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# **GEOSX Documentation**

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## Contents

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<b>1</b>	<b>Quick Start Guide</b>	<b>3</b>
1.1	Frequently Asked Questions . . . . .	3
1.2	Repository Organization . . . . .	4
1.3	Username and Authentication . . . . .	5
1.4	Download . . . . .	5
1.5	Configuration . . . . .	7
1.6	Compilation . . . . .	8
1.7	Running . . . . .	9
1.8	Testing . . . . .	10
<b>2</b>	<b>Tutorials</b>	<b>11</b>
2.1	Tutorial 1: First Steps . . . . .	11
2.2	Tutorial 2: External Meshes . . . . .	24
2.3	Tutorial 3: Regions and Property Specifications . . . . .	33
2.4	Tutorial 4: Boundary Conditions and Time-Dependent Functions . . . . .	44
<b>3</b>	<b>Basic Examples</b>	<b>49</b>
3.1	Multiphase Flow . . . . .	49
3.2	Multiphase Flow with Wells . . . . .	58
3.3	CO <sub>2</sub> Injection . . . . .	71
3.4	Poromechanics . . . . .	80
3.5	Hydraulic Fracturing . . . . .	85
3.6	Triaxial Driver . . . . .	99
<b>4</b>	<b>Advanced Examples</b>	<b>107</b>
4.1	Validation and Verification Studies . . . . .	107
4.2	Performance Benchmarks . . . . .	275
4.3	Application Studies . . . . .	275
4.4	pygeosx Examples . . . . .	275
<b>5</b>	<b>User Guide</b>	<b>287</b>
5.1	Input Files . . . . .	287
5.2	Meshes . . . . .	294
5.3	Physics Solvers . . . . .	301
5.4	Constitutive Models . . . . .	329
5.5	Initial and Boundary Conditions . . . . .	385
5.6	Event Management . . . . .	396

5.7	Tasks Manager	402
5.8	Functions	403
5.9	Linear Solvers	408
5.10	Numerical Methods	412
5.11	Parallel Partitioning	414
5.12	Outputs	415
5.13	pygeosx — GEOSX in Python	420
5.14	Indices and tables	422
<b>6</b>	<b>Developer Guide</b>	<b>423</b>
6.1	Contributing	423
6.2	Code Components	461
<b>7</b>	<b>Doxygen</b>	<b>527</b>
<b>8</b>	<b>Python Tools</b>	<b>529</b>
8.1	Python Tools Setup	529
8.2	Packages	530
<b>9</b>	<b>Build Guide</b>	<b>549</b>
9.1	System prerequisites	549
9.2	Third-party dependencies	550
9.3	Building GEOSX	553
9.4	Spack and Uberenv	554
9.5	Continuous Integration process	557
<b>10</b>	<b>Datastructure Index</b>	<b>561</b>
10.1	Input Schema Definitions	561
10.2	Element: AcousticSEM	562
10.3	Element: Aquifer	564
10.4	Element: Benchmarks	565
10.5	Element: BiotPorosity	565
10.6	Element: BlackOilFluid	566
10.7	Element: Blueprint	567
10.8	Element: BoundedPlane	567
10.9	Element: Box	567
10.10	Element: BrooksCoreyBakerRelativePermeability	568
10.11	Element: BrooksCoreyCapillaryPressure	569
10.12	Element: BrooksCoreyRelativePermeability	569
10.13	Element: CO2BrineEzrokhiFluid	569
10.14	Element: CO2BrineEzrokhiThermalFluid	570
10.15	Element: CO2BrinePhillipsFluid	570
10.16	Element: CO2BrinePhillipsThermalFluid	570
10.17	Element: CarmanKozenyPermeability	571
10.18	Element: CellElementRegion	571
10.19	Element: ChomboIO	571
10.20	Element: CompositeFunction	572
10.21	Element: CompositionalMultiphaseFVM	574
10.22	Element: CompositionalMultiphaseFluid	575
10.23	Element: CompositionalMultiphaseHybridFVM	577
10.24	Element: CompositionalMultiphaseReservoir	578
10.25	Element: CompositionalMultiphaseStatistics	578
10.26	Element: CompositionalMultiphaseWell	579
10.27	Element: CompressibleSinglePhaseFluid	580
10.28	Element: CompressibleSolidCarmanKozenyPermeability	580



10.29 Element: CompressibleSolidConstantPermeability . . . . .	581
10.30 Element: CompressibleSolidParallelPlatesPermeability . . . . .	581
10.31 Element: CompressibleSolidSlipDependentPermeability . . . . .	581
10.32 Element: ConstantPermeability . . . . .	581
10.33 Element: Constitutive . . . . .	581
10.34 Element: Coulomb . . . . .	583
10.35 Element: Cylinder . . . . .	584
10.36 Element: DamageElasticIsotropic . . . . .	584
10.37 Element: DamageSpectralElasticIsotropic . . . . .	584
10.38 Element: DamageVolDevElasticIsotropic . . . . .	585
10.39 Element: DeadOilFluid . . . . .	586
10.40 Element: DelftEgg . . . . .	587
10.41 Element: Dirichlet . . . . .	587
10.42 Element: DruckerPrager . . . . .	588
10.43 Element: ElasticIsotropic . . . . .	588
10.44 Element: ElasticIsotropicPressureDependent . . . . .	588
10.45 Element: ElasticOrthotropic . . . . .	589
10.46 Element: ElasticSEM . . . . .	590
10.47 Element: ElasticTransverseIsotropic . . . . .	591
10.48 Element: ElementRegions . . . . .	591
10.49 Element: EmbeddedSurfaceGenerator . . . . .	592
10.50 Element: Events . . . . .	592
10.51 Element: ExtendedDruckerPrager . . . . .	593
10.52 Element: FieldSpecification . . . . .	593
10.53 Element: FieldSpecifications . . . . .	594
10.54 Element: File . . . . .	594
10.55 Element: FiniteElementSpace . . . . .	594
10.56 Element: FiniteElements . . . . .	594
10.57 Element: FiniteVolume . . . . .	595
10.58 Element: FlowProppantTransport . . . . .	595
10.59 Element: FrictionlessContact . . . . .	596
10.60 Element: Functions . . . . .	596
10.61 Element: Geometry . . . . .	596
10.62 Element: HaltEvent . . . . .	597
10.63 Element: HybridMimeticDiscretization . . . . .	597
10.64 Element: Hydrofracture . . . . .	599
10.65 Element: HydrostaticEquilibrium . . . . .	600
10.66 Element: Included . . . . .	600
10.67 Element: InternalMesh . . . . .	601
10.68 Element: InternalWell . . . . .	602
10.69 Element: InternalWellbore . . . . .	603
10.70 Element: JFunctionCapillaryPressure . . . . .	605
10.71 Element: LagrangianContact . . . . .	606
10.72 Element: LaplaceFEM . . . . .	608
10.73 Element: LinearSolverParameters . . . . .	610
10.74 Element: Mesh . . . . .	611
10.75 Element: ModifiedCamClay . . . . .	611
10.76 Element: MultiPhaseConstantThermalConductivity . . . . .	611
10.77 Element: MultiPhaseVolumeWeightedThermalConductivity . . . . .	612
10.78 Element: MultiphasePoromechanics . . . . .	612
10.79 Element: MultiphasePoromechanicsReservoir . . . . .	613
10.80 Element: MultivariableTableFunction . . . . .	613
10.81 Element: NonlinearSolverParameters . . . . .	615
10.82 Element: NullModel . . . . .	616

10.83	Element: NumericalMethods	616
10.84	Element: Outputs	616
10.85	Element: PAMELAMesh	616
10.86	Element: PML	617
10.87	Element: PVTDriver	617
10.88	Element: PackCollection	618
10.89	Element: ParallelPlatesPermeability	618
10.90	Element: Parameter	618
10.91	Element: Parameters	618
10.92	Element: ParticleFluid	619
10.93	Element: Perforation	619
10.94	Element: PeriodicEvent	620
10.95	Element: PermeabilityBase	620
10.96	Element: PhaseFieldDamageFEM	621
10.97	Element: PhaseFieldFracture	623
10.98	Element: PorousDelftEgg	624
10.99	Element: PorousDruckerPrager	624
10.100	Element: PorousElasticIsotropic	624
10.101	Element: PorousElasticOrthotropic	624
10.102	Element: PorousElasticTransverseIsotropic	625
10.103	Element: PorousExtendedDruckerPrager	625
10.104	Element: PorousModifiedCamClay	625
10.105	Element: PressurePorosity	625
10.106	Element: Problem	626
10.107	Element: ProppantPermeability	626
10.108	Element: ProppantPorosity	626
10.109	Element: ProppantSlurryFluid	627
10.110	Element: ProppantSolidProppantPermeability	627
10.111	Element: ProppantTransport	629
10.112	Element: Python	630
10.113	Element: ReactiveCompositionalMultiphaseOBL	632
10.114	Element: Restart	633
10.115	Element: Run	633
10.116	Element: Silo	634
10.117	Element: SinglePhaseConstantThermalConductivity	634
10.118	Element: SinglePhaseFVM	635
10.119	Element: SinglePhaseHybridFVM	636
10.120	Element: SinglePhasePoromechanics	637
10.121	Element: SinglePhasePoromechanicsEmbeddedFractures	638
10.122	Element: SinglePhasePoromechanicsReservoir	639
10.123	Element: SinglePhaseProppantFVM	640
10.124	Element: SinglePhaseReservoir	641
10.125	Element: SinglePhaseStatistics	641
10.126	Element: SinglePhaseWell	642
10.127	Element: SlipDependentPermeability	642
10.128	Element: SolidInternalEnergy	642
10.129	Element: SolidMechanicsEmbeddedFractures	643
10.130	Element: SolidMechanicsLagrangianSSLE	645
10.131	Element: SolidMechanicsStateReset	646
10.132	Element: SolidMechanicsStatistics	646
10.133	Element: SolidMechanics_LagrangianFEM	648
10.134	Element: SoloEvent	649
10.135	Element: Solvers	649
10.136	Element: SourceFlux	650

10.137Element: SurfaceElementRegion . . . . .	651
10.138Element: SurfaceGenerator . . . . .	652
10.139Element: SymbolicFunction . . . . .	653
10.140Element: TableCapillaryPressure . . . . .	655
10.141Element: TableFunction . . . . .	656
10.142Element: TableRelativePermeability . . . . .	658
10.143Element: TableRelativePermeabilityHysteresis . . . . .	660
10.144Element: Tasks . . . . .	661
10.145Element: ThermalCompressibleSinglePhaseFluid . . . . .	662
10.146Element: ThickPlane . . . . .	663
10.147Element: TimeHistory . . . . .	663
10.148Element: Traction . . . . .	664
10.149Element: TriaxialDriver . . . . .	665
10.150Element: TwoPointFluxApproximation . . . . .	665
10.151Element: VTK . . . . .	665
10.152Element: VTKMesh . . . . .	666
10.153Element: VanGenuchtenBakerRelativePermeability . . . . .	667
10.154Element: VanGenuchtenCapillaryPressure . . . . .	668
10.155Element: WellControls . . . . .	670
10.156Element: WellElementRegion . . . . .	671
10.157Element: lassen . . . . .	671
10.158Element: quartz . . . . .	671
10.159Datastructure: AcousticSEM . . . . .	672
10.160Datastructure: Aquifer . . . . .	673
10.161Datastructure: Benchmarks . . . . .	673
10.162Datastructure: BiotPorosity . . . . .	673
10.163Datastructure: BlackOilFluid . . . . .	675
10.164Datastructure: Blueprint . . . . .	676
10.165Datastructure: BoundedPlane . . . . .	676
10.166Datastructure: Box . . . . .	676
10.167Datastructure: BrooksCoreyBakerRelativePermeability . . . . .	676
10.168Datastructure: BrooksCoreyCapillaryPressure . . . . .	677
10.169Datastructure: BrooksCoreyRelativePermeability . . . . .	677
10.170Datastructure: CO2BrineEzrokhiFluid . . . . .	679
10.171Datastructure: CO2BrineEzrokhiThermalFluid . . . . .	681
10.172Datastructure: CO2BrinePhillipsFluid . . . . .	683
10.173Datastructure: CO2BrinePhillipsThermalFluid . . . . .	685
10.174Datastructure: CarmanKozenyPermeability . . . . .	686
10.175Datastructure: CellElementRegion . . . . .	686
10.176Datastructure: ChomboIO . . . . .	686
10.177Datastructure: CompositeFunction . . . . .	686
10.178Datastructure: CompositionalMultiphaseFVM . . . . .	687
10.179Datastructure: CompositionalMultiphaseFluid . . . . .	689
10.180Datastructure: CompositionalMultiphaseHybridFVM . . . . .	690
10.181Datastructure: CompositionalMultiphaseReservoir . . . . .	691
10.182Datastructure: CompositionalMultiphaseStatistics . . . . .	691
10.183Datastructure: CompositionalMultiphaseWell . . . . .	692
10.184Datastructure: CompressibleSinglePhaseFluid . . . . .	693
10.185Datastructure: CompressibleSolidCarmanKozenyPermeability . . . . .	693
10.186Datastructure: CompressibleSolidConstantPermeability . . . . .	693
10.187Datastructure: CompressibleSolidParallelPlatesPermeability . . . . .	693
10.188Datastructure: CompressibleSolidSlipDependentPermeability . . . . .	693
10.189Datastructure: ConstantPermeability . . . . .	694
10.190Datastructure: Constitutive . . . . .	694

10.191	Datastructure: ConstitutiveModels	695
10.192	Datastructure: Coulomb	695
10.193	Datastructure: Cylinder	695
10.194	Datastructure: DamageElasticIsotropic	696
10.195	Datastructure: DamageSpectralElasticIsotropic	696
10.196	Datastructure: DamageVolDevElasticIsotropic	696
10.197	Datastructure: DeadOilFluid	698
10.198	Datastructure: DelftEgg	699
10.199	Datastructure: Dirichlet	699
10.200	Datastructure: DruckerPrager	699
10.201	Datastructure: ElasticIsotropic	699
10.202	Datastructure: ElasticIsotropicPressureDependent	700
10.203	Datastructure: ElasticOrthotropic	700
10.204	Datastructure: ElasticSEM	701
10.205	Datastructure: ElasticTransverseIsotropic	702
10.206	Datastructure: ElementRegions	702
10.207	Datastructure: EmbeddedSurfaceGenerator	703
10.208	Datastructure: Events	703
10.209	Datastructure: ExtendedDruckerPrager	704
10.210	Datastructure: FieldSpecification	704
10.211	Datastructure: FieldSpecifications	704
10.212	Datastructure: File	704
10.213	Datastructure: FiniteElementSpace	704
10.214	Datastructure: FiniteElements	705
10.215	Datastructure: FiniteVolume	705
10.216	Datastructure: FlowProppantTransport	705
10.217	Datastructure: FrictionlessContact	706
10.218	Datastructure: Functions	706
10.219	Datastructure: Geometry	706
10.220	Datastructure: HaltEvent	706
10.221	Datastructure: HybridMimeticDiscretization	706
10.222	Datastructure: Hydrofracture	707
10.223	Datastructure: HydrostaticEquilibrium	707
10.224	Datastructure: Included	707
10.225	Datastructure: InternalMesh	708
10.226	Datastructure: InternalWell	708
10.227	Datastructure: InternalWellbore	708
10.228	Datastructure: JFunctionCapillaryPressure	708
10.229	Datastructure: LagrangianContact	709
10.230	Datastructure: LaplaceFEM	709
10.231	Datastructure: Level0	709
10.232	Datastructure: LinearSolverParameters	710
10.233	Datastructure: Mesh	710
10.234	Datastructure: MeshBodies	710
10.235	Datastructure: ModifiedCamClay	710
10.236	Datastructure: MultiPhaseConstantThermalConductivity	711
10.237	Datastructure: MultiPhaseVolumeWeightedThermalConductivity	711
10.238	Datastructure: MultiphasePoromechanics	711
10.239	Datastructure: MultiphasePoromechanicsReservoir	712
10.240	Datastructure: MultivariableTableFunction	712
10.241	Datastructure: NonlinearSolverParameters	712
10.242	Datastructure: NullModel	713
10.243	Datastructure: NumericalMethods	713
10.244	Datastructure: Outputs	713

10.245	Datastructure: PAMELAMesh . . . . .	713
10.246	Datastructure: PML . . . . .	713
10.247	Datastructure: PVTDriver . . . . .	713
10.248	Datastructure: PackCollection . . . . .	714
10.249	Datastructure: ParallelPlatesPermeability . . . . .	714
10.250	Datastructure: Parameter . . . . .	714
10.251	Datastructure: Parameters . . . . .	714
10.252	Datastructure: ParticleFluid . . . . .	714
10.253	Datastructure: Perforation . . . . .	714
10.254	Datastructure: PeriodicEvent . . . . .	715
10.255	Datastructure: PermeabilityBase . . . . .	715
10.256	Datastructure: PhaseFieldDamageFEM . . . . .	715
10.257	Datastructure: PhaseFieldFracture . . . . .	716
10.258	Datastructure: PorousDelftEgg . . . . .	716
10.259	Datastructure: PorousDruckerPrager . . . . .	716
10.260	Datastructure: PorousElasticIsotropic . . . . .	716
10.261	Datastructure: PorousElasticOrthotropic . . . . .	716
10.262	Datastructure: PorousElasticTransverseIsotropic . . . . .	717
10.263	Datastructure: PorousExtendedDruckerPrager . . . . .	717
10.264	Datastructure: PorousModifiedCamClay . . . . .	717
10.265	Datastructure: PressurePorosity . . . . .	717
10.266	Datastructure: Problem . . . . .	718
10.267	Datastructure: ProppantPermeability . . . . .	718
10.268	Datastructure: ProppantPorosity . . . . .	718
10.269	Datastructure: ProppantSlurryFluid . . . . .	719
10.270	Datastructure: ProppantSolidProppantPermeability . . . . .	719
10.271	Datastructure: ProppantTransport . . . . .	720
10.272	Datastructure: Python . . . . .	720
10.273	Datastructure: ReactiveCompositionalMultiphaseOBL . . . . .	720
10.274	Datastructure: Restart . . . . .	720
10.275	Datastructure: Run . . . . .	721
10.276	Datastructure: Silo . . . . .	721
10.277	Datastructure: SinglePhaseConstantThermalConductivity . . . . .	721
10.278	Datastructure: SinglePhaseFVM . . . . .	721
10.279	Datastructure: SinglePhaseHybridFVM . . . . .	722
10.280	Datastructure: SinglePhasePoromechanics . . . . .	723
10.281	Datastructure: SinglePhasePoromechanicsEmbeddedFractures . . . . .	724
10.282	Datastructure: SinglePhasePoromechanicsReservoir . . . . .	725
10.283	Datastructure: SinglePhaseProppantFVM . . . . .	725
10.284	Datastructure: SinglePhaseReservoir . . . . .	726
10.285	Datastructure: SinglePhaseStatistics . . . . .	726
10.286	Datastructure: SinglePhaseWell . . . . .	727
10.287	Datastructure: SlipDependentPermeability . . . . .	727
10.288	Datastructure: SolidInternalEnergy . . . . .	728
10.289	Datastructure: SolidMechanicsEmbeddedFractures . . . . .	728
10.290	Datastructure: SolidMechanicsLagrangianSSLE . . . . .	729
10.291	Datastructure: SolidMechanicsStateReset . . . . .	729
10.292	Datastructure: SolidMechanicsStatistics . . . . .	729
10.293	Datastructure: SolidMechanics_LagrangianFEM . . . . .	730
10.294	Datastructure: SoloEvent . . . . .	730
10.295	Datastructure: SolverStatistics . . . . .	730
10.296	Datastructure: Solvers . . . . .	731
10.297	Datastructure: SourceFlux . . . . .	731
10.298	Datastructure: SurfaceElementRegion . . . . .	732

10.299	Datastructure: SurfaceGenerator . . . . .	733
10.300	Datastructure: SymbolicFunction . . . . .	733
10.301	Datastructure: TableCapillaryPressure . . . . .	734
10.302	Datastructure: TableFunction . . . . .	734
10.303	Datastructure: TableRelativePermeability . . . . .	734
10.304	Datastructure: TableRelativePermeabilityHysteresis . . . . .	735
10.305	Datastructure: Tasks . . . . .	736
10.306	Datastructure: ThermalCompressibleSinglePhaseFluid . . . . .	736
10.307	Datastructure: ThickPlane . . . . .	736
10.308	Datastructure: TimeHistory . . . . .	736
10.309	Datastructure: Traction . . . . .	737
10.310	Datastructure: TriaxialDriver . . . . .	737
10.311	Datastructure: TwoPointFluxApproximation . . . . .	737
10.312	Datastructure: VTK . . . . .	737
10.313	Datastructure: VTKMesh . . . . .	737
10.314	Datastructure: VanGenuchtenBakerRelativePermeability . . . . .	738
10.315	Datastructure: VanGenuchtenCapillaryPressure . . . . .	738
10.316	Datastructure: WellControls . . . . .	738
10.317	Datastructure: WellElementRegion . . . . .	739
10.318	Datastructure: WellElementRegionUniqueSubRegion . . . . .	740
10.319	Datastructure: commandLine . . . . .	741
10.320	Datastructure: domain . . . . .	741
10.321	Datastructure: edgeManager . . . . .	742
10.322	Datastructure: elementRegionsGroup . . . . .	742
10.323	Datastructure: elementSubRegions . . . . .	742
10.324	Datastructure: embeddedSurfacesEdgeManager . . . . .	743
10.325	Datastructure: embeddedSurfacesNodeManager . . . . .	744
10.326	Datastructure: faceManager . . . . .	745
10.327	Datastructure: lassen . . . . .	746
10.328	Datastructure: meshLevels . . . . .	746
10.329	Datastructure: neighborData . . . . .	746
10.330	Datastructure: nodeManager . . . . .	746
10.331	Datastructure: quartz . . . . .	747
10.332	Datastructure: sets . . . . .	747
10.333	Datastructure: wellElementSubRegion . . . . .	747
<b>11</b>	<b>Contributors</b>	<b>749</b>
<b>12</b>	<b>Publications</b>	<b>751</b>
12.1	Preprints and Early-Views . . . . .	751
12.2	2022 . . . . .	753
12.3	2021 . . . . .	755
12.4	2020 . . . . .	756
12.5	2019 . . . . .	757
<b>13</b>	<b>Acknowledgements</b>	<b>759</b>
<b>14</b>	<b>Indices and tables</b>	<b>761</b>
	<b>Python Module Index</b>	<b>763</b>
	<b>Index</b>	<b>765</b>

Welcome to our documentation pages!

GEOSX is a code framework focused on enabling streamlined development of physics simulations on high performance computing platforms. Our documentation is organized into several separate guides, given that different users will have different needs.

We recommend all users begin with the Quick Start guide, which will walk you through downloading and compiling the code. Application focused users may then want to explore our Tutorials, which provide an introduction to the basic capabilities of the code. More detailed descriptions of these capabilities can then be found in the User Guide.

For those interested in developing new capabilities in GEOSX, we provide a Developer Guide. The code itself is also documented inline using doxygen. The Build Guide contains more detailed information about third-party dependencies, the build system, and the continuous integration system. Finally, GEOSX has a self-documenting data structure. The Datastructure Index is an automatically generated list of all available input parameters and data structures in the code. This is a comprehensive resource, but probably not the place to start.

High quality documentation is a critical component of a successful code. If you have suggestions for improving the guides below, please post an issue on our [issue tracker](#).





The goal of this page is to get you started as quickly as possible using GEOSX. We will walk you through downloading the source, compiling the code, and testing the installation.

Before jumping to the installation process, we want to first address some frequently asked questions we get from new users. If you are itching to get started, feel free to jump ahead to the relevant sections.

## 1.1 Frequently Asked Questions

### 1.1.1 Does GEOSX have a graphical user interface?:

Given the focus on rapid development and HPC environments, GEOSX does not have a graphical user interface. This is consistent with many other high performance computing packages, but we recognize it can be a deal-breaker for certain users. For those who can get past this failing, we promise we still have a lot to offer. In a typical workflow, you will prepare an XML-based input file describing your problem. You may also prepare a mesh file containing geometric and property information describing, say, a reservoir you would like to simulate. There is no shortage of GUI tools that can help you in this model building stage. The resulting input deck is then consumed by GEOSX to run the simulation and produce results. This may be done in a terminal of your local machine or by submitting a job to a remote server. The resulting output files can then be visualized by any number of graphical visualization programs (typically [VisIt](#) or [paraview](#)). Thus, while GEOSX is GUI free, the typical workflow is not.

### 1.1.2 Do I need to be a code developer to use GEOSX?:

For the moment, most users will need to download and compile the code from source, which we readily admit this requires a certain level of development expertise. We try to make this process as easy as possible, and we are working on additional deployment options to make this process easier. Once installed, however, our goal is to make GEOSX accessible to developers and non-developers alike. Our target audience includes engineers and scientists who want to solve tough application problems, but could care less about the insides of the tool. For those of you who *are* interested in scientific computing, however, GEOSX is an open source project and we welcome external contributions.

### 1.1.3 What are the system requirements?:

GEOSX is primarily written in C++, with a focus on standards compliance and platform-to-platform portability. It is designed to run on everything from commodity laptops to the world's most powerful supercomputers. We regularly test the code across a variety of operating systems and compilers. Most of these operating systems are Linux/UNIX based (e.g. Ubuntu, CentOS, Mac OSX). We do have developers working in Windows environments, but they use a Virtual Machine or work within a docker image rather than directly in the Windows environment. In the instructions below, we assume you have access to fairly standard development tools. Using advanced features of GEOSX, like GPU-acceleration, will of course introduce additional hardware and software requirements.

### 1.1.4 Help, I get errors while trying to download/compile/run!:

Unfortunately, no set of instructions is foolproof. It is simply impossible to anticipate every system configuration or user. If you run into problems during the installation, we recommend the following five-step process:

1. Take a moment to relax, and then re-read the instructions carefully. Perhaps you overlooked a key step? Re-read the error message(s) closely. Modern compilation tools are often quite helpful in reporting exactly why things fail.
2. Type a few keywords from your error into a search engine. It is possible someone else out there has encountered your problem before, and a well-chosen keyword can often produce an instant solution. Note that when a compilation fails, you may get pages and pages of errors. Try to identify the *first* one to occur and fix that. One error will often trigger subsequent errors, and looking at the *last* error on the screen may not be so helpful.
3. If you encounter problems building one of the third-party libraries we depend on, check out their support pages. They may be able to help you more directly than we can.
4. Still stuck? Check out our [issues tracker](#), searching current or closed issues that may address your problem. Perhaps someone has had an identical issue, or something close. The issue tracker has a convenient search bar where you can search for relevant keywords. Remember to remove the default `is:open` keyword to search both open and closed issues.
5. If you have exhausted the options above, it is time to seek help from the developers. Post an issue on our issue tracker. Be specific, providing as much information as possible about your system setup and the error you are encountering. Please be patient in this process, as we may need to correspond a few times and ask you to run additional tests. Most of the time, users have a slightly unusual system configuration that we haven't encountered yet, such as an older version of a particular library. Other times there is a legitimate bug in GEOSX to be addressed. Take pride in the fact that you may be saving the next user from wasted time and frustration.

## 1.2 Repository Organization

The source for GEOSX and related tools are hosted on [Github](#). We use [Git workflows](#) to version control our code and manage the entire development process. On Github, we have a [GEOSX Organization](#) that hosts several related repositories.

You should sign up for a free Github account, particularly if you are interested in posting issues to our issue tracker and communicating with the developers. The main repository of interest is obviously GEOSX itself: [GEOSX](#)

We also rely on two types of dependencies: first-party and third-party. First-party dependencies are projects directly associated with the GEOSX effort, but kept in separate repositories because they form stand-alone tools. For example, there is a geologic mesh handling package called [PAMELA](#) and an equation-of-state package called [PVTPackage](#). These packages are handled as [Git Submodules](#), which provides a transparent way of coordinating multiple code development projects. Most users will never have to worry that these modules are in fact separate projects from GEOSX.

We also rely on several open-source Third-Party Libraries (TPLs) (see [thirdPartyLibs](#)). These are well-respected projects developed externally to GEOSX. We have found, however, that many compilation issues stem from version incompatibilities between different packages. To address this, we provide a mirror of these TPLs, with version combinations we know play nicely together. We also provide a build script that conveniently and consistently builds those dependencies.

Our build system will automatically use the mirror package versions by default. You are welcome to tune your configuration, however, to point to different versions installed on your system. If you work on an HPC platform, for example, common packages may already be available and optimized for platform hardware. For new users, however, it may be safer to begin with the TPL mirror.

---

**Note:** If you are working on an HPC platform with several other GEOSX users, we often compile the TPLs in a shared location so individual users don't have to waste their storage quota. Inquire with your institution's point-of-contact whether this option already exists. For all LLNL systems, the answer is yes.

---

Finally, there are also several private repositories only accessible to the core development team, which we use for behind-the-scene testing and maintenance of the code.

## 1.3 Username and Authentication

New users should sign up for a free [Github account](#).

If you intend to develop in the GEOSX codebase, you may benefit from setting up your git credentials (see [Git Workflow](#)).

## 1.4 Download

It is possible to directly download the source code as a zip file. We strongly suggest, however, that users don't rely on this option. Instead, most users should use Git to either *clone* or *fork* the repository. This makes it much easier to stay up to date with the latest releases and bug fixes. If you are not familiar with the basics of Git, [here is a helpful resource](#) to get you started.

The tutorial here assumes you will use a https clone with no specific credentials. Using an ssh connection pattern requires a very slight modification. See the **Additional Notes** at the end of this section for details.

If you do not already have Git installed on your system, you will need to install it. We recommend using a relatively recent version of Git, as there have been some notable improvements over the past few years. You can check if Git is already available by opening a terminal and typing

```
git --version
```

You'll also need the `git-lfs` large file extension.

The first task is to clone the GEOSX and `thirdPartyLibs` repositories. If you do not tell it otherwise, the build system will expect the GEOSX and `thirdPartyLibs` to be parallel to each other in the directory structure. For example,

```
codes/
├── GEOSX/
└── thirdPartyLibs/
```

where the toplevel `codes` directory can be re-named and located wherever you like. It is possible to customize the build system to expect a different structure, but for now let us assume you take the simplest approach.

First, using a terminal, create the `codes` directory wherever you like.

```
cd /insert/your/desired/path/  
mkdir codes  
cd codes
```

Inside this directory, we can clone the GEOSX repository. We will also use some Git commands to initialize and download the submodules (e.g. PAMELA). Note that most users will not have access to our integrated tests repository, and so we “deinit” (deactivate) this submodule. Developers who will be working with the integratedTests repository should skip this line.

```
git clone https://github.com/GEOSX/GEOSX.git  
cd GEOSX  
git lfs install  
git submodule init  
git submodule deinit integratedTests  
git submodule update  
cd ..
```

If all goes well, you should have a complete copy of the GEOSX source at this point. The most common errors people encounter here have to do with Github not recognizing their authentication settings and/or repository permissions. See the previous section for tips on ensuring your SSH is working properly.

*Note:* The integratedTests submodule is not publicly available, with access limited to the core development team. This may cause the `git submodule update` command to fail if you forget the `git submodule deinit integratedTests` step above. This submodule is not required for building GEOSX. If you see an error message here, however, you may need to initialize and update the submodules manually:

```
cd GEOSX  
git submodule update --init src/cmake/blt  
git submodule update --init src/coreComponents/LvArray  
git submodule update --init src/coreComponents/fileIO/coupling/hdf5_interface  
git submodule update --init src/coreComponents/mesh/PAMELA  
git submodule update --init src/coreComponents/constitutive/PVTPackage  
cd ..
```

Once we have grabbed GEOSX, we do the same for the thirdPartyLibs repository. From the `codes` directory, type

```
git clone https://github.com/GEOSX/thirdPartyLibs.git  
cd thirdPartyLibs  
git lfs install  
git pull  
git submodule init  
git submodule update  
cd ..
```

Again, if all goes well you should now have a copy of all necessary TPL packages.

### Additional Notes:

#. `git-lfs` may not function properly (or may be very slow) if your version of `git` and `git-lfs` are not current. If you are using an older version, you may need to add `git lfs pull` after `git pull` in the above procedures.

#. You can adapt the commands if you use an ssh connection instead. The `clone https://github.com/GEOSX/GEOSX.git` becomes `git clone git@github.com:GEOSX/GEOSX.git`. You may also be willing to insert your credentials in the command line (less secure) `git clone https://${USER}:${TOKEN}@github.com/GEOSX/GEOSX.git`.

## 1.5 Configuration

At a minimum, you will need a relatively recent compiler suite installed on your system (e.g. [GCC](#), [Clang](#)) as well as [CMake](#). If you want to run jobs using MPI-based parallelism, you will also need an MPI implementation (e.g. [OpenMPI](#), [MVAPICH](#)). Note that GEOSX supports a variety of parallel computing models, depending on the hardware and software environment. Advanced users are referred to the [Build Guide](#) for a discussion of the available configuration options.

Before beginning, it is a good idea to have a clear idea of the flavor and version of the build tools you are using. If something goes wrong, the first thing the support team will ask you for is this information.

```
cpp --version
mpic++ --version
cmake --version
```

Here, you may need to replace `cpp` with the full path to the C++ compiler you would like to use, depending on how your path and any aliases are configured.

GEOSX compilations are driven by a `cmake host-config` file, which tells the build system about the compilers you are using, where various packages reside, and what options you want to enable. We have created a number of default `hostconfig` files for common systems. You should browse them to see if any are close to your needs:

```
cd GEOSX/host-configs
```

We maintain `host configs` (ending in `.cmake`) for HPC systems at various institutions, as well as ones for common personal systems. If you cannot find one that matches your needs, we suggest beginning with one of the shorter ones and modifying as needed. A typical one may look like:

```
# file: your-platform.cmake

# detect host and name the configuration file
site_name(HOST_NAME)
set(CONFIG_NAME "your-platform" CACHE PATH "")
message("CONFIG_NAME = ${CONFIG_NAME}")

# set paths to C, C++, and Fortran compilers. Note that while GEOSX does not contain
↳ any Fortran code,
# some of the third-party libraries do contain Fortran code. Thus a Fortran compiler
↳ must be specified.
set(CMAKE_C_COMPILER "/usr/bin/clang" CACHE PATH "")
set(CMAKE_CXX_COMPILER "/usr/bin/clang++" CACHE PATH "")
set(CMAKE_Fortran_COMPILER "/usr/local/bin/gfortran" CACHE PATH "")
set(ENABLE_Fortran OFF CACHE BOOL "" FORCE)

# enable MPI and set paths to compilers and executable.
# Note that the MPI compilers are wrappers around standard serial compilers.
# Therefore, the MPI compilers must wrap the appropriate serial compilers specified
# in CMAKE_C_COMPILER, CMAKE_CXX_COMPILER, and CMAKE_Fortran_COMPILER.
set(ENABLE_MPI ON CACHE BOOL "")
set(MPI_C_COMPILER "/usr/local/bin/mpicc" CACHE PATH "")
set(MPI_CXX_COMPILER "/usr/local/bin/mpicxx" CACHE PATH "")
set(MPI_Fortran_COMPILER "/usr/local/bin/mpifort" CACHE PATH "")
set(MPIEXEC "/usr/local/bin/mpirun" CACHE PATH "")

# disable CUDA and OpenMP
set(CUDA_ENABLED OFF CACHE BOOL "" FORCE)
set(ENABLE_OPENMP OFF CACHE BOOL "" FORCE)
```

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```
# enable PAMELA and PVTPackage
set(ENABLE_PAMELA ON CACHE BOOL "" FORCE)
set(ENABLE_PVTPackage ON CACHE BOOL "" FORCE)

# enable tests
set(ENABLE_GTEST_DEATH_TESTS ON CACHE BOOL "" FORCE )

# define the path to your compiled installation directory
set(GEOSX_TPL_DIR "/path/to/your/TPL/installation/dir" CACHE PATH "")
# let GEOSX define some third party libraries information for you
include(${CMAKE_CURRENT_LIST_DIR}/tpls.cmake)
```

The various `set()` commands are used to set environment variables that control the build. You will see in the above example that we set the C++ compiler to `/user/bin/clang++` and so forth. We also disable CUDA and OpenMP, but enable PAMELA and PVTPackage. The final line is related to our unit test suite. See the [Build Guide](#) for more details on available options.

---

**Note:** If you develop a new `host-config` for a particular platform that may be useful for other users, please consider sharing it with the developer team.

---

## 1.6 Compilation

We will begin by compiling the TPLs, followed by the main code. If you work on an HPC system with other GEOSX developers, check with them to see if the TPLs have already been compiled in a shared directory. If this is the case, you can skip ahead to just compiling the main code. If you are working on your own machine, you will need to compile both.

We strongly suggest that GEOSX and TPLs be built with the same `hostconfig` file. Below, we assume that you keep it in, say, `GEOSX/host-configs/your-platform.cmake`, but this is up to you.

We begin with the third-party libraries, and use a `python config-build.py` script to configure and build all of the TPLs. Note that we will request a Release build type, which will enable various optimizations. The other option is a Debug build, which allows for debugging but will be much slower in production mode. The TPLs will then be built in a build directory named consistently with your `hostconfig` file.

```
cd thirdPartyLibs
python scripts/config-build.py -hc ../GEOSX/host-configs/your-platform.cmake -bt Release
cd build-your-platform-release
make
```

Note that building all of the TPLs can take quite a while, so you may want to go get a cup of coffee at this point. Also note that you should *not* use a `parallel make -j N` command to try and speed up the build time.

The next step is to compile the main code. Again, the `config-build.py` sets up `cmake` for you, so the process is very similar.

```
cd ../../GEOSX
python scripts/config-build.py -hc host-configs/your-platform.cmake -bt Release
cd build-your-platform-release
make -j4
make install
```

The host-config file is the place to set all relevant configuration options. Note that the path to the previously installed third party libraries is typically specified within this file. An alternative is to set the path `GEOSX_TPL_DIR` via a `cmake` command line option, e.g.

```
python scripts/config-build.py -hc host-configs/your-platform.cmake -bt Release -D_
↳GEOSX_TPL_DIR=/full/path/to/thirdPartyLibs
```

We highly recommend using full paths, rather than relative paths, whenever possible. The parallel make `-j 4` will use four processes for compilation, which can substantially speed up the build if you have a multi-processor machine. You can adjust this value to match the number of processors available on your machine. The `make install` command then installs GEOSX to a default location unless otherwise specified.

If all goes well, a `geosx` executable should now be available:

```
GEOSX/install-your-platform-release/bin/geosx
```

## 1.7 Running

We can do a quick check that the `geosx` executable is working properly by calling the executable with our help flag

```
./bin/geosx --help
```

This should print out a brief summary of the available command line arguments:

```
USAGE: geosx -i input.xml [options]

Options:
-?, --help
-i, --input,           Input xml filename (required)
-r, --restart,         Target restart filename
-x, --x-partitions,    Number of partitions in the x-direction
-y, --y-partitions,    Number of partitions in the y-direction
-z, --z-partitions,    Number of partitions in the z-direction
-s, --schema,          Name of the output schema
-b, --use-nonblocking, Use non-blocking MPI communication
-n, --name,            Name of the problem, used for output
-s, --suppress-pinned, Suppress usage of pinned memory for MPI communication buffers
-o, --output,          Directory to put the output files
-t, --timers,          String specifying the type of timer output
--trace-data-migration, Trace host-device data migration
--pause-for,           Pause geosx for a given number of seconds before starting_
↳execution
```

Obviously this doesn't do much interesting, but it will at least confirm that the executable runs. In typical usage, an input XML must be provided describing the problem to be run, e.g.

```
./bin/geosx -i your-problem.xml
```

In a parallel setting, the command might look something like

```
mpirun -np 8 ./bin/geosx -i your-problem.xml -x 2 -y 2 -z 2
```

Note that we provide a series of [Tutorials](#) to walk you through the actual usage of the code, with several input examples. Once you are comfortable the build is working properly, we suggest new users start working through these tutorials.

## 1.8 Testing

It is wise to run our unit test suite as an additional check that everything is working properly. You can run them in the build folder you just created.

```
cd GEOSX/build-your-platform-release
ctest -V
```

This will run a large suite of simple tests that check various components of the code. If you have access, you may also consider running the integrated tests. Please refer to [Integrated Tests](#) for further information.

---

**Note:** If *all* of the unit tests fail, there is likely something wrong with your installation. Refer to the FAQs above for how best to proceed in this situation. If only a few tests fail, it is possible that your platform configuration has exposed some issue that our existing platform tests do not catch. If you suspect this is the case, please consider posting an issue to our issue tracker (after first checking whether other users have encountered a similar issue).

---



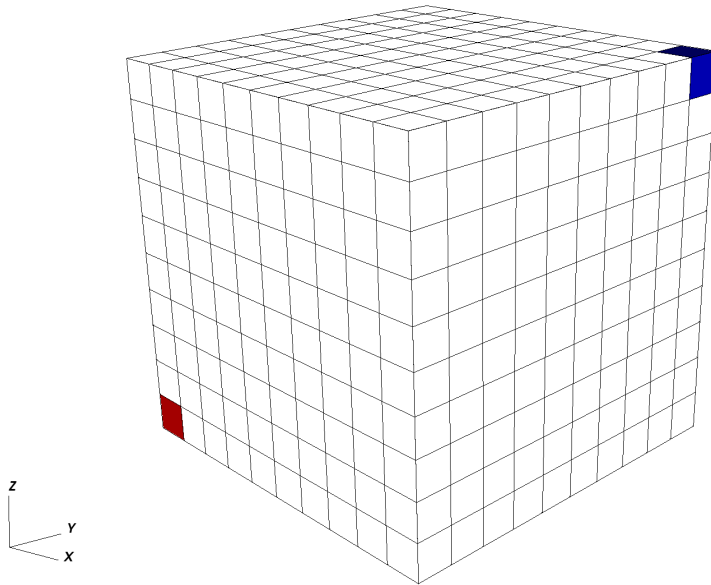
The easiest way to learn to use GEOSX is through worked examples. Here, we have included tutorials showing how to run some common problems. After working through these examples, you should have a good understanding of how to set up and solve your own models.

Note that these tutorials are intended to be followed in sequence, as each step introduces a few new skills. Most of the tutorial models are also quite small, so that large computational resources are not required.

## 2.1 Tutorial 1: First Steps

### Context

In this tutorial, we use a single-phase flow solver (see [Singlephase Flow Solver](#)) to solve for pressure propagation on a 10x10x10 cube mesh with anisotropic permeability values. The pressure source is the lowest-left corner element, and the pressure sink sits at the opposite top corner.



## Objectives

At the end of this tutorial you will know:

- the basic structure of XML input files used by GEOSX,
- how to run GEOSX on a simple case requiring no external input files,
- the basic syntax of a solver block for single-phase problems,
- how to control output and visualize results.

## Input file

GEOSX runs by reading user input information from one or more XML files. For this tutorial, we only need a single GEOSX input file located at:

```
inputFiles/singlePhaseFlow/3D_10x10x10_compressible_smoke.xml
```

## Running GEOSX

If our XML input file is called `my_input.xml`, GEOSX runs this file by executing:

```
/path/to/geosx -i /path/to/my_input.xml
```

The `-i` flag indicates the path to the XML input file. To get help on what other command line input flags GEOSX supports, run `geosx --help`.

## Input file structure

XML files store information in a tree-like structure using nested blocks of information called *elements*. In GEOSX, the root of this tree structure is the element called `Problem`. All elements in an XML file are defined by an opening *tag* (`<ElementName>`) and end by a corresponding closing tag (`</ElementName>`). Elements can have properties defined as *attributes* with `key="value"` pairs. A typical GEOSX input file contains the following tags:

1. *Solver*
2. *Mesh*

3. *Geometry*
4. *Events*
5. *NumericalMethods*
6. *ElementRegions*
7. *Constitutive*
8. *FieldSpecifications*
9. *Outputs*

### XML validation tools

If you have not already done so, please use or enable an XML validation tool (see **User Guide/Input Files/Input Validation**). Such tools will help you identify common issues that may occur when working with XML files.

---

**Note:** Common errors come from the fact that XML is case-sensitive, and all opened tags must be properly closed.

---

## 2.1.1 Single-phase solver

GEOSX is a multiphysics simulator. To find the solution to different physical problems such as diffusion or mechanical deformation, GEOSX uses one or more physics solvers. The `Solvers` tag is used to define and parameterize these solvers. Different combinations of solvers can be applied in different regions of the domain at different moments of the simulation.

In this first example, we use one type of solver in the entire domain and for the entire duration of the simulation. The solver we are specifying here is a single-phase flow solver. In GEOSX, such a solver is created using a `SinglePhaseFVM` element. This type of solver is one among several cell-centered single-phase finite volume methods.

The XML block used to define this single-phase finite volume solver is shown here:

```
<Solvers>
  <SinglePhaseFVM
    name="SinglePhaseFlow"
    logLevel="1"
    discretization="singlePhaseTPFA"
    targetRegions="{ mainRegion }">
      <NonlinearSolverParameters
        newtonTol="1.0e-6"
        newtonMaxIter="8"/>
      <LinearSolverParameters
        solverType="gmres"
        preconditionerType="amg"
        krylovTol="1.0e-10"/>
    </SinglePhaseFVM>
  </Solvers>
```

Each type of solver has a specific set of parameters that are required and some parameters that are optional. Optional values are usually set with sensible default values.

### name

First, we register a solver of type `SinglePhaseFVM` with a user-chosen name, here `SinglePhaseFlow`. This unique user-defined name can be almost anything. However, some symbols are known to cause issues in names : avoid commas, slashes, curly braces. GEOSX is case-sensitive: it makes a distinction between two `SinglePhaseFVM`

solvers called `mySolver` and `MySolver`. Giving elements a name is a common practice in GEOSX: users need to give unique identifiers to objects they define. That name is the handle to this instance of a solver class.

### logLevel

Then, we set a solver-specific level of console logging (`logLevel` set to 1 here). Notice that the value (1) is between double-quotes. This is a general convention for all attributes: we write `key="value"` regardless of the value type (integers, strings, lists, etc.).

For `logLevel`, higher values lead to more console output or intermediate results saved to files. When debugging, higher `logLevel` values is often convenient. In production runs, you may want to suppress most console output.

### discretization

For solvers of the `SinglePhaseFVM` family, one required attribute is a discretization scheme. Here, we use a Two-Point Flux Approximation (TPFA) finite volume discretization scheme called `singlePhaseTPFA`. To know the list of admissible values of an attribute, please see GEOSX's XML schema. This discretization type must know how to find permeability values that it uses internally to compute transmissibilities. The `permeabilityNames` attribute tells the solver the user-defined name (the *handle*) of the permeability values that will be defined elsewhere in the input file. Note that the order of attributes inside an element is not important.

### fluidNames, solidNames, targetRegions

Here, we specify a collection of fluids, rocks, and target regions of the mesh on which the solver will apply. Curly brackets are used in GEOSX inputs to indicate collections of values (sets or lists). The curly brackets used here are necessary, even if the collection contains a single value. Commas are used to separate members of a set.

### Nested elements

Finally, note that other XML elements can be nested inside the `Solvers` element. Here, we use specific XML elements to set values for numerical tolerances. The solver stops when numerical residuals are smaller than the specified tolerances (convergence is achieved) or when the maximum number of iterations allowed is exceeded (convergence not achieved).

## 2.1.2 Mesh

To solve this problem, we need to define a mesh for our numerical calculations. This is the role of the **Mesh** element.

There are two approaches to specifying meshes in GEOSX: internal or external.

- The external approach allows to import mesh files created outside GEOSX, such as a corner-point grid or an unstructured grid representing complex shapes and structures.
- The internal approach uses GEOSX's built-in capability to create simple meshes from a small number of parameters. It does not require any external file information. The geometric complexity of internal meshes is limited, but many practical problems can be solved on such simple grids.

In this tutorial, to keep things self-contained, we use the internal mesh generator. We parameterize it with the **InternalMesh** element.

```
<Mesh>
  <InternalMesh
    name="mesh"
    elementTypes="{ C3D8 }"
    xCoords="{ 0, 10 }"
    yCoords="{ 0, 10 }"
    zCoords="{ 0, 10 }"
    nx="{ 10 }"
    ny="{ 10 }"
    nz="{ 10 }"
```

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```
cellBlockNames="{ cellBlock }"/>
</Mesh>
```

**name**

Just like for solvers, we register the `InternalMesh` element using a unique **name** attribute. Here the `InternalMesh` object is instantiated with the name `mesh`.

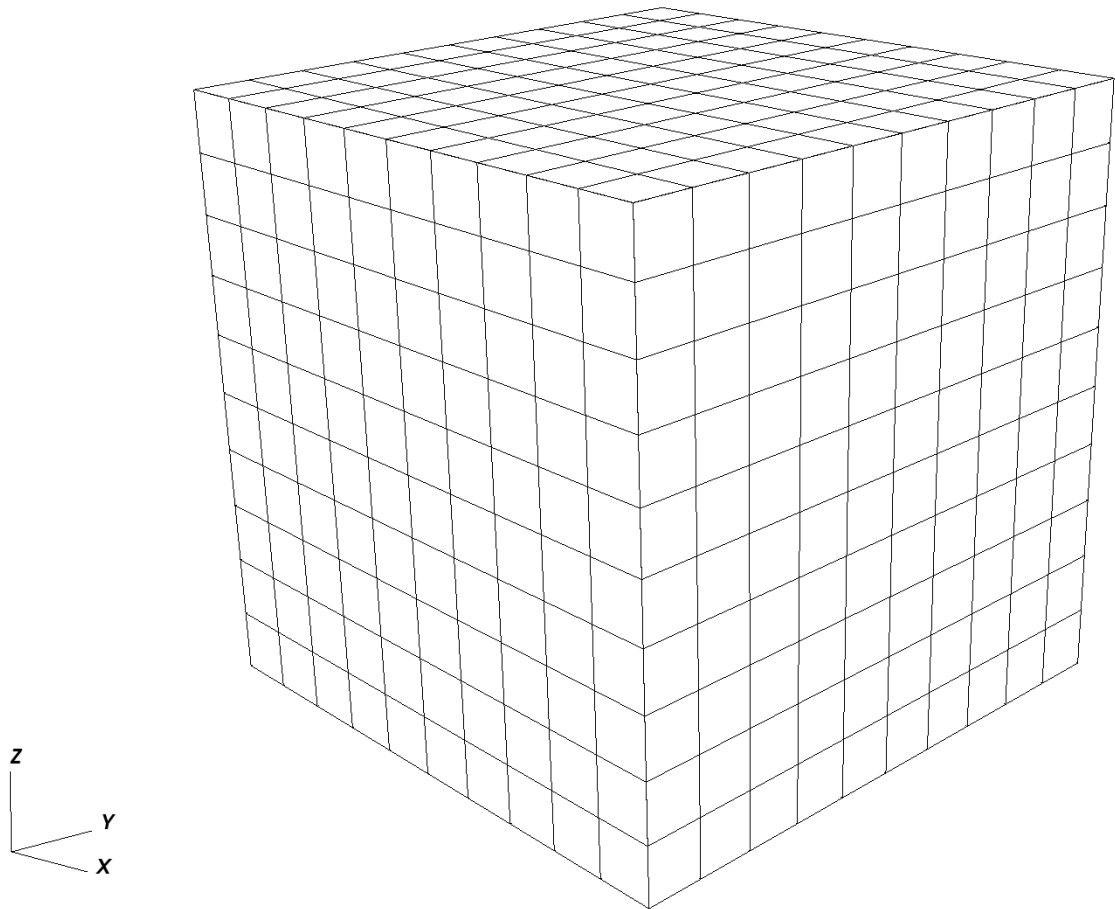
**elementTypes**

We specify the collection of elements types that this mesh contains. Tetrahedra, hexahedra, and wedges are examples of element types. If a mesh contains different types of elements (a hybrid mesh), we should indicate this here by listing all unique types of elements in curly brackets. Keeping things simple, our element collection has only one type of element: a `C3D8` type representing a hexahedral element (linear 8-node brick).

A mesh can contain several geometrical types of elements. For numerical convenience, elements are aggregated by types into `cellBlocks`. Here, we only linear 8-node brick elements, so the entire domain is one object called `cellBlock`.

**xCoords, yCoords, zCoords, nx, ny, nz**

This specifies the spatial arrangement of the mesh elements. The mesh defined here goes from coordinate  $x=0$  to  $x=10$  in the  $x$ -direction, with  $nx=10$  subdivisions along this segment. The same is true for the  $y$ -dimension and the  $z$ -dimension. Our mesh is a cube of  $10 \times 10 \times 10 = 1,000$  elements with a bounding box defined by corner coordinates  $(0,0,0)$  and  $(10,10,10)$ .



### 2.1.3 Geometry

The `Geometry` tag allows users to capture subregions of a mesh and assign them a unique name. Here, we name two `Box` elements, one for the location of the `source` and one for the `sink`. Pressure values are assigned to these named regions elsewhere in the input file.

The pressure source is the element in the (0,0,0) corner of the domain, and the sink is the element in the (10,10,10) corner.

For an element to be inside a geometric region, it must have all its vertices strictly inside that region. Consequently, we need to extend the geometry limits a small amount beyond the actual coordinates of the elements to catch all vertices. Here, we use a safety padding of 0.01.

```
<Geometry>
  <Box
    name="source"
    xMin="{ -0.01, -0.01, -0.01 }"
    xMax="{ 1.01, 1.01, 1.01 }"/>

  <Box
```

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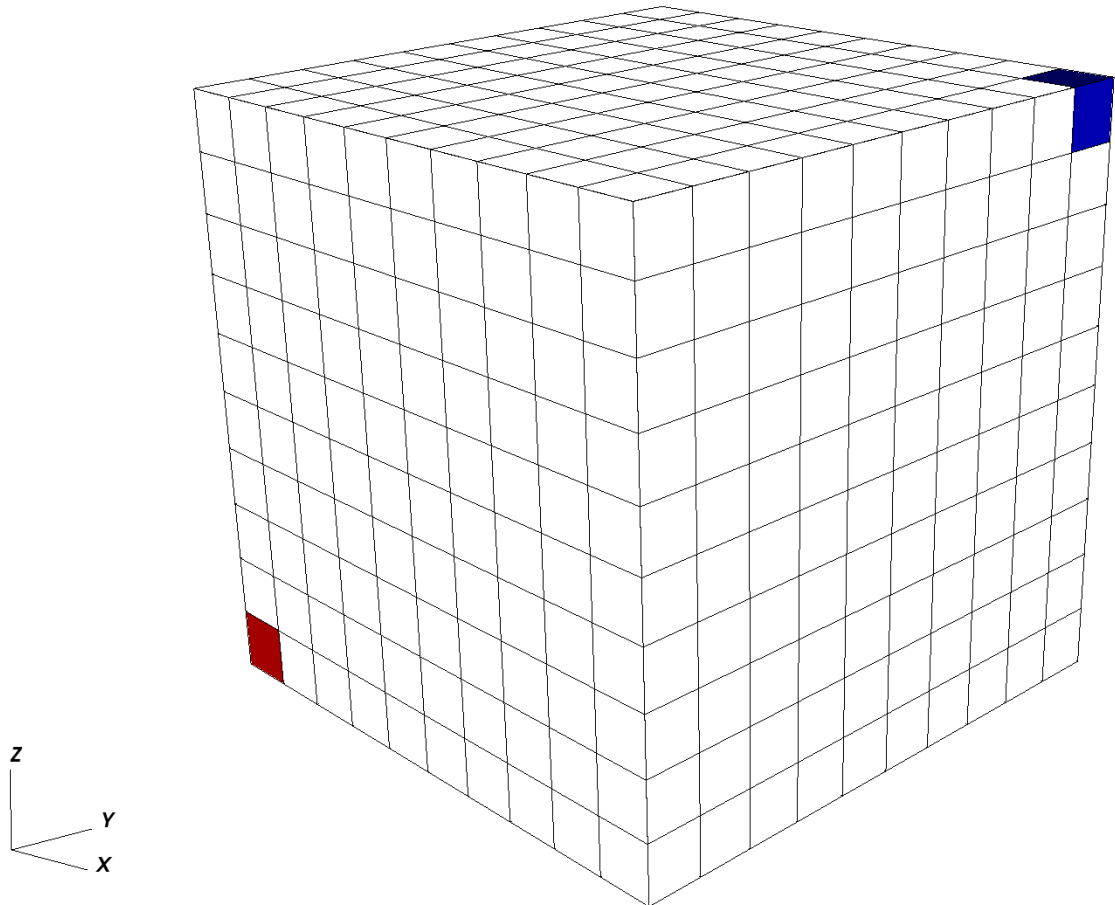
(continued from previous page)

```

name="sink"
xMin="{ 8.99, 8.99, 8.99 }"
xMax="{ 10.01, 10.01, 10.01 }"/>
</Geometry>

```

There are several methods to achieve similar conditions (Dirichlet boundary condition on faces, etc.). The `Box` defined here is one of the simplest approaches.



## 2.1.4 Events

In GEOSX, we call `Events` anything that happens at a set time or frequency. Events are a central element for time-stepping in GEOSX, and a dedicated section just for events is necessary to give them the treatment they deserve.

For now, we focus on three simple events: the time at which we wish the simulation to end (`maxTime`), the times at which we want the solver to perform updates, and the times we wish to have simulation output values reported.

In GEOSX, all times are specified in **seconds**, so here `maxTime=5000.0` means that the simulation will run from time 0 to time 5,000 seconds.

If we focus on the `PeriodicEvent` elements, we see :

1. A **periodic solver** application: this event is named `solverApplications`. With the attribute `forceDt=20`, it tells the solver to compute results at 20-second time intervals. We know what this event does by looking at its `target` attribute: here, from time 0 to `maxTime` and with a forced time step of 20 seconds, we instruct GEOSX to call the solver registered as `SinglePhaseFlow`. Note the hierarchical structure of the target formulation, using `'/'` to indicate a specific named instance (`SinglePhaseFlow`) of an element (`Solvers`). If the solver needs to take smaller time steps, it is allowed to do so, but it will have to compute results for every 20-second increment between time zero and `maxTime` regardless of possible intermediate time steps.
2. An **output event**: this event is used for reporting purposes and instructs GEOSX to write out results at specific frequencies. Here, we need to see results at every 100-second increment. This event triggers a full application of solvers, even if solvers were not summoned by the previous event. In other words, an output event will force an application of solvers, possibly in addition to the periodic events requested directly.

```
<Events maxTime="5000.0">
  <PeriodicEvent
    name="solverApplications"
    forceDt="20.0"
    target="/Solvers/SinglePhaseFlow"/>
  <PeriodicEvent
    name="outputs"
    timeFrequency="100.0"
    target="/Outputs/siloOutput"/>
</Events>
```

### 2.1.5 Numerical methods

GEOSX comes with several useful numerical methods. In the `Solvers` elements, for instance, we had specified to use a two-point flux approximation as discretization scheme for the finite volume single-phase solver. Now to use this scheme, we need to supply more details in the `NumericalMethods` element.

```
<NumericalMethods>
  <FiniteVolume>
    <TwoPointFluxApproximation
      name="singlePhaseTPFA"
    />
  </FiniteVolume>
</NumericalMethods>
```

The `fieldName` attribute specifies which property will be used for flux computations, and also specifies that for Dirichlet boundary conditions, the pressure value at the element face is used. The `coefficientName` attribute is used for the stencil transmissibility computations.

Note that in GEOSX, there is a difference between physics solvers and numerical methods. Their parameterizations are thus independent. We can have multiple solvers using the same numerical scheme but with different tolerances, for instance.

### 2.1.6 Regions

In GEOSX, `ElementsRegions` are used to attach material properties to regions of elements. Here, we use only one **CellElementRegion** to represent the entire domain (user name: `mainRegion`). It contains all the blocks called `cellBlock` defined in the mesh section. We specify the materials contained in that region using a `materialList`. Several materials coexist in `cellBlock`, and we list them using their user-defined names: `water`, `rockPorosity`, and `rockPerm`, etc. What these names mean, and the physical properties that they are attached to are defined next.



```

<ElementRegions>
  <CellElementRegion
    name="mainRegion"
    cellBlocks="{ cellBlock }"
    materialList="{ water, rock }"/>
</ElementRegions>

```

## 2.1.7 Constitutive models

The Constitutive element attaches physical properties to all materials contained in the domain.

The physical properties of the materials defined as water, rockPorosity, and rockPerm are provided here, each material being derived from a different material type: CompressibleSinglePhaseFluid for the water, PressurePorosity for the rock porosity, and ConstantPermeability for rock permeability. The list of attributes differs between these constitutive materials.

```

<Constitutive>
  <CompressibleSinglePhaseFluid
    name="water"
    defaultDensity="1000"
    defaultViscosity="0.001"
    referencePressure="0.0"
    compressibility="5e-10"
    viscosibility="0.0"/>

  <CompressibleSolidConstantPermeability
    name="rock"
    solidModelName="nullSolid"
    porosityModelName="rockPorosity"
    permeabilityModelName="rockPerm"/>

  <NullModel
    name="nullSolid"/>

  <PressurePorosity
    name="rockPorosity"
    defaultReferencePorosity="0.05"
    referencePressure="0.0"
    compressibility="1.0e-9"/>

  <ConstantPermeability
    name="rockPerm"
    permeabilityComponents="{ 1.0e-12, 1.0e-12, 1.0e-15 }"/>
</Constitutive>

```

The names water, rockPorosity and rockPerm are defined by the user as handles to specific instances of physical materials. GEOSX uses S.I. units throughout, not field units. Pressures, for instance, are in Pascal, not psia. The x- and y-permeability are set to 1.0e-12 m<sup>2</sup> corresponding to approximately 1 Darcy.

We have used the handles water, rockPorosity and rockPerm in the input file in the ElementRegions section of the XML file, before the registration of these materials took place here, in Constitutive element.

**Note:** This highlights an important aspect of using XML in GEOSX: the order in which objects are registered and used in the XML file is not important.

### 2.1.8 Defining properties

In the `FieldSpecifications` section, properties such as source and sink pressures are set. GEOSX offers a lot of flexibility to specify field values through space and time.

Spatially, in GEOSX, all field specifications are associated to a target object on which the field values are mounted. This allows for a lot of freedom in defining fields: for instance, one can have volume property values attached to a subset of volume elements of the mesh, or surface properties attached to faces of a subset of elements.

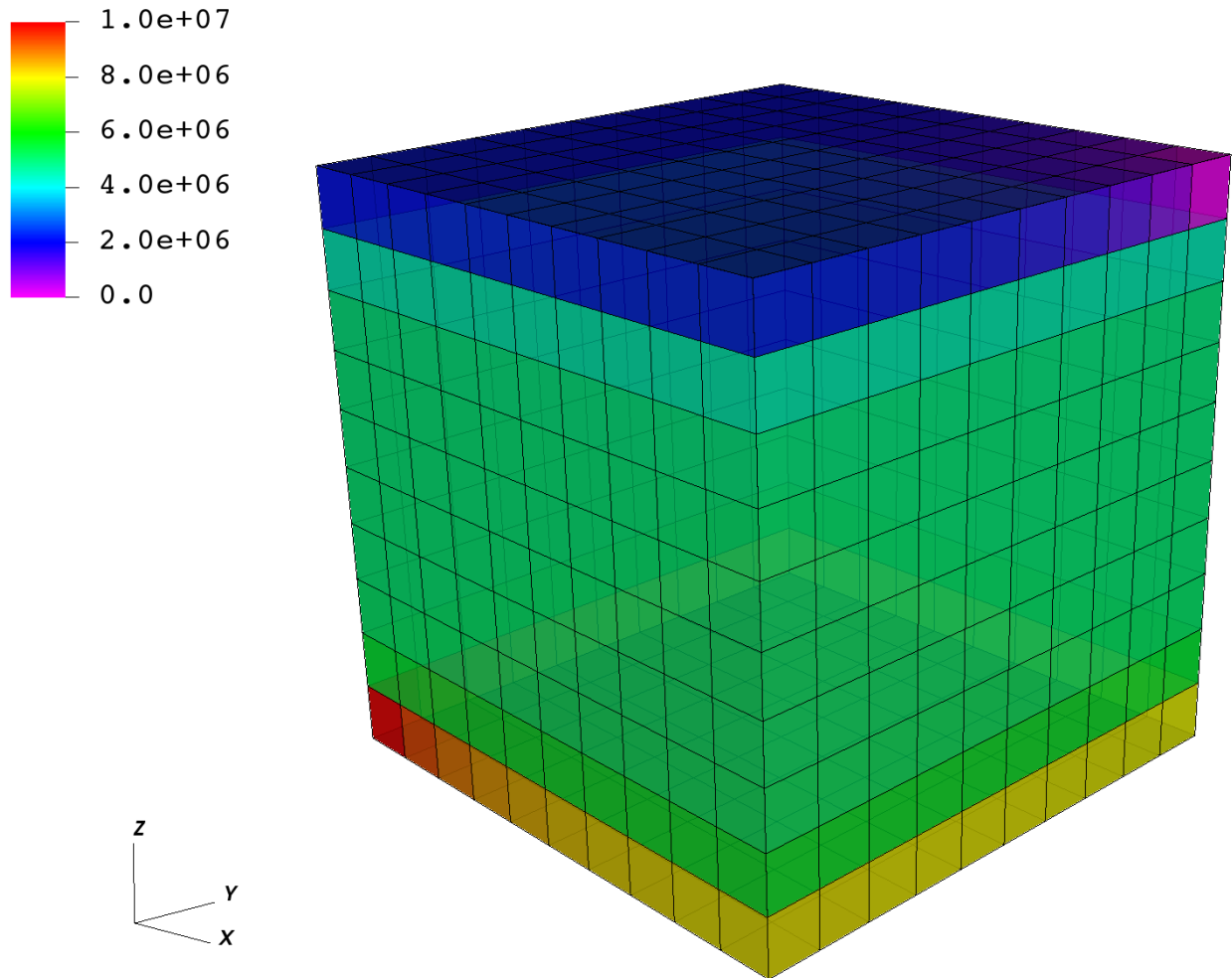
For each `FieldSpecification`, we specify a name, a `fieldName` (this name is used by solvers or numerical methods), an `objectPath`, `setNames` and a `scale`. The `ObjectPath` is important and it reflects the internal class hierarchy of the code. Here, for the `fieldName` `pressure`, we assign the value defined by `scale` (5e6 Pascal) to one of the `ElementRegions` (class) called `mainRegions` (instance). More specifically, we target the `elementSubRegions` called `cellBlock` (this contains all the C3D8 elements, effectively all the domain). The `setNames` allows to use the elements defined in `Geometry`, or use everything in the object path (using the `all`).

```
<FieldSpecifications>
  <FieldSpecification
    name="initialPressure"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions/mainRegion/cellBlock"
    fieldName="pressure"
    scale="5e6"/>

  <FieldSpecification
    name="sourceTerm"
    objectPath="ElementRegions/mainRegion/cellBlock"
    fieldName="pressure"
    scale="1e7"
    setNames="{ source }"/>

  <FieldSpecification
    name="sinkTerm"
    objectPath="ElementRegions/mainRegion/cellBlock"
    fieldName="pressure"
    scale="0.0"
    setNames="{ sink }"/>
</FieldSpecifications>
```

The image below shows the pressures after the very first time step, with the domain initialized at 5 MPa, the sink at 0 MPa on the top right, and the source in the lower left corner at 10 MPa.



### 2.1.9 Output

In order to retrieve results from a simulation, we need to instantiate one or multiple `Outputs`.

Here, we define a single object of type `Silo`. `Silo` is a library and a format for reading and writing a wide variety of scientific data. Data in `Silo` format can be read by `VisIt`.

This `Silo` output object is called `siloOutput`. We had referred to this object already in the `Events` section: it was the target of a periodic event named `outputs`. You can verify that the `Events` section is using this object as a target. It does so by pointing to `/Outputs/siloOutput`.

```
<Outputs>
  <Silo
    name="siloOutput"/>
</Outputs>
```

GEOSX currently supports outputs that are readable by `VisIt` and Kitware's `Paraview`, as well as other visualization tools. In this example, we only request a `Silo` format compatible with `VisIt`.

All elements are now in place to run GEOSX.

## 2.1.10 Running GEOSX

The command to run GEOSX is

```
path/to/geosx -i path/to/this/xml_file.xml
```

Note that all paths for files included in the XML file are relative to this XML file.

While running GEOSX, it logs status information on the console output with a verbosity that is controlled at the object level, and that can be changed using the `logLevel` flag.

The first few lines appearing to the console are indicating that the XML elements are read and registered correctly:

```
Adding Solver of type SinglePhaseFVM, named SinglePhaseFlow
Adding Mesh: InternalMesh, mesh
Adding Geometric Object: Box, source
Adding Geometric Object: Box, sink
Adding Event: PeriodicEvent, solverApplications
Adding Event: PeriodicEvent, outputs
Adding Output: Silo, siloOutput
Adding Object CellElementRegion named mainRegion from ObjectManager::Catalog.
  mainRegion/cellBlock/water is allocated with 1 quadrature points.
  mainRegion/cellBlock/rock is allocated with 1 quadrature points.
  mainRegion/cellBlock/rockPerm is allocated with 1 quadrature points.
  mainRegion/cellBlock/rockPorosity is allocated with 1 quadrature points.
  mainRegion/cellBlock/nullSolid is allocated with 1 quadrature points.
```

Then, we go into the execution of the simulation itself:

```
Time: 0s, dt:20s, Cycle: 0
  Attempt: 0, NewtonIter: 0
    ( R ) = ( 5.65e+00 ) ;
  Attempt: 0, NewtonIter: 1
    ( R ) = ( 2.07e-04 ) ;
  Last LinSolve(iter,res) = ( 63, 8.96e-11 ) ;
  Attempt: 0, NewtonIter: 2
    ( R ) = ( 9.86e-11 ) ;
  Last LinSolve(iter,res) = ( 70, 4.07e-11 ) ;
```

Each time iteration at every 20s interval is logged to console, until the end of the simulation at `maxTime=5000`:

```
Time: 4980s, dt:20s, Cycle: 249
  Attempt: 0, NewtonIter: 0
    ( R ) = ( 4.74e-09 ) ;
  Attempt: 0, NewtonIter: 1
    ( R ) = ( 2.05e-14 ) ;
  Last LinSolve(iter,res) = ( 67, 5.61e-11 ) ;
SinglePhaseFlow: Newton solver converged in less than 4 iterations, time-step_
→required will be doubled.
Cleaning up events
Umpire          HOST sum across ranks: 14.8 MB
Umpire          HOST          rank max: 14.8 MB
total time      5.658s
initialization time 0.147s
run time        3.289s
```

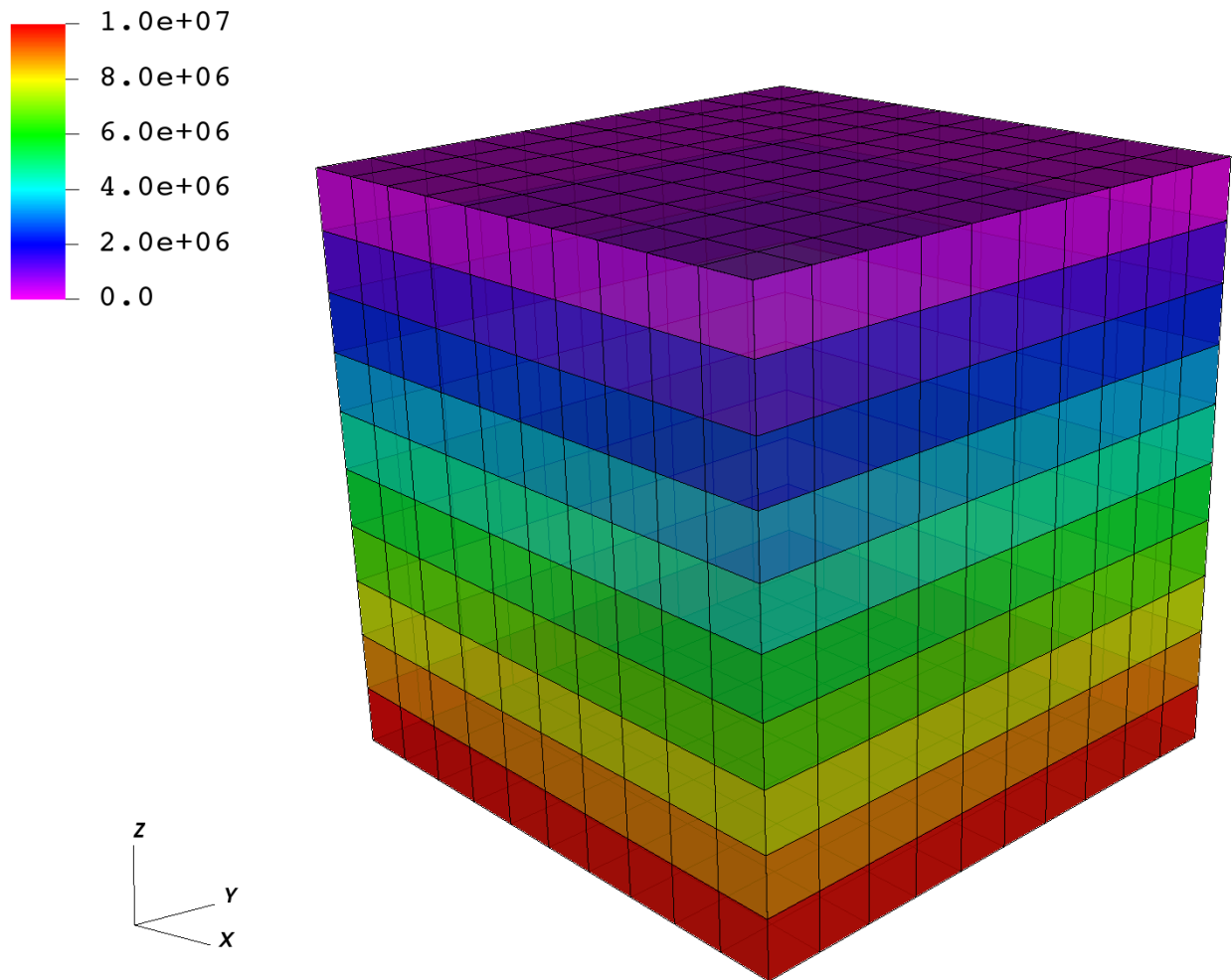
All newton iterations are logged along with corresponding nonlinear residuals for each time iteration. In turn, for each newton iteration, `LinSolve` provides the number of linear iterations and the final residual reached by the linear solver. Information on run times, initialization times, and maximum amounts of memory (high water mark) are given at the end of the simulation, if successful.

Congratulations on completing this first run!

### 2.1.11 Visualization

Here, we have requested results to be written in Silo, a format compatible with [VisIt](#). To visualize results, open VisIt and directly load the database of simulation output files.

After a few time step, pressure between the source and sink are in equilibrium, as shown on the representation below.



### 2.1.12 To go further

#### Feedback on this tutorial

This concludes the single-phase internal mesh tutorial. For any feedback on this tutorial, please submit a [GitHub issue](#) on the project's [GitHub](#) page.

#### For more details

- More on single-phase flow solvers, please see *Singlephase Flow Solver*.
- More on meshes, please see *Meshes*.

- More on events, please see [Event Management](#).

## 2.2 Tutorial 2: External Meshes

### Context

In this tutorial, we use a simple single-phase flow solver (see [Singlephase Flow Solver](#)) to solve for pressure propagation on a mesh that is imported into GEOSX. The main goal of this tutorial is to learn how to work with external meshes, and to learn how easy it is to swap meshes on the same physical problem in GEOSX. This makes GEOSX a powerful tool to solve real field applications with complex geometries and perform assessments of mesh geometry and resolution effects.

### Objectives

At the end of this tutorial you will know:

- the syntax and format of input meshes,
- how to input external files into a GEOSX input XML file,
- how to run the same physical problem with two different meshes,
- how to use and visualize hexahedral and tetrahedral meshes.

### Input Files

This tutorial uses an XML file containing the main input for GEOSX and a separate file with all the mesh information. As we will see later, the main XML file points to the external mesh file with an `include` statement. The XML input file for this test case is located at:

```
inputFiles/singlePhaseFlow/pamela_test/3D_10x10x10_compressible_pamela_hex_gravity_
↪ smoke.xml
```

The mesh file format used in this tutorial is called **MSH**. This format is a standard scientific meshing format not specific to GEOSX. It is maintained as the native format of the meshing tool **Gmsh**. MSH is designed for unstructured meshes and contains a compact and complete representation of the mesh geometry and of its properties. The mesh file used here is human-readable ASCII. It contains a list of nodes with their (x,y,z) coordinates, and a list of elements that are constructed from these nodes.

### 2.2.1 Hexahedral elements

In the first part of the tutorial, we will run flow simulations on a mesh made of hexahedral elements. These types of elements are used in classical cartesian grids (sugar cubes) or corner-point grids or pillar grids.

#### Brief discussion about hexahedral meshes in GEOSX

Although closely related, the hexahedral grids that GEOSX can process are slightly different than either structured grid or corner-point grids. The differences are worth pointing out here. In GEOSX:

- **hexahedra can have irregular shapes:** no pillars are needed and vertices can be anywhere in space. This is useful for grids that turn, fold, or are heavily bent. Hexahedral blocks should nevertheless have 8 distinct vertices that are not coalesced. Some tolerance exists for degeneration to wedges in some solvers (finite element solvers), but it is best to avoid such situations and label elements according to their actual shape. Butterfly cells, flat cells, negative or zero volume cells will cause problems.

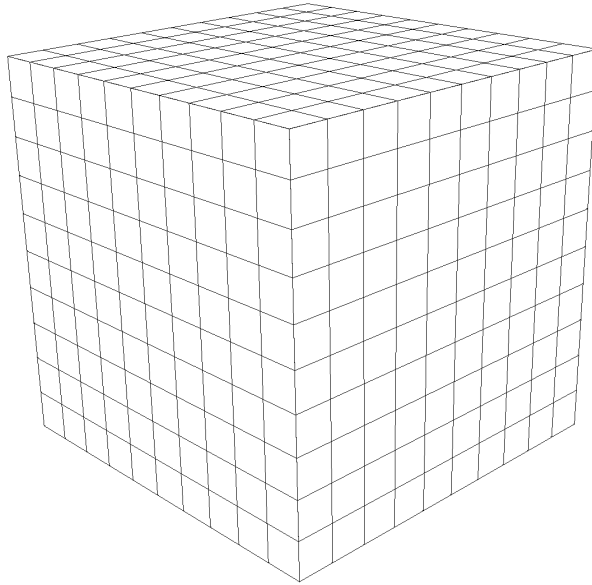
- **the mesh needs to be conformal:** in 3D, this means that neighboring grid blocks have to share exactly a complete face. Note that corner-point grids do not have this requirement and neighboring blocks can be offset. When importing grids from commonly-used ge modeling packages, this is an important consideration. This problem is solved by splitting shifted grid blocks to restore conformity. While it may seem convenient to be able to have offset grid blocks at first, the advantages of conformal grids used in GEOSX are worth the extra meshing effort: by using conformal grids, GEOSX can run finite element and finite volume simulations on the same mesh without problems, going seamlessly from one numerical method to the other. This is key to enabling multiphysics simulation.
- **there is no assumption of overall structure:** GEOSX does not need to know a number of block in the X, Y, Z direction (no NX, NY, NZ) and does not assume that the mesh is a full cartesian domain that the interesting parts of the reservoir must be carved out from. Blocks are numbered by indices that assume nothing about spatial positioning and there is no concept of (i,j,k). This approach also implies that no “masks” are needed to remove inactive or dead cells, as often done in cartesian grids to get the actual reservoir contours from a bounding box, and here we only need to specify grid blocks that are active. For performance and flexibility, this lean approach to meshes is important.

### Importing an external mesh with PAMELA

In this first part of the tutorial, we use a hexahedral mesh provided with GEOSX. This hexahedral mesh is strictly identical to the grid used in the first tutorial (*Tutorial 1: First Steps*), but instead of using the internal grid generator GEOSX, we specify it with spatial node coordinates in MSH format.

The process by which grids are imported into GEOSX is worth explaining. To import external grid into GEOSX, we use an external component (submodule) called **PAMELA**. PAMELA (Parallel Meshing Library) was developed as a stand-alone utility to import grids in multiple formats and write them into memory for GEOSX. Although PAMELA is not necessary to run GEOSX (the internal grid generator of GEOSX has plenty of interesting features), you need PAMELA if you want to import external grids.

So here, our mesh consists of a simple sugar-cube stack of size 10x10x10. We inject fluid from one vertical face of a cube (the face corresponding to  $x=0$ ), and we let the pressure equilibrate in the closed domain. The displacement is a single-phase, compressible fluid subject to gravity forces, so we expect the pressure to be constant on the injection face, and to be close to hydrostatic on the opposite plane ( $x=10$ ). We use GEOSX to compute the pressure inside each grid block over a period of time of 100 seconds.



To see how to import such a mesh, we inspect the following XML file:

```
inputFiles/singlePhaseFlow/pamela_test/3D_10x10x10_compressible_pamela_hex_gravity_
↪ smoke.xml
```

In the XML Mesh tag, instead of an InternalMesh tag, we have a PAMELAMesh tag. We see that a file called cube\_10x10x10\_hex.msh is imported using PAMELA, and this object is instantiated with a user-defined name value. The file here contains geometric information in MSH format (it can also contain properties, as we will see in the next tutorial).

```
<Mesh>
  <PAMELAMesh
    name="CubeHex"
    file="cube_10x10x10_hex.msh"/>
</Mesh>
```

Here are the first few lines of the msh file :

Listing 2.1: cube\_10x10x10\_hex.msh

```
$MeshFormat
2.2 0 8
$EndMeshFormat
$Nodes
1331
1      0      0      0
2      10     0      0
3      10     10     0
4      0      10     0
5      0      0      10
6      10     0      10
7      10     10     10
8      0      10     10
9      1      0      0
```

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10	2	0	0
11	3	0	0
12	4	0	0
13	5	0	0
14	6	0	0
15	7	0	0

GEOSX can run different physical solvers on different regions of the mesh at different times. Here, to keep things simple, we run one solver (single-phase flow) on the entire domain throughout the simulation. Even this is trivial, we need to define and name a region encompassing the entire domain and assign it to the single-phase flow solver. We also need to provide material properties to the regions. This is done by specifying `ElementRegions`. Here, the entire field is one region called `Domain`, and contains multiple constitutive models, including `water`, `rockPorosity`, and `rockPerm`.

```
<ElementRegions>
  <CellElementRegion
    name="Domain"
    cellBlocks="{ DEFAULT_HEX }"
    materialList="{ water, rock }"/>
</ElementRegions>
```

## Running GEOSX

The command to run GEOSX is

```
path/to/geosx -i ../../../../inputFiles/singlePhaseFlow/pamela_test/3D_10x10x10_
↳compressible_pamela_hex_gravity_smoke.xml
```

Note that all paths for files included in the XML file are relative to this XML file, not to the GEOSX executable. When running GEOSX, console messages will provide indications regarding the status of the simulation.

In our case, the first lines are:

```
Adding Solver of type SinglePhaseFlow, named SinglePhaseFlow
Adding Mesh: PAMELAMesh, CubeHex
Adding Geometric Object: Box, all
Adding Geometric Object: Box, left
Adding Event: PeriodicEvent, solverApplications
Adding Event: PeriodicEvent, outputs
Adding Event: PeriodicEvent, restarts
Adding Output: Silo, siloWellPump
Adding Output: Restart, restartOutput
Adding Object CellElementRegion named Domain from ObjectManager::Catalog.
```

This indicates initialization of GEOSX. The mesh preprocessing tool PAMELA is launched next, with console messages as follows.

```
0 >>> *****
0 >>> PAMELA Library Import tool
0 >>> *****
0 >>> GMSH FORMAT IDENTIFIED
0 >>> *** Importing Gmsh mesh format...
0 >>> Reading nodes...
0 >>> Done0
0 >>> Reading elements...
```

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```
0 >>> Number of nodes = 1331
0 >>> Number of triangles = 0
0 >>> Number of quadrilaterals = 0
0 >>> Number of tetrahedra = 0
0 >>> Number of hexahedra = 1000
0 >>> Number of pyramids = 0
0 >>> Number of wedges = 0
0 >>> *** Done
0 >>> *** Creating Polygons from Polyhedra...
0 >>> 3300 polygons have been created
0 >>> *** Done
0 >>> *** Perform partitioning...
0 >>> TRIVIAL partitioning...
0 >>> Ghost elements...
0 >>> Clean mesh...
0 >>> *** Done...
0 >>> Clean Adjacency...
0 >>> *** Done...
Writing into the GEOSX mesh data structure
Running simulation
```

Notice the specification of the number of nodes (1331), and hexahedra (1000). After the adjacency calculations, GEOSX starts the simulation itself, with the time-step increments specified in the XML file.

At the end of your simulation, you should see something like:

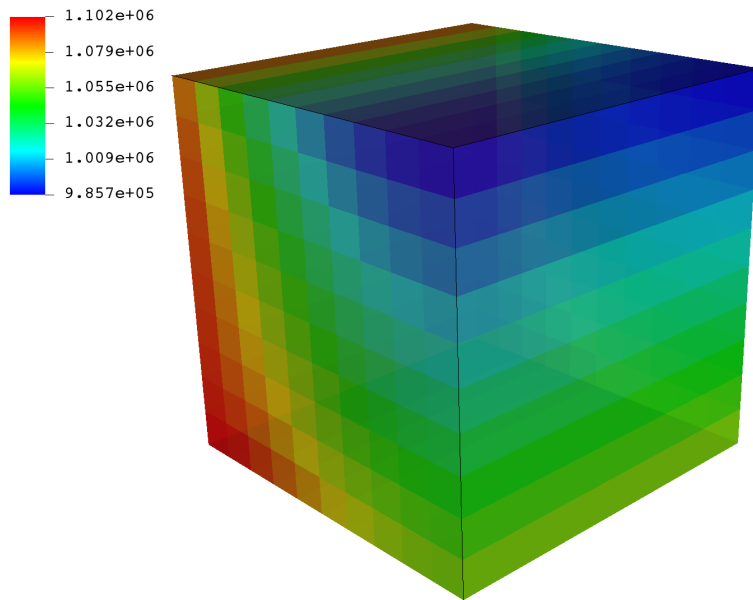
```
Time: 99s, dt:1s, Cycle: 99
Cleaning up events
Writing out restart file at 3D_10x10x10_compressible_pamela_hex_gravity_smoke_restart_
→000000100/rank_0000000.hdf5

init time = 0.081181s, run time = 5.4595s
```

Once this is done, GEOSX is finished and we can inspect the outcome.

### Visualization of results in VisIt

All results are written in a format compatible with [VisIt](#). To load the results, point VisIt to the database file written in the Silo output folder.



We see that the face  $x=0$  shown here in the back of the illustration applies a constant pressure boundary condition (colored in red), whereas the face across from it displays a pressure field under gravity effect, equilibrated and hydrostatic. These results are consistent with what we expect.

Let us now see if a tetrahedral mesh, under the same exact physical conditions, can reproduce these results.

### 2.2.2 Externally Generated Tetrahedral Elements

In the second part of the tutorial, we discretize the same cubic domain but with tetrahedral elements. Tetrahedral meshes are not yet common in geomodeling but offer tremendous flexibility in modeling fracture planes, faults, complex reservoir horizons and boundaries. Just like for hexahedral meshes, and for the same reasons (compatibility with finite volume and finite element methods), tetrahedral meshes in GEOSX must be conformal.

As stated previously, the problem we wish to solve here is the exact same physical problem as with hexahedral grid blocks. We apply a constant pressure condition (injection) from the  $x=0$  vertical face of the domain, and we let pressure equilibrate over time. We observe the opposite side of the cube and expect to see hydrostatic pressure profiles because of the gravitational effect. The displacement is a single phase, compressible flow subject to gravity forces. We use GEOSX to compute the pressure inside each grid block.

The set-up for this problem is almost identical to the hexahedral mesh set-up. We simply point our `Mesh` tag to include a tetrahedral grid. The beauty of not relying on I,J,K indices for any property specification or well trajectory makes it **easy to try different meshes for the same physical problems with GEOSX**. Swapping out meshes without requiring other modifications to the input files makes mesh refinement studies easy to perform with GEOSX.

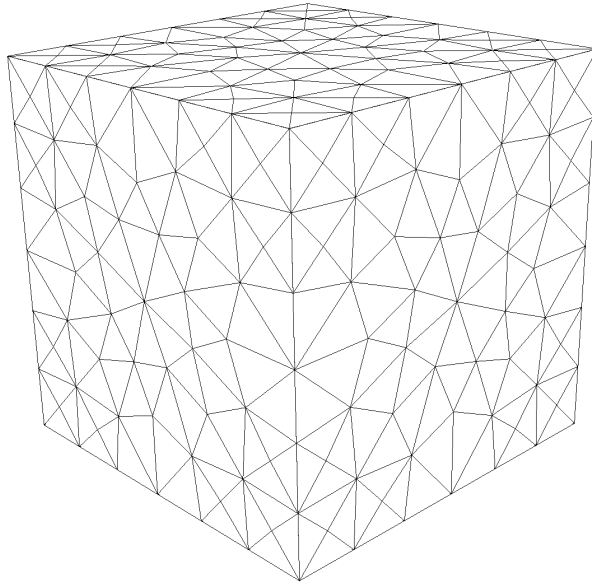
Like before, the XML file for this problem is the following:

```
inputFiles/singlePhaseFlow/pamela_test/3D_10x10x10_compressible_pamela_tetra_gravity_
↪ smoke.xml
```

The only difference, is that now, the `Mesh` tag points GEOSX to a different mesh file called `cube_10x10x10_tet.msh`. This file contains nodes and tetrahedral elements in `Gmsh` format, representing a different discretization of the exact same 10x10x10 cubic domain.

```
<Mesh>
  <PAMELAMesh
    name="CubeTetra"
    file="cube_10x10x10_tet.msh"/>
</Mesh>
```

The mesh now looks like this:



And the MSH file starts as follows (notice the tetrahedral point coordinates as real numbers):

Listing 2.2: cube\_10x10x10\_tet.msh

```
$MeshFormat
2.2 0 8
$EndMeshFormat
$Nodes
366
1 0 0 10
2 0 0 0
3 0 10 10
4 0 10 0
5 10 0 10
6 10 0 0
7 10 10 10
8 10 10 0
9 0 0 1.6666666666666662
10 0 0 3.3333333333333323
11 0 0 4.9999999999999986
12 0 0 6.6666666666666647
13 0 0 8.3333333333333321
14 0 1.6666666666666662 10
15 0 3.3333333333333323 10
```

Again, the entire field is one region called `Domain` and contains `water` and `rock` only.

```
<ElementRegions>
  <CellElementRegion
    name="Domain"
    cellBlocks="{ DEFAULT_TETRA }"
    materialList="{ water, rock }"/>
</ElementRegions>
```

## Running GEOSX

The command to run GEOSX is

```
path/to/geosx -i ../../../../inputFiles/singlePhaseFlow/pamela_test/3D_10x10x10_
↳compressible_pamela_tetra_gravity_smoke.xml
```

Again, all paths for files included in the XML file are relative to this XML file, not to the GEOSX executable. When running GEOSX, console messages will provide indications regarding the status of the simulation. In our case, the first lines are:

```
Adding Solver of type SinglePhaseFVM, named SinglePhaseFlow
Adding Mesh: PAMELAMesh, CubeTetra
Adding Geometric Object: Box, left
Adding Event: PeriodicEvent, solverApplications
Adding Event: PeriodicEvent, outputs
Adding Event: PeriodicEvent, restarts
Adding Output: Silo, siloWellPump
Adding Output: Restart, restartOutput
Adding Object CellElementRegion named Domain from ObjectManager::Catalog.
Reading external mesh from /****/inputFiles/singlePhaseFlow/pamela_test/cube_10x10x10_
↳tet.msh
```

Followed by:

```
0 >>> *****
0 >>> PAMELA Library Import tool
0 >>> *****
0 >>> GMSH FORMAT IDENTIFIED
0 >>> *** Importing Gmsh mesh format...
0 >>> Reading nodes...
0 >>> Done0
0 >>> Reading elements...
0 >>> Number of nodes = 366
0 >>> Number of triangles = 624
0 >>> Number of quadrilaterals = 0
0 >>> Number of tetrahedra = 1153
0 >>> Number of hexahedra = 0
0 >>> Number of pyramids = 0
0 >>> Number of wedges = 0
0 >>> *** Done
0 >>> *** Creating Polygons from Polyhedra...
0 >>> 1994 polygons have been created
0 >>> *** Done
0 >>> *** Perform partitioning...
0 >>> TRIVIAL partitioning...
0 >>> Ghost elements...
0 >>> Clean mesh...
0 >>> *** Done...
```

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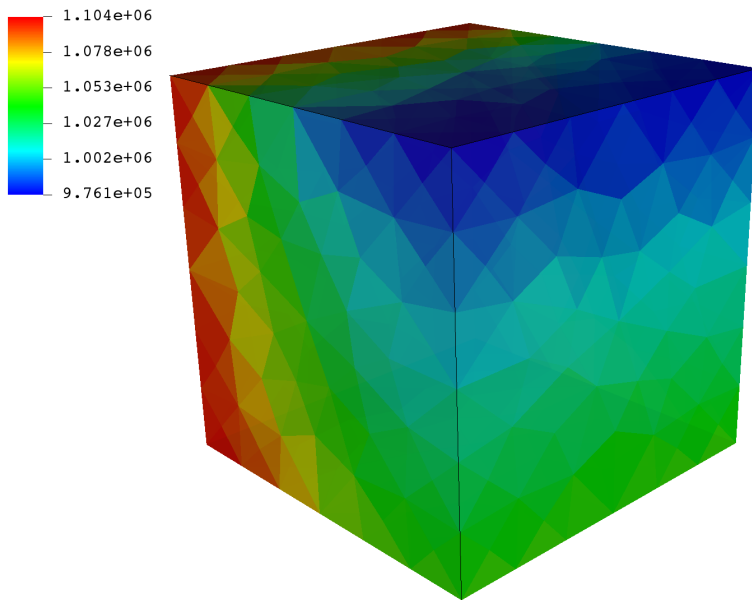
```
0 >>> Clean Adjacency...
0 >>> *** Done...
Writing into the GEOSX mesh data structure
  Domain/DEFAULT_TETRA/water is allocated with 1 quadrature points.
  Domain/DEFAULT_TETRA/rock is allocated with 1 quadrature points.
  Domain/DEFAULT_TETRA/rockPerm is allocated with 1 quadrature points.
  Domain/DEFAULT_TETRA/rockPorosity is allocated with 1 quadrature points.
  Domain/DEFAULT_TETRA/nullSolid is allocated with 1 quadrature points.
PAMELAMesh CubeTetra: importing field data from mesh dataset
```

We see that we have now 366 nodes and 1153 tetrahedral elements. And finally, when the simulation is successfully done we see:

```
Time: 0s, dt:1s, Cycle: 0
Time: 1s, dt:1s, Cycle: 1
Time: 2s, dt:1s, Cycle: 2
Time: 3s, dt:1s, Cycle: 3
Time: 4s, dt:1s, Cycle: 4
Time: 5s, dt:1s, Cycle: 5
...
Time: 95s, dt:1s, Cycle: 95
Time: 96s, dt:1s, Cycle: 96
Time: 97s, dt:1s, Cycle: 97
Time: 98s, dt:1s, Cycle: 98
Time: 99s, dt:1s, Cycle: 99
Cleaning up events
Umpire          HOST sum across ranks:    4.9 MB
Umpire          HOST          rank max:    4.9 MB
total time                      3.164s
initialization time             0.178s
run time                       2.659s
```

## Visualization of results in VisIt

All results are written in a format compatible with [VisIt](#) by default. If we load into VisIt the *.database* file found in the Silo folder, we observe the following results:



Here, we can see that despite the different mesh sizes and shapes, we are able to recover our pressure profile without any problems, or degradation in runtime performance.

### 2.2.3 To go further

#### Feedback on this tutorial

This concludes the single-phase external mesh tutorial. For any feedback on this tutorial, please submit a [GitHub issue](#) on the project's [GitHub page](#).

#### For more details

- A complete description of the Internal Mesh generator is found here [Meshes](#).
- PAMELA being an external submodule has less documentation, but the same [Meshes](#) page may get you started.
- GEOSX can handle tetrahedra, hexahedra, pyramids, wedges, and any combination thereof in one mesh. For more information on how MSH formats can help you specify these mesh types, see the [Gmsh](#) website.

## 2.3 Tutorial 3: Regions and Property Specifications

### Context

In this tutorial, we set up a simple field case for single-phase flow simulation (see [Singlephase Flow Solver](#)). We demonstrate how to run a basic flow simulation in the reservoir layer. We do not consider any coupling with wells. Injection and production will be specified by imposing a high pressure in the cells close to the injection area and a low pressure in the cells close to the production area.

### Objectives

At the end of this tutorial you will know:

- how to import external mesh information and properties,
- how to run a specific solver (here, flow) in a specific region only,

- the basic method of using boxes to set up boundary conditions,
- how to use *TableFunction* to import fields varying in time and/or space,
- how to control output frequency and export results for visualization.

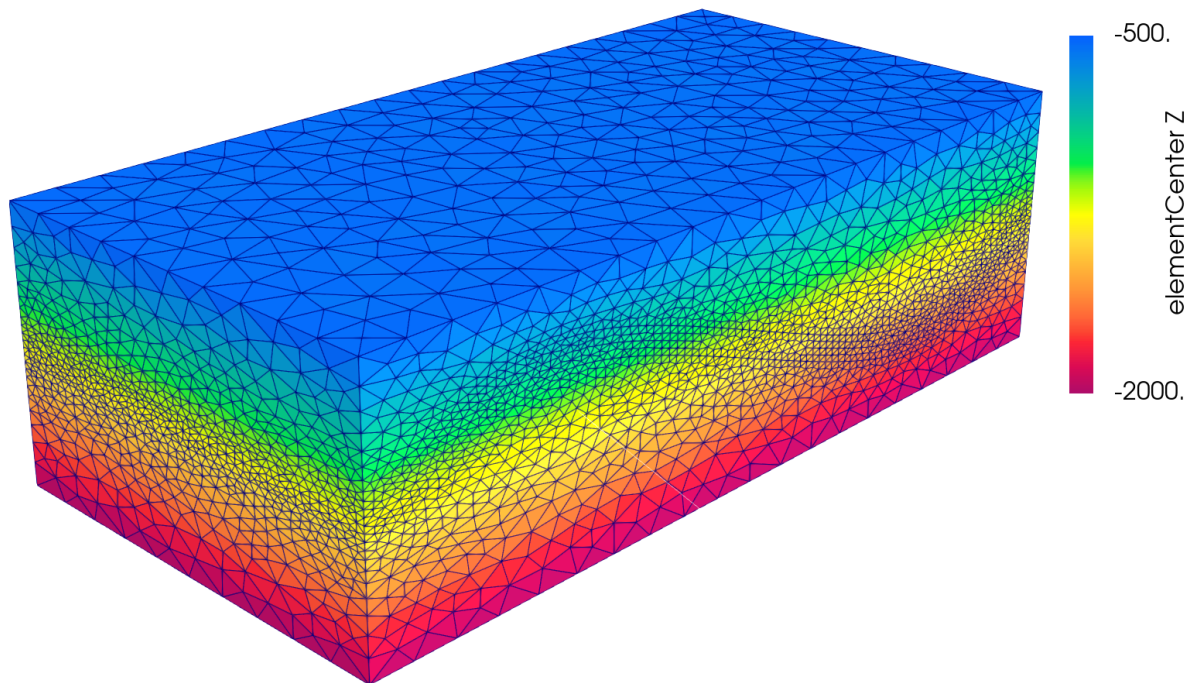
### Input file

The XML input file for this test case is located at:

```
inputFiles/singlePhaseFlow/FieldCaseTutorial3_base.xml
```

```
inputFiles/singlePhaseFlow/FieldCaseTutorial3_smoke.xml
```

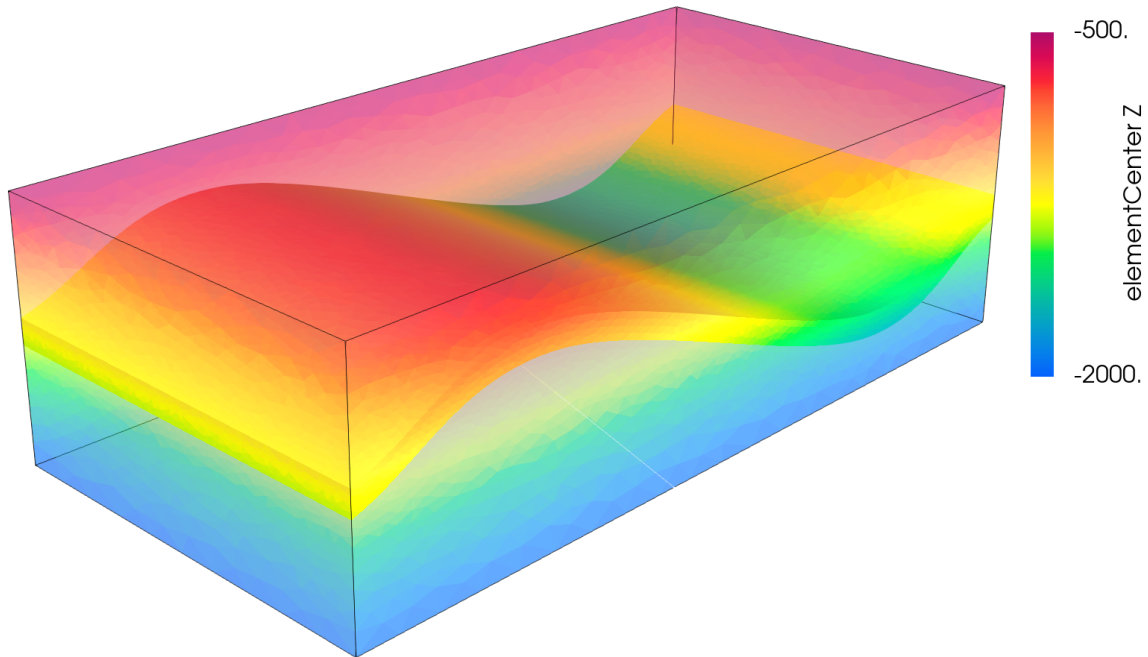
We consider the following mesh as a numerical support to the simulations in this tutorial:



This mesh contains three continuous regions:

- a Top region (overburden, elementary tag = *Overburden*)
- a Middle region (reservoir layer, elementary tag = *Reservoir*)
- a Bottom region (underburden, elementary tag = *Underburden*)





The mesh is defined using the VTK file format (see [Meshes](#) for more information on the supported mesh file format). Each tetrahedron is associated to a unique tag.

The XML file considered here follows the typical structure of the GEOSX input files:

1. *Solver*
2. *Mesh*
3. *Geometry*
4. *Events*
5. *NumericalMethods*
6. *ElementRegions*
7. *Constitutive*
8. *FieldSpecifications*
9. *Outputs*
10. *Functions*

### 2.3.1 Single-phase solver

Let us inspect the **Solver** XML tags.

```
<Solvers>
  <SinglePhaseFVM
    name="SinglePhaseFlow"
    discretization="singlePhaseTPFA"
    targetRegions="{ Reservoir }">
    <NonlinearSolverParameters
      newtonTol="1.0e-6"
      newtonMaxIter="8"/>
  </SinglePhaseFVM>
</Solvers>
```

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```

<LinearSolverParameters
  solverType="gmres"
  preconditionerType="amg"
  amgSmootherType="lljacobi"
  krylovTol="1.0e-10"/>
</SinglePhaseFVM>
</Solvers>

```

This node gathers all the information previously defined. We use a classical `SinglePhaseFVM` Finite Volume Method, with the two-point flux approximation as will be defined in the **NumericalMethods** tag. The `targetRegions` refers only to the Reservoir region because we only solve for flow in this region.

The `NonlinearSolverParameters` and `LinearSolverParameters` are used to set usual numerical solver parameters such as the linear and nonlinear tolerances, the preconditioner and solver types or the maximum number of nonlinear iterations.

### 2.3.2 Mesh

Here, we use the `VTKMesh` to load the mesh (see *Importing the Mesh*). The syntax to import external meshes is simple : in the XML file, the mesh `file` is included with its relative or absolute path to the location of the GEOSX XML file and a user-specified name label for the mesh object.

```

<Mesh>
  <VTKMesh name="SyntheticMesh"
    file="synthetic.vtu" />
</Mesh>

```

### 2.3.3 Geometry

Here, we are using definition of `source` and `sink` boxes in addition to the `all` box in order to flag sets of nodes or cells which will act as injection or production.

```

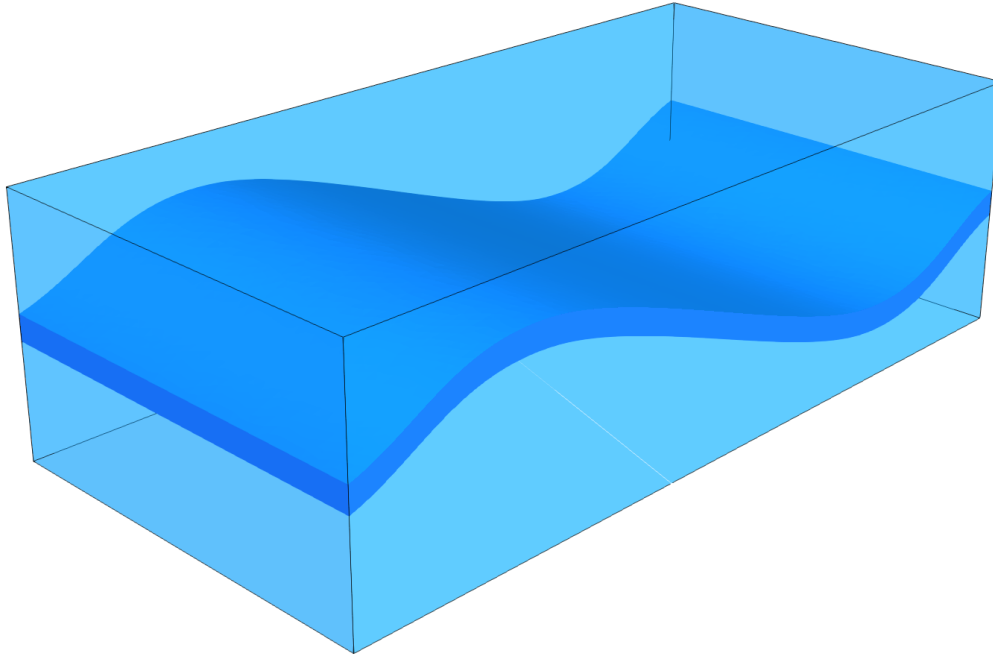
<Geometry>
  <Box
    name="all"
    xMin="{ -1e9, -1e9, -1e9 }"
    xMax="{ 1e9, 1e9, 1e9 }"/>

  <Box
    name="source"
    xMin="{ 15500, 7000, -5000 }"
    xMax="{ 16000, 7500, 0 }"/>

  <Box
    name="sink"
    xMin="{ 6500, 1500, -5000 }"
    xMax="{ 7000, 2000, 0 }"/>
</Geometry>

```

In order to define a box, the user defines `xMax` and `xMin`, two diagonally opposite nodes of the box.



### 2.3.4 Events

The events are used here to guide the simulation through time, and specify when outputs must be triggered.

```
<Events maxTime="100.0e6">
  <PeriodicEvent name="solverApplications"
    forceDt="10.0e6"
    target="/Solvers/SinglePhaseFlow" />

  <PeriodicEvent name="outputs"
    timeFrequency="10.0e6"
    target="/Outputs/reservoir_with_properties" />
</Events>
```

The **Events** tag is associated with the `maxTime` keyword defining the maximum time. If this time is ever reached or exceeded, the simulation ends.

Two `PeriodicEvent` are defined. - The first one, `solverApplications`, is associated with the solver. The `forceDt` keyword means that there will always be time-steps of 23 days (2 000 000 seconds). - The second, `outputs`, is associated with the output. The `timeFrequency` keyword means that it will be executed every 116 days (10 000 000 seconds).

### 2.3.5 Numerical methods

Defining the numerical method used in the solver, we will provide information on how to discretize our equations. Here a classical two-point flux approximation (TPFA) scheme is used to discretize water fluxes over faces.

```
<NumericalMethods>
  <FiniteVolume>
    <TwoPointFluxApproximation
      name="singlePhaseTPFA"
```

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```

    />
  </FiniteVolume>
</NumericalMethods>

```

### 2.3.6 Regions

Assuming that the overburden and the underburden are impermeable, and flow only takes place in the reservoir, we need to define regions.

There are two methods to achieve this regional solve.

- The first solution is to define a unique `CellElementRegion` corresponding to the reservoir.

```

<ElementRegions>
  <CellElementRegion
    name="Reservoir"
    cellBlocks="{ 2_tetrahedra }"
    materialList="{ water, rock, rockPerm, rockPorosity, nullSolid }"/>
</ElementRegions>

```

- The second solution is to define all the `CellElementRegions` as they are in the VTK file, but defining the solvers only on the reservoir layer. In this case, the **ElementRegions** tag is :

```

<ElementRegions>
  <CellElementRegion
    name="Reservoir"
    cellBlocks="{ 2_tetrahedra }"
    materialList="{ water, rock }"/>

  <CellElementRegion
    name="Burden"
    cellBlocks="{ 1_tetrahedra, 3_tetrahedra }"
    materialList="{ water, rock }"/>
</ElementRegions>

```

We opt for the latest as it allows to visualize over- and underburdens and to change regions handling in their tag without needing to amend the **ElementRegion** tag.

---

**Note:** The material list here was set for a single-phase flow problem. This list is subject to change if the problem is not a single-phase flow problem.

---

### 2.3.7 Constitutive models

We simulate a single-phase flow in the reservoir layer, hence with multiple types of materials, a fluid (water) and solid (rock permeability and porosity).

```

<Constitutive>
  <CompressibleSinglePhaseFluid
    name="water"
    defaultDensity="1000"
    defaultViscosity="0.001"
    referencePressure="0.0"

```

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```

compressibility="1e-9"
viscosity="0.0"/>

<CompressibleSolidConstantPermeability
  name="rock"
  solidModelName="nullSolid"
  porosityModelName="rockPorosity"
  permeabilityModelName="rockPerm"/>

<NullModel
  name="nullSolid"/>

<PressurePorosity
  name="rockPorosity"
  defaultReferencePorosity="0.05"
  referencePressure="10e7"
  compressibility="1.0e-9"/>

<ConstantPermeability
  name="rockPerm"
  permeabilityComponents="{ 1.0e-13, 1.0e-13, 1.0e-16 }"/>
</Constitutive>

```

The constitutive parameters such as the density, the viscosity, and the compressibility are specified in the International System of Units.

---

**Note:** To consider an incompressible fluid, the user has to set the compressibility to 0.

---



---

**Note:** GEOSX handles permeability as a diagonal matrix, so the three values of the permeability tensor are set individually using the component field.

---

### 2.3.8 Defining properties

The next step is to specify fields, including:

- The initial value (here, the pressure has to be initialized)
- The static properties (here, we have to define the permeability tensor and the porosity)
- The boundary conditions (here, the injection and production pressure have to be set)

```

<FieldSpecifications>

  <FieldSpecification
    name="permx"
    initialCondition="1"
    component="0"
    setNames="{ all }"
    objectPath="ElementRegions/Reservoir"
    fieldName="rockPerm_permeability"
    scale="1e-15"
    functionName="permxFunc"/>

```

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```

<FieldSpecification
  name="permy"
  initialCondition="1"
  component="1"
  setNames="{ all }"
  objectPath="ElementRegions/Reservoir"
  fieldName="rockPerm_permeability"
  scale="1e-15"
  functionName="permyFunc"/>

<FieldSpecification
  name="permz"
  initialCondition="1"
  component="2"
  setNames="{ all }"
  objectPath="ElementRegions/Reservoir"
  fieldName="rockPerm_permeability"
  scale="3e-15"
  functionName="permzFunc"/>

<FieldSpecification
  name="initialPressure"
  initialCondition="1"
  setNames="{ all }"
  objectPath="ElementRegions/Reservoir/2_tetrahedra"
  fieldName="pressure"
  scale="1e7"
/>

<FieldSpecification
  name="sourceTerm"
  objectPath="ElementRegions/Reservoir/2_tetrahedra"
  fieldName="pressure"
  scale="15e7"
  setNames="{ source }"
/>

<FieldSpecification
  name="sinkTerm"
  objectPath="ElementRegions/Reservoir/2_tetrahedra"
  fieldName="pressure"
  scale="5e7"
  setNames="{ sink }"/>
</FieldSpecifications>

```

You may note :

- All static parameters and initial value fields must have `initialCondition` field set to 1.
- The `objectPath` refers to the `ElementRegion` in which the field has its value,
- The `setName` field points to the box previously defined to apply the fields,
- `name` and `fieldName` have a different meaning: `name` is used to give a name to the XML block. This name must be unique. `fieldName` is the name of the field registered in GEOSX. This value has to be set according to the expected input fields of each solver.

## 2.3.9 Output

The **Outputs** XML tag is used to trigger the writing of visualization files. Here, we write files in a format natively readable by Paraview under the tag *VTK*:

```
<Outputs>
  <VTK
    name="reservoir_with_properties"/>
</Outputs>
```

**Note:** The name keyword defines the name of the output directory.

## 2.3.10 Using functions to specify properties

Eventually, one can define varying properties using `TableFunction` (*Functions*) under the **Functions** tag:

```
<Functions>
  <TableFunction
    name="timeInj"
    inputVarNames="{ time }"
    coordinates="{ 1e6, 10e6, 50e6 }"
    values="{ 1, 0.01, 0.00001 }"/>

  <TableFunction
    name="initialPressureFunc"
    inputVarNames="{ elementCenter }"
    coordinateFiles="{ tables_FieldCaseTuto/xlin.geos, tables_FieldCaseTuto/ylin.
→geos, tables_FieldCaseTuto/zlin.geos }"
    voxelFile="tables_FieldCaseTuto/pressure.geos"/>

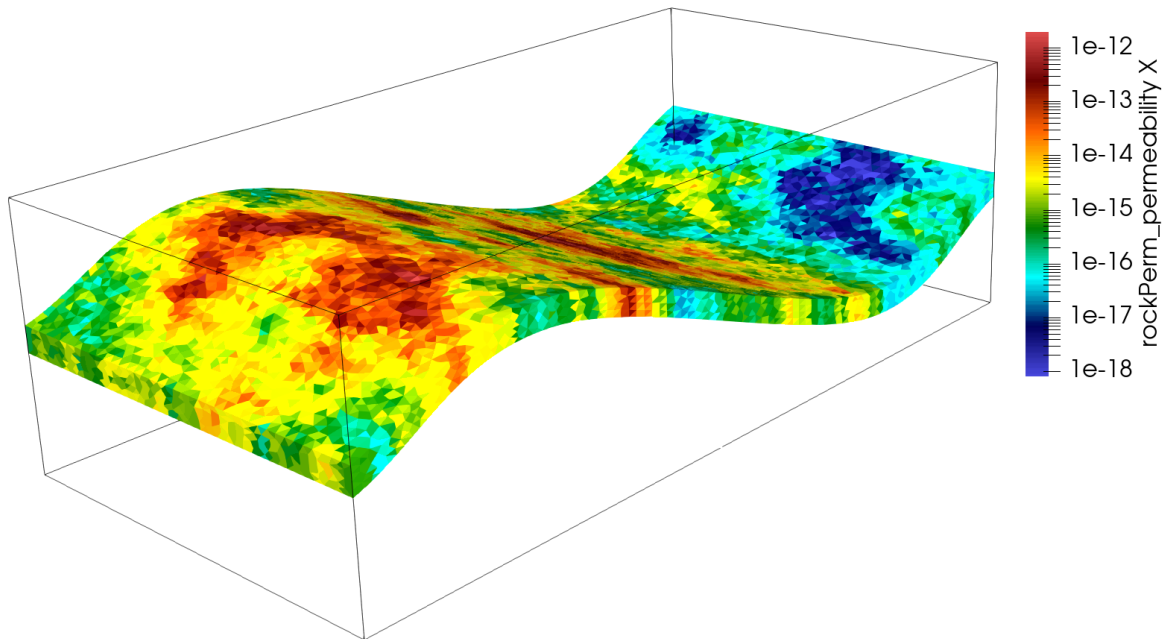
  <TableFunction
    name="permxFunc"
    inputVarNames="{ elementCenter }"
    coordinateFiles="{ tables_FieldCaseTuto/xlin.geos, tables_FieldCaseTuto/ylin.
→geos, tables_FieldCaseTuto/zlin.geos }"
    voxelFile="tables_FieldCaseTuto/permx.geos"
    interpolation="nearest"/>

  <TableFunction
    name="permyFunc"
    inputVarNames="{ elementCenter }"
    coordinateFiles="{ tables_FieldCaseTuto/xlin.geos, tables_FieldCaseTuto/ylin.
→geos, tables_FieldCaseTuto/zlin.geos }"
    voxelFile="tables_FieldCaseTuto/permy.geos"
    interpolation="nearest"/>

  <TableFunction
    name="permzFunc"
    inputVarNames="{ elementCenter }"
    coordinateFiles="{ tables_FieldCaseTuto/xlin.geos, tables_FieldCaseTuto/ylin.
→geos, tables_FieldCaseTuto/zlin.geos }"
    voxelFile="tables_FieldCaseTuto/permz.geos"
    interpolation="nearest"/>
</Functions>
```

Here, the injection pressure is set to vary with time. Attentive reader might have noticed that `sourceTerm` was bound to a `TableFunction` named *timeInj* under **FieldSpecifications** tag definition. The initial pressure is set based on the values contained in the table formed by the files which are specified. In particular, the files *xlin.geos*, *ylin.geos* and *zlin.geos* define a regular meshing of the bounding box containing the reservoir. The *pressure.geos* file then defines the values of the pressure at those points.

We proceed in a similar manner as for *pressure.geos* to map a heterogeneous permeability field (here the 5th layer of the SPE 10 test case) onto our unstructured grid. This mapping will use a nearest point interpolation rule.



**Note:** The varying values imposed in *values* or passed through *voxelFile* are premultiplied by the *scale* attribute from **FieldSpecifications**.

### 2.3.11 Running GEOSX

The simulation can be launched with:

```
geosx -i FieldCaseTutorial13_smoke.xml
```

One can notice the correct load of the field function among the starting output messages

```
Adding Mesh: VTKMesh, SyntheticMesh
Adding Event: PeriodicEvent, solverApplications
Adding Event: PeriodicEvent, outputs
Adding Solver of type SinglePhaseFVM, named SinglePhaseFlow
Adding Geometric Object: Box, all
Adding Geometric Object: Box, source
Adding Geometric Object: Box, sink
Adding Output: VTK, reservoir_with_properties
    TableFunction: timeInj
    TableFunction: initialPressureFunc
    TableFunction: permxFunc
```

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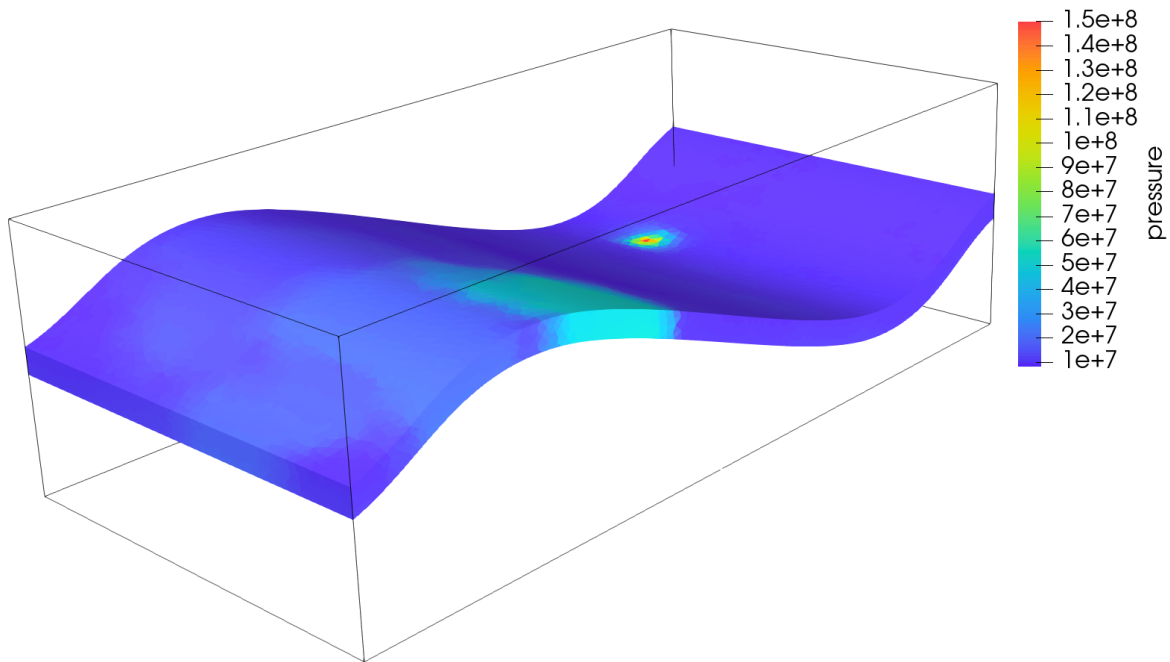


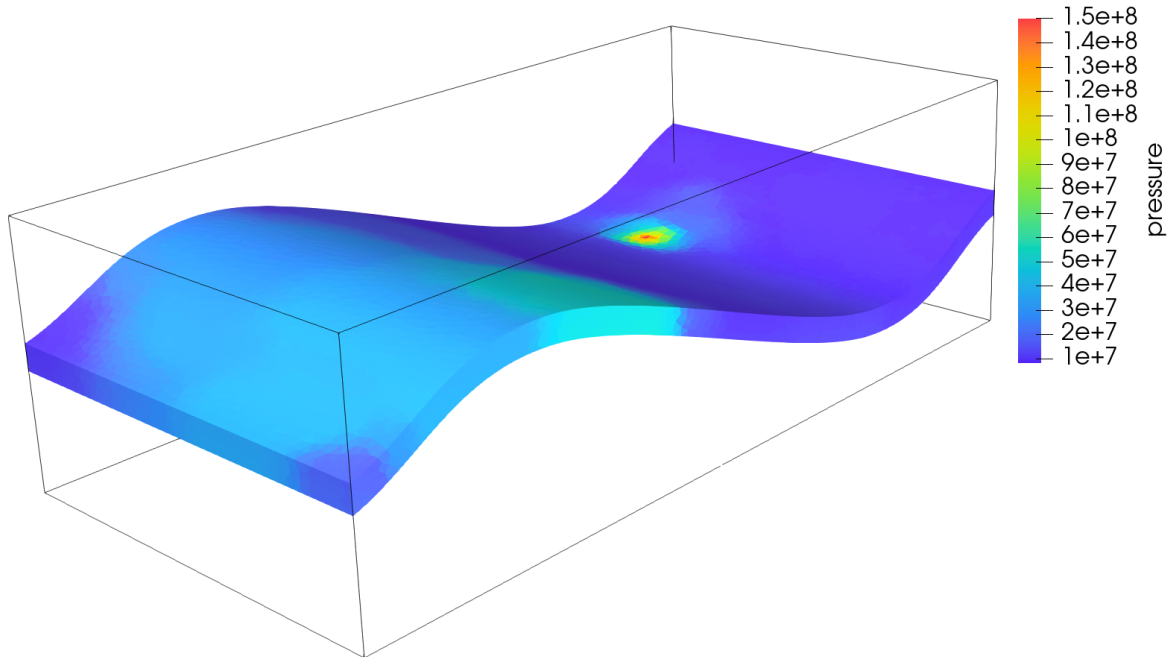
(continued from previous page)

```
TableFunction: permyFunc
TableFunction: permzFunc
Adding Object CellElementRegion named Reservoir from ObjectManager::Catalog.
Adding Object CellElementRegion named Burden from ObjectManager::Catalog.
```

### 2.3.12 Visualization of results

We can open the file *syntheticReservoirVizFile.pvd* with Paraview to visualize the simulation results. In the event block, we have asked for the output to be generated at regular intervals throughout the simulation, we can thus visualize the pressure distribution at different simulation times, showing the variation in the injection control.





### 2.3.13 To go further

#### Feedback on this tutorial

This concludes this tutorial. For any feedback, please submit a [GitHub issue](#) on the project's GitHub page.

#### For more details

- More on meshes, please see [Meshes](#).
- More on events, please see [Event Management](#).

## 2.4 Tutorial 4: Boundary Conditions and Time-Dependent Functions

### Context

In this tutorial, we use a small strain linear elastic based solid mechanics solver (see [Solid Mechanics Solver](#)) from GEOSX to solve for the bending problem of a three-dimensional cantilever beam. The beam is fixed at one end, and subjects to a traction force pointing to the y-positive direction on the other end. The beam is deformed in the x-y plane.

### Objectives

At the end of this tutorial, you will know:

- how to use the solid mechanics solver to solve a quasistatic problem,
- how to set up displacement boundary condition at element nodes,
- how to set up traction boundary condition on element surfaces,
- how to use a table function to control time-dependent loading.

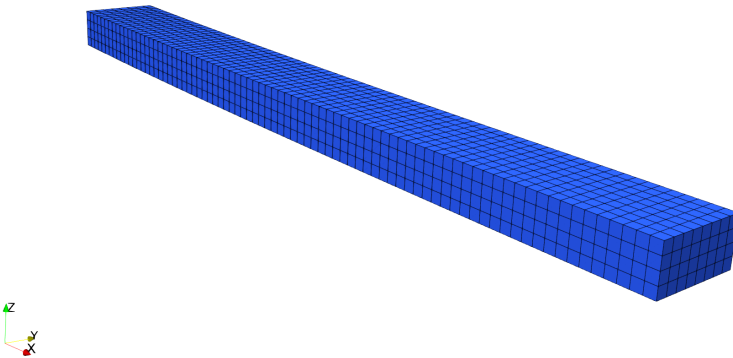
### Input file

This tutorial uses no external input files and everything required is contained within a single GEOSX input file. The xml input file for this test case is located at:

```
inputFiles/solidMechanics/beamBending_base.xml
inputFiles/solidMechanics/beamBending_benchmark.xml
```

### 2.4.1 Discretized computational domain

The following mesh is used in this tutorial:



This mesh contains 80 x 8 x 4 eight-node brick elements in the x, y and z directions, respectively. Here, the `InternalMesh` is used to generate a structured three-dimensional mesh with `C3D8` as the `elementTypes`. This mesh is defined as a cell block with the name `cb1`.

```
<Mesh>
  <InternalMesh
    name="mesh1"
    elementTypes="{ C3D8 }"
    xCoords="{ 0, 80 }"
    yCoords="{ 0, 8 }"
    zCoords="{ 0, 4 }"
    nx="{ 160 }"
    ny="{ 16 }"
    nz="{ 8 }"
    cellBlockNames="{ cb1 }"/>
  </InternalMesh>
</Mesh>
```

### 2.4.2 Gravity

The gravity is turned off explicitly at the beginning of the input file:

```
<Solvers
  gravityVector="{ 0.0, 0.0, 0.0 }">
```

### 2.4.3 Solid mechanics solver

The solid mechanics solver is based on the small strain Lagrangian finite element formulation. The problem is run as `QuasiStatic` without considering the beam inertial. The computational domain is discretized by `FE1`, which is defined in the `NumericalMethods` block. The material is designated as `shale`, whose properties are defined in the `Constitutive` block.

```
<SolidMechanicsLagrangianSSLE
  name="lagsolve"
  timeIntegrationOption="QuasiStatic"
  discretization="FE1"
  targetRegions="{ Region2 }"
  logLevel="1">
```

## 2.4.4 Finite element discretization

The computational domain is discretized by C3D8 elements with the first order interpolation functions at each direction in the parent domain. The 2 x 2 x 2 Gauss quadrature rule is adopted to be compatible with the first order interpolation functions.

```
<NumericalMethods>
  <FiniteElements>
    <FiniteElementSpace
      name="FE1"
      order="1"/>
    </FiniteElements>
  </NumericalMethods>
```

## 2.4.5 Constitutive model

Recall that in the `SolidMechanicsLagrangianSSLE` block, `shale` is designated as the material in the computational domain. Here, the material is defined as linear isotropic.

```
<ElasticIsotropic
  name="shale"
  defaultDensity="2700"
  defaultBulkModulus="5.5556e9"
  defaultShearModulus="4.16667e9"/>
```

## 2.4.6 Boundary conditions

As aforementioned, the beam is fixed on one end, and subjects to surface traction on the other end. These boundary conditions are set up through the `FieldSpecifications` block. Here, `nodeManager` and `faceManager` in the `objectPath` indicate that the boundary conditions are applied to the element nodes and faces, respectively. Component 0, 1, and 2 refer to the x, y, and z direction, respectively. And the non-zero values given by `Scale` indicate the magnitude of the loading. Some shorthands, such as `xneg` and `xpos`, are used as the locations where the boundary conditions are applied in the computational domain. For instance, `xneg` means the portion of the computational domain located at the left-most in the x-axis, while `xpos` refers to the portion located at the right-most area in the x-axis. Similar shorthands include `ypos`, `yneg`, `zpos`, and `zneg`. Particularly, the time-dependent loading applied at the beam tip is defined through a function with the name `timeFunction`.

```
<FieldSpecifications>
  <FieldSpecification
    name="xnegconstraint"
    objectPath="nodeManager"
    fieldName="totalDisplacement"
    component="0"
    scale="0.0"
```

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```

    setNames="{ xneg }"/>

    <FieldSpecification
      name="yconstraint"
      objectPath="nodeManager"
      fieldName="totalDisplacement"
      component="1"
      scale="0.0"
      setNames="{ xneg }"/>

    <FieldSpecification
      name="zconstraint"
      objectPath="nodeManager"
      fieldName="totalDisplacement"
      component="2"
      scale="0.0"
      setNames="{ zneg, zpos }"/>

    <Traction
      name="xposconstraint"
      objectPath="faceManager"
      scale="1.0e6"
      direction="{ 0, 1, 0 }"
      functionName="timeFunction"
      setNames="{ xpos }"/>
  </FieldSpecifications>

```

### 2.4.7 Table function

A table function is used to define the time-dependent loading at the beam tip. The `coordinates` and `values` form a time-magnitude pair for the loading time history. In this case, the loading magnitude increases linearly as the time evolves.

```

<Functions>
  <TableFunction
    name="timeFunction"
    inputVarNames="{ time }"
    coordinates="{ 0.0, 10.0 }"
    values="{ 0.0, 10.0 }"/>
</Functions>

```

### 2.4.8 Execution

Finally, the execution of the simulation is set up in the `Events` block, where `target` points to the solid mechanics solver defined in the `Solvers` block, and the time increment `forceDt` is set as 1.0s.

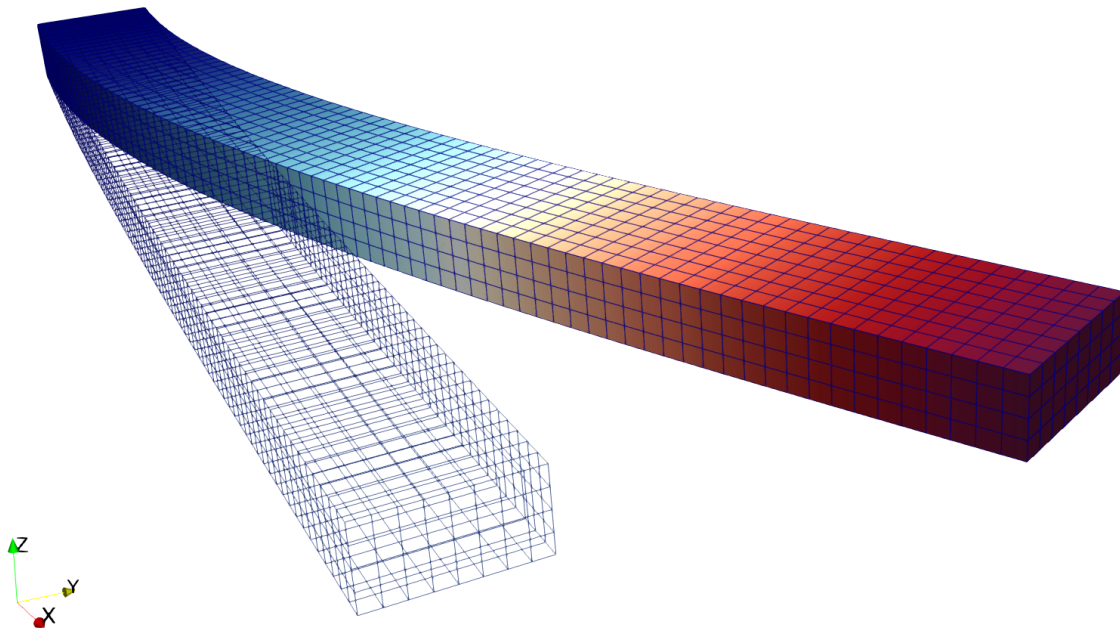
```

<PeriodicEvent
  name="solverApplications"
  forceDt="1.0"
  target="/Solvers/lagsolve"/>

```

### 2.4.9 Result

The deformed beam is shown as following (notice that the displacement is visually magnified):



### 2.4.10 To go further

#### Feedback on this tutorial

This concludes the solid mechanics for small-strain linear elasticity tutorial. For any feedback on this tutorial, please submit a [GitHub issue](#) on the project's [GitHub](#) page.

#### For more details

- More on meshes, please see *[Meshes](#)*.
- More on events, please see *[Event Management](#)*.

## 3.1 Multiphase Flow

### Context

In this example, we set up a multiphase, multicomponent test case (see *Compositional Multiphase Flow Solver*). The permeability field corresponds to the two bottom layers (layers 84 and 85) of the SPE10 test case. The thermodynamic behavior of the fluid mixture is specified using a simple immiscible two-phase (Dead-Oil) model. Injection and production are simulated using boundary conditions.

### Objective

The main objective of this example is to review the main elements of a simple two-phase simulation in GEOSX, including:

- the compositional multiphase flow solver,
- the multiphase constitutive models,
- the specifications of multiphase boundary conditions.

### Input file

This example is based on the XML file located at

```
inputFiles/compositionalMultiphaseFlow/benchmarks/SPE10/deadOilSpe10Layers84_85_base_  
↪iterative.xml
```

The XML file considered here follows the typical structure of the GEOSX input files:

1. *Solver*
2. *Mesh*
3. *Geometry*
4. *Events*
5. *NumericalMethods*

6. *ElementRegions*
7. *Constitutive*
8. *FieldSpecifications*
9. *Outputs*

### 3.1.1 Multiphase flow solver

In GEOSX, the setup of a multiphase simulation starts in the **Solvers** XML block of the input file. This example relies on a solver of type **CompositionalMultiphaseFVM** that implements a fully implicit finite-volume scheme based on the standard two-point approximation of the flux (TPFA). More information on this solver can be found at [Compositional Multiphase Flow Solver](#).

Let us have a closer look at the **Solvers** XML block displayed below. The solver has a name (here, `compflow`) that can be chosen by the user and is not imposed by GEOSX. Note that this name is used in the **Events** XML block to trigger the application of the solver. Using the `targetRegions` attribute, the solver defines the target regions on which it is applied. In this example, there is only one region, named `reservoir`.

The **CompositionalMultiphaseFVM** block contains two important sub-blocks, namely **NonlinearSolverParameters** and **LinearSolverParameters**. In **NonlinearSolverParameters**, one can finely tune the nonlinear tolerance, the application of the linear search algorithm, and the heuristics used to increase the time step size. In **LinearSolverParameters**, the user can specify the linear tolerance, the type of (direct or iterative) linear solver, and the type of preconditioner, if any. For large multiphase flow problems, we recommend using an iterative linear solver (`solverType="gmres"` or `solverType="fgmres"`) combined with the multigrid reduction (MGR) preconditioner (`preconditionerType="mgr"`). More information about the MGR preconditioner can be found in [Linear Solvers](#).

---

**Note:** For non-trivial simulations, we recommend setting the `initialDt` attribute to a small value (relative to the time scale of the problem) in seconds. If the simulation appears to be slow, use `logLevel="1"` in **CompositionalMultiphaseFVM** to detect potential Newton convergence problems. If the Newton solver struggles, please set `lineSearchAction="Attempt"` in **NonlinearSolverParameters**. If the Newton convergence is good, please add `logLevel="1"` in the **LinearSolverParameters** block to detect linear solver problems, especially if an iterative linear solver is used.

---

---

**Note:** To use the linear solver options of this example, you need to ensure that GEOSX is configured to use the Hypre linear solver package.

---

```
<Solvers>

  <CompositionalMultiphaseFVM
    name="compflow"
    logLevel="1"
    discretization="fluidTPFA"
    targetRegions="{ reservoir }"
    temperature="300"
    useMass="1"
    initialDt="1e3"
    maxCompFractionChange="0.1">
    <NonlinearSolverParameters
      newtonTol="1.0e-4"
      newtonMaxIter="40"
```

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```

    maxTimeStepCuts="10"
    lineSearchAction="None"/>
    <LinearSolverParameters
      solverType="fgmres"
      preconditionerType="mgr"
      krylovTol="1.0e-5"/>
    </CompositionalMultiphaseFVM>
  </Solvers>

```

### 3.1.2 Mesh

In this simulation, we define a simple mesh generated internally using the **InternalMesh** generator, as illustrated in the previous examples. The mesh dimensions and cell sizes are chosen to be those specified in the SPE10 test case, but are limited to the two bottom layers. The mesh description must be done in meters.

```

<Mesh>
  <InternalMesh
    name="mesh"
    elementTypes="{ C3D8 }"
    xCoords="{ 0, 365.76 }"
    yCoords="{ 0, 670.56 }"
    zCoords="{ 0, 1.22 }"
    nx="{ 60 }"
    ny="{ 220 }"
    nz="{ 2 }"
    cellBlockNames="{ block }"/>
  </Mesh>

```

### 3.1.3 Geometry

As in the previous examples, the **Geometry** XML block is used to select the cells in which the boundary conditions are applied. To mimic the setup of the original SPE10 test case, we place a source term in the middle of the domain, and a sink term in each corner. The specification of the boundary conditions applied to the selected mesh cells is done in the **FieldSpecifications** block of the XML file using the names of the boxes defined here.

```

<Geometry>

  <Box
    name="source"
    xMin="{ 182.85, 335.25, -0.01 }"
    xMax="{ 189.00, 338.35, 2.00 }"/>

  <Box
    name="sink1"
    xMin="{ -0.01, -0.01, -0.01 }"
    xMax="{ 6.126, 3.078, 2.00 }"/>

  <Box
    name="sink2"
    xMin="{ -0.01, 667.482, -0.01 }"
    xMax="{ 6.126, 670.60, 2.00 }"/>

  <Box
    name="sink3"
    xMin="{ 359.634, -0.01, -0.01 }"

```

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```

    xMax="{ 365.8, 3.048, 2.00 }"/>
  <Box
    name="sink4"
    xMin="{ 359.634, 667.482, -0.01 }"
    xMax="{ 365.8, 670.60, 2.00 }"/>

</Geometry>

```

### 3.1.4 Events

In the **Events** XML block of this example, we specify two types of **PeriodicEvents** serving different purposes, namely solver application and result output.

The periodic event named `solverApplications` triggers the application of the solver on its target region. This event must point to the solver by name. In this example, the name of the solver is `compflow` and was defined in the **Solvers** block. The time step is initialized using the `initialDt` attribute of the flow solver. Then, if the solver converges in less than a certain number of nonlinear iterations (by default, 40% of the maximum number of nonlinear iterations), the time step will be increased until it reaches the maximum time step size specified with `maxEventDt`. If the time step fails, the time step will be cut. The parameters defining the time stepping strategy can be finely tuned by the user in the flow solver block. Note that all times are in seconds.

The output event forces GEOSX to write out the results at the frequency specified by the attribute `timeFrequency`. Here, we choose to output the results using the VTK format (see [Tutorial 2: External Meshes](#) for a tutorial that uses the Silo output file format). Using `targetExactTimestep=1` in this XML block forces GEOSX to adapt the time stepping to ensure that an output is generated exactly at the time frequency requested by the user. In the `target` attribute, we must use the name defined in the **VTK** XML tag inside the **Output** XML section, as documented at the end of this example (here, `vtkOutput`).

More information about events can be found at [Event Management](#).

```

<Events
  maxTime="2e6">

  <PeriodicEvent
    name="outputs"
    timeFrequency="5e5"
    targetExactTimestep="1"
    target="/Outputs/vtkOutput"/>

  <PeriodicEvent
    name="solverApplications"
    maxEventDt="5e5"
    target="/Solvers/compflow"/>

  <PeriodicEvent
    name="restarts"
    timeFrequency="1e6"
    targetExactTimestep="0"
    target="/Outputs/restartOutput"/>

</Events>

```

### 3.1.5 Numerical methods

In the **NumericalMethods** XML block, we select a two-point flux approximation (TPFA) finite-volume scheme to discretize the governing equations on the reservoir mesh. TPFA is currently the only numerical scheme that can be used with a flow solver of type **CompositionalMultiphaseFVM**.

```
<NumericalMethods>
  <FiniteVolume>
    <TwoPointFluxApproximation
      name="fluidTPFA"/>
    </FiniteVolume>
  </NumericalMethods>
```

### 3.1.6 Reservoir region

In the **ElementRegions** XML block, we define a **CellElementRegion** named `reservoir` corresponding to the reservoir mesh. The attribute `cellBlocks` is set to `block` to point this element region to the hexahedral mesh defined internally.

The **CellElementRegion** must also point to the constitutive models that are used to update the dynamic rock and fluid properties in the cells of the reservoir mesh. The names `fluid`, `rock`, and `relperm` used for this in the `materialList` correspond to the attribute name of the **Constitutive** block.

```
<ElementRegions>
  <CellElementRegion
    name="reservoir"
    cellBlocks="{ block }"
    materialList="{ fluid, rock, relperm }"/>
  </ElementRegions>
```

### 3.1.7 Constitutive models

For a simulation performed with the **CompositionalMultiphaseFVM** physics solver, at least four types of constitutive models must be specified in the **Constitutive** XML block:

- a fluid model describing the thermodynamics behavior of the fluid mixture,
- a relative permeability model,
- a rock permeability model,
- a rock porosity model.

All these models use SI units exclusively. A capillary pressure model can also be specified in this block but is omitted here for simplicity.

Here, we introduce a fluid model describing a simplified mixture thermodynamic behavior. Specifically, we use an immiscible two-phase (Dead Oil) model by placing the XML tag **DeadOilFluid**. Other fluid models can be used with the **CompositionalMultiphaseFVM** solver, as explained in *Fluid Models*.

With the tag **BrooksCoreyRelativePermeability**, we define a relative permeability model. A list of available relative permeability models can be found at *Relative Permeability Models*.

The properties are chosen to match those of the original SPE10 test case.

**Note:** The names and order of the phases listed for the attribute `phaseNames` must be identical in the fluid model (here, **DeadOilFluid**) and the relative permeability model (here, **BrooksCoreyRelativePermeability**). Otherwise, GEOSX will throw an error and terminate.

---

We also introduce models to define rock compressibility and permeability. This step is similar to what is described in the previous examples (see for instance *Tutorial 1: First Steps*).

We remind the reader that the attribute name of the constitutive models defined here must be used in the **ElementRegions** and **Solvers** XML blocks to point the element regions and the physics solvers to their respective constitutive models.

```
<Constitutive>

  <DeadOilFluid
    name="fluid"
    phaseNames="{ oil, water }"
    surfaceDensities="{ 800.0, 1022.0 }"
    componentMolarWeight="{ 114e-3, 18e-3 }"
    hydrocarbonFormationVolFactorTableNames="{ B_o_table }"
    hydrocarbonViscosityTableNames="{ visc_o_table }"
    waterReferencePressure="30600000.1"
    waterFormationVolumeFactor="1.03"
    waterCompressibility="0.00000000041"
    waterViscosity="0.0003"/>

  <CompressibleSolidConstantPermeability
    name="rock"
    solidModelName="nullSolid"
    porosityModelName="rockPorosity"
    permeabilityModelName="rockPerm"/>

  <NullModel
    name="nullSolid"/>

  <PressurePorosity
    name="rockPorosity"
    defaultReferencePorosity="0.1"
    referencePressure="1.0e7"
    compressibility="1e-10"/>

  <BrooksCoreyRelativePermeability
    name="relperm"
    phaseNames="{ oil, water }"
    phaseMinVolumeFraction="{ 0.0, 0.0 }"
    phaseRelPermExponent="{ 2.0, 2.0 }"
    phaseRelPermMaxValue="{ 1.0, 1.0 }"/>

  <ConstantPermeability
    name="rockPerm"
    permeabilityComponents="{ 1.0e-14, 1.0e-14, 1.0e-18 }"/>

</Constitutive>
```

### 3.1.8 Initial and boundary conditions

In the **FieldSpecifications** section, we define the initial and boundary conditions as well as the geological properties (porosity, permeability). All this is done using SI units. Here, we focus on the specification of the initial and boundary conditions for a simulation performed with the **CompositionalMultiphaseFVM** solver. We refer to [Tutorial 1: First Steps](#) for a more general discussion on the **FieldSpecification** XML blocks.

For a simulation performed with the **CompositionalMultiphaseFVM** solver, we have to set the initial pressure as well as the initial global component fractions (in this case, the oil and water component fractions). The `component` attribute of the **FieldSpecification** XML block must use the order in which the `phaseNames` have been defined in the **DeadOilFluid** XML block. In other words, `component=0` is used to initialize the oil global component fraction and `component=1` is used to initialize the water global component fraction, because we previously set `phaseNames="{oil, water}"` in the **DeadOilFluid** XML block.

To specify the sink terms, we use the **FieldSpecification** mechanism in a similar fashion to impose the sink pressure and composition. This is done to mimic a pressure-controlled well (before breakthrough). To specify the source term, we use a **SourceFlux** block to impose a fixed mass injection rate of component 1 (water) to mimic a rate-controlled well.

```
<FieldSpecifications>
  <FieldSpecification
    name="permx"
    component="0"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions/reservoir/block"
    fieldName="rockPerm_permeability"
    functionName="permxFunc"
    scale="9.869233e-16"/>
  <FieldSpecification
    name="permy"
    component="1"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions/reservoir/block"
    fieldName="rockPerm_permeability"
    functionName="permyFunc"
    scale="9.869233e-16"/>
  <FieldSpecification
    name="permz"
    component="2"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions/reservoir/block"
    fieldName="rockPerm_permeability"
    functionName="permzFunc"
    scale="9.869233e-16"/>

  <FieldSpecification
    name="referencePorosity"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions/reservoir/block"
    fieldName="rockPorosity_referencePorosity"
    functionName="poroFunc"
    scale="1.0"/>

  <FieldSpecification
```

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```

    name="initialPressure"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions/reservoir/block"
    fieldName="pressure"
    scale="4.1369e+7"/>
<FieldSpecification
  name="initialComposition_oil"
  initialCondition="1"
  setNames="{ all }"
  objectPath="ElementRegions/reservoir/block"
  fieldName="globalCompFraction"
  component="0"
  scale="0.9995"/>
<FieldSpecification
  name="initialComposition_water"
  initialCondition="1"
  setNames="{ all }"
  objectPath="ElementRegions/reservoir/block"
  fieldName="globalCompFraction"
  component="1"
  scale="0.0005"/>

<SourceFlux
  name="sourceTerm"
  objectPath="ElementRegions/reservoir/block"
  scale="-0.07279"
  component="1"
  setNames="{ source }"/>

<FieldSpecification
  name="sinkPressure"
  setNames="{ sink1, sink2, sink3, sink4 }"
  objectPath="ElementRegions/reservoir/block"
  fieldName="pressure"
  scale="2.7579e+7"/>
<FieldSpecification
  name="sinkComposition_oil"
  setNames="{ sink1, sink2, sink3, sink4 }"
  objectPath="ElementRegions/reservoir/block"
  fieldName="globalCompFraction"
  component="0"
  scale="0.9995"/>
<FieldSpecification
  name="sinkComposition_water"
  setNames="{ sink1, sink2, sink3, sink4 }"
  objectPath="ElementRegions/reservoir/block"
  fieldName="globalCompFraction"
  component="1"
  scale="0.0005"/>

</FieldSpecifications>

```

### 3.1.9 Output

In this section, we request an output of the results in VTK format. Note that the name defined here must match the names used in the **Events** XML block to define the output frequency.

```
<Outputs>
  <VTK
    name="vtkOutput"/>

  <Restart
    name="restartOutput"/>

</Outputs>
```

All elements are now in place to run GEOSX.

### 3.1.10 Running GEOSX

The first few lines appearing to the console are indicating that the XML elements are read and registered correctly:

```
Adding Solver of type CompositionalMultiphaseFVM, named compflow
Adding Mesh: InternalMesh, mesh
Adding Geometric Object: Box, source
Adding Geometric Object: Box, sink1
Adding Geometric Object: Box, sink2
Adding Geometric Object: Box, sink3
Adding Geometric Object: Box, sink4
Adding Event: PeriodicEvent, outputs
Adding Event: PeriodicEvent, solverApplications
TableFunction: permxFunc
TableFunction: permyFunc
TableFunction: permzFunc
TableFunction: poroFunc
TableFunction: B_o_table
TableFunction: visc_o_table
Adding Output: VTK, vtkOutput
Adding Object CellElementRegion named region from ObjectManager::Catalog.
region/block/fluid is allocated with 1 quadrature points.
region/block/rock is allocated with 1 quadrature points.
aaregion/block/relperm is allocated with 1 quadrature points.
```

At this point, we are done with the case set-up and the code steps into the execution of the simulation itself:

```
Time: 0s, dt:1000s, Cycle: 0

  Attempt: 0, NewtonIter: 0
  ( Rfluid ) = (2.28e+00) ;      ( R ) = ( 2.28e+00 ) ;
  Attempt: 0, NewtonIter: 1
  ( Rfluid ) = (8.83e-03) ;      ( R ) = ( 8.83e-03 ) ;
  Last LinSolve(iter,res) = ( 2, 2.74e-03 ) ;
  Attempt: 0, NewtonIter: 2
  ( Rfluid ) = (8.86e-05) ;      ( R ) = ( 8.86e-05 ) ;
  Last LinSolve(iter,res) = ( 2, 8.92e-03 ) ;

compflow: Max phase CFL number: 0.00399585
```

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```
compflow: Max component CFL number: 0.152466
compflow: Newton solver converged in less than 16 iterations, time-step required will
↳be doubled.
```

### 3.1.11 Visualization

A file compatible with Paraview is produced in this example. It is found in the output folder, and usually has the extension *.pvd*. More details about this file format can be found [here](#). We can load this file into Paraview directly and visualize results:

### 3.1.12 To go further

#### Feedback on this example

This concludes the example on setting up an immiscible two-phase flow simulation in a channelized permeability field. For any feedback on this example, please submit a [GitHub issue on the project's GitHub page](#).

#### For more details

- A complete description of the reservoir flow solver is found here: *Compositional Multiphase Flow Solver*.
- The available constitutive models are listed at *Constitutive Models*.

## 3.2 Multiphase Flow with Wells

### Context

In this example, we build on the concepts presented in *Multiphase Flow* to show how to set up a multiphase water injection problem with wells in the three-dimensional *Egg model*. The twelve wells (four producers and eight injectors) are placed according to the description of the original test case.

### Objectives

In this example, we re-use many GEOSX features already presented in *Multiphase Flow*, but we now focus on:

- how to import an external mesh with embedded geological properties (permeability) in the VTK format (*.vtu*),
- how to set up the wells.

### Input file

This example is based on the XML file located at

```
../../../../inputFiles/compositionalMultiphaseWell/benchmarks/Egg/deadOilEgg_base_
↳iterative.xml
```

The mesh file corresponding to the Egg model is stored in the GEOSXDATA repository. Therefore, you must first download the GEOSXDATA repository in the same folder as the GEOSX repository to run this test case.

---

**Note:** GEOSXDATA is a separate repository in which we store large mesh files in order to keep the main GEOSX repository lightweight.

---

The XML file considered here follows the typical structure of the GEOSX input files:



1. *Solver*
2. *Mesh*
3. *Events*
4. *NumericalMethods*
5. *ElementRegions*
6. *Constitutive*
7. *FieldSpecifications*
8. *Outputs*
9. *Tasks*

### 3.2.1 Coupling the flow solver with wells

In GEOSX, the simulation of reservoir flow with wells is set up by combining three solvers listed and parameterized in the **Solvers** XML block of the input file. We introduce separately a flow solver and a well solver acting on different regions of the domain—respectively, the reservoir region and the well regions. To drive the simulation and bind these single-physics solvers, we also specify a *coupling solver* between the reservoir flow solver and the well solver. This coupling of single-physics solvers is the generic approach used in GEOSX to define multiphysics problems. It is illustrated in *Poromechanics* for a poroelastic test case.

The three solvers employed in this example are:

- the single-physics reservoir flow solver, a solver of type **CompositionalMultiphaseFVM** named `compositionalMultiphaseFlow` (more information on this solver at [Compositional Multiphase Flow Solver](#)),
- the single-physics well solver, a solver of type **CompositionalMultiphaseWell** named `compositionalMultiphaseWell` (more information on this solver at [Compositional Multiphase Well Solver](#)),
- the coupling solver that binds the two single-physics solvers above, an object of type **CompositionalMultiphaseReservoir** named `coupledFlowAndWells`.

The **Solvers** XML block is shown below. The coupling solver points to the two single-physics solvers using the attributes `flowSolverName` and `wellSolverName`. These names can be chosen by the user and are not imposed by GEOSX. The flow solver is applied to the reservoir and the well solver is applied to the wells, as specified by their respective `targetRegions` attributes.

The simulation is fully coupled and driven by the coupled solver. Therefore, the time stepping information (here, `initialDt`, but there may be other parameters used to fine-tune the time stepping strategy), the nonlinear solver parameters, and the linear solver parameters must be specified at the level of the coupling solver. There is no need to specify these parameters at the level of the single-physics solvers. Any solver information specified in the single-physics XML blocks will not be taken into account.

---

**Note:** It is worth repeating the `logLevel="1"` parameter at the level of the well solver to make sure that a notification is issued when the well control is switched (from rate control to BHP control, for instance).

---

Here, we instruct GEOSX to perform at most `newtonMaxIter = "10"` Newton iterations. GEOSX will adjust the time step size as follows:

- if the Newton solver converges in `timeStepIncreaseIterLimit × newtonMaxIter = 5` iterations or fewer, GEOSX will double the time step size for the next time step,

- if the Newton solver converges in `timeStepDecreaseIterLimit` x `newtonMaxIter` = 8 iterations or more, GEOSX will reduce the time step size for the next time step by a factor `timestepCutFactor` = 0.1,
- if the Newton solver fails to converge in `newtonMaxIter` = 10, GEOSX will cut the time step size by a factor `timestepCutFactor` = 0.1 and restart from the previous converged time step.

The maximum number of time step cuts is specified by the attribute `maxTimeStepCuts`. Note that a backtracking line search can be activated by setting the attribute `lineSearchAction` to `Attempt` or `Require`. If `lineSearchAction` = "Attempt", we accept the nonlinear iteration even if the line search does not reduce the residual norm. If `lineSearchAction` = "Require", we cut the time step if the line search does not reduce the residual norm.

**Note:** To use the linear solver options of this example, you need to ensure that GEOSX is configured to use the Hypr linear solver package.

```
<Solvers>
  <CompositionalMultiphaseReservoir
    name="coupledFlowAndWells"
    flowSolverName="compositionalMultiphaseFlow"
    wellSolverName="compositionalMultiphaseWell"
    logLevel="1"
    initialDt="1e4"
    targetRegions="{ reservoir, wellRegion1, wellRegion2, wellRegion3, wellRegion4,
↪wellRegion5, wellRegion6, wellRegion7, wellRegion8, wellRegion9, wellRegion10,
↪wellRegion11, wellRegion12 }">
    <NonlinearSolverParameters
      newtonTol="1.0e-4"
      newtonMaxIter="25"
      timeStepDecreaseIterLimit="0.9"
      timeStepIncreaseIterLimit="0.6"
      timeStepCutFactor="0.1"
      maxTimeStepCuts="10"
      lineSearchAction="None"/>
    <LinearSolverParameters
      solverType="fgmres"
      preconditionerType="mgr"
      krylovTol="1e-4"
      krylovAdaptiveTol="1"
      krylovWeakestTol="1e-2"/>
  </CompositionalMultiphaseReservoir>

  <CompositionalMultiphaseFVM
    name="compositionalMultiphaseFlow"
    targetRegions="{ reservoir }"
    discretization="fluidTPFA"
    temperature="297.15"
    maxCompFractionChange="0.3"
    logLevel="1"
    useMass="1"/>

  <CompositionalMultiphaseWell
    name="compositionalMultiphaseWell"
    targetRegions="{ wellRegion1, wellRegion2, wellRegion3, wellRegion4,
↪wellRegion5, wellRegion6, wellRegion7, wellRegion8, wellRegion9, wellRegion10,
↪wellRegion11, wellRegion12 }"
```

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```

maxCompFractionChange="0.5"
logLevel="1"
useMass="1">
<WellControls
  name="wellControls1"
  type="producer"
  control="BHP"
  referenceElevation="28"
  targetBHP="3.9e7"
  targetPhaseRate="1e6"
  targetPhaseName="oil"/>
<WellControls
  name="wellControls2"
  type="producer"
  control="BHP"
  referenceElevation="28"
  targetBHP="3.9e7"
  targetPhaseRate="1e6"
  targetPhaseName="oil"/>

```

### 3.2.2 Mesh definition and well geometry

In the presence of wells, the **Mesh** block of the XML input file includes two parts:

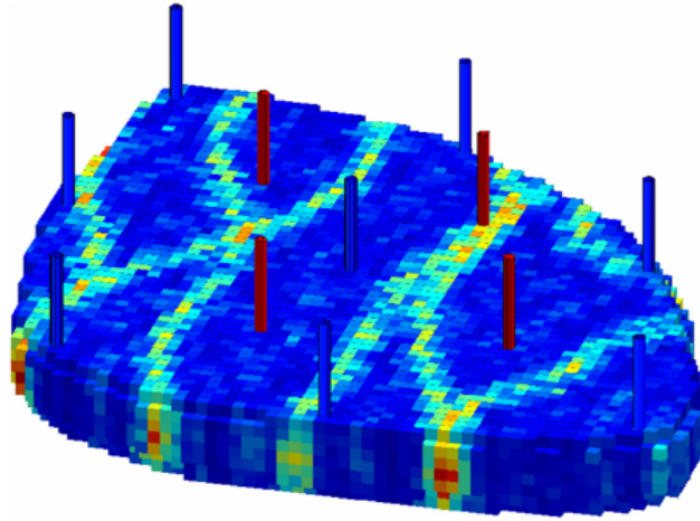
- a sub-block **VTKMesh** defining the reservoir mesh (see [Tutorial 2: External Meshes](#) for more on this),
- a collection of sub-blocks **InternalWell** defining the geometry of the wells.

The reservoir mesh is imported from a `.vtu` file that contains the mesh geometry and also includes the permeability values in the x, y, and z directions. These quantities must be specified using the metric unit system, i.e., in meters for the well geometry and square meters for the permeability field. We note that the mesh file only contains the active cells, so there is no keyword needed in the XML file to define them.

Each well is defined internally (i.e., not imported from a file) in a separate **InternalWell** XML sub-block. An **InternalWell** sub-block must point to the reservoir mesh that the well perforates using the attribute `meshName`, to the region corresponding to this well using the attribute `wellRegionName`, and to the control of this well using the attribute `wellControl`. Each block **InternalWell** must point to the reservoir mesh (using the attribute `meshName`), the corresponding well region (using the attribute `wellRegionName`), and the corresponding well control (using the attribute `wellControlName`).

Each well is defined using a vertical polyline going through the seven layers of the mesh, with a perforation in each layer. The well placement implemented here follows the pattern of the original test case. The well geometry must be specified in meters.

The location of the perforations is found internally using the linear distance along the wellbore from the top of the well, specified by the attribute `distanceFromHead`. It is the responsibility of the user to make sure that there is a perforation in the bottom cell of the well mesh otherwise an error will be thrown and the simulation will terminate. For each perforation, the well transmissibility factors employed to compute the perforation rates are calculated internally using the Peaceman formulation.



```

<Mesh>
  <VTKMesh
    name="mesh"
    file="../../../../../../GEOSXDATA/DataSets/Egg/egg.vtu"
    fieldsToImport="{ PERM }"
    fieldNamesInGEOSX="{ rockPerm_permeability }"/>

  <InternalWell
    name="wellProducer1"
    wellRegionName="wellRegion1"
    wellControlsName="wellControls1"
    meshName="mesh"
    polylineNodeCoords="{ { 124, 340, 28 },
                          { 124, 340, 0 } }"
    polylineSegmentConn="{ { 0, 1 } }"
    radius="0.1"
    numElementsPerSegment="7">
    <Perforation
      name="producer1_perf1"
      distanceFromHead="2"/>
    <Perforation
      name="producer1_perf2"
      distanceFromHead="6"/>
    <Perforation
      name="producer1_perf3"
      distanceFromHead="10"/>
    <Perforation
      name="producer1_perf4"
      distanceFromHead="14"/>
    <Perforation
      name="producer1_perf5"
      distanceFromHead="18"/>
    <Perforation
      name="producer1_perf6"
      distanceFromHead="22"/>
    <Perforation
      name="producer1_perf7"
      distanceFromHead="26"/>
  </InternalWell>

```

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```

</InternalWell>

<InternalWell
  name="wellProducer2"
  wellRegionName="wellRegion2"
  wellControlsName="wellControls2"
  meshName="mesh"
  polylineNodeCoords="{ { 276, 316, 28 },
                        { 276, 316, 0 } }"
  polylineSegmentConn="{ { 0, 1 } }"
  radius="0.1"
  numElementsPerSegment="7">
  <Perforation
    name="producer2_perf1"
    distanceFromHead="2"/>
  <Perforation
    name="producer2_perf2"
    distanceFromHead="6"/>
  <Perforation
    name="producer2_perf3"
    distanceFromHead="10"/>
  <Perforation
    name="producer2_perf4"
    distanceFromHead="14"/>
  <Perforation
    name="producer2_perf5"
    distanceFromHead="18"/>
  <Perforation
    name="producer2_perf6"
    distanceFromHead="22"/>
  <Perforation
    name="producer2_perf7"
    distanceFromHead="26"/>
</InternalWell>

```

### 3.2.3 Events

In the **Events** XML block, we specify four types of **PeriodicEvents**.

The periodic event named `solverApplications` notifies GEOSX that the coupled solver `coupledFlowAndWells` has to be applied to its target regions (here, reservoir and wells) at every time step. The time stepping strategy has been fully defined in the **CompositionalMultiphaseReservoir** coupling block using the `initialDt` attribute and the **NonlinearSolverParameters** nested block.

We also define an output event instructing GEOSX to write out `.vtk` files at the time frequency specified by the attribute `timeFrequency`. Here, we choose to output the results using the VTK format (see [Tutorial 2: External Meshes](#) for an example that uses the Silo output file format). The `target` attribute must point to the **VTK** sub-block of the **Outputs** block (defined at the end of the XML file) by name (here, `vtkOutput`).

We define the events involved in the collection and output of the well production rates following the procedure defined in [Tasks Manager](#). The time history collection events trigger the collection of the well rates at the desired frequency, while the time history output events trigger the output of the HDF5 files containing the time series. These events point by name to the corresponding blocks of the **Tasks** and **Outputs** XML blocks, respectively. Here, these names are `wellRateCollection1` and `timeHistoryOutput1`.

```
<Events
  maxTime="1.5e7">
  <PeriodicEvent
    name="vtk"
    timeFrequency="2e6"
    target="/Outputs/vtkOutput"/>

  <PeriodicEvent
    name="timeHistoryOutput1"
    timeFrequency="1.5e7"
    target="/Outputs/timeHistoryOutput1"/>

  <PeriodicEvent
    name="timeHistoryOutput2"
    timeFrequency="1.5e7"
    target="/Outputs/timeHistoryOutput2"/>

  <PeriodicEvent
    name="timeHistoryOutput3"
    timeFrequency="1.5e7"
    target="/Outputs/timeHistoryOutput3"/>

  <PeriodicEvent
    name="timeHistoryOutput4"
    timeFrequency="1.5e7"
    target="/Outputs/timeHistoryOutput4"/>

  <PeriodicEvent
    name="solverApplications"
    maxEventDt="5e5"
    target="/Solvers/coupledFlowAndWells"/>

  <PeriodicEvent
    name="timeHistoryCollection1"
    timeFrequency="1e6"
    target="/Tasks/wellRateCollection1"/>

  <PeriodicEvent
    name="timeHistoryCollection2"
    timeFrequency="1e6"
    target="/Tasks/wellRateCollection2"/>

  <PeriodicEvent
    name="timeHistoryCollection3"
    timeFrequency="1e6"
    target="/Tasks/wellRateCollection3"/>

  <PeriodicEvent
    name="timeHistoryCollection4"
    timeFrequency="1e6"
    target="/Tasks/wellRateCollection4"/>

  <PeriodicEvent
    name="restarts"
    timeFrequency="7.5e6"
    targetExactTimestep="0"
    target="/Outputs/restartOutput"/>
```

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`</Events>`

### 3.2.4 Numerical methods

In the `NumericalMethods` XML block, we instruct GEOSX to use a TPFA finite-volume numerical scheme. This part is similar to the corresponding section of *Multiphase Flow*, and has been adapted to match the specifications of the Egg model.

```
<NumericalMethods>
  <FiniteVolume>
    <TwoPointFluxApproximation
      name="fluidTPFA"/>
    </FiniteVolume>
  </NumericalMethods>
```

### 3.2.5 Reservoir and well regions

In this section of the input file, we follow the procedure already described in *Multiphase Flow* for the definition of the reservoir region with multiphase constitutive models.

We associate a **CellElementRegion** named `reservoir` to the reservoir mesh. Since we have imported a mesh with one region consisting of hexahedral cells, we must set the attribute `cellBlocks` to `hexahedra`.

**Note:** If you use a name that is not `hexahedra` for this attribute, GEOSX will throw an error at the beginning of the simulation.

We also associate a **WellElementRegion** to each well. As the **CellElementRegion**, it contains a `materialList` that must point (by name) to the constitutive models defined in the **Constitutive** XML block.

```
<ElementRegions>
  <CellElementRegion
    name="reservoir"
    cellBlocks="{ hexahedra }"
    materialList="{ fluid, rock, relperm }"/>

  <WellElementRegion
    name="wellRegion1"
    materialList="{ fluid, relperm }"/>

  <WellElementRegion
    name="wellRegion2"
    materialList="{ fluid, relperm }"/>
```

### 3.2.6 Constitutive models

The **CompositionalMultiphaseFVM** physics solver relies on at least four types of constitutive models listed in the **Constitutive** XML block:

- a fluid model describing the thermodynamics behavior of the fluid mixture,

- a relative permeability model,
- a rock permeability model,
- a rock porosity model.

All the parameters must be provided using the SI unit system.

This part is identical to that of *Multiphase Flow*.

```
<Constitutive>
  <DeadOilFluid
    name="fluid"
    phaseNames="{ oil, water }"
    surfaceDensities="{ 848.9, 1025.2 }"
    componentMolarWeight="{ 114e-3, 18e-3 }"
    tableFiles="{ pvdo.txt, pvtw.txt }"/>

  <BrooksCoreyRelativePermeability
    name="relperm"
    phaseNames="{ oil, water }"
    phaseMinVolumeFraction="{ 0.1, 0.2 }"
    phaseRelPermExponent="{ 4.0, 3.0 }"
    phaseRelPermMaxValue="{ 0.8, 0.75 }"/>

  <CompressibleSolidConstantPermeability
    name="rock"
    solidModelName="nullSolid"
    porosityModelName="rockPorosity"
    permeabilityModelName="rockPerm"/>

  <NullModel
    name="nullSolid"/>

  <PressurePorosity
    name="rockPorosity"
    defaultReferencePorosity="0.2"
    referencePressure="0.0"
    compressibility="1.0e-13"/>

  <ConstantPermeability
    name="rockPerm"
    permeabilityComponents="{ 1.0e-12, 1.0e-12, 1.0e-12 }"/>
</Constitutive>
```

### 3.2.7 Initial conditions

We are ready to specify the reservoir initial conditions of the problem in the **FieldSpecifications** XML block. The well variables do not have to be initialized here since they will be defined internally.

The formulation of the **CompositionalMultiphaseFVM** physics solver (documented at *Compositional Multiphase Flow Solver*) requires the definition of the initial pressure field and initial global component fractions. We define here a uniform pressure field that does not satisfy the hydrostatic equilibrium, but a hydrostatic initialization of the pressure field is possible using *Functions*:. For the initialization of the global component fractions, we remind the user that their component attribute (here, 0 or 1) is used to point to a specific entry of the phaseNames attribute in the **DeadOilFluid** block.

Note that we also define the uniform porosity field here since it is not included in the mesh file imported by the **VTKMesh**.



```

<FieldSpecifications>
  <FieldSpecification
    name="initialPressure"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions/reservoir/hexahedra"
    fieldName="pressure"
    scale="4e7"/>

  <FieldSpecification
    name="initialComposition_oil"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions/reservoir/hexahedra"
    fieldName="globalCompFraction"
    component="0"
    scale="0.9"/>

  <FieldSpecification
    name="initialComposition_water"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions/reservoir/hexahedra"
    fieldName="globalCompFraction"
    component="1"
    scale="0.1"/>
</FieldSpecifications>

```

### 3.2.8 Outputs

In this section, we request an output of the results in VTK format and an output of the rates for each producing well. Note that the name defined here must match the name used in the **Events** XML block to define the output frequency.

```

<Outputs>
  <VTK
    name="vtkOutput"/>

  <TimeHistory
    name="timeHistoryOutput1"
    sources="{ /Tasks/wellRateCollection1 }"
    filename="wellRateHistory1"/>

  <TimeHistory
    name="timeHistoryOutput2"
    sources="{ /Tasks/wellRateCollection2 }"
    filename="wellRateHistory2"/>

  <TimeHistory
    name="timeHistoryOutput3"
    sources="{ /Tasks/wellRateCollection3 }"
    filename="wellRateHistory3"/>

  <TimeHistory
    name="timeHistoryOutput4"
    sources="{ /Tasks/wellRateCollection4 }"
    filename="wellRateHistory4"/>

```

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```

<Restart
  name="restartOutput" />

</Outputs>

```

### 3.2.9 Tasks

In the **Events** block, we have defined four events requesting that a task periodically collects the rate for each producing well. This task is defined here, in the **PackCollection** XML sub-block of the **Tasks** block. The task contains the path to the object on which the field to collect is registered (here, a `WellElementSubRegion`) and the name of the field (here, `wellElementMixtureConnectionRate`). The details of the history collection mechanism can be found in *Tasks Manager*.

```

<Tasks>
  <PackCollection
    name="wellRateCollection1"
    objectPath="ElementRegions/wellRegion1/wellRegion1UniqueSubRegion"
    fieldName="wellElementMixtureConnectionRate"/>

  <PackCollection
    name="wellRateCollection2"
    objectPath="ElementRegions/wellRegion2/wellRegion2UniqueSubRegion"
    fieldName="wellElementMixtureConnectionRate"/>

  <PackCollection
    name="wellRateCollection3"
    objectPath="ElementRegions/wellRegion3/wellRegion3UniqueSubRegion"
    fieldName="wellElementMixtureConnectionRate"/>

  <PackCollection
    name="wellRateCollection4"
    objectPath="ElementRegions/wellRegion4/wellRegion4UniqueSubRegion"
    fieldName="wellElementMixtureConnectionRate"/>
</Tasks>

```

All elements are now in place to run GEOSX.

### 3.2.10 Running GEOSX

The first few lines appearing to the console are indicating that the XML elements are read and registered correctly:

```

Adding Mesh: VTKMesh, mesh
Adding Mesh: InternalWell, wellProducer1
Adding Mesh: InternalWell, wellProducer2
Adding Mesh: InternalWell, wellProducer3
Adding Mesh: InternalWell, wellProducer4
Adding Mesh: InternalWell, wellInjector1
Adding Mesh: InternalWell, wellInjector2
Adding Mesh: InternalWell, wellInjector3
Adding Mesh: InternalWell, wellInjector4
Adding Mesh: InternalWell, wellInjector5
Adding Mesh: InternalWell, wellInjector6

```

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```

Adding Mesh: InternalWell, wellInjector7
Adding Mesh: InternalWell, wellInjector8
Adding Solver of type CompositionalMultiphaseReservoir, named coupledFlowAndWells
Adding Solver of type CompositionalMultiphaseFVM, named compositionalMultiphaseFlow
Adding Solver of type CompositionalMultiphaseWell, named compositionalMultiphaseWell
Adding Event: PeriodicEvent, vtk
Adding Event: PeriodicEvent, timeHistoryOutput1
Adding Event: PeriodicEvent, timeHistoryOutput2
Adding Event: PeriodicEvent, timeHistoryOutput3
Adding Event: PeriodicEvent, timeHistoryOutput4
Adding Event: PeriodicEvent, solverApplications
Adding Event: PeriodicEvent, timeHistoryCollection1
Adding Event: PeriodicEvent, timeHistoryCollection2
Adding Event: PeriodicEvent, timeHistoryCollection3
Adding Event: PeriodicEvent, timeHistoryCollection4
Adding Event: PeriodicEvent, restarts
Adding Output: VTK, vtkOutput
Adding Output: TimeHistory, timeHistoryOutput1
Adding Output: TimeHistory, timeHistoryOutput2
Adding Output: TimeHistory, timeHistoryOutput3
Adding Output: TimeHistory, timeHistoryOutput4
Adding Output: Restart, restartOutput
Adding Object CellElementRegion named reservoir from ObjectManager::Catalog.
Adding Object WellElementRegion named wellRegion1 from ObjectManager::Catalog.
Adding Object WellElementRegion named wellRegion2 from ObjectManager::Catalog.
Adding Object WellElementRegion named wellRegion3 from ObjectManager::Catalog.
Adding Object WellElementRegion named wellRegion4 from ObjectManager::Catalog.
Adding Object WellElementRegion named wellRegion5 from ObjectManager::Catalog.
Adding Object WellElementRegion named wellRegion6 from ObjectManager::Catalog.
Adding Object WellElementRegion named wellRegion7 from ObjectManager::Catalog.
Adding Object WellElementRegion named wellRegion8 from ObjectManager::Catalog.
Adding Object WellElementRegion named wellRegion9 from ObjectManager::Catalog.
Adding Object WellElementRegion named wellRegion10 from ObjectManager::Catalog.
Adding Object WellElementRegion named wellRegion11 from ObjectManager::Catalog.
Adding Object WellElementRegion named wellRegion12 from ObjectManager::Catalog.

```

This is followed by the creation of the 18553 hexahedral cells of the imported mesh. At this point, we are done with the case set-up and the code steps into the execution of the simulation itself:

```

Time: 0s, dt:10000s, Cycle: 0
  Attempt: 0, ConfigurationIter: 0, NewtonIter: 0
    ( Rflow ) = ( 1.01e+01 ) ;      ( Rwell ) = ( 4.96e+00 ) ;      ( R ) = ( 1.13e+01
↪ ) ;
  Attempt: 0, ConfigurationIter: 0, NewtonIter: 1
    ( Rflow ) = ( 1.96e+00 ) ;      ( Rwell ) = ( 8.07e-01 ) ;      ( R ) = ( 2.12e+00
↪ ) ;
  Last LinSolve(iter,res) = ( 44, 8.96e-03 ) ;
  Attempt: 0, ConfigurationIter: 0, NewtonIter: 2
    ( Rflow ) = ( 4.14e-01 ) ;      ( Rwell ) = ( 1.19e-01 ) ;      ( R ) = ( 4.31e-01
↪ ) ;
  Last LinSolve(iter,res) = ( 44, 9.50e-03 ) ;
  Attempt: 0, ConfigurationIter: 0, NewtonIter: 3
    ( Rflow ) = ( 1.77e-02 ) ;      ( Rwell ) = ( 9.38e-03 ) ;      ( R ) = ( 2.00e-02
↪ ) ;
  Last LinSolve(iter,res) = ( 47, 8.69e-03 ) ;
  Attempt: 0, ConfigurationIter: 0, NewtonIter: 4

```

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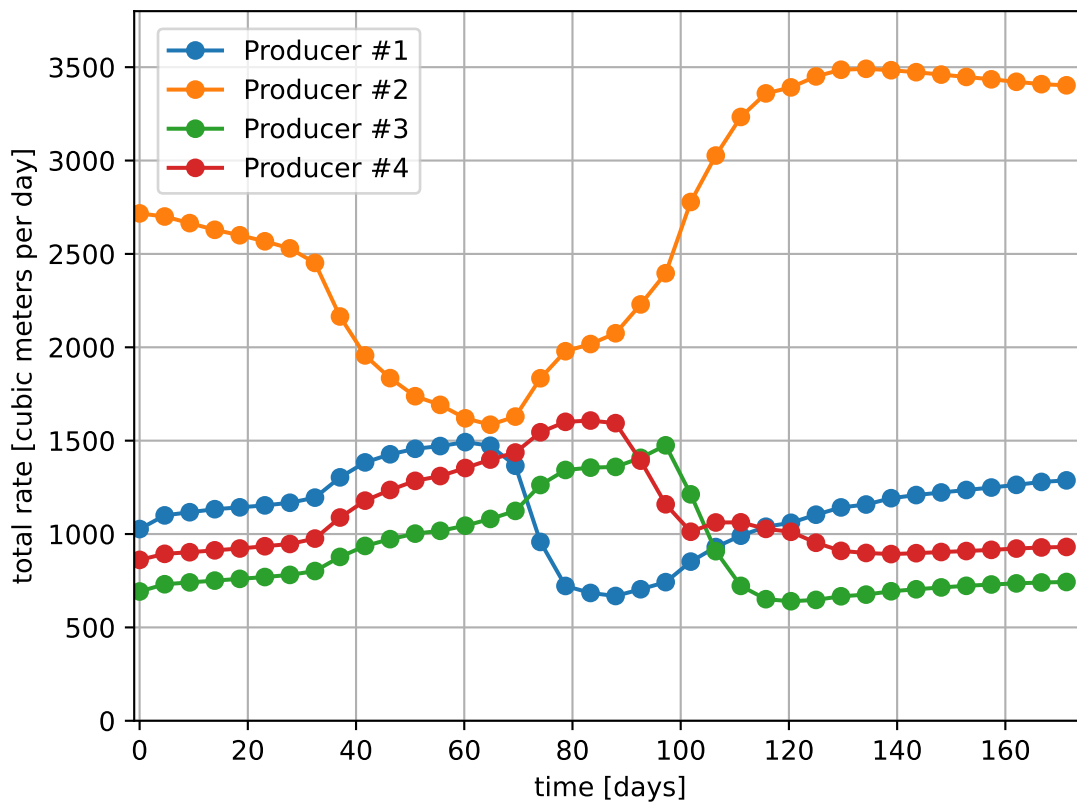
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```
( Rflow ) = ( 1.13e-04 ) ;      ( Rwell ) = ( 5.09e-05 ) ;      ( R ) = ( 1.24e-04
↪ ) ;
Last LinSolve(iter,res) = ( 50, 9.54e-03 ) ;
Attempt: 0, ConfigurationIter: 0, NewtonIter: 5
( Rflow ) = ( 2.17e-08 ) ;      ( Rwell ) = ( 1.15e-07 ) ;      ( R ) = ( 1.17e-07
↪ ) ;
Last LinSolve(iter,res) = ( 55, 2.71e-04 ) ;
coupledFlowAndWells: Newton solver converged in less than 15 iterations, time-step
↪ required will be doubled.
```

### 3.2.11 Visualization

A file compatible with Paraview is produced in this example. It is found in the output folder, and usually has the extension *.pvd*. More details about this file format can be found [here](#). We can load this file into Paraview directly and visualize results:

We have instructed GEOSX to output the time series of rates for each producer. The data contained in the corresponding hdf5 files can be extracted and plotted as shown below.



### 3.2.12 To go further

#### Feedback on this example

This concludes the example on setting up a Dead-Oil simulation in the Egg model. For any feedback on this example, please submit a [GitHub issue on the project's GitHub page](#).

#### For more details

- A complete description of the reservoir flow solver is found here: *Compositional Multiphase Flow Solver*.
- The well solver is description at *Compositional Multiphase Well Solver*.
- The available constitutive models are listed at *Constitutive Models*.

## 3.3 CO<sub>2</sub> Injection

### Context

In this example, we show how to set up a multiphase simulation of CO<sub>2</sub> injection.

### Objectives

At the end of this example you will know:

- how to set up a CO<sub>2</sub> injection scenario with a well,
- how to run a case using MPI-parallelism.

### Input file

The XML file for this test case is located at :

```
inputFiles/compositionalMultiphaseWell/simpleCo2InjTutorial_base.xml
```

```
inputFiles/compositionalMultiphaseWell/simpleCo2InjTutorial_smoke.xml
```

This mesh is a simple internally generated regular grid (50 x 1 x 150). A single CO<sub>2</sub> injection well is at the center of the reservoir.

The XML file considered here follows the typical structure of the GEOSX input files:

1. *Solver*
2. *Mesh*
3. *Events*
4. *NumericalMethods*
5. *ElementRegions*
6. *Constitutive*
7. *FieldSpecifications*
8. *Outputs*
9. *Tasks*

### 3.3.1 Multiphase flow and well solvers

Let us inspect the **Solver** XML tags. They consist of three blocks **CompositionalMultiphaseFVM**, **CompositionalMultiphaseWell** and **CompositionalMultiphaseReservoir**, which are respectively handling the solution from multiphase flow in the reservoir, multiphase flow in the wells, and coupling between those two parts.

```
<Solvers>
  <CompositionalMultiphaseReservoir
    name="coupledFlowAndWells"
    flowSolverName="compositionalMultiphaseFlow"
    wellSolverName="compositionalMultiphaseWell"
    logLevel="1"
    initialDt="1e2"
    targetRegions="{ reservoir, wellRegion }">
    <NonlinearSolverParameters
      newtonTol="1.0e-4"
      lineSearchAction="None"
      maxTimeStepCuts="10"
      newtonMaxIter="40"/>
    <LinearSolverParameters
      solverType="fgmres"
      preconditionerType="mgr"
      krylovTol="1e-5"/>
    </CompositionalMultiphaseReservoir>

    <CompositionalMultiphaseFVM
      name="compositionalMultiphaseFlow"
      targetRegions="{ reservoir }"
      discretization="fluidTPFA"
      temperature="368.15"
      maxCompFractionChange="0.2"
      logLevel="1"
      useMass="1"/>

    <CompositionalMultiphaseWell
      name="compositionalMultiphaseWell"
      targetRegions="{ wellRegion }"
      maxCompFractionChange="0.2"
      logLevel="1"
      useMass="1">
      <WellControls
        name="wellControls"
        type="injector"
        control="totalVolRate"
        enableCrossflow="0"
        referenceElevation="6650"
        useSurfaceConditions="1"
        surfacePressure="101325"
        surfaceTemperature="288.71"
        targetBHP="5e7"
        targetTotalRate="1.5"
        injectionTemperature="368.15"
        injectionStream="{ 1, 0 }"/>
      </CompositionalMultiphaseWell>
    </Solvers>
```

In the **CompositionalMultiphaseFVM** (*Compositional Multiphase Flow Solver*), a classical multiphase compositional solver with a TPFA discretization is described.

The **CompositionalMultiphaseWell** (*Compositional Multiphase Well Solver*) consists of wellbore specifications (see *Multiphase Flow with Wells* for detailed example). As its reservoir counterpart, it includes references to fluid and relative permeability models, but also defines a **WellControls** sub-tag. This sub-tag specifies the CO<sub>2</sub> injector control mode: the well is initially rate-controlled, with a rate specified in `targetTotalRate` and a maximum pressure specified in `targetBHP`. The injector-specific attribute, `injectionStream`, describes the composition of the injected mixture (here, pure CO<sub>2</sub>).

The **CompositionalMultiphaseReservoir** coupling section describes the binding between those two previous elements (see *Poromechanics* for detailed example on coupling physics in GEOSX). In addition to being bound to the previously described blocks through `flowSolverName` and `wellSolverName` sub-tags, it contains the `initialDt` starting time-step size value and defines the **NonlinearSolverParameters** and **LinearSolverParameters** that are used to control Newton-loop and linear solver behaviors (see *Linear Solvers* for a detailed description of linear solver attributes).

---

**Note:** To use the linear solver options of this example, you need to ensure that GEOSX is configured to use the Hypr linear solver package.

---

### 3.3.2 Mesh and well geometry

In this example, the **Mesh** tag is used to generate the reservoir mesh internally (*Tutorial 1: First Steps*). The internal generation of well is defined with the **InternalWell** sub-tag. Apart from the name identifier attribute and their `wellRegionName` (*ElementRegions*) and `wellControlsName` (*Solver*) binding attributes, `polylineNodeCoords` and `polylineSegmentConn` attributes are used to define the path of the wellbore and connections between its nodes. The `numElementsPerSegment` discretizes the wellbore segments while the `radius` attribute specifies the wellbore radius (*Multiphase Flow with Wells* for details on wells). Once the wellbore is defined and discretized, the position of **Perforations** is defined using the linear distance from the head of the wellbore (`distanceFromHead`).

```
<Mesh>
  <InternalMesh
    name="cartesianMesh"
    elementTypes="{ C3D8 }"
    xCoords="{ 0, 1000 }"
    yCoords="{ 450, 550 }"
    zCoords="{ 6500, 7700 }"
    nx="{ 50 }"
    ny="{ 1 }"
    nz="{ 150 }"
    cellBlockNames="{ cellBlock }"/>

  <InternalWell
    name="wellInjector1"
    wellRegionName="wellRegion"
    wellControlsName="wellControls"
    meshName="cartesianMesh"
    polylineNodeCoords="{ { 500.0, 500.0, 6650.00 },
                          { 500.0, 500.0, 6600.00 } }"
    polylineSegmentConn="{ { 0, 1 } }"
    radius="0.1"
    numElementsPerSegment="2">
    <Perforation
      name="injector1_perf1"
      distanceFromHead="45"/>
```

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```
</InternalWell>
</Mesh>
```

**Note:** It is the responsibility of the user to make sure that there is a perforation in the bottom cell of the well mesh, otherwise an error will be thrown and the simulation will terminate.

### 3.3.3 Events

The solver is applied as a periodic event whose target is referred to as `coupledFlowAndWells` nametag. Using the `maxEventDt` attribute, we specify a max time step size of  $5 \times 10^6$  seconds.

The output event triggers a VTK output every  $10^7$  seconds, constraining the solver schedule to match exactly these dates. The output path to data is specified as a `target` of this **PeriodicEvent**.

Another periodic event is defined under the name `restarts`. It consists of saved checkpoints every  $5 \times 10^7$  seconds, whose physical output folder name is defined under the **Output** tag.

Finally, the time history collection and output events are used to trigger the mechanisms involved in the generation of a time series of well pressure (see the procedure outlined in *Tasks Manager*, and the example in *Multiphase Flow with Wells*).

```
<Events
  maxTime="5e8">

  <PeriodicEvent
    name="outputs"
    timeFrequency="1e7"
    targetExactTimestep="1"
    target="/Outputs/simpleReservoirViz"/>

  <PeriodicEvent
    name="restarts"
    timeFrequency="5e7"
    targetExactTimestep="1"
    target="/Outputs/restartOutput"/>

  <PeriodicEvent
    name="timeHistoryCollection"
    timeFrequency="1e7"
    targetExactTimestep="1"
    target="/Tasks/wellPressureCollection" />

  <PeriodicEvent
    name="timeHistoryOutput"
    timeFrequency="2e8"
    targetExactTimestep="1"
    target="/Outputs/timeHistoryOutput" />

  <PeriodicEvent
    name="solverApplications"
    maxEventDt="5e5"
    target="/Solvers/coupledFlowAndWells"/>

</Events>
```



### 3.3.4 Numerical methods

The **TwoPointFluxApproximation** is chosen for the fluid equation discretization. The tag specifies:

- A primary field to solve for as `fieldName`. For a flow problem, this field is pressure.
- A set of target regions in `targetRegions`.
- A `coefficientName` pointing to the field used for TPFA transmissibilities construction.
- A `coefficientModelNames` used to specify the permeability constitutive model(s).

```
<NumericalMethods>
  <FiniteVolume>
    <TwoPointFluxApproximation
      name="fluidTPFA"/>
    </FiniteVolume>
  </NumericalMethods>
```

### 3.3.5 Element regions

We define a **CellElementRegion** pointing to the cell block defining the reservoir mesh, and a **WellElementRegion** for the well. The two regions contain a list of constitutive model names.

```
<ElementRegions>
  <CellElementRegion
    name="reservoir"
    cellBlocks="{ cellBlock }"
    materialList="{ fluid, rock, relperm }"/>

  <WellElementRegion
    name="wellRegion"
    materialList="{ fluid, relperm, rockPerm }"/>
</ElementRegions>
```

### 3.3.6 Constitutive laws

Under the **Constitutive** tag, four items can be found:

- **CO2BrinePhillipsFluid** : this tag defines phase names, component molar weights, and fluid behaviors such as CO<sub>2</sub> solubility in brine and viscosity/density dependencies on pressure and temperature.
- **PressurePorosity** : this tag contains all the data needed to model rock compressibility.
- **BrooksCoreyRelativePermeability** : this tag defines the relative permeability model for each phase, its end-point values, residual volume fractions (saturations), and the Corey exponents.
- **ConstantPermeability** : this tag defines the permeability model that is set to a simple constant diagonal tensor, whose values are defined in `permeabilityComponent`. Note that these values will be overwritten by the permeability field imported in **FieldSpecifications**.

```
<Constitutive>
  <CO2BrinePhillipsFluid
    name="fluid"
    phaseNames="{ gas, water }"
    componentNames="{ co2, water }"
    componentMolarWeight="{ 44e-3, 18e-3 }"
```

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```

phasePVTParaFiles="{ pvtgas.txt, pvtliquid.txt }"
flashModelParaFile="co2flash.txt"/>

<CompressibleSolidConstantPermeability
  name="rock"
  solidModelName="nullSolid"
  porosityModelName="rockPorosity"
  permeabilityModelName="rockPerm"/>

<NullModel
  name="nullSolid"/>

<PressurePorosity
  name="rockPorosity"
  defaultReferencePorosity="0.1"
  referencePressure="1.0e7"
  compressibility="4.5e-10"/>

<BrooksCoreyRelativePermeability
  name="relperm"
  phaseNames="{ gas, water }"
  phaseMinVolumeFraction="{ 0.05, 0.30 }"
  phaseRelPermExponent="{ 2.0, 2.0 }"
  phaseRelPermMaxValue="{ 1.0, 1.0 }"/>

<ConstantPermeability
  name="rockPerm"
  permeabilityComponents="{ 1.0e-17, 1.0e-17, 3.0e-17 }"/>

</Constitutive>

```

The PVT data specified by **CO2BrinePhillipsFluid** is set to model the behavior of the CO<sub>2</sub>-brine system as a function of pressure, temperature, and salinity. We currently rely on a two-phase, two-component (CO<sub>2</sub> and H<sub>2</sub>O) model in which salinity is a constant parameter in space and in time. The model is described in detail in *CO<sub>2</sub>-brine model*. The model definition requires three text files:

In *co2flash.txt*, we define the CO<sub>2</sub> solubility model used to compute the amount of CO<sub>2</sub> dissolved in the brine phase as a function of pressure (in Pascal), temperature (in Kelvin), and salinity (in units of molality):

```
FlashModel CO2Solubility 1e6 1.5e7 5e4 367.15 369.15 1 0
```

The first keyword is an identifier for the model type (here, a flash model). It is followed by the model name. Then, the lower, upper, and step increment values for pressure and temperature ranges are specified. The trailing 0 defines a zero-salinity in the model. Note that the water component is not allowed to evaporate into the CO<sub>2</sub>-rich phase.

The *pvtgas.txt* and *pvtliquid.txt* files define the models used to compute the density and viscosity of the two phases, as follows:

```
DensityFun SpanWagnerCO2Density 1e6 1.5e7 5e4 94 96 1
ViscosityFun FenghourCO2Viscosity 1e6 1.5e7 5e4 94 96
```

```
DensityFun PhillipsBrineDensity 1e6 1.5e7 5e4 94 96 1 0
ViscosityFun PhillipsBrineViscosity 0
```

In these files, the first keyword of each line is an identifier for the model type (either a density or a viscosity model). It is followed by the model name. Then, the lower, upper, and step increment values for pressure and temperature

ranges are specified. The trailing 0 for `PhillipsBrineDensity` and `PhillipsBrineViscosity` entry is the salinity of the brine, set to zero.

**Note:** It is the responsibility of the user to make sure that the pressure and temperature values encountered in the simulation (in the reservoir and in the well) are within the bounds specified in the PVT files. GEOSX will not throw an error if a value outside these bounds is encountered, but the (nonlinear) behavior of the simulation and the quality of the results will likely be negatively impacted.

### 3.3.7 Property specification

The **FieldSpecifications** tag is used to declare fields such as directional permeability, reference porosity, initial pressure, and compositions. Here, these fields are homogeneous, except for the permeability field that is taken as an heterogeneous log-normally distributed field and specified in **Functions** as in *Tutorial 3: Regions and Property Specifications*.

```
<FieldSpecifications>
  <FieldSpecification
    name="permx"
    initialCondition="1"
    component="0"
    setNames="{ all }"
    objectPath="ElementRegions"
    fieldName="rockPerm_permeability"
    scale="1e-15"
    functionName="permxFunc"/>

  <FieldSpecification
    name="permy"
    initialCondition="1"
    component="1"
    setNames="{ all }"
    objectPath="ElementRegions"
    fieldName="rockPerm_permeability"
    scale="1e-15"
    functionName="permyFunc"/>

  <FieldSpecification
    name="permz"
    initialCondition="1"
    component="2"
    setNames="{ all }"
    objectPath="ElementRegions"
    fieldName="rockPerm_permeability"
    scale="3e-15"
    functionName="permzFunc"/>

  <FieldSpecification
    name="initialPressure"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions/reservoir"
    fieldName="pressure"
    scale="1.25e7"/>
```

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```

<FieldSpecification
  name="initialComposition_co2"
  initialCondition="1"
  setNames="{ all }"
  objectPath="ElementRegions/reservoir"
  fieldName="globalCompFraction"
  component="0"
  scale="0.0"/>

<FieldSpecification
  name="initialComposition_water"
  initialCondition="1"
  setNames="{ all }"
  objectPath="ElementRegions/reservoir"
  fieldName="globalCompFraction"
  component="1"
  scale="1.0"/>
</FieldSpecifications>

```

**Note:** In this case, we are using the same permeability field (*perm.geos*) for all the directions. Note also that the `fieldName` values are set to *rockPerm\_permeability* to access the permeability field handled as a **Constitutive** law. These permeability values will overwrite the values already set in the **Constitutive** block.

### 3.3.8 Output

The **Outputs** XML tag is used to write visualization, restart, and time history files.

Here, we write visualization files in a format natively readable by Paraview under the tag **VTK**. A **Restart** tag is also be specified. In conjunction with a **PeriodicEvent**, a restart file allows to resume computations from a set of checkpoints in time. Finally, we require an output of the well pressure history using the **TimeHistory** tag.

```

<Outputs>
  <VTK
    name="simpleReservoirViz"/>

  <Restart
    name="restartOutput"/>

  <TimeHistory
    name="timeHistoryOutput"
    sources="{/Tasks/wellPressureCollection}"
    filename="wellPressureHistory" />

</Outputs>

```

### 3.3.9 Tasks

In the **Events** block, we have defined an event requesting that a task periodically collects the pressure at the well. This task is defined here, in the **PackCollection** XML sub-block of the **Tasks** block. The task contains the path to the object on which the field to collect is registered (here, a *WellElementSubRegion*) and the name of the field (here, *pressure*). The details of the history collection mechanism can be found in *Tasks Manager*.

```

<Tasks>
  <PackCollection
    name="wellPressureCollection"
    objectPath="ElementRegions/wellRegion/wellRegionUniqueSubRegion"
    fieldName="pressure" />
  </Tasks>

```

### 3.3.10 Running GEOSX

The simulation can be launched with 4 cores using MPI-parallelism:

```
mpirun -np 4 geosx -i SimpleCo2InjTutorial.xml -x 1 -y 1 -z 4
```

A restart from a checkpoint file *SimpleCo2InjTutorial\_restart\_000000024.root* is always available thanks to the following command line :

```
mpirun -np 4 geosx -i SimpleCo2InjTutorial.xml -r SimpleCo2InjTutorial_restart_
↪000000024 -x 1 -y 1 -z 4
```

The output then shows the loading of HDF5 restart files by each core.

```

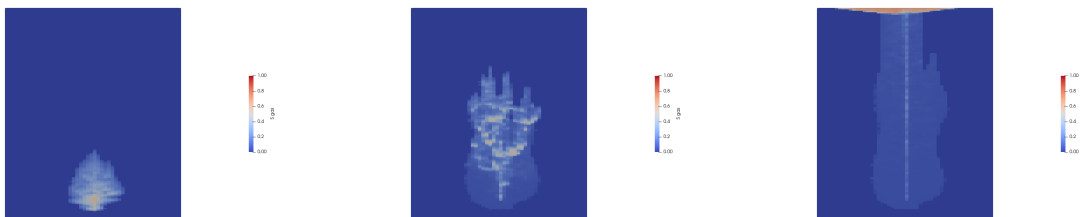
Loading restart file SimpleCo2InjTutorial_restart_000000024
Rank 0: rankFilePattern = SimpleCo2InjTutorial_restart_000000024/rank_%07d.hdf5
Rank 0: Reading in restart file at SimpleCo2InjTutorial_restart_000000024/rank_
↪00000000.hdf5
Rank 1: Reading in restart file at SimpleCo2InjTutorial_restart_000000024/rank_
↪00000001.hdf5
Rank 3: Reading in restart file at SimpleCo2InjTutorial_restart_000000024/rank_
↪00000003.hdf5
Rank 2: Reading in restart file at SimpleCo2InjTutorial_restart_000000024/rank_
↪00000002.hdf5

```

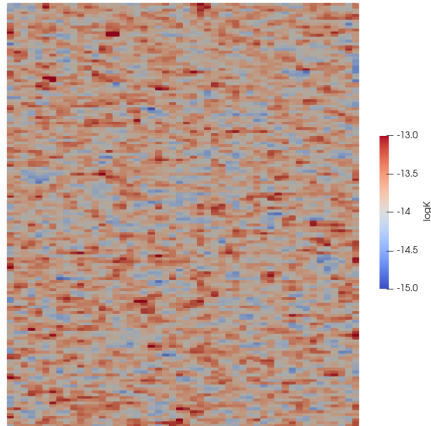
and the simulation restarts from this point in time.

### 3.3.11 Visualization

Using Paraview, we can observe the CO<sub>2</sub> plume moving upward under buoyancy effects and forming a gas cap at the top of the domain,



The heterogeneous values of the log permeability field can also be visualized in Paraview as shown below:



### 3.3.12 To go further

#### Feedback on this example

This concludes the CO<sub>2</sub> injection field case example. For any feedback on this example, please submit a [GitHub issue](#) on the project's [GitHub page](#).

#### For more details

- A complete description of the reservoir flow solver is found here: *Compositional Multiphase Flow Solver*.
- The well solver is described at *Compositional Multiphase Well Solver*.
- The available fluid constitutive models are listed at *Fluid Models*.

## 3.4 Poromechanics

### Context

In this example, we use a coupled solver to solve a poroelastic Terzaghi-type problem, a classic benchmark in poroelasticity. We do so by coupling a single phase flow solver with a small-strain Lagrangian mechanics solver.

### Objectives

At the end of this example you will know:

- how to use multiple solvers for poromechanical problems,
- how to define finite elements and finite volume numerical methods.

### Input file

This example uses no external input files and everything required is contained within two GEOSX input files located at:

```
inputFiles/poromechanics/PoroElastic_Terzaghi_base_direct.xml
```

```
inputFiles/poromechanics/PoroElastic_Terzaghi_smoke.xml
```

### 3.4.1 Description of the case

We simulate the consolidation of a poroelastic fluid-saturated column of height  $L$  having unit cross-section. The column is instantaneously loaded at time  $t = 0$  s with a constant compressive traction  $w$  applied on the face highlighted in red in the figure below. Only the loaded face is permeable to fluid flow, with the remaining parts of the boundary subject to roller constraints and impervious.

Fig. 3.1: Sketch of the setup for Terzaghi's problem.

GEOSX will calculate displacement and pressure fields along the column as a function of time. We will use the analytical solution for pressure to check the accuracy of the solution obtained with GEOSX, namely

$$p(x, t) = \frac{4}{\pi} p_0 \sum_{m=0}^{\infty} \frac{1}{2m+1} \exp \left[ -\frac{(2m+1)^2 \pi^2 c_c t}{4L^2} \right] \sin \left[ \frac{(2m+1)\pi x}{2L} \right],$$

where  $p_0 = \frac{b}{K_v S_e + b^2} |w|$  is the initial pressure, constant throughout the column, and  $c_c = \frac{\kappa}{\mu} \frac{K_v}{K_v S_e + b^2}$  is the consolidation coefficient (or diffusion coefficient), with

- $b$  Biot's coefficient
- $K_v = \frac{E(1-\nu)}{(1+\nu)(1-2\nu)}$  the uniaxial bulk modulus,  $E$  Young's modulus, and  $\nu$  Poisson's ratio
- $S_e = \frac{(b-\phi)(1-b)}{K} + \phi c_f$  the constrained specific storage coefficient,  $\phi$  porosity,  $K = \frac{E}{3(1-2\nu)}$  the bulk modulus, and  $c_f$  the fluid compressibility
- $\kappa$  the isotropic permeability
- $\mu$  the fluid viscosity

The characteristic consolidation time of the system is defined as  $t_c = \frac{L^2}{c_c}$ . Knowledge of  $t_c$  is useful for choosing appropriately the timestep sizes that are used in the discrete model.

### 3.4.2 Coupled solvers

GEOSX is a multi-physics tool. Different combinations of physics solvers available in the code can be applied in different regions of the mesh at different moments of the simulation. The `XML Solvers` tag is used to list and parameterize these solvers.

We define and characterize each single-physics solver separately. Then, we define a *coupling solver* between these single-physics solvers as another, separate, solver. This approach allows for generality and flexibility in our multi-physics resolutions. The order in which these solver specifications is done is not important. It is important, though, to instantiate each single-physics solver with meaningful names. The names given to these single-physics solver instances will be used to recognize them and create the coupling.

To define a poromechanical coupling, we will effectively define three solvers:

- the single-physics flow solver, a solver of type `SinglePhaseFVM` called here `SinglePhaseFlowSolver` (more information on these solvers at [Singlephase Flow Solver](#)),
- the small-stress Lagrangian mechanics solver, a solver of type `SolidMechanicsLagrangianSSLE` called here `LinearElasticitySolver` (more information here: [Solid Mechanics Solver](#)),
- the coupling solver that will bind the two single-physics solvers above, an object of type `SinglePhasePoromechanics` called here `PoroelasticitySolver` (more information at [Poromechanics Solver](#)).

Note that the name attribute of these solvers is chosen by the user and is not imposed by GEOSX.

The two single-physics solvers are parameterized as explained in their respective documentation.

Let us focus our attention on the coupling solver. This solver (`PoroelasticitySolver`) uses a set of attributes that specifically describe the coupling for a poromechanical framework. For instance, we must point this solver to the correct fluid solver (here: `SinglePhaseFlowSolver`), the correct solid solver (here: `LinearElasticitySolver`). Now that these two solvers are tied together inside the coupling solver, we have a coupled multiphysics problem defined. More parameters are required to characterize a coupling. Here, we specify the discretization method (`FE1`, defined further in the input file), and the target regions (here, we only have one, `Domain`).

```
<SinglePhasePoromechanics
  name="PoroelasticitySolver"
  solidSolverName="LinearElasticitySolver"
  flowSolverName="SinglePhaseFlowSolver"
  logLevel="1"
  targetRegions="{ Domain }">
  <LinearSolverParameters
    directParallel="0"/>
</SinglePhasePoromechanics>

<SolidMechanicsLagrangianSSLE
  name="LinearElasticitySolver"
  timeIntegrationOption="QuasiStatic"
  logLevel="1"
  discretization="FE1"
  targetRegions="{ Domain }"/>

<SinglePhaseFVM
  name="SinglePhaseFlowSolver"
  logLevel="1"
  discretization="singlePhaseTPFA"
  targetRegions="{ Domain }"/>
</Solvers>
```

### 3.4.3 Multiphysics numerical methods

Numerical methods in multiphysics settings are similar to single physics numerical methods. All can be defined under the same `NumericalMethods` XML tag. In this problem, we use finite volume for flow and finite elements for solid mechanics. Both methods require additional parameterization attributes to be defined here.

As we have seen before, the coupling solver and the solid mechanics solver require the specification of a discretization method called `FE1`. This discretization method is defined here as a finite element method using linear basis functions and Gaussian quadrature rules. For more information on defining finite elements numerical schemes, please see the dedicated *FiniteElement* section.

The finite volume method requires the specification of a discretization scheme. Here, we use a two-point flux approximation as described in the dedicated documentation (found here: *FiniteVolume*).

```
<NumericalMethods>
  <FiniteElements>
    <FiniteElementSpace
      name="FE1"
      order="1"/>
    </FiniteElements>

    <FiniteVolume>
```

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```

<TwoPointFluxApproximation
  name="singlePhaseTPFA"/>
</FiniteVolume>
</NumericalMethods>

```

### 3.4.4 Mesh, material properties, and boundary conditions

Last, let us take a closer look at the geometry of this simple problem. We use the internal mesh generator to create a beam-like mesh, with one single element along the Y and Z axes, and 21 elements along the X axis. All the elements are hexahedral elements (C3D8) of the same dimension (1x1x1 meters).

```

<Mesh>
  <InternalMesh
    name="mesh1"
    elementTypes="{ C3D8 }"
    xCoords="{ 0, 10 }"
    yCoords="{ 0, 1 }"
    zCoords="{ 0, 1 }"
    nx="{ 400 }"
    ny="{ 16 }"
    nz="{ 16 }"
    cellBlockNames="{ cb1 }"/>
  </InternalMesh>
</Mesh>

```

The parameters used in the simulation are summarized in the following table.

Symbol	Parameter	Units	Value
$E$	Young's modulus	[Pa]	$10^4$
$\nu$	Poisson's ratio	[-]	0.2
$b$	Biot's coefficient	[-]	1.0
$\phi$	Porosity	[-]	0.3
$\rho_f$	Fluid density	[kg/m <sup>3</sup> ]	1.0
$c_f$	Fluid compressibility	[Pa <sup>-1</sup> ]	0.0
$\kappa$	Permeability	[m <sup>2</sup> ]	$10^{-4}$
$\mu$	Fluid viscosity	[Pa s]	1.0
$ w $	Applied compression	[Pa]	1.0
$L$	Column length	[m]	10.0

Material properties and boundary conditions are specified in the `Constitutive` and `FieldSpecifications` sections. For such set of parameters we have  $p_0 = 1.0$  Pa,  $c_c = 1.111 \text{ m}^2 \text{ s}^{-1}$ , and  $t_c = 90$  s. Therefore, as shown in the `Events` section, we run this simulation for 90 seconds.

### 3.4.5 Running GEOSX

To run the case, use the following command:

```
path/to/geosx -i inputFiles/poromechanics/PoroElastic_Terzaghi_smoke.xml
```

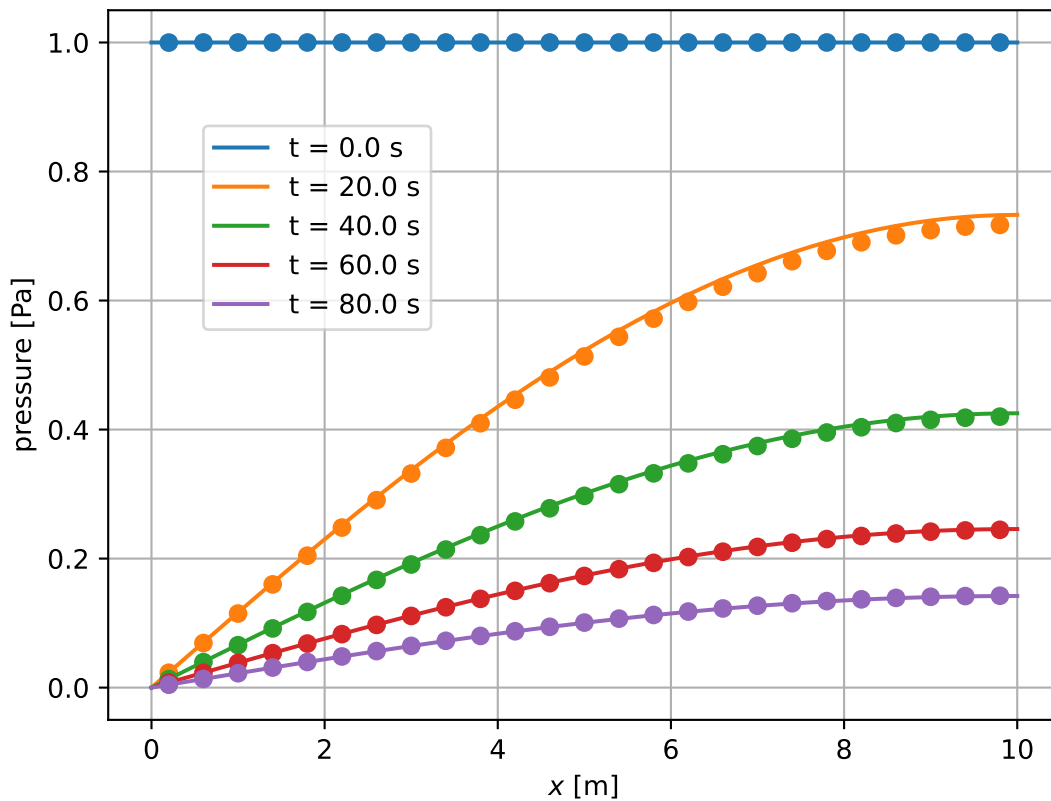
Here, we see for instance the `RSolid` and `RFluid` at a representative timestep (residual values for solid and fluid mechanics solvers, respectively)

```
Attempt: 0, NewtonIter: 0
( RSolid ) = ( 5.00e-01 ) ;      ( Rsolid, Rfluid ) = ( 5.00e-01, 0.00e+00 )
( R ) = ( 5.00e-01 ) ;
Attempt: 0, NewtonIter: 1
( RSolid ) = ( 4.26e-16 ) ;      ( Rsolid, Rfluid ) = ( 4.26e-16, 4.22e-17 )
( R ) = ( 4.28e-16 ) ;
```

As expected, since we are dealing with a linear problem, the fully implicit solver converges in a single iteration.

### 3.4.6 Inspecting results

This plot compares the analytical pressure solution (continuous lines) at selected times with the numerical solution (markers).



### 3.4.7 To go further

#### Feedback on this example

This concludes the poroelastic example. For any feedback on this example, please submit a [GitHub issue](#) on the project's GitHub page.

#### For more details

- More on poroelastic multiphysics solvers, please see [Poromechanics Solver](#).
- More on numerical methods, please see [Numerical Methods](#).
- More on functions, please see [Functions](#).

## 3.5 Hydraulic Fracturing

### Context

In this example, we use a fully coupled hydrofracture solver from GEOSX to solve for the propagation of a single fracture within a reservoir with heterogeneous in-situ properties. Advanced xml features will be used throughout the example.

### Objectives

At the end of this example you will know:

- how to use multiple solvers for hydraulic fracturing problems,
- how to specify pre-existing fractures and where new fractures can develop,
- how to construct a mesh with bias,
- how to specify heterogeneous in-situ properties and initial conditions,
- how to use parameters, symbolic math, and units in xml files.

### Input files

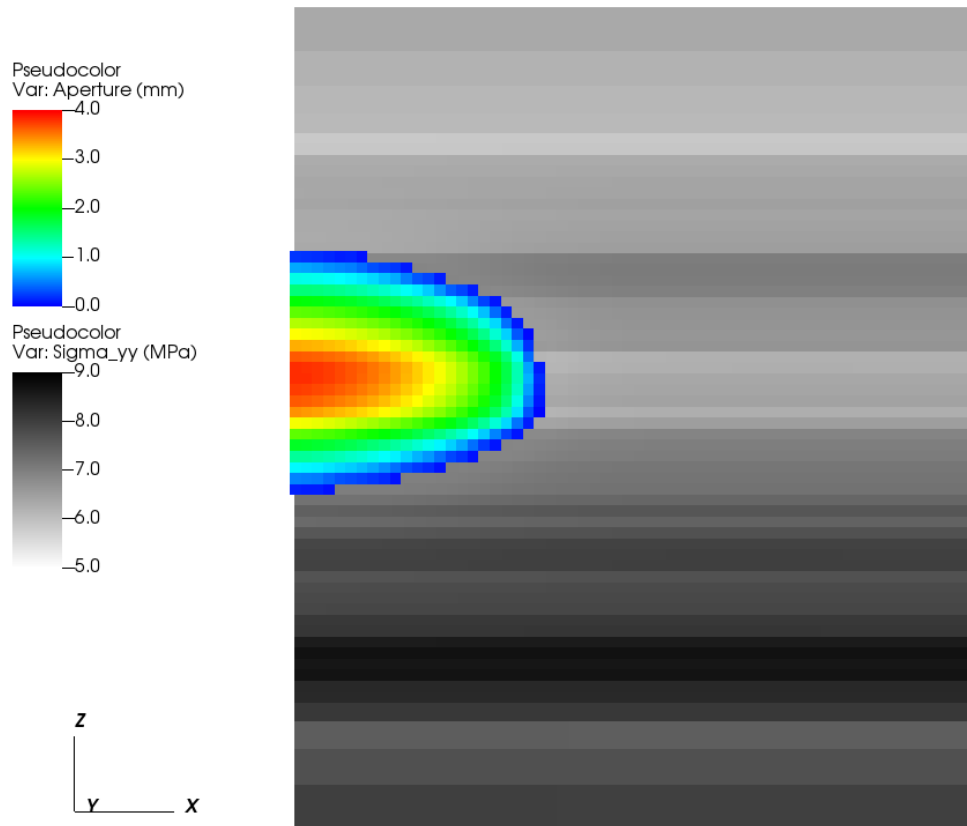
This example uses a set of input files and table files located at:

```
inputFiles/hydraulicFracturing
```

Because the input files use the advanced xml features, they must be preprocessed using the `geosx_xml_tools` package. If you have not already done so, setup these features by following the instructions here: [Advanced XML Features](#).

### 3.5.1 Description of the case

Here, our goal is to demonstrate how hydraulic fractures are modeled in a typical environment. The in-situ properties and initial conditions are based upon a randomly generated, fractal, 1D layer-cake model.



The inputs for this case are contained inside a case-specific (`heterogeneousInSitu_benchmark.xml`) and base (`heterogeneousInSitu_base.xml`) XML files. The `tables` directory contains the pre-constructed geologic model. This example will first focus on the case-specific input file, which contains the key parameter definitions, then consider the base xml file.

### 3.5.2 Included: including external xml files

At the head of the case-specific xml file is a block that will instruct GEOSX to include an external file. In our case, this points to the base hydraulic fracturing input file.

```
<Included>
  <File
    name="./heterogeneousInSitu_base.xml"/>
  </Included>
```

### 3.5.3 Parameters: defining variables to be used throughout the file

The `Parameters` block defines a series of variables that can be used throughout the input file. These variables allow a given input file to be easily understood and/or modified for a specific environment, even by non-expert users. Parameters are specified in pairs of names and values. The names should only contain alphanumeric characters and underlines. The values can contain any type (strings, doubles, etc.).

Parameters can be used throughout the input file (or an included input file) by placing them in-between dollar signs. Barring any circular-definition errors, parameters can be used within other parameters. For example, see the parameter `mu_upscaled`. The value of this parameter is a symbolic expression, which is denoted by the surrounding back-ticks, and is dependent upon two other parameters. During pre-processing, `geosx_xml_tools` will substitute the parameter definitions, and evaluate the symbolic expression using a python-derived syntax.

A number of the input parameters include optional unit definitions, which are denoted by the square brackets following a value. For example, the parameter `t_max` is used to set the maximum time for the simulation to 20 minutes.

```
<Parameters>
  <!-- Use the swarm upscaling law -->
  <Parameter
    name="Nperf"
    value="5"/>

  <Parameter
    name="Nswarm"
    value="5"/>

  <Parameter
    name="mu_init"
    value="0.001"/>

  <Parameter
    name="K_init"
    value="1e6"/>

  <Parameter
    name="mu_upscaled"
    value="`$mu_init$*($Nswarm$**2)`"/>

  <Parameter
    name="K_upscaled"
    value="`$K_init$*($Nswarm$**0.5)`"/>

  <Parameter
    name="ContactStiffness"
    value="1e10"/>

  <!-- Event timing -->
  <Parameter
    name="pump_start"
    value="1 [min]"/>

  <Parameter
    name="pump_ramp"
    value="5 [s]"/>

  <Parameter
    name="pump_ramp_dt_limit"
    value="0.2 [s]"/>

  <Parameter
    name="dt_max"
    value="30 [s]"/>

  <Parameter
    name="t_max"
```

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```

    value="20 [min]"/>

    <!-- Etc. -->
    <Parameter
      name="table_root"
      value="./tables"/>

    <Parameter
      name="t_allocation"
      value="28 [min]"/>
  </Parameters>

```

### 3.5.4 Mesh with biased boundaries

The mesh block for this example uses a biased mesh along the simulation boundaries to reduce the size of the problem, while maintaining the desired spatial extents. For a given element region with bias, the left-hand element will be  $x\%$  smaller and the right-hand element will be  $x\%$  larger than the average element size. Along the  $x$ -axis of the mesh, we select a value of zero for the first region to indicate that we want a uniform-sized mesh, and we select a bias of  $-0.6$  for the second region to indicate that we want the element size to smoothly increase in size as it moves in the  $+x$  direction. The other dimensions of the mesh follow a similar pattern.

```

<Mesh>
  <InternalMesh
    name="mesh1"
    xCoords="{ 0, 200, 250 }"
    yCoords="{ -100, 0, 100 }"
    zCoords="{ -150, -100, 0, 100, 150 }"
    nx="{ 50, 5 }"
    ny="{ 10, 10 }"
    nz="{ 5, 25, 25, 5 }"
    xBias="{ 0, -0.6 }"
    yBias="{ 0.6, -0.6 }"
    zBias="{ 0.6, 0, 0, -0.6 }"
    cellBlockNames="{ cb1 }"
    elementTypes="{ C3D8 }"/>
  </InternalMesh>
</Mesh>

```

### 3.5.5 Defining a fracture nodeset

For this example, we want to propagate a single hydraulic fracture along the plane defined by  $y = 0$ . To achieve this, we need to define three nodesets:

- `source_a`: The location where we want to inject fluid. Typically, we want this to be a single face in the  $x$ - $z$  plane.
- `perf_a`: This is the initial fracture for the simulation. This nodeset needs to be at least two-faces wide in the  $x$ - $z$  plane (to represent the fracture at least one internal node needs to be open).
- `fracturable_a`: This is the set of faces where we will allow the fracture to grow. For a problem where we expect the fracture to curve out of the plane defined by  $y = 0$ , this could be replaced.

```

<Geometry>
  <Box
    name="source_a"
    xMin="{ -0.1, -0.1, -0.1 }"

```

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```

xMax="{ 4.1, 0.1, 4.1 }"/>

<Box
  name="perf_a"
  xMin="{ -4.1, -0.1, -4.1 }"
  xMax="{ 4.1, 0.1, 4.1 }"/>

<ThickPlane
  name="fracturable_a"
  normal="{ 0, 1, 0 }"
  origin="{ 0, 0, 0 }"
  thickness="0.1"/>
</Geometry>

```

### 3.5.6 Boundary conditions

The boundary conditions for this problem are defined in the case-specific and the base xml files. The case specific block includes four instructions:

- `frac`: this marks the initial perforation.
- `separableFace`: this marks the set of faces that are allowed to break during the simulation.
- `waterDensity`: this initializes the fluid in the perforation.
- `sourceTerm`: this instructs the code to inject fluid into the `source_a` nodeset. Note the usage of the symbolic expression and parameters in the scale. This boundary condition is also driven by a function, which we will define later.

```

<FieldSpecifications>
  <!-- Fracture-related nodesets -->
  <FieldSpecification
    name="frac"
    fieldName="ruptureState"
    initialCondition="1"
    objectPath="faceManager"
    scale="1"
    setNames="{ perf_a }"/>

  <FieldSpecification
    name="separableFace"
    fieldName="isFaceSeparable"
    initialCondition="1"
    objectPath="faceManager"
    scale="1"
    setNames="{ fracturable_a }"/>

  <!-- Fluid Initial Conditions -->
  <FieldSpecification
    name="waterDensity"
    fieldName="water_density"
    initialCondition="1"
    objectPath="ElementRegions"
    scale="1000"
    setNames="{ perf_a }"/>

```

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```

<!-- Fluid BC's -->
<!-- Note: the units of the source flux BC are in kg/s not m3/s -->
<SourceFlux
  name="sourceTerm"
  objectPath="ElementRegions/Fracture"
  scale="-1000.0/$Nperf$"
  functionName="flow_rate"
  setNames="{ source_a }"/>
</FieldSpecifications>

```

The base block includes instructions to set the initial in-situ properties and stresses. It is also used to specify the external mechanical boundaries on the system. In this example, we are using roller-boundary conditions (zero normal-displacement). Depending upon how close they are to the fracture, they can significantly affect its growth. Therefore, it is important to test whether the size of the model is large enough to avoid this.

```

<FieldSpecifications>
  <FieldSpecification
    name="bulk_modulus"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions"
    fieldName="rock_bulkModulus"
    functionName="bulk_modulus"
    scale="1.0"/>

  <FieldSpecification
    name="shear_modulus"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions"
    fieldName="rock_shearModulus"
    functionName="shear_modulus"
    scale="1.0"/>

  <FieldSpecification
    name="sigma_xx"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions"
    fieldName="rock_stress"
    component="0"
    functionName="sigma_xx"
    scale="1.0"/>

  <FieldSpecification
    name="sigma_yy"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions"
    fieldName="rock_stress"
    component="1"
    functionName="sigma_yy"
    scale="1.0"/>

  <FieldSpecification
    name="sigma_zz"

```

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```

        initialCondition="1"
        setNames="{ all }"
        objectPath="ElementRegions"
        fieldName="rock_stress"
        component="2"
        functionName="sigma_zz"
        scale="1.0"/>

<!-- Mechanical BC's -->
<FieldSpecification
  name="x_constraint"
  component="0"
  fieldName="totalDisplacement"
  objectPath="nodeManager"
  scale="0.0"
  setNames="{ xneg, xpos }"/>

<FieldSpecification
  name="y_constraint"
  component="1"
  fieldName="totalDisplacement"
  objectPath="nodeManager"
  scale="0.0"
  setNames="{ yneg, ypos }"/>

<FieldSpecification
  name="z_constraint"
  component="2"
  fieldName="totalDisplacement"
  objectPath="nodeManager"
  scale="0.0"
  setNames="{ zneg, zpos }"/>
</FieldSpecifications>

```

### 3.5.7 Coupled hydraulic fracturing solver

The Solvers block is located in the base xml file. Note that the `gravityVector` attribute indicates that we are applying gravity in the z-direction in this problem.

Similar to other coupled physics solvers, the Hydrofracture solver is specified in three parts:

- Hydrofracture: this is the primary solver, which will be called by the event manager. Two of its key attributes are the names of the dependent solid and fluid solvers.
- SolidMechanicsLagrangianSSLE: this is the solid mechanics solver.
- SinglePhaseFVM: this is the fluid solver.

The final solver present in this example is the SurfaceGenerator, which manages how faces in the model break.

```

<Solvers
  gravityVector="{ 0.0, 0.0, -9.81 }">
  <Hydrofracture
    name="hydrofracture"
    solidSolverName="lagsolve"
    flowSolverName="SinglePhaseFlow"
    surfaceGeneratorName="SurfaceGen"

```

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```

couplingTypeOption="FIM"
logLevel="1"
targetRegions="{ Fracture }"
contactRelationName="fractureContact"
maxNumResolves="2">
  <NonlinearSolverParameters
    newtonTol="1.0e-6"
    newtonMaxIter="40"
    lineSearchMaxCuts="3"/>
  <LinearSolverParameters
    solverType="gmres"
    preconditionerType="amg"/>
</Hydrofracture>

<SolidMechanicsLagrangianSSLE
  name="lagsolve"
  timeIntegrationOption="QuasiStatic"
  discretization="FE1"
  targetRegions="{ Domain, Fracture }"
  contactRelationName="fractureContact">
  <NonlinearSolverParameters
    newtonTol="1.0e-6"/>
  <LinearSolverParameters
    solverType="gmres"
    krylovTol="1.0e-10"/>
</SolidMechanicsLagrangianSSLE>

<SinglePhaseFVM
  name="SinglePhaseFlow"
  discretization="singlePhaseTPFA"
  targetRegions="{ Fracture }">
  <NonlinearSolverParameters
    newtonTol="1.0e-5"
    newtonMaxIter="10"/>
  <LinearSolverParameters
    solverType="gmres"
    krylovTol="1.0e-12"/>
</SinglePhaseFVM>

<SurfaceGenerator
  name="SurfaceGen"
  targetRegions="{ Domain }"
  rockToughness="$K_upscaled$"
  nodeBasedSIF="1"
  mpiCommOrder="1"/>
</Solvers>

```

### 3.5.8 Events

Rather than explicitly specify the desired timestep behavior, this example uses a flexible approach for timestepping. The hydrofracture solver is applied in the `solverApplications` event, which request a `maxEventDt = 30` s. To maintain stability during the critical early phase of the model, we delay turning on the pump by `pump_start`. We then use the `pumpStart` event to limit the further limit the timestep to `pump_ramp_dt_limit` as the fracture experiences rapid development (`pump_start` to `pump_start + pump_ramp`). Note that while this event does not have a target, it can still influence the time step behavior. After this period, the hydraulic fracture solver will

attempt to increase / decrease the requested timestep to maintain stability.

Other key events in this problem include:

- `preFracture`: this calls the surface generator at the beginning of the problem and helps to initialize the fracture.
- `outputs_vtk` and `outputs_silo`: these produces output vtk and silo files.
- `restarts` (inactive): this is a `HaltEvent`, which tracks the external clock. When the runtime exceeds the specified value (here `$t_allocation$=28` minutes), the code will call the target (which writes a restart file) and instruct the code to exit.

```
<Events
  maxTime="$t_max$"
  logLevel="1">
  <!-- Generate the initial fractures -->
  <SoloEvent
    name="preFracture"
    target="/Solvers/SurfaceGen"/>

  <!-- Primary outputs -->
  <PeriodicEvent
    name="outputs_vtk"
    timeFrequency="1 [min]"
    targetExactTimestep="0"
    target="/Outputs/vtkOutput"/>

  <PeriodicEvent
    name="outputs_silo"
    timeFrequency="1 [min]"
    targetExactTimestep="0"
    target="/Outputs/siloOutput"/>

  <!-- Apply the hydrofracture solver -->
  <PeriodicEvent
    name="solverApplications"
    maxEventDt="$dt_max$"
    target="/Solvers/hydrofracture"/>

  <!-- Limit dt during the pump ramp-up -->
  <PeriodicEvent
    name="pumpStart"
    beginTime="$pump_start$"
    endTime="`$pump_start$+$pump_ramp$`"
    maxEventDt="$pump_ramp_dt_limit$"/>

  <!-- Watch the wall-clock, write a restart, and exit gracefully if necessary -->
  <!-- <HaltEvent
    name="restarts"
    maxRuntime="$t_allocation$"
    target="/Outputs/restartOutput"/> -->
</Events>
```

### 3.5.9 Functions to set in-situ properties

The function definitions are in the base xml file, and rely upon the files in the tables directory. The functions in this example include the flow rate over time, the in-situ principal stress, and the bulk/shear moduli of the rock. Note the use of the `table_root` parameter, which contains the root path to the table files.

The `flow_rate` TableFunction is an example of a 1D function. It has a single input, which is time. The table is defined using a single coordinateFile:

```
0.00000e+00
6.00000e+01
1.20000e+02
3.72000e+03
3.78000e+03
1.00000e+09
```

And a single voxelFile:

```
0.00000e+00
0.00000e+00
5.00000e-02
5.00000e-02
0.00000e+00
0.00000e+00
```

Given the specified linear interpolation method, these values define a simple trapezoidal function. Note: since this is a 1D table, these values could alternately be given within the xml file using the `coordinates` and `values` attributes.

The `sigma_xx` TableFunction is an example of a 3D function. It uses `elementCenter` as its input, which is a vector. It is specified using a set of three coordinate files (one for each axis), and a single voxel file. The geologic model in this example is a layer-cake, which was randomly generated, so the size of the x and y axes are 1. The interpolation method used here is upper, so the values in the table indicate those at the top of each layer.

```
<Functions>
  <!-- Pumping Schedule -->
  <TableFunction
    name="flow_rate"
    inputVarNames="{ time }"
    coordinateFiles="{ $table_root$/flowRate_time.csv }"
    voxelFile="$table_root$/flowRate.csv"/>

  <!-- Geologic Model -->
  <TableFunction
    name="sigma_xx"
    inputVarNames="{ elementCenter }"
    coordinateFiles="{ $table_root$/x.csv, $table_root$/y.csv, $table_root$/z.csv }"
    voxelFile="$table_root$/sigma_xx.csv"
    interpolation="upper"/>

  <TableFunction
    name="sigma_yy"
    inputVarNames="{ elementCenter }"
    coordinateFiles="{ $table_root$/x.csv, $table_root$/y.csv, $table_root$/z.csv }"
    voxelFile="$table_root$/sigma_yy.csv"
    interpolation="upper"/>

  <TableFunction
    name="sigma_zz"
    inputVarNames="{ elementCenter }"
    coordinateFiles="{ $table_root$/x.csv, $table_root$/y.csv, $table_root$/z.csv }"
    voxelFile="$table_root$/sigma_zz.csv"
    interpolation="upper"/>

  <TableFunction
```

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```

name="init_pressure"
inputVarNames="{ elementCenter }"
coordinateFiles="{ $table_root$/x.csv, $table_root$/y.csv, $table_root$/z.csv }"
voxelFile="$table_root$/porePressure.csv"
interpolation="upper"/>

<TableFunction
name="bulk_modulus"
inputVarNames="{ elementCenter }"
coordinateFiles="{ $table_root$/x.csv, $table_root$/y.csv, $table_root$/z.csv }"
voxelFile="$table_root$/bulkModulus.csv"
interpolation="upper"/>

<TableFunction
name="shear_modulus"
inputVarNames="{ elementCenter }"
coordinateFiles="{ $table_root$/x.csv, $table_root$/y.csv, $table_root$/z.csv }"
voxelFile="$table_root$/shearModulus.csv"
interpolation="upper"/>

<TableFunction
name="apertureTable"
coordinates="{ -1.0e-3, 0.0 }"
values="{ 1.0e-6, 1.0e-4 }"/>

</Functions>

```

### 3.5.10 Running GEOSX

Assuming that the preprocessing tools have been correctly installed (see [Advanced XML Features](#)), there will be a script in the GEOSX build/bin directory called *geosx\_preprocessed*. Replacing *geosx* with *geosx\_preprocessed* in an input command will automatically apply the preprocessor and send the results to GEOSX.

Before beginning, we recommend that you make a local copy of the example and its tables. Because we are using advanced xml features in this example, the input file must be pre-processed before running. For example, this will run the code on a debug partition using a total of 36 cores.

```

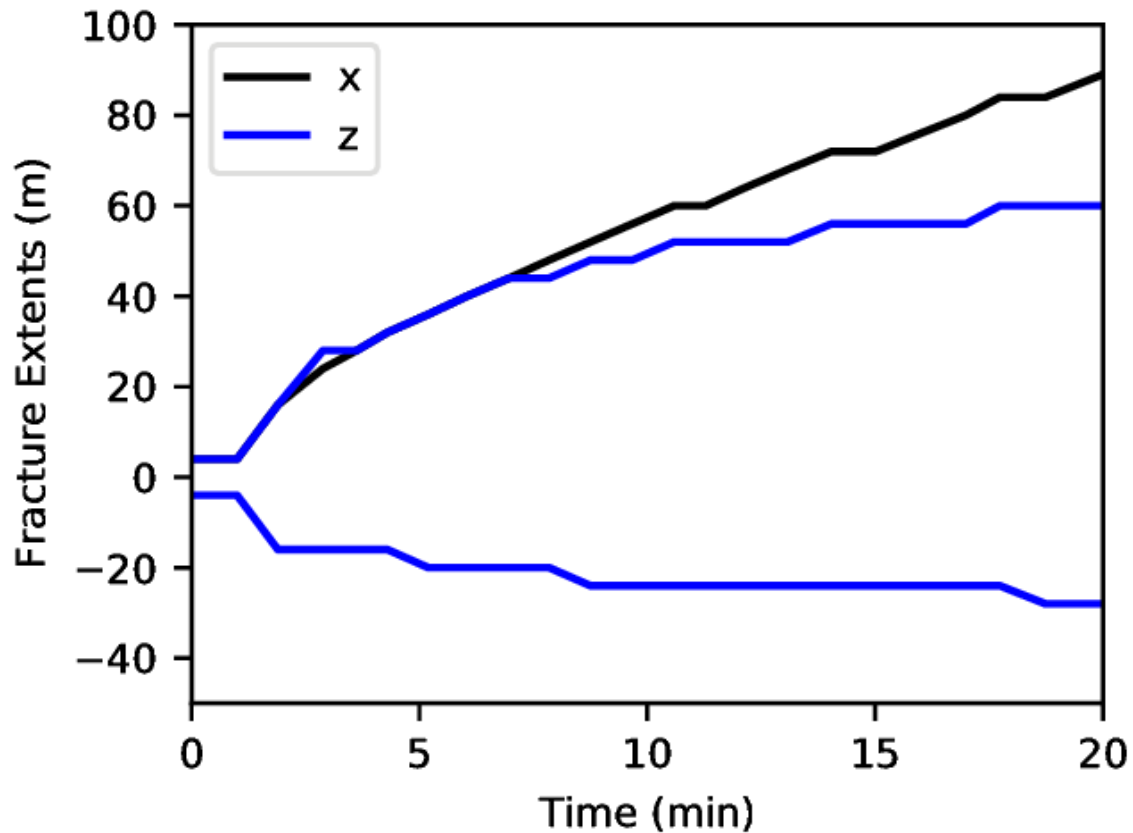
cp -r examples/hydraulicFracturing ./hf_example
cd hf_example
srun -n 36 -ppdebug geosx_preprocessed -i heterogeneousInSitu_benchmark.xml -x 6 -y 2_
↩-z 3 -o hf_results

```

Note that as part of the automatic preprocessing step a compiled xml file is written to the disk (by default '[input\_name].preprocessed'). When developing an xml with advanced features, we recommend that you check this file to ensure its accuracy.

### 3.5.11 Inspecting results

In the above example, we requested vtk- and silo-format output files every minute. We can therefore import these into VisIt, Paraview, or python and visualize the outcome. The following figure shows the extents of the generated fracture over time:

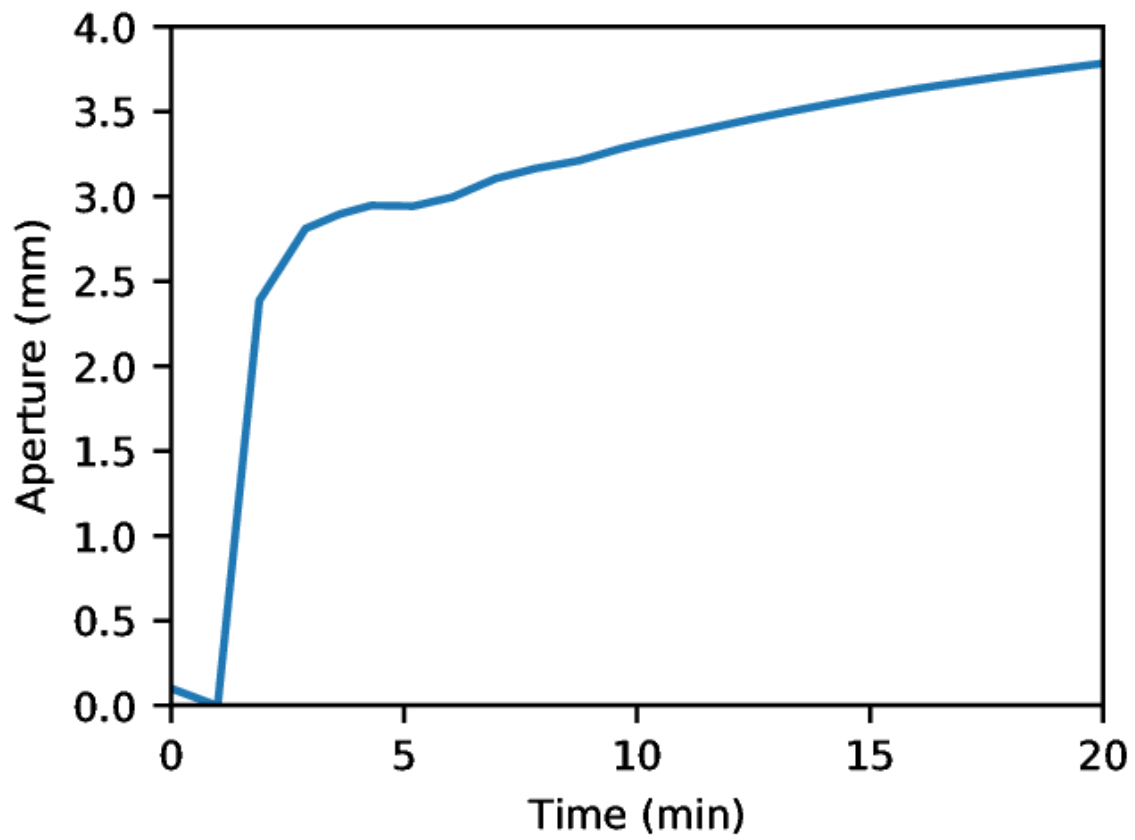


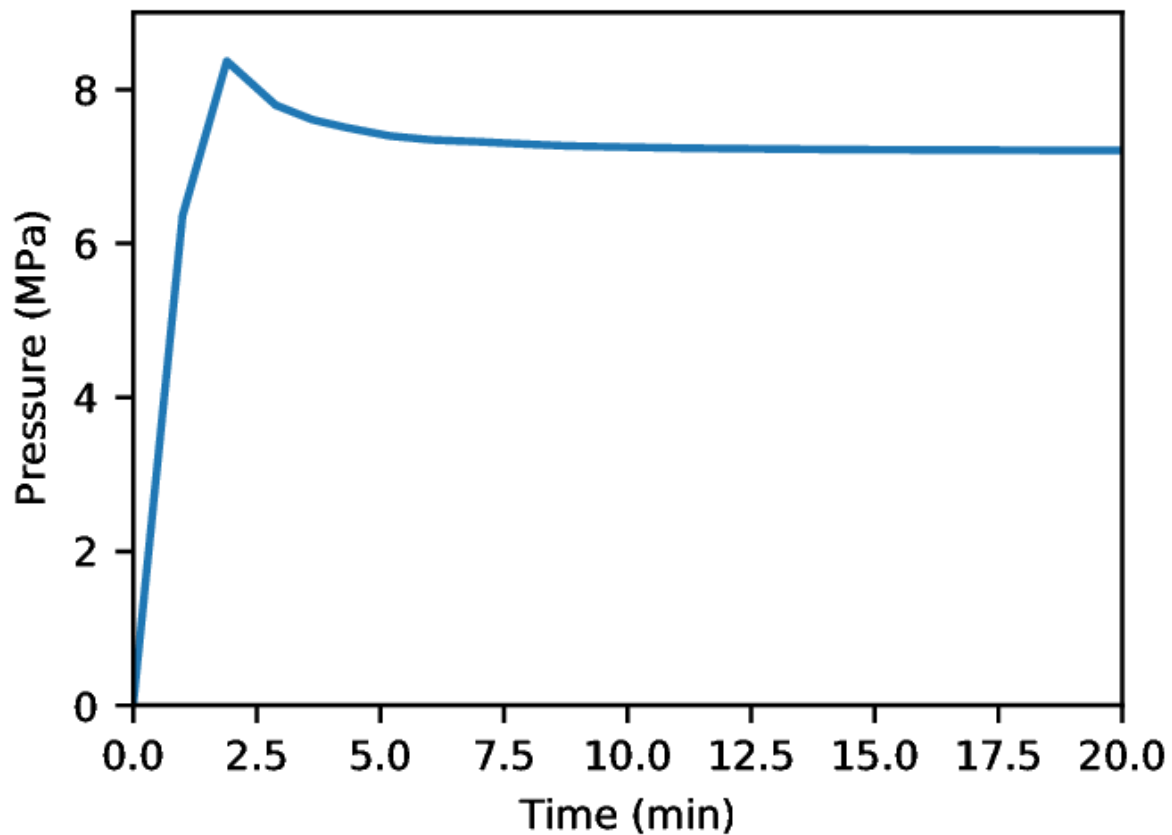
Notes for visualization tools:

- 1) In Visit, we currently recommend that you look at the silo-format files (due to a compatibility issue with vtk)
- 2) In Paraview, you may need to use the Multi-block Inspector (on the right-hand side of the screen by default) to limit the visualization to the fracture. In addition, the Properties inspector (on the left-hand side of the screen by default) may not include some of the parameters present on the fracture. Instead, we recommend that you use the property dropdown box at the top of the screen.

Because we did not explicitly specify any fracture barriers in this example, the fracture dimensions are controlled by the in-situ stresses. During the first couple of minutes of growth, the fracture quickly reaches its maximum/minimum height, which corresponds to a region of low in-situ minimum stress.

The following figures show the aperture and pressure of the hydraulic fracture (near the source) over time:





### 3.5.12 Modifying Parameters Via the Command-Line

The advanced xml feature preprocessor allows parameters to be set or overridden by specifying any number of *-p name value* arguments on the command-line. Note that if the parameter value has spaces, it needs to be enclosed by quotation marks.

To illustrate this feature, we can re-run the previous analysis with viscosity increased from 1 cP to 5 cP:

```
srun -n 36 -ppdebug geosx_preprocessed -i heterogeneousInSitu_benchmark.xml -p mu 0.
↪005 -x 6 -y 2 -z 3 -o hf_results_lower_mu
```

### 3.5.13 To go further

#### Feedback on this example

This concludes the hydraulic fracturing example. For any feedback on this example, please submit a [GitHub issue](#) on the project's [GitHub](#) page.

#### For more details

- More on advanced xml features, please see *Advanced XML Features*.
- More on functions, please see *Functions*.
- More on biased meshes, please see *Mesh Bias*.



## 3.6 Triaxial Driver

### Context

In this example, we use the `TriaxialDriver` inside GEOSX to simulate rock mechanics experiments, such as triaxial tests. The triaxial driver allows to calibrate properties and model parameters against experimental data before using them in field-scale simulations.

### Objectives

At the end of this example, you will know:

- how to set up a triaxial test scenario with the `TriaxialDriver`,
- how to define a material model,
- how to specify loading conditions with `TableFunction`,
- how to save and postprocess simulation results.

### Input file

The XML file for this test case is located at:

```
inputFiles/triaxialDriver/triaxialDriver_ExtendedDruckerPrager.xml
```

This example also uses a set of table files located at:

```
inputFiles/triaxialDriver/tables/
```

Last, a Python script for post-processing the results is provided:

```
src/docs/sphinx/basicExamples/triaxialDriver/triaxialDriverFigure.py
```

### 3.6.1 Description of the case

Instead of launching a full finite-element simulation to mimic experimental loading conditions, GEOSX provides a `TriaxialDriver` to investigate constitutive behaviors and simulate laboratory tests. The triaxial driver makes it easy to interpret the mechanical response and calibrate the constitutive models against experimental data.

In this example, the Extended Drucker-Prager model (see *Model: Extended Drucker-Prager*) is used to solve elastoplastic deformations of rock samples when subject to controlled loading conditions. The strain-hardening Extended Drucker-Prager model with an associated plastic flow rule is tested in this example. To replicate a typical triaxial test, we use a table function to specify loading conditions in axial and radial directions. The resulting strains and stresses in both directions are numerically calculated and saved into a simple ASCII output file.

For this example, we focus on the `Task`, the `Constitutive`, and the `Functions` tags.

### 3.6.2 Task

In GEOSX, the `TriaxialDriver` is defined with a dedicated XML structure. The `TriaxialDriver` is added to a standard XML input deck as a solo task to the `Tasks` queue and added as a `SoloEvent` to the event queue.

For this example, we simulate the elastoplastic deformation of a confined specimen caused by external load. A homogeneous domain with one solid material is assumed. The material is named `ExtendedDruckerPrager`, and its mechanical properties are specified in the `Constitutive` section.

Different testing modes are available in the `TriaxialDriver` to mimic different laboratory loading conditions:

mode	axial loading	radial loading	initial stress
strainControl	axial strain controlled with axialControl	radial strain controlled with radialControl	isotropic stress using initialStress
stressControl	axial stress controlled with axialControl	radial stress controlled with radialControl	isotropic stress using initialStress
mixedControl	axial strain controlled with axialControl	radial stress controlled with radialControl	isotropic stress using initialStress

A triaxial test is usually conducted on a confined rock sample to determine material properties. As shown, a conventional triaxial test is described using the mode="mixedControl" testing mode in the TriaxialDriver.

In a triaxial test, the testing sample is under confining pressure (radial stresses) and subject to increased axial load. Therefore, a stress control radialControl="stressFunction" is defined in the radial direction to impose confining pressure. A strain control axialControl="strainFunction" is applied in the axial direction to represent axial compression.

The initial stress is specified by initialStress="-10.e6". To ensure static equilibrium at the first timestep, its value should be consistent with the initial set of applied stresses defined in axial or radial loading functions. This stress has a negative value due to the negative sign convention for compressive stress in GEOSX.

Then, steps="200" defines the number of load steps and output="simulationResults.txt" specifies an output file to which the simulation results will be written.

```
<Tasks>
  <TriaxialDriver
    name="triaxialDriver"
    material="ExtendedDruckerPrager"
    mode="mixedControl"
    axialControl="strainFunction"
    radialControl="stressFunction"
    initialStress="-10.e6"
    steps="200"
    output="simulationResults.txt" />
  </TriaxialDriver>
</Tasks>
```

In addition to triaxial tests, volumetric and oedometer tests can be simulated by changing the strainControl mode, and by defining loading controls in axial and radial direction accordingly. A volumetric test can be modelled by setting the axial and radial control functions to the same strain function, whereas an oedometer test runs by setting the radial strain to zero (see *Triaxial Driver*).

### 3.6.3 Constitutive laws

Any solid material model implemented in GEOSX can be called by the TriaxialDriver.

For this problem, Extended Drucker Prager model ExtendedDruckerPrager is used to describe the mechanical behavior of an isotropic material, when subject to external loading. As for the material parameters, defaultInitialFrictionAngle, defaultResidualFrictionAngle and defaultCohesion denote the initial friction angle, the residual friction angle, and cohesion, respectively, as defined by the Mohr-Coulomb failure envelope. As the residual friction angle defaultResidualFrictionAngle is larger than the initial one defaultInitialFrictionAngle, a strain hardening model is adopted, whose hardening rate is given as defaultHardening="0.0001". If the residual friction angle is set to be less than the initial one, strain weakening will take place. Setting defaultDilationRatio="1.0" corresponds to an associated flow rule.

```
<Constitutive>
  <ExtendedDruckerPrager
```

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```

    name="ExtendedDruckerPrager"
    defaultDensity="2700"
    defaultBulkModulus="10.0e9"
    defaultShearModulus="6.0e9"
    defaultCohesion="6.0e6"
    defaultInitialFrictionAngle="16.0"
    defaultResidualFrictionAngle="20.0"
    defaultDilationRatio="1.0"
    defaultHardening="0.0001"
  />
</Constitutive>

```

All constitutive parameters such as density, viscosity, bulk modulus, and shear modulus are specified in the International System of Units.

### 3.6.4 Functions

The `TriaxialDriver` uses a simple form of time-stepping to advance through the loading steps, allowing for simulating both rate-dependent and rate-independent models.

In this case, users should define two different time history functions (`strainFunction` and `stressFunction`) to describe loading conditions in axial and radial direction respectively. More specifically, the table functions in this example include the temporal variations of radial stress and axial strain, which rely upon the external files in the table directory (see [Functions](#)). Note that for standard tests with simple loading history, functions can be embedded directly in the XML file without using external tables.

```

<Functions>
  <TableFunction
    name="strainFunction"
    inputVarNames="{ time }"
    coordinateFiles="{ tables/time.geos }"
    voxelFile="tables/axialStrain.geos"/>

  <TableFunction
    name="stressFunction"
    inputVarNames="{ time }"
    coordinateFiles="{ tables/time.geos }"
    voxelFile="tables/radialStress.geos"/>
</Functions>

```

The `strainFunction` `TableFunction` is an example of a 1D interpolated function, which describes the strain as a function of time `inputVarNames="{ time }"`. This table is defined using a single coordinate file:

```

0
1
2
3
4
5

```

And a single voxel file:

```

0.0
-0.004
-0.002

```

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```
-0.005
-0.003
-0.006
```

Similarly, the correlation between the confining stress and time is given through the `stressFunction` defined using the same coordinate file:

```
0
1
2
3
4
5
```

And a different voxel file:

```
-10.0e6
-10.0e6
-10.0e6
-10.0e6
-10.0e6
-10.0e6
```

For this specific test, the confining stress is kept constant and equal to the `initialStress`. Instead of monotonic changing the axial load, two loading/unloading cycles are specified in the `strainFunction`. This way, both plastic loading and elastic unloading can be modeled.

Note that by convention in GEOSX, `stressFunction` and `strainFunction` have negative values for a compressive test.

### 3.6.5 Mesh

Even if discretization is not required for the `TriaxialDriver`, a dummy mesh should be defined to pass all the necessary checks when initializing GEOSX and running the module. A dummy mesh should be created in the `Mesh` section and assigned to the `cellBlocks` in the `ElementRegions` section.

```
<Mesh>
  <InternalMesh
    name="mesh1"
    elementTypes="{ C3D8 }"
    xCoords="{ 0, 1 }"
    yCoords="{ 0, 1 }"
    zCoords="{ 0, 1 }"
    nx="{ 1 }"
    ny="{ 1 }"
    nz="{ 1 }"
    cellBlockNames="{ cellBlock01 }"/>
  </Mesh>

  <ElementRegions>
    <CellElementRegion
      name="dummy"
      cellBlocks="{ cellBlock01 }"
      materialList="{ dummy }"/>
    </ElementRegions>
```

Once calibrated, the testing constitutive models can be easily used in full field-scale simulation by adding solver, discretization, and boundary condition blocks to the xml file. Also, it is possible to run a full GEOSX model and generate identical results as those provided by the `TriaxialDriver`.

### 3.6.6 Running TriaxialDriver

The `TriaxialDriver` is launched like any other GEOSX simulation by using the following command:

```
path/to/geosx -i triaxialDriver_ExtendedDruckerPrager.xml
```

The running log appears to the console to indicate if the case can be successfully executed or not:

```
Max threads: 32
MKL max threads: 16
GEOSX version 0.2.0 (HEAD, sha1: bb16d72)
Adding Event: SoloEvent, triaxialDriver
    TableFunction: strainFunction
    TableFunction: stressFunction
Adding Mesh: InternalMesh, mesh1
Adding Object CellElementRegion named dummy from ObjectManager::Catalog.
Total number of nodes:8
Total number of elems:1
Rank 0: Total number of nodes:8
        dummy/cellBlock01 does not have a discretization associated with it.
Time: 0s, dt:1s, Cycle: 0
Cleaning up events
Umpire          HOST sum across ranks: 23.2 KB
Umpire          HOST          rank max: 23.2 KB
total time      0.435s
initialization time 0.053s
run time        0.004s
```

### 3.6.7 Inspecting results

The simulation results are saved in a text file, named `simulationResults.txt`. This output file has a brief header explaining the meaning of each column. Each row corresponds to one timestep of the driver, starting from initial conditions in the first row.

```
# column 1 = time
# column 2 = axial_strain
# column 3 = radial_strain_1
# column 4 = radial_strain_2
# column 5 = axial_stress
# column 6 = radial_stress_1
# column 7 = radial_stress_2
# column 8 = newton_iter
# column 9 = residual_norm
0.0000e+00 0.0000e+00 0.0000e+00 0.0000e+00 -1.0000e+07 -1.0000e+07 -1.0000e+07 0.
→0000e+00 0.0000e+00
2.5000e-02 -1.0000e-04 2.5000e-05 2.5000e-05 -1.1500e+07 -1.0000e+07 -1.0000e+07 1.
→0000e+00 0.0000e+00
5.0000e-02 -2.0000e-04 5.0000e-05 5.0000e-05 -1.3000e+07 -1.0000e+07 -1.0000e+07 1.
→0000e+00 0.0000e+00
7.5000e-02 -3.0000e-04 7.5000e-05 7.5000e-05 -1.4500e+07 -1.0000e+07 -1.0000e+07 1.
→0000e+00 0.0000e+00
```

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```
1.0000e-01 -4.0000e-04 1.0000e-04 1.0000e-04 -1.6000e+07 -1.0000e+07 -1.0000e+07 1.  
↪0000e+00 0.0000e+00  
... 
```

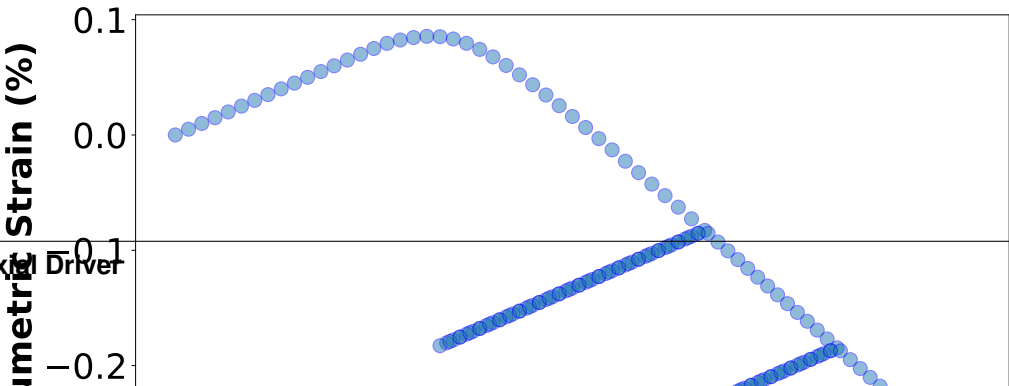
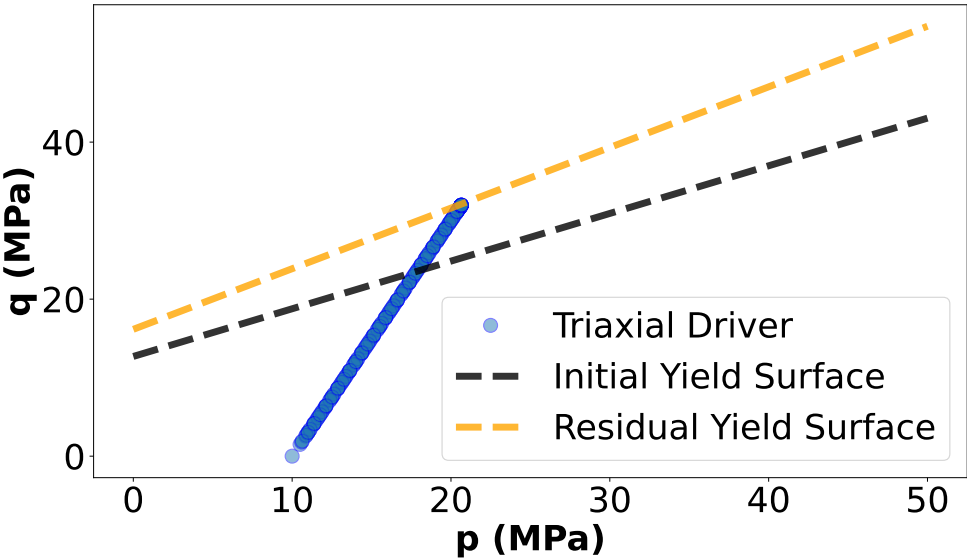
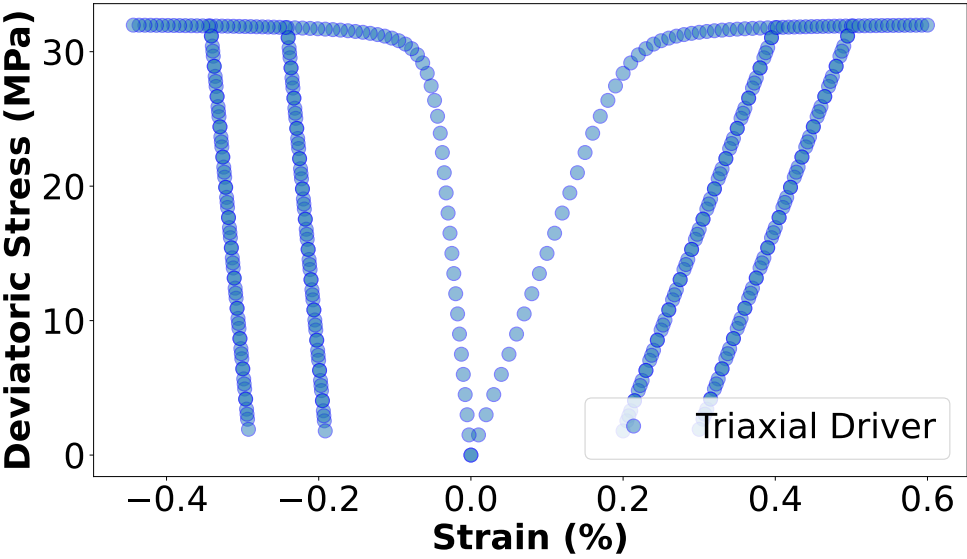
Note that the file contains two columns for radial strains (`radial_strain_1` and `radial_strain_2`) and two columns for radial stresses (`radial_stress_1` and `radial_stress_2`). For isotropic materials, the stresses and strains along the two radial axes would be the same. However, the stresses and strains in the radial directions can differ for cases with anisotropic materials and true-triaxial loading conditions.

This output file can be processed and visualized using any tool. As an example here, with the provided python script, the simulated stress-strain curve, p-q diagram and relationship between volumetric strain and axial strain are plotted, and used to validate results against experimental observations:

### 3.6.8 To go further

#### Feedback on this example

For any feedback on this example, please submit a [GitHub issue on the project's GitHub page](#).







## 4.1 Validation and Verification Studies

### 4.1.1 Carbon Storage

#### Verification of CO<sub>2</sub> Core Flood Experiment with Buckley-Leverett Solution

##### Context

In this example, we simulate a CO<sub>2</sub> core flood experiment representing immiscible transport of two-phase flow (CO<sub>2</sub> and water) through porous media (Ekechukwu et al., 2022). This problem is solved using the multiphase flow solver in GEOSX to obtain the temporal evolution of saturation along the flow direction, and verified against the Buckley-Leverett analytical solutions (Buckley and Leverett, 1942; Arabzai and Honma, 2013).

##### Input file

The xml input files for the test case are located at:

```
inputFiles/compositionalMultiphaseFlow/benchmarks/buckleyLeverettProblem/  
↪buckleyLeverett_base.xml  
inputFiles/compositionalMultiphaseFlow/benchmarks/buckleyLeverettProblem/  
↪buckleyLeverett_benchmark.xml
```

Table files and a Python script for post-processing the simulation results are provided:

```
inputFiles/compositionalMultiphaseFlow/benchmarks/buckleyLeverettProblem/  
↪buckleyLeverett_table
```

```
src/docs/sphinx/advancedExamples/validationStudies/carbonStorage/buckleyLeverett/  
↪buckleyLeverettFigure.py
```

## Description of the case

We model the immiscible displacement of brine by CO<sub>2</sub> in a quasi one-dimensional domain that mimics a CO<sub>2</sub> core flood experiment, as shown below. The domain is horizontal, homogeneous, isotropic and isothermal. Prior to injection, the domain is fully saturated with brine. To match the analytical example configuration, supercritical CO<sub>2</sub> is injected from the inlet and a constant flow rate is imposed. To meet the requirements of immiscible transport in one-dimensional domain, we assume linear and horizontal flow, incompressible and immiscible phases, negligible capillary pressure and gravitational forces, and no poromechanical effects. Upon injection, the saturation front of the injected phase (supercritical CO<sub>2</sub>) forms a sharp leading edge and advances with time.



Fig. 4.1: Sketch of the problem

We set up and solve a multiphase flow model to obtain the spatial and temporal solutions of phase saturations and pore pressures across the domain upon injection. Saturation profiles along the flow direction are evaluated and compared with their corresponding analytical solutions (Arabzai and Honma, 2013).

A power-law Brooks-Corey relation is used here to describe gas  $k_{rg}$  and water  $k_{rw}$  relative permeabilities:

$$k_{rg} = k_{rg}^0 (S_g^*)^{n_g}$$

$$k_{rw} = k_{rw}^0 (S_w^*)^{n_w}$$

where  $k_{rg}^0$  and  $k_{rw}^0$  are the maximum relative permeability of gas and water phase respectively;  $n_g$  and  $n_w$  are the Corey exponents; dimensionless volume fraction (saturation) of gas phase  $S_g^*$  and water phase  $S_w^*$  are given as:

$$S_g^* = \frac{S_g - S_{gr}}{1 - S_{gr} - S_{wr}}$$

$$S_w^* = \frac{S_w - S_{wr}}{1 - S_{gr} - S_{wr}}$$

where  $S_{gr}$  and  $S_{wr}$  are the residual gas and water saturation;

According to the Buckley–Leverett theory with constant fluid viscosity, the fractional flow of gas phase  $f_g$  can be expressed as:

$$f_g = \frac{\frac{k_{rg}}{\mu_g}}{\frac{k_{rg}}{\mu_g} + \frac{k_{rw}}{\mu_w}}$$

where  $\mu_g$  and  $\mu_w$  represent the viscosity of gas and water phase respectively, assumed to be constant in this study.

The position of a particular saturation is given as a function of the injection time  $t$  and the value of the derivative  $\frac{df_g}{dS_g}$  at that saturation:

$$x_{S_g} = \frac{Q_T t}{A\phi} \left( \frac{df_g}{dS_g} \right)$$

where  $Q_T$  is the total flow rate,  $A$  is the area of the cross-section in the core sample,  $\phi$  is the rock porosity. In addition, the abrupt saturation front is determined based on the tangent point on the fractional flow curve.

For this example, we focus on the `Mesh`, the `Constitutive`, and the `FieldSpecifications` tags.

## Mesh

The mesh was created with the internal mesh generator and parametrized in the `InternalMesh` XML tag. It contains  $1000 \times 1 \times 1$  eight-node brick elements in the x, y, and z directions respectively. Such eight-node hexahedral elements are defined as `C3D8` `elementTypes`, and their collection forms a mesh with one group of cell blocks named here `cellBlock`. The width of the domain should be large enough to ensure the formation of a one-dimension flow.

```
<Mesh>
  <InternalMesh
    name="mesh"
    elementTypes="{ C3D8 }"
    xCoords="{ 0, 0.1 }"
    yCoords="{ 0, 0.00202683 }"
    zCoords="{ 0, 1 }"
    nx="{ 1000 }"
    ny="{ 1 }"
    nz="{ 1 }"
    cellBlockNames="{ cellBlock }"/>
  </InternalMesh>
</Mesh>
```

## Flow solver

The isothermal immiscible simulation is performed using the GEOSX general-purpose multiphase flow solver. The multiphase flow solver, a solver of type `CompositionalMultiphaseFVM` called here `compflow` (more information on these solvers at *Compositional Multiphase Flow Solver*) is defined in the XML block **CompositionalMultiphaseFVM**:

```
<Solvers>
  <CompositionalMultiphaseFVM
    name="compflow"
    logLevel="1"
    discretization="fluidTPFA"
    temperature="300"
    initialDt="0.001"
    useMass="1"
    targetRegions="{ region }">
    <NonlinearSolverParameters
      newtonTol="1.0e-6"
      newtonMaxIter="50"
      maxTimeStepCuts="2"
      lineSearchMaxCuts="2"/>
    <LinearSolverParameters
      solverType="direct"
      directParallel="0"
      logLevel="0"/>
  </CompositionalMultiphaseFVM>
</Solvers>
```

We use the `targetRegions` attribute to define the regions where the flow solver is applied. Here, we only simulate fluid flow in one region named as `region`. We specify the discretization method (`fluidTPFA`, defined in the

NumericalMethods section), and the initial reservoir temperature (temperature="300").

## Constitutive laws

This benchmark example involves an immiscible, incompressible, two-phase model, whose fluid rheology and permeability are specified in the Constitutive section. The best approach to represent this fluid behavior in GEOSX is to use the **DeadOilFluid** model in GEOSX.

```
<Constitutive>
  <CompressibleSolidConstantPermeability
    name="rock"
    solidModelName="nullSolid"
    porosityModelName="rockPorosity"
    permeabilityModelName="rockPerm"/>

  <NullModel
    name="nullSolid"/>

  <PressurePorosity
    name="rockPorosity"
    defaultReferencePorosity="0.2"
    referencePressure="1e7"
    compressibility="1.0e-15"/>

  <ConstantPermeability
    name="rockPerm"
    permeabilityComponents="{ 9.0e-13, 9.0e-13, 9.0e-13 }"/>

  <BrooksCoreyRelativePermeability
    name="relperm"
    phaseNames="{ gas, water }"
    phaseMinVolumeFraction="{ 0.0, 0.0 }"
    phaseRelPermExponent="{ 3.5, 3.5 }"
    phaseRelPermMaxValue="{ 1.0, 1.0 }"/>

  <DeadOilFluid
    name="fluid"
    phaseNames="{ gas, water }"
    surfaceDensities="{ 280.0, 992.0 }"
    componentMolarWeight="{ 44e-3, 18e-3 }"
    tableFiles="{ buckleyLeverett_table/pvdg.txt, buckleyLeverett_table/pvtw.txt }"/
  >
</Constitutive>
```

Constant fluid densities and viscosities are given in the external tables for both phases. The formation volume factors are set to 1 for incompressible fluids. The relative permeability for both phases are modeled with the power-law correlations BrooksCoreyRelativePermeability (more information at [Brooks-Corey relative permeability model](#)), as shown below. Capillary pressure is assumed to be negligible.

All constitutive parameters such as density, viscosity, and permeability are specified in the International System of Units.

## Time history function

In the Tasks section, PackCollection tasks are defined to collect time history information from fields. Either the entire field or specified named sets of indices in the field can be collected. In this exam-

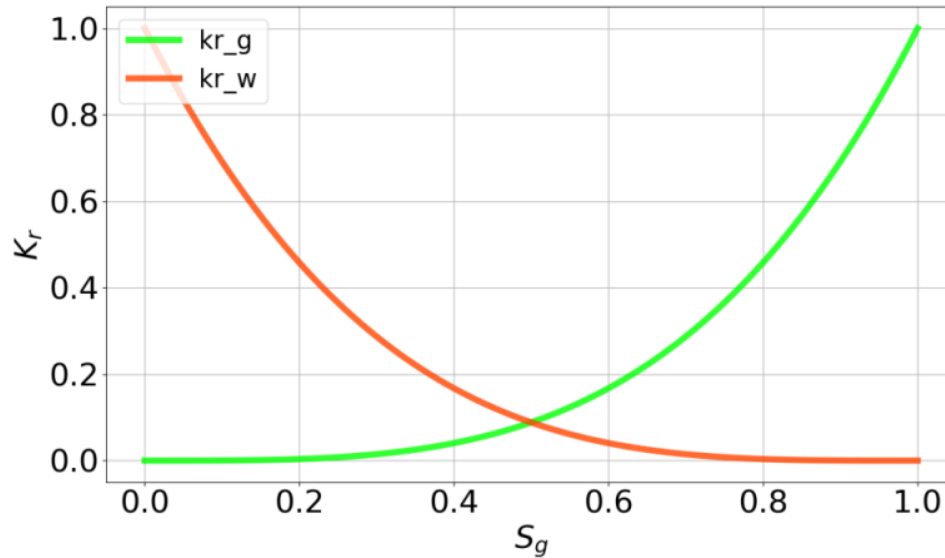


Fig. 4.2: Relative permeabilities of both phases

ple, `phaseVolumeFractionCollection` is specified to output the time history of phase saturations `fieldName="phaseVolumeFraction"` across the computational domain.

```
<Tasks>
  <PackCollection
    name="phaseVolumeFractionCollection"
    objectPath="ElementRegions/region/cellBlock"
    fieldName="phaseVolumeFraction"/>
</Tasks>
```

This task is triggered using the Event manager with a `PeriodicEvent` defined for the recurring tasks. GEOSX writes one file named after the string defined in the `filename` keyword, formatted as a HDF5 file (`saturationHistory.hdf5`). The `TimeHistory` file contains the collected time history information from the specified time history collector. This file includes datasets for the simulation time, element center, and the time history information for both phases. A Python script is prepared to read and plot any specified subset of the time history data for verification and visualization.

## Initial and boundary conditions

The next step is to specify fields, including:

- The initial value (the pore pressure and phase saturations have to be initialized)
- The boundary conditions (fluid injection rates and controls on the fluid outlet have to be set)

In this example, the domain is initially saturated with brine in a uniform pressure field. The `component` attribute of the **FieldSpecification** XML block must use the order in which the `phaseNames` have been defined in the **DeadOilFluid** XML block. In other words, `component=0` is used to initialize the gas global component fraction and `component=1` is used to initialize the water global component fraction, because we previously set `phaseNames="{gas, water}"` in the **DeadOilFluid** XML block.

A mass injection rate `SourceFlux` (`scale="-0.00007"`) of pure CO<sub>2</sub> (`component="0"`) is applied at the fluid inlet, named `source`. The value given for `scale` is  $Q_T \rho_g$ . Pressure and composition controls at the fluid outlet

(sink) are also specified. The `setNames="{ source }"` and `setNames="{ sink }"` are defined using the **Box XML** tags of the **Geometry** section.

These boundary conditions are set up through the `FieldSpecifications` section.

```
<FieldSpecifications>
  <FieldSpecification
    name="initialPressure"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions"
    fieldName="pressure"
    scale="1e7"/>

  <FieldSpecification
    name="initialComposition_gas"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions"
    fieldName="globalCompFraction"
    component="0"
    scale="0.001"/>

  <FieldSpecification
    name="initialComposition_water"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions"
    fieldName="globalCompFraction"
    component="1"
    scale="0.999"/>

  <SourceFlux
    name="sourceTerm"
    objectPath="ElementRegions"
    scale="-0.00007"
    component="0"
    setNames="{ source }"/>

  <FieldSpecification
    name="sinkTermPressure"
    objectPath="faceManager"
    fieldName="pressure"
    scale="1e7"
    setNames="{ sink }"/>

  <FieldSpecification
    name="sinkTermTemperature"
    objectPath="faceManager"
    fieldName="temperature"
    scale="300"
    setNames="{ sink }"/>

  <FieldSpecification
    name="sinkTermComposition_gas"
    setNames="{ sink }"
    objectPath="faceManager"
    fieldName="globalCompFraction"
```

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```

component="0"
scale="0.001"/>

<FieldSpecification
  name="sinkTermComposition_water"
  setNames="{ sink }"
  objectPath="faceManager"
  fieldName="globalCompFraction"
  component="1"
  scale="0.999"/>
</FieldSpecifications>

```

The parameters used in the simulation are summarized in the following table, and specified in the Constitutive and FieldSpecifications sections.

Symbol	Parameter	Unit	Value
$k_{rg}^0$	Max Relative Permeability of Gas	[-]	1.0
$k_{rw}^0$	Max Relative Permeability of Water	[-]	1.0
$n_g$	Corey Exponent of Gas	[-]	3.5
$n_w$	Corey Exponent of Water	[-]	3.5
$S_{gr}$	Residual Gas Saturation	[-]	0.0
$S_{wr}$	Residual Water Saturation	[-]	0.0
$\phi$	Porosity	[-]	0.2
$\kappa$	Matrix Permeability	[m <sup>2</sup> ]	$9.0 \cdot 10^{-13}$
$\mu_g$	Gas Viscosity	[Pa s]	$2.3 \cdot 10^{-5}$
$\mu_w$	Water Viscosity	[Pa s]	$5.5 \cdot 10^{-4}$
$Q_T$	Total Flow Rate	[m <sup>3</sup> /s]	$2.5 \cdot 10^{-7}$
$D_L$	Domain Length	[m]	0.1
$D_W$	Domain Width	[m]	1.0
$D_T$	Domain Thickness	[m]	0.002

## Inspecting results

We request VTK-format output files and use Paraview to visualize the results. The following figure shows the distribution of phase saturations and pore pressures in the computational domain at  $t = 70s$ .

Two following dimensionless terms are defined when comparing the numerical solution with the analytical solutions:

$$t^* = \frac{Q_T t}{AD_L \phi}$$

$$x_d = \frac{x_{S_g}}{D_L}$$

The figure below compares the results from GEOSX (dashed curves) and the corresponding analytical solution (solid curves) for the change of gas saturation ( $S_g$ ) and water saturation ( $S_w$ ) along the flow direction.

GEOSX reliably captures the immiscible transport of two phase flow (CO<sub>2</sub> and water). GEOSX matches the analytical solutions in the formation and the progress of abrupt fronts in the saturation profiles.

## To go further

### Feedback on this example

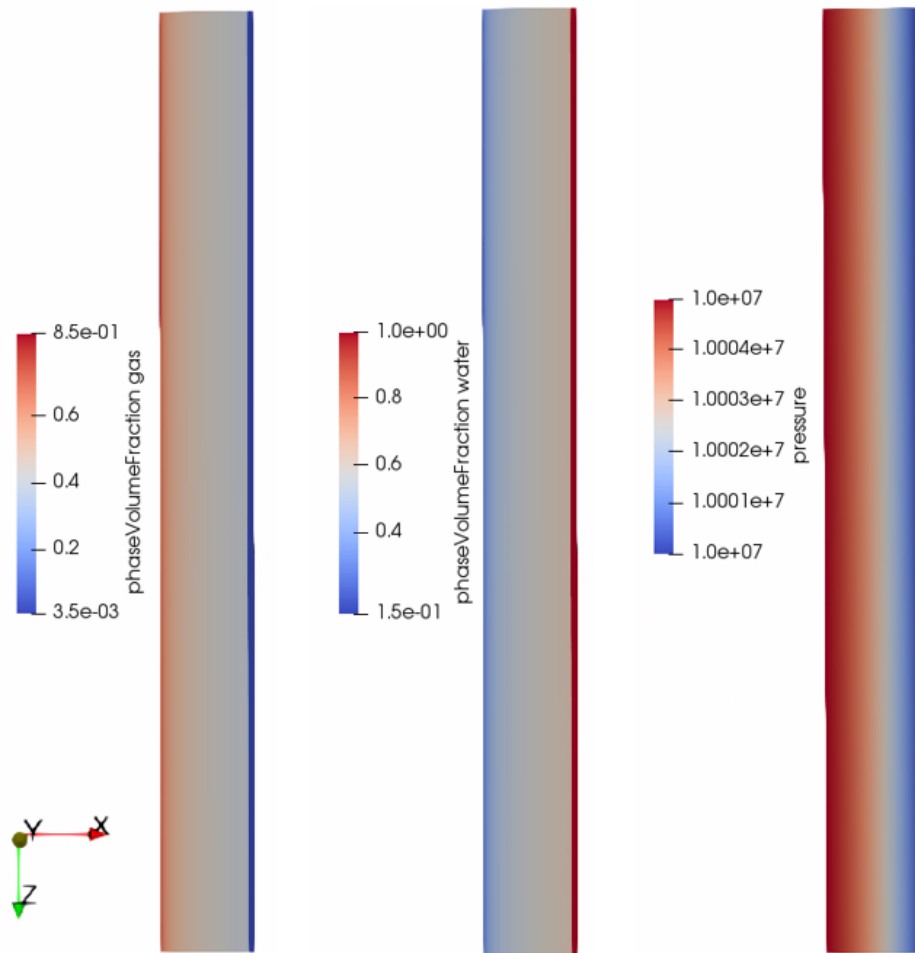
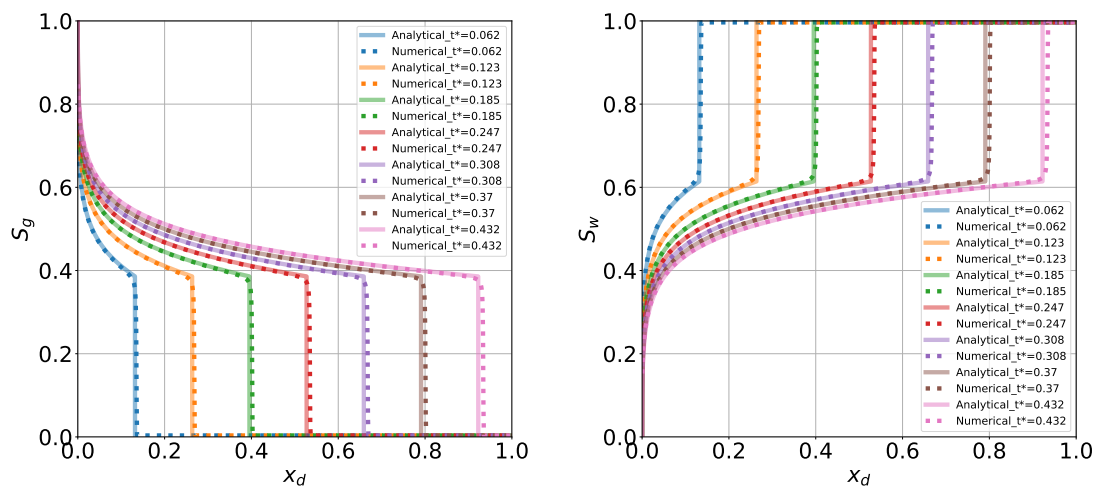


Fig. 4.3: Simulation results of phase saturations and pore pressure





For any feedback on this example, please submit a [GitHub issue](#) on the project's [GitHub page](#).

## CO2 Plume Evolution and Leakage Through an Abandoned Well

### Context

We consider a benchmark problem used in (Class et al., 2009) to compare a number of numerical models applied to CO2 storage in geological formations. Using a simplified isothermal and immiscible two-phase setup, this test case addresses the simulation of the advective spreading of CO2 injected into an aquifer and the leakage of CO2 from the aquifer through an abandoned, leaky well.

Our goal is to review the different sections of the XML file reproducing the benchmark configuration and to demonstrate that the GEOSX results (i.e., arrival time of the CO2 plume at the leaky well and leakage rate through the abandoned well) are in agreement with the reference results published in (Class et al., 2009).

The GEOSX results obtained for the non-isothermal version of this test case (referred to as Problem 1.2 in (Class et al., 2009)) will be presented in a separate documentation page.

### Input file

This benchmark test is based on the XML file located below:

```
inputFiles/compositionalMultiphaseFlow/benchmarks/isothermalLeakyWell/
→ isothermalLeakyWell_benchmark.xml
```

### Problem description

The following text is adapted from the detailed description of the benchmark test case presented in (Ebigbo, Class, Helmig, 2007) and (Class et al., 2009).

The leakage scenario considered here involves one CO2 injection well, one leaky well, two aquifers and an aquitard. The setup is illustrated in the figure below. The leaky well connects the two aquifers. CO2 is injected into the lower aquifer, comes in contact with the leaky well and rises to the higher aquifer. The advective flow (including the buoyancy effects) of CO2 in the initially brine-saturated aquifers and through the leaky well is the most important process in this problem.

The model domain is located 2840 to 3000 m below the surface and has the following dimensions: 1000 x 1000 x 160 m. The distance between the injection and the leaky well is 100 m, and the injection rate of CO2 into the lower aquifer is constant (equal to 8.87 kg/s). The wells are described as cylinders with a 0.15 m radius and are treated as a porous medium with a higher permeability than the aquifers (i.e., this problem does not require a well model).

The authors have used the following simplifying assumptions:

- The formation is isotropic.
- All processes are isothermal.
- CO2 and brine are two separate and immiscible phases. Mutual dissolution is neglected.
- Fluid properties such as density and viscosity are constant.
- The pressure conditions at the lateral boundaries are constant over time, and equal to the initial hydrostatic condition.
- Relative permeabilities are linear functions of saturation.
- Capillary pressure is negligible.

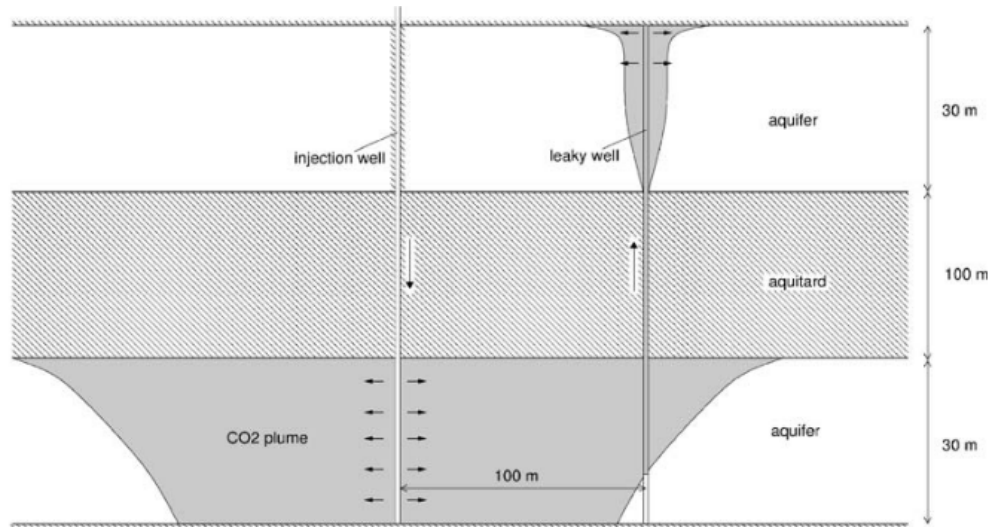


Fig. 4.4: Leakage scenario (image taken from (Ebigbo, Class, Helmig, 2007)).

### Mesh and element regions

The structured mesh is generated using the internal mesh generator as parameterized in the **InternalMesh** block of the XML file. The mesh contains 112 x 101 x 60 hexahedral elements (C3D8 element type) in the x, y, and z directions respectively.

The attributes `nx`, `ny`, `nz`, and `cellBlockNames` are used to define  $5 \times 3 \times 3 = 45$  cell blocks. Note that the cell block names listed in `cellBlockNames` are mapped to their corresponding cell block using an k-j-i logic in which the k index is the fastest index, and the i index is the slowest index.

```
<Mesh>
  <InternalMesh
    name="mesh"
    elementTypes="{ C3D8 }"
    xCoords="{ -500, -0.1329, 0.1329, 99.8671, 100.1329, 500 }"
    yCoords="{ -500, -0.1329, 0.1329, 500 }"
    zCoords="{ -3000, -2970, -2870, -2840 }"
    nx="{ 50, 1, 20, 1, 40 }"
    ny="{ 50, 1, 50 }"
    nz="{ 20, 30, 10 }"
    cellBlockNames="{ aquiferBottom00, aquitard00, aquiferTop00,
                      aquiferBottom01, aquitard01, aquiferTop01,
                      aquiferBottom02, aquitard02, aquiferTop02,
                      aquiferBottom10, aquitard10, aquiferTop10,
                      aquiferBottom11, aquitard11, aquiferTop11,
                      aquiferBottom12, aquitard12, aquiferTop12,
                      aquiferBottom20, aquitard20, aquiferTop20,
                      aquiferBottom21, aquitard21, aquiferTop21,
                      aquiferBottom22, aquitard22, aquiferTop22,
                      aquiferBottom30, aquitard30, aquiferTop30,
                      aquiferBottom31, aquitard31, aquiferTop31,
                      aquiferBottom32, aquitard32, aquiferTop32,
                      aquiferBottom40, aquitard40, aquiferTop40,
                      aquiferBottom41, aquitard41, aquiferTop41,
                      aquiferBottom42, aquitard42, aquiferTop42 }"/>
  </InternalMesh>
</Mesh>
```

**Note:** In the GEOSX input file, the two wells are defined as columns of hexahedral elements of individual size 0.2658 x 0.2658 x 1 m. The coefficient 0.2858 is chosen to ensure that the cross-section of the wells is equal to the one defined in the benchmark, in which wells are defined as cylinders with a radius of 0.15.

The cell block names are used in the **ElementRegions** block to define element regions in the aquifers and in the wells. In the benchmark definition, the wells are treated as a porous medium with a higher permeability and therefore the rock models are not the same in the aquifers (rock) and in the wells (rockWell). These names are defined in the **Constitutive** block.

```
<ElementRegions>
  <CellElementRegion
    name="aquiferBottom"
    cellBlocks="{ aquiferBottom00, aquiferBottom01, aquiferBottom02,
                  aquiferBottom10,          aquiferBottom12,
                  aquiferBottom20, aquiferBottom21, aquiferBottom22,
                  aquiferBottom30,          aquiferBottom32,
                  aquiferBottom40, aquiferBottom41, aquiferBottom42 }"
    materialList="{ fluid, rock, relperm }"/>
  <CellElementRegion
    name="aquiferTop"
    cellBlocks="{ aquiferTop00, aquiferTop01, aquiferTop02,
                  aquiferTop10,          aquiferTop12,
                  aquiferTop20, aquiferTop21, aquiferTop22,
                  aquiferTop30,          aquiferTop32,
                  aquiferTop40, aquiferTop41, aquiferTop42 }"
    materialList="{ fluid, rock, relperm }"/>
  <CellElementRegion
    name="injectionWell"
    cellBlocks="{ aquiferBottom31 }"
    materialList="{ fluid, rockWell, relperm }"/>
  <CellElementRegion
    name="leakyWell"
    cellBlocks="{ aquiferTop11, aquitard11, aquiferBottom11 }"
    materialList="{ fluid, rockWell, relperm }"/>
  <CellElementRegion
    name="Barrier"
    cellBlocks="{ aquitard00, aquitard01, aquitard02,
                  aquitard10,          aquitard12,
                  aquitard20, aquitard21, aquitard22,
                  aquitard30, aquitard31, aquitard32,
                  aquitard40, aquitard41, aquitard42,
                  aquiferTop31 }"
    materialList="{ }"/>
</ElementRegions>
```

Defining these element regions allows the flow solver to be applied only to the top and bottom aquifers (and wells), since we follow the approach of (Class et al., 2009) and do not simulate flow in the aquitard, as we will see in the next section.

**Note:** Since the two aquifer regions share the same material properties, they could have been defined as a single **CellElementRegion** for a more concise XML file producing the same results. The same is true for the two well regions.

## Flow solver

The isothermal immiscible simulation is performed by the GEOSX general-purpose multiphase flow solver based on a TPFA discretization defined in the XML block **CompositionalMultiphaseFVM**:

```
<Solvers>
  <CompositionalMultiphaseFVM
    name="compflow"
    logLevel="1"
    discretization="fluidTPFA"
    temperature="313.15"
    useMass="1"
    targetRegions="{ aquiferTop, aquiferBottom, injectionWell, leakyWell }">
    <NonlinearSolverParameters
      newtonTol="1.0e-3"
      newtonMaxIter="100"
      maxTimeStepCuts="5"
      lineSearchAction="Attempt"/>
    <LinearSolverParameters
      solverType="fgmres"
      preconditionerType="mgr"
      krylovTol="1e-4"/>
    </CompositionalMultiphaseFVM>
  </Solvers>
```

We use the `targetRegions` attribute to define the regions where the flow solver is applied. Here, we follow the approach described in (Class et al., 2009) and do not simulate flow in the aquitard, considered as impermeable to flow. We only simulate flow in the two aquifers (`aquiferTop` and `aquiferBottom`), in the bottom part of the injection well (`injectionWell`), and in the leaky well connecting the two aquifers (`leakyWell`).

## Constitutive laws

This benchmark test involves an immiscible, incompressible, two-phase model. The best approach to represent this fluid behavior in GEOSX is to use the **DeadOilFluid** model, although this model was initially developed for simple isothermal oil-water or oil-gas systems (note that this is also the approach used for the Eclipse simulator, as documented in (Class et al., 2009)).

```
<DeadOilFluid
  name="fluid"
  phaseNames="{ oil, water }"
  surfaceDensities="{ 479.0, 1045.0 }"
  componentMolarWeight="{ 114e-3, 18e-3 }"
  hydrocarbonFormationVolFactorTableNames="{ B_o_table }"
  hydrocarbonViscosityTableNames="{ visc_o_table }"
  waterReferencePressure="1e7"
  waterFormationVolumeFactor="1.0"
  waterCompressibility="1e-15"
  waterViscosity="2.535e-4"/>
```

The fluid properties (constant densities and constant viscosities) are those given in the article documenting the benchmark. To define an incompressible fluid, we set all the formation volume factors to 1.

---

**Note:** For now, we have to use the label `oil` to denote the CO<sub>2</sub> phase in the Dead-Oil model, but this will be changed in future GEOSX releases.

---

The rock model defines an incompressible porous medium with a porosity equal to 0.15. The relative permeability model is linear. Capillary pressure is neglected.

### Initial and boundary conditions

The domain is initially saturated with brine with a hydrostatic pressure field. This is specified using the **HydrostaticEquilibrium** XML tag in the **FieldSpecifications** block. The datum pressure and elevation used below are defined in (Class et al., 2009)).

```
<HydrostaticEquilibrium
  name="equil"
  objectPath="ElementRegions"
  datumElevation="-3000"
  datumPressure="3.086e7"
  initialPhaseName="water"
  componentNames="{ oil, water }"
  componentFractionVsElevationTableNames="{ initOilCompFracTable,
                                             initWaterCompFracTable }"
  temperatureVsElevationTableName="initTempTable"/>
```

In the **Functions** block, the **TableFunction** s named `initOilCompFracTable` and `initWaterCompFracTable` define the brine-saturated initial state, while the **TableFunction** named `initTempTable` defines the homogeneous temperature field (temperature is not used in this benchmark).

Since the fluid densities are constant for this simple incompressible fluid model, the same result could have been achieved using a simpler table in a **FieldSpecification** tag, as explained in *Hydrostatic Equilibrium Initial Condition*. To impose the Dirichlet boundary conditions on the four sides of the aquifers, we use this simple table-based approach as shown below:

```
<FieldSpecification
  name="bcPressureAquiferBottom"
  objectPath="ElementRegions/aquiferBottom"
  setNames="{ east, west, south, north }"
  fieldName="pressure"
  functionName="pressureFunction"
  scale="1"/>
<FieldSpecification
  name="bcPressureAquiferTop"
  objectPath="ElementRegions/aquiferTop"
  setNames="{ east, west, south, north }"
  fieldName="pressure"
  functionName="pressureFunction"
  scale="1"/>
<FieldSpecification
  name="bcPressureInjectionWell"
  objectPath="ElementRegions/injectionWell"
  setNames="{ east, west, south, north }"
  fieldName="pressure"
  functionName="pressureFunction"
  scale="1"/>
<FieldSpecification
  name="bcPressureLeakyWell"
  objectPath="ElementRegions/leakyWell"
  setNames="{ east, west, south, north }"
  fieldName="pressure"
  functionName="pressureFunction"
```

(continues on next page)

```

    scale="1"/>
<FieldSpecification
  name="bcCompositionOilAquiferBottom"
  setNames="{ east, west, south, north }"
  objectPath="ElementRegions/aquiferBottom"
  fieldName="globalCompFraction"
  component="0"
  scale="0.000001"/>
<FieldSpecification
  name="bcCompositionOilAquiferTop"
  setNames="{ east, west, south, north }"
  objectPath="ElementRegions/aquiferTop"
  fieldName="globalCompFraction"
  component="0"
  scale="0.000001"/>
<FieldSpecification
  name="bcCompositionOilInjectionWell"
  setNames="{ east, west, south, north }"
  objectPath="ElementRegions/injectionWell"
  fieldName="globalCompFraction"
  component="0"
  scale="0.000001"/>
<FieldSpecification
  name="bcCompositionOilLeakyWell"
  setNames="{ east, west, south, north }"
  objectPath="ElementRegions/leakyWell"
  fieldName="globalCompFraction"
  component="0"
  scale="0.000001"/>
<FieldSpecification
  name="bcCompositionWaterAquiferBottom"
  setNames="{ east, west, south, north }"
  objectPath="ElementRegions/aquiferBottom"
  fieldName="globalCompFraction"
  component="1"
  scale="0.999999"/>
<FieldSpecification
  name="bcCompositionWaterAquiferTop"
  setNames="{ east, west, south, north }"
  objectPath="ElementRegions/aquiferTop"
  fieldName="globalCompFraction"
  component="1"
  scale="0.999999"/>
<FieldSpecification
  name="bcCompositionWaterInjectionWell"
  setNames="{ east, west, south, north }"
  objectPath="ElementRegions/injectionWell"
  fieldName="globalCompFraction"
  component="1"
  scale="0.999999"/>
<FieldSpecification
  name="bcCompositionWaterLeakyWell"
  setNames="{ east, west, south, north }"
  objectPath="ElementRegions/leakyWell"
  fieldName="globalCompFraction"
  component="1"
  scale="0.999999"/>

```

where the `setNames = "{ east, west, south, north }"` are defined using the **Box** XML tags of the **Geometry** section, and where the tables are defined as **TableFunction** in the **Functions** section.

To reproduce the behavior of a rate-controlled well, we use the **SourceFlux** tag on the `source` set (located in the `injectionWell` cell element regions), with the injection rate specified in the benchmark description (8.87 kg/s):

```
<SourceFlux
  name="sourceTerm"
  objectPath="ElementRegions/injectionWell"
  component="0"
  scale="-8.87"
  setNames="{ source }"/>
```

**Note:** If the `setNames` attribute of **SourceFlux** contains multiple cells (which is the case here), then the amount injected in each cell is equal to the total injection rate (specified with `scale`) divided by the number of cells.

## Inspecting results

We request VTK-format output files and use Paraview to visualize the results. The following figure shows the distribution of CO<sub>2</sub> saturation and pressure along the slice defined by `x = 0` at `t = 200` days.

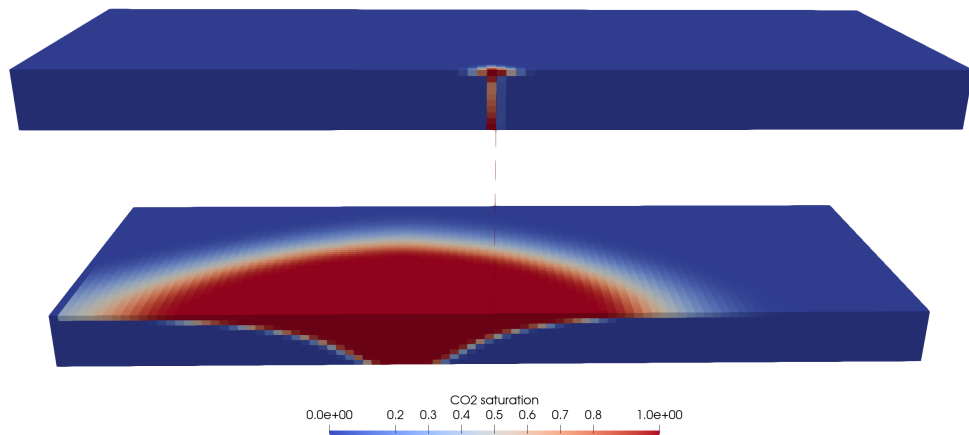


Fig. 4.5: CO<sub>2</sub> saturation after 200 days

To validate the GEOSX results, we consider the metrics used in (Class et al., 2009).

First, we consider the arrival time of the CO<sub>2</sub> plume at the leaky well. As in (Class et al., 2009), we use the leakage rate threshold of 0.005% to detect the arrival time. Although the arrival time is highly dependent on the degree of spatial refinement in the vicinity of the wells (not documented in (Class et al., 2009)), the next table shows that the GEOSX arrival time at the leaky well (9.6 days) is in agreement with the values published in (Class et al., 2009).

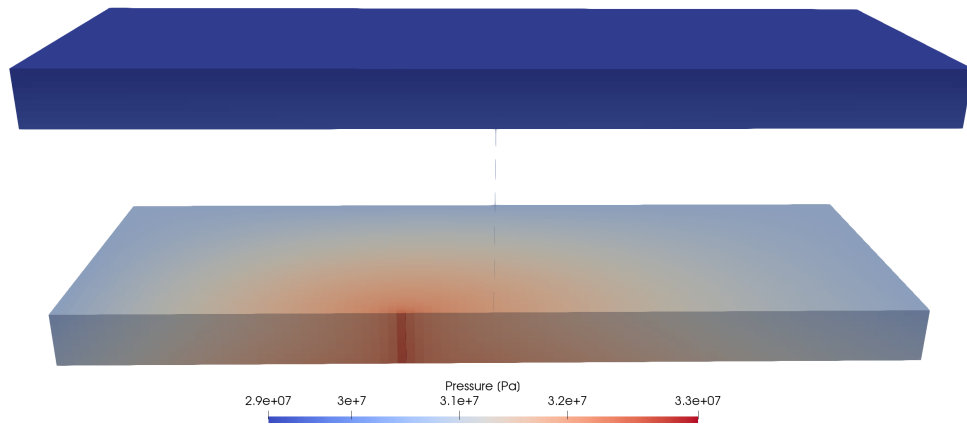


Fig. 4.6: Pressure after 200 days

Code	Arrival time [day]
GEOSX	9.6
COORES	8
DuMux	6
ECLIPSE	8
FEHM	4
IPARS-CO2	10
MUFTE	8
RockFlow	19
ELSA	14
TOUGH2/ECO2N	4
TOUGH2/ECO2N (2)	10
TOUGH2 (3)	9
VESA	7

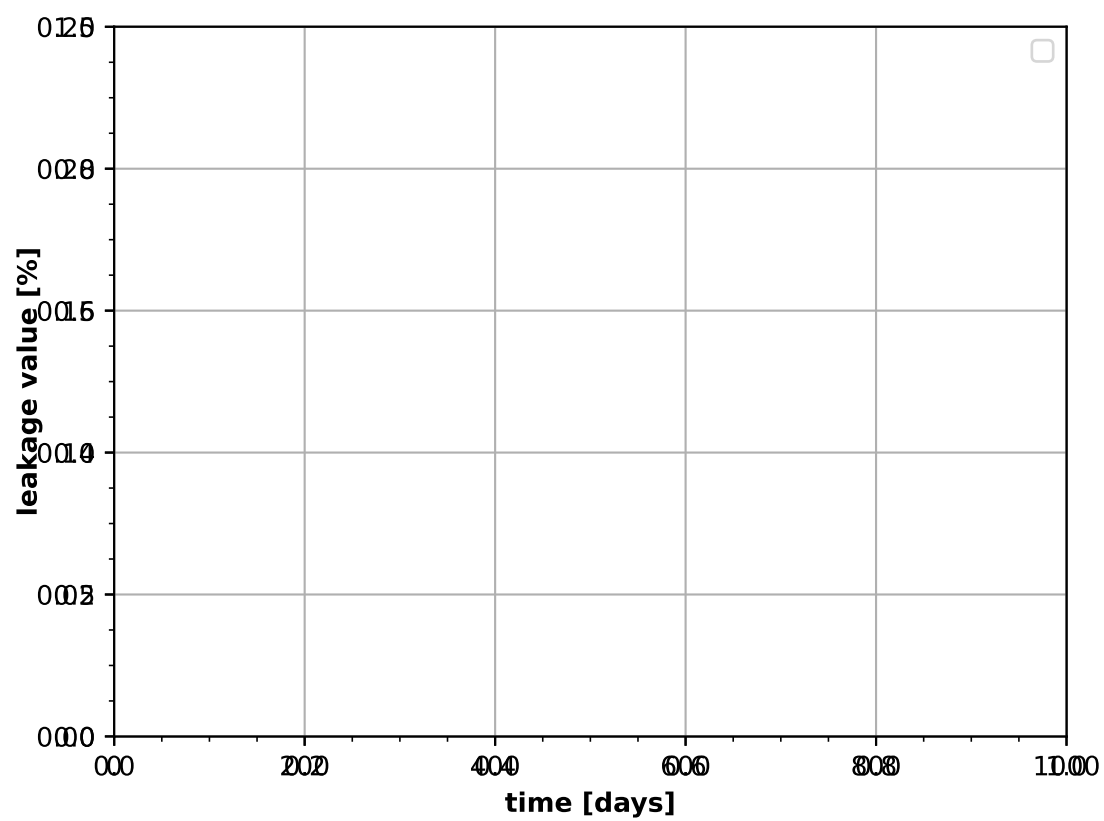
**Note:** The time-stepping strategy used by the codes considered in (Class et al., 2009), as well as the meshes that have been used, are not documented in the benchmark description. Therefore, even on this simple test case, we cannot expect to obtain an exact match with the published results.

Next, we measure the CO<sub>2</sub> leakage rate through the leaky well, defined by the authors as the CO<sub>2</sub> mass flow at midway between top and bottom aquifers divided by the injection rate (8.87 kg/s), in percent. The GEOSX leakage rate is shown in the figure below:

The leakage rates computed by the codes considered in (Class et al., 2009) are shown in the figure below.

The comparison between the previous two figures shows that GEOSX can successfully reproduce the trend in leakage rate observed in the published benchmark results. To further validate the GEOSX results, we reproduce below Table 8 of (Class et al., 2009) to compare the maximum leakage rate, the time at which this maximum leakage rate is attained, and the leakage rate at 1000 days.





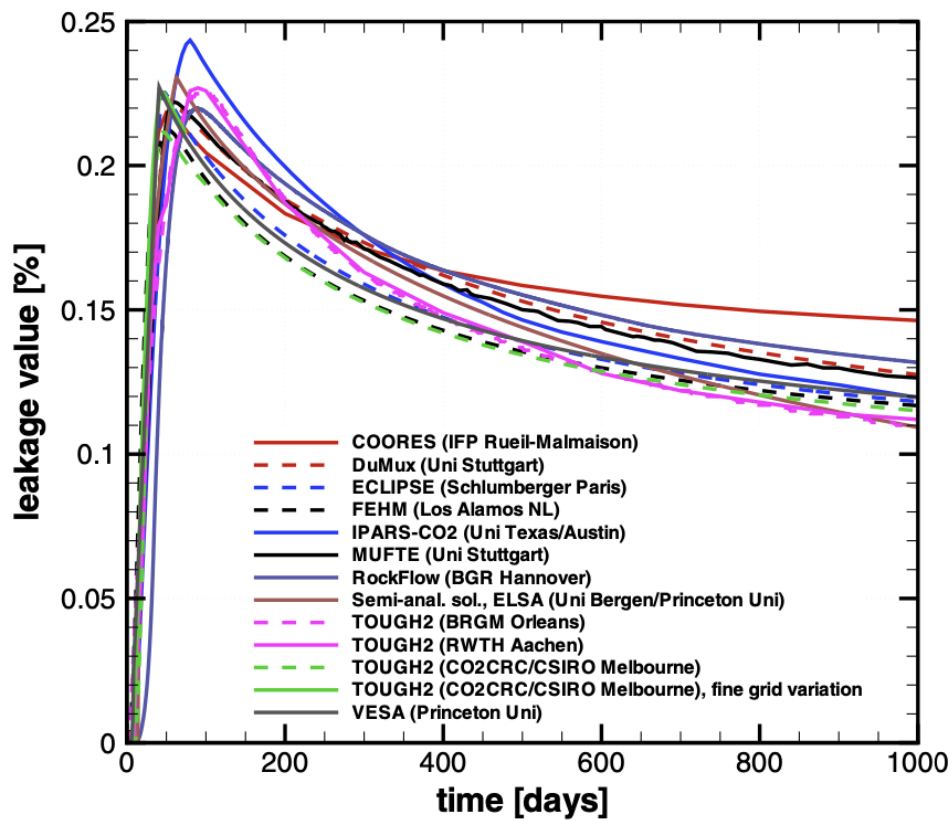


Fig. 4.7: Leakage rates [%] obtained with the simulators considered in (Class et al., 2009).

Code	Max leakage [%]	Time at max leakage [day]	Leakage at 1000 days [%]
GEOSX	0.219	50.6	0.1172
COORES	0.219	50	0.146
DuMux	0.220	61	0.128
ECLIPSE	0.225	48	0.118
FEHM	0.216	53	0.119
IPARS-CO2	0.242	80	0.120
MUFTE	0.222	58	0.126
RockFlow	0.220	74	0.132
ELSA	0.231	63	0.109
TOUGH2/ECO2N	0.226	93	0.110
TOUGH2/ECO2N (2)	0.212	46	0.115
TOUGH2 (3)	0.227	89	0.112
VESA	0.227	41	0.120

This table confirms the agreement between GEOSX and the results of (Class et al., 2009). A particularly good match is obtained with Eclipse, FEHM, and TOUGH2/ECO2N (2).

### To go further

The more complex, non-isothermal version of this test will be added to this suite of validation examples soon.

### Feedback on this example

For any feedback on this example, please submit a [GitHub issue on the project's GitHub page](#).

## 4.1.2 Fault Mechanics

### Single Fracture Under Shear Compression

#### Context

In this example, a single fracture is simulated using a Lagrange contact model in a 2D infinite domain and subjected to a constant uniaxial compressive remote stress (Franceschini et al., 2020). An analytical solution (Phan et al., 2003) is available for verifying the accuracy of the numerical results, providing an analytical form for the normal traction and slip on the fracture surface due to frictional contact. In this example, the `TimeHistory` function and a Python script are used to output and postprocess multi-dimensional data (traction and displacement on the fracture surface).

#### Input file

Everything required is contained within two GEOSX input files and one mesh file located at:

```
inputFiles/lagrangianContactMechanics/ContactMechanics_SingleFracCompression_base.xml
```

```
inputFiles/lagrangianContactMechanics/ContactMechanics_SingleFracCompression_
↪benchmark.xml
```

```
inputFiles/lagrangianContactMechanics/crackInPlane_benchmark.vtu
```

## Description of the case

We simulate an inclined fracture under a compressive horizontal stress ( $\sigma$ ), as shown below. This fracture is placed in an infinite, homogeneous, isotropic, and elastic medium. Uniaxial compression and frictional contact on the fracture surface cause mechanical deformation to the surrounding rock and sliding along the fracture plane. For verification purposes, plane strain deformation and Coulomb failure criterion are considered in this numerical model.

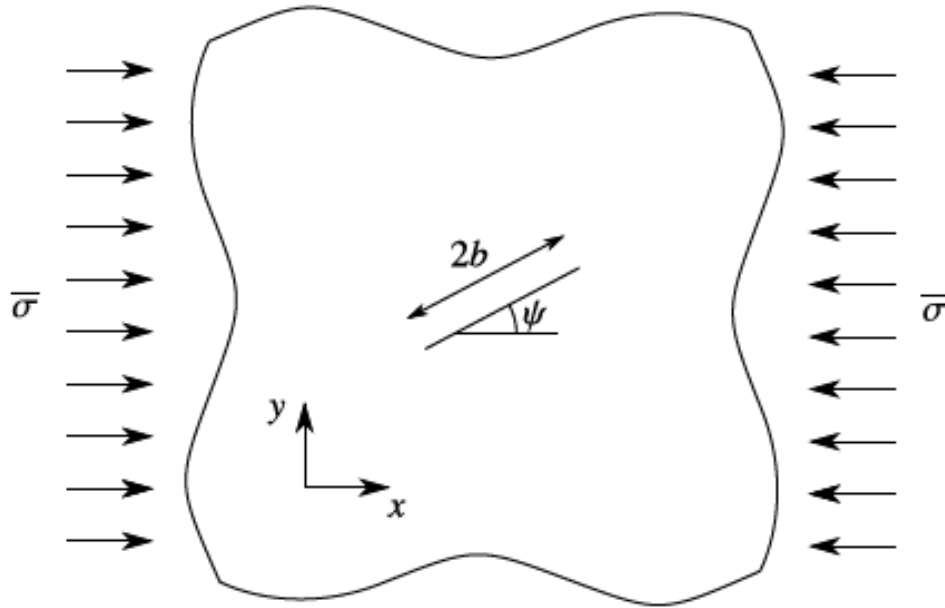


Fig. 4.8: Sketch of the problem

To simulate this phenomenon, we use a Lagrange contact model. Displacement and stress fields on the fracture plane are calculated numerically. Predictions of the normal traction ( $t_N$ ) and slip ( $g_T$ ) on the fracture surface are compared with the corresponding analytical solution (Phan et al., 2003).

$$t_N = -\sigma(\sin(\psi))^2$$

$$g_T = \frac{4(1-\nu^2)}{E}(\sigma \sin(\psi)(\cos(\psi) - \sin(\psi) \tan(\theta))) \sqrt{b^2 - (b-\xi)^2}$$

where  $\psi$  is the inclination angle,  $\nu$  is Poisson's ratio,  $E$  is Young's modulus,  $\theta$  is the friction angle,  $b$  is the fracture half-length,  $\xi$  is a local coordinate on the fracture varying in the range  $[0, 2b]$ .

In this example, we focus our attention on the Mesh tags, the Constitutive tags, and the FieldSpecifications tags.

## Mesh

The following figure shows the mesh used in this problem.

Here, we load the mesh with PAMELAMesh (see [Importing the Mesh](#)). The syntax to import external meshes is simple: in the XML file, the mesh file `crackInPlane_benchmark.vtu` is included with its relative or absolute path to the location of the GEOSX XML file and a user-specified label (here `CubeHex`) is given to the mesh object. This unstructured mesh contains quadrilaterals elements and interface elements. Refinement is performed to conform with the fracture geometry specified in the `Geometry` section.

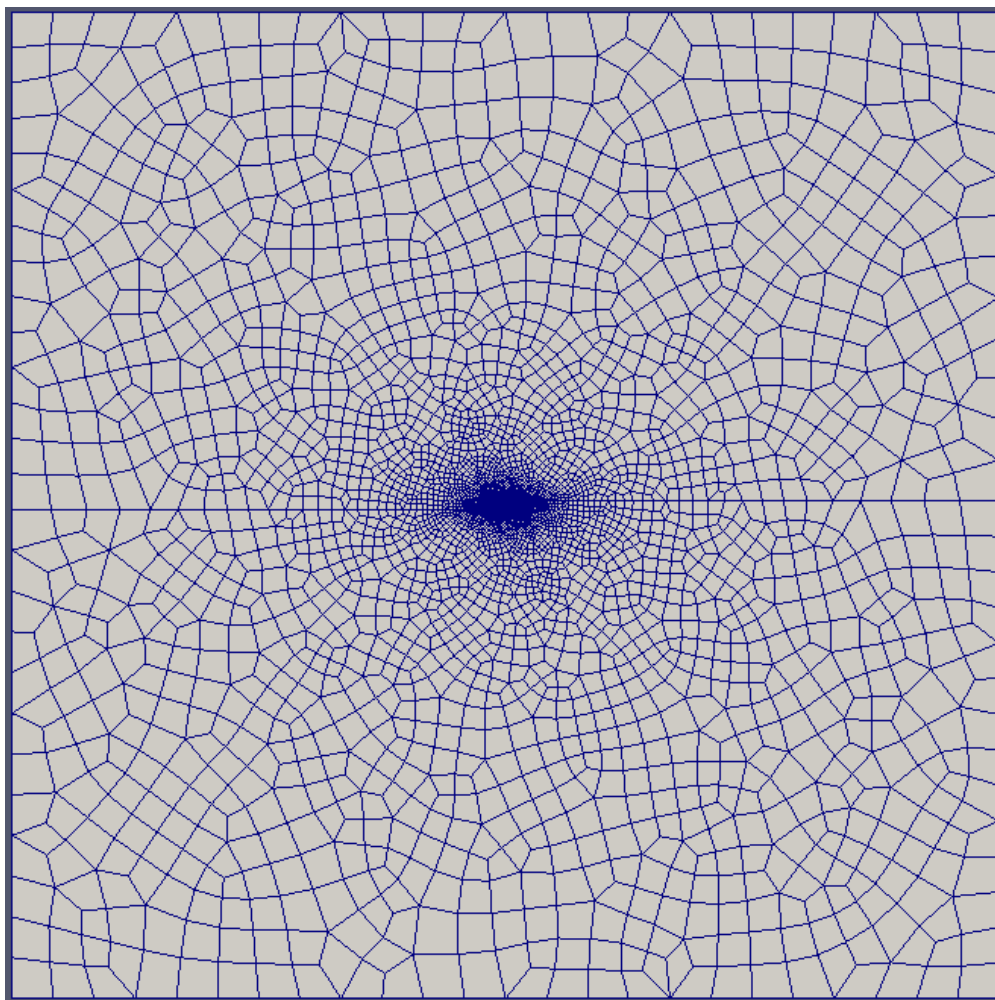


Fig. 4.9: Imported mesh

```

<Mesh>
  <VTKMesh
    name="CubeHex"
    file="crackInPlane_benchmark.vtu"/>
  </Mesh>

<Geometry>
  <BoundedPlane
    name="fracture"
    normal="{ -0.342020143325669, 0.939692620785908, 0.0 }"
    origin="{ 0.0, 0.0, 0.0 }"
    lengthVector="{ 0.939692620785908, 0.342020143325669, 0.0 }"
    widthVector="{ 0.0, 0.0, 1.0 }"
    dimensions="{ 2, 10 }"/>

  <BoundedPlane
    name="core"
    normal="{ -0.342020143325669, 0.939692620785908, 0.0 }"
    origin="{ 0.0, 0.0, 0.0 }"
    lengthVector="{ 0.939692620785908, 0.342020143325669, 0.0 }"
    widthVector="{ 0.0, 0.0, 1.0 }"
    dimensions="{ 2, 10 }"/>

  <Box
    name="rightPoint"
    xMin="{ 39.9, -40.1, -0.001 }"
    xMax="{ 40.1, 40.1, 0.051 }"/>

  <Box
    name="leftPoint"
    xMin="{ -40.1, -40.1, -0.001 }"
    xMax="{ -39.9, 40.1, 0.051 }"/>

  <Box
    name="topPoint"
    xMin="{ -40.1, 39.9, -0.001 }"
    xMax="{ 40.1, 40.1, 0.051 }"/>

  <Box
    name="bottomPoint"
    xMin="{ -40.1, -40.1, -0.001 }"
    xMax="{ 40.1, -39.9, 0.051 }"/>

  <Box
    name="front"
    xMin="{ -40.1, -40.1, -0.001 }"
    xMax="{ 40.1, 40.1, 0.001 }"/>

  <Box
    name="rear"
    xMin="{ -40.1, -40.1, 0.049 }"
    xMax="{ 40.1, 40.1, 0.051 }"/>

  <Box
    name="xmin"
    xMin="{ -40.1, -40.1, -0.001 }"
    xMax="{ -39.9, 40.1, 0.051 }"/>

```

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```

<Box
  name="xmax"
  xMin="{39.9, -40.1, -0.001}"
  xMax="{40.1, 40.1, 0.051}" />
</Geometry>

```

## Solid mechanics solver

GEOSX is a multi-physics platform. Different combinations of physics solvers available in the code can be applied in different regions of the domain and be functional at different stages of the simulation. The `Solvers` tag in the XML file is used to list and parameterize these solvers.

To specify a coupling between two different solvers, we define and characterize each single-physics solver separately. Then, we customize a *coupling solver* between these single-physics solvers as an additional solver. This approach allows for generality and flexibility in constructing multi-physics solvers. Each single-physics solver should be given a meaningful and distinct name because GEOSX recognizes these single-physics solvers based on their given names to create the coupling.

To setup a coupling between rock and fracture deformations, we define three different solvers:

- For solving the frictional contact, we define a Lagrangian contact solver, called here `lagrangiancontact`. In this solver, we specify `targetRegions` that includes both the continuum region `Region` and the discontinuum region `Fracture` where the solver is applied to couple rock and fracture deformation. The contact constitutive law used for the fracture elements is named `fractureMaterial`, and defined later in the Constitutive section.
- Rock deformations are handled by a solid mechanics solver `SolidMechanics_LagrangianFEM`. This solid mechanics solver (see *Solid Mechanics Solver*) is based on the Lagrangian finite element formulation. The problem is run as `QuasiStatic` without considering inertial effects. The computational domain is discretized by `FE1`, which is defined in the `NumericalMethods` section. The solid material is named `rock`, and its mechanical properties are specified later in the `Constitutive` section.
- The solver `SurfaceGenerator` defines the fracture region and rock toughness.

```

<Solvers
  gravityVector="{0.0, 0.0, 0.0}">
  <LagrangianContact
    name="lagrangiancontact"
    solidSolverName="lagsolve"
    stabilizationName="TPFAstabilization"
    logLevel="1"
    discretization="FE1"
    targetRegions="{ Region, Fracture }"
    contactRelationName="fractureMaterial"
    fractureRegionName="Fracture">
    <NonlinearSolverParameters
      newtonTol="1.0e-8"
      logLevel="2"
      newtonMaxIter="10"
      maxNumConfigurationAttempts="10"
      lineSearchAction="Require"
      lineSearchMaxCuts="2"
      maxTimeStepCuts="2" />
    <LinearSolverParameters

```

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```

        solverType="direct"
        directParallel="0"
        logLevel="0"/>
    </LagrangianContact>

    <SolidMechanics_LagrangianFEM
        name="lagsolve"
        timeIntegrationOption="QuasiStatic"
        logLevel="0"
        discretization="FE1"
        targetRegions="{ Region, Fracture }">
        <NonlinearSolverParameters
            newtonTol="1.0e-6"
            newtonMaxIter="5"/>
        <LinearSolverParameters
            krylovTol="1.0e-10"
            logLevel="0"/>
    </SolidMechanics_LagrangianFEM>

    <SurfaceGenerator
        name="SurfaceGen"
        logLevel="0"
        fractureRegion="Fracture"
        targetRegions="{ Region }"
        rockToughness="1.0e6"
        mpiCommOrder="1"/>
</Solvers>

```

## Constitutive laws

For this specific problem, we simulate the elastic deformation and fracture slippage caused by uniaxial compression. A homogeneous and isotropic domain with one solid material is assumed, with mechanical properties specified in the Constitutive section.

Fracture surface slippage is assumed to be governed by the Coulomb failure criterion. The contact constitutive behavior is named `fractureMaterial` in the `Coulomb` block, where cohesion `cohesion="0.0"` and friction coefficient `frictionCoefficient="0.577350269"` are specified.

```

<Constitutive>
    <ElasticIsotropic
        name="rock"
        defaultDensity="2700"
        defaultBulkModulus="16.666666666666666e9"
        defaultShearModulus="1.0e10"/>

    <Coulomb
        name="fractureMaterial"
        cohesion="0.0"
        frictionCoefficient="0.577350269"
        apertureTableName="apertureTable"/>
</Constitutive>

```

Recall that in the `SolidMechanics_LagrangianFEM` section, `rock` is the material of the computational domain. Here, the isotropic elastic model `ElasticIsotropic` is used to simulate the mechanical behavior of `rock`.

All constitutive parameters such as density, bulk modulus, and shear modulus are specified in the International System



of Units.

## Time history function

In the Tasks section, PackCollection tasks are defined to collect time history information from fields. Either the entire field or specified named sets of indices in the field can be collected. In this example, tractionCollection and displacementJumpCollection tasks are specified to output the local traction `fieldName="traction"` and relative displacement `fieldName="displacementJump"` on the fracture surface.

```
<Tasks>
  <PackCollection
    name="tractionCollection"
    objectPath="ElementRegions/Fracture/faceElementSubRegion"
    fieldName="traction"/>

  <PackCollection
    name="displacementJumpCollection"
    objectPath="ElementRegions/Fracture/faceElementSubRegion"
    fieldName="displacementJump"/>
</Tasks>
```

These two tasks are triggered using the Event management, with PeriodicEvent defined for these recurring tasks. GEOSX writes two files named after the string defined in the filename keyword and formatted as HDF5 files (displacementJump\_history.hdf5 and traction\_history.hdf5). The TimeHistory file contains the collected time history information from each specified time history collector. This information includes datasets for the simulation time, element center defined in the local coordinate system, and the time history information. Then, a Python script is used to access and plot any specified subset of the time history data for verification and visualization.

## Initial and boundary conditions

The next step is to specify fields, including:

- The initial value (the remote compressive stress needs to be initialized),
- The boundary conditions (the constraints of the outer boundaries have to be set).

In this tutorial, we specify an uniaxial horizontal stress ( $\sigma_x = -1.0e8$  Pa). The remaining parts of the outer boundaries are subjected to roller constraints. These boundary conditions are set up through the FieldSpecifications section.

```
<FieldSpecifications>
  <FieldSpecification
    name="frac"
    initialCondition="1"
    setNames="{ fracture }"
    objectPath="faceManager"
    fieldName="ruptureState"
    scale="1"/>

  <FieldSpecification
    name="separableFace"
    initialCondition="1"
    setNames="{ core }"
    objectPath="faceManager"
```

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```

    fieldName="isFaceSeparable"
    scale="1"/>

<FieldSpecification
  name="xconstraint"
  objectPath="nodeManager"
  fieldName="totalDisplacement"
  component="0"
  scale="0.0"
  setNames="{ leftPoint, rightPoint }"/>

<FieldSpecification
  name="yconstraint"
  objectPath="nodeManager"
  fieldName="totalDisplacement"
  component="1"
  scale="0.0"
  setNames="{ bottomPoint, topPoint }"/>

<FieldSpecification
  name="zconstraint"
  objectPath="nodeManager"
  fieldName="totalDisplacement"
  component="2"
  scale="0.0"
  setNames="{ front, rear }"/>

<FieldSpecification
  name="Sigmax"
  initialCondition="1"
  setNames="{ all }"
  objectPath="ElementRegions/Region"
  fieldName="rock_stress"
  component="0"
  scale="-1.0e8"/>
</FieldSpecifications>

```

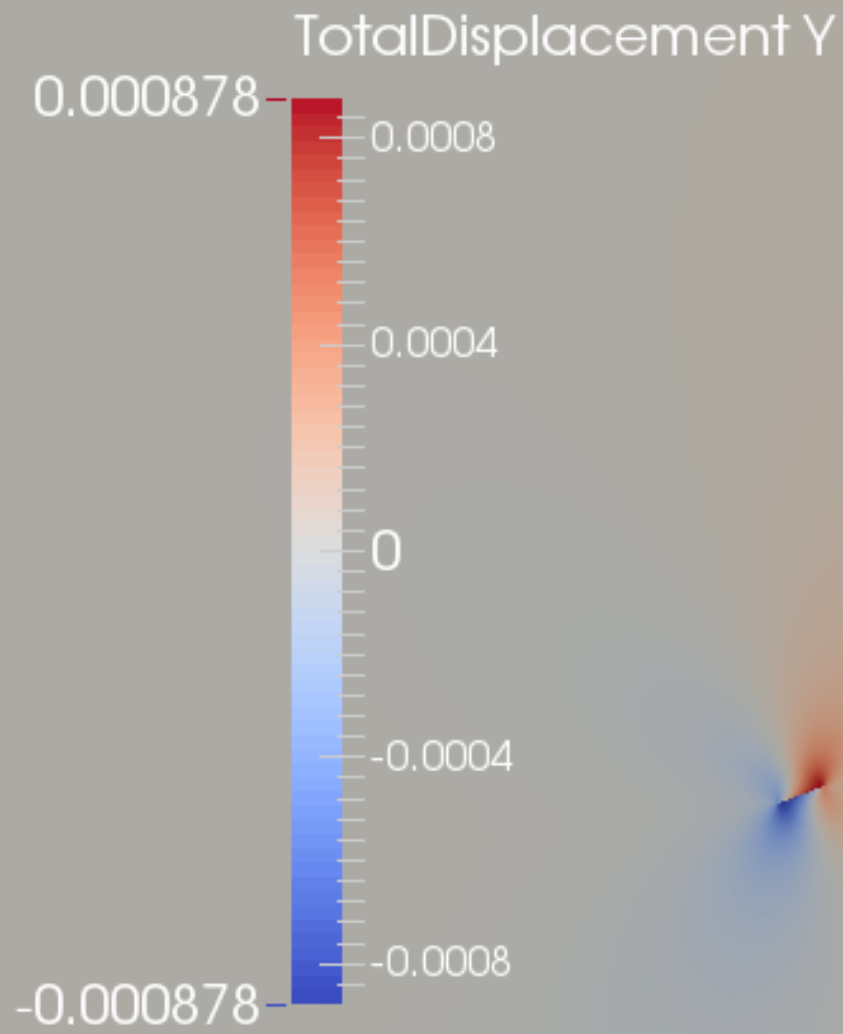
Note that the remote stress has a negative value, due to the negative sign convention for compressive stresses in GEOSX.

The parameters used in the simulation are summarized in the following table.

Symbol	Parameter	Unit	Value
$K$	Bulk Modulus	[GPa]	16.67
$G$	Shear Modulus	[GPa]	10.0
$\sigma$	Compressive Stress	[MPa]	-100.0
$\theta$	Friction Angle	[Degree]	30.0
$\psi$	Inclination Angle	[Degree]	20.0
$b$	Fracture Half Length	[m]	1.0

## Inspecting results

We request VTK-format output files and use Paraview to visualize the results. The following figure shows the distribution of  $u_y$  in the computational domain.



The next figure shows the distribution of relative shear displacement values on the fracture surface.

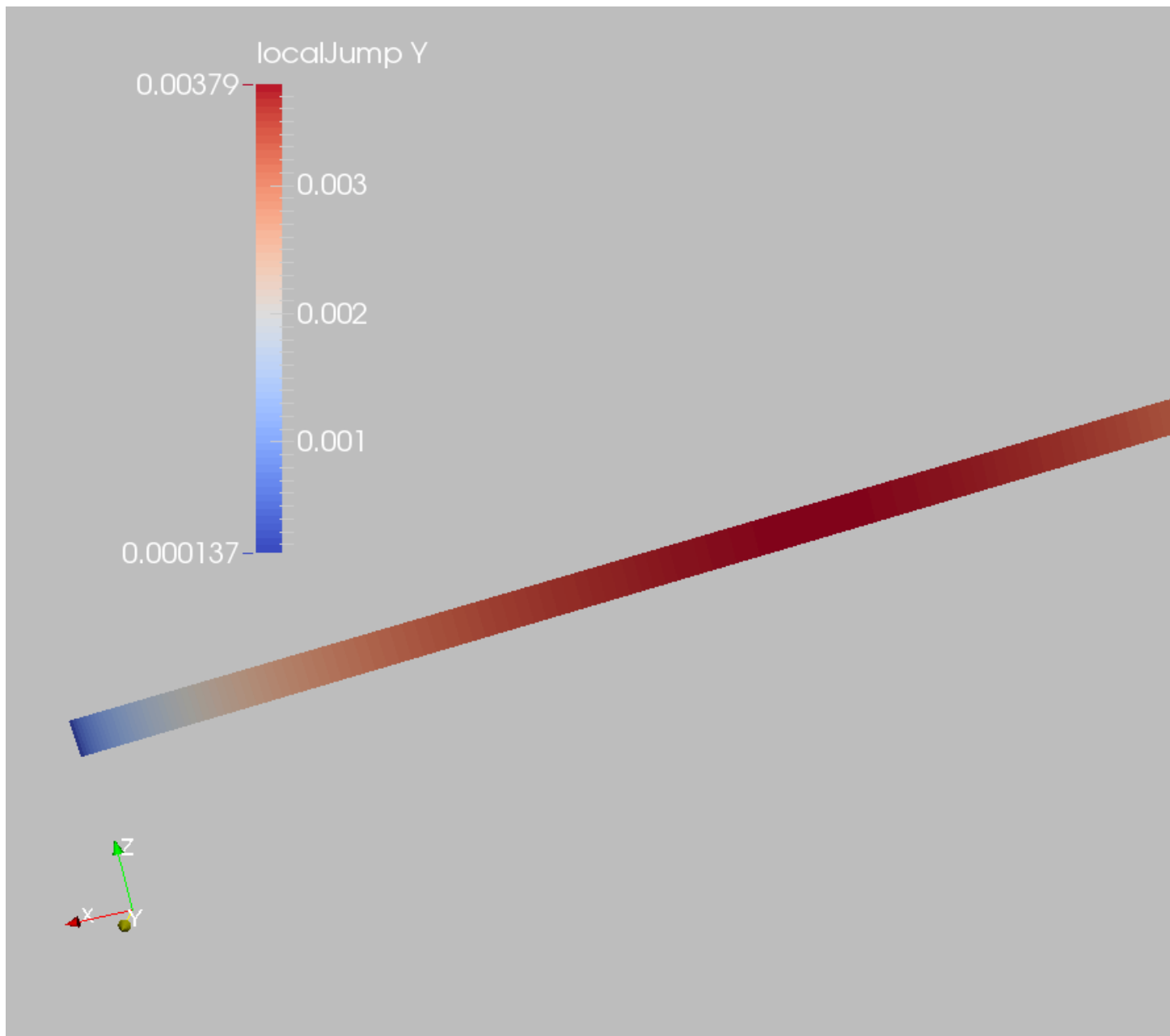


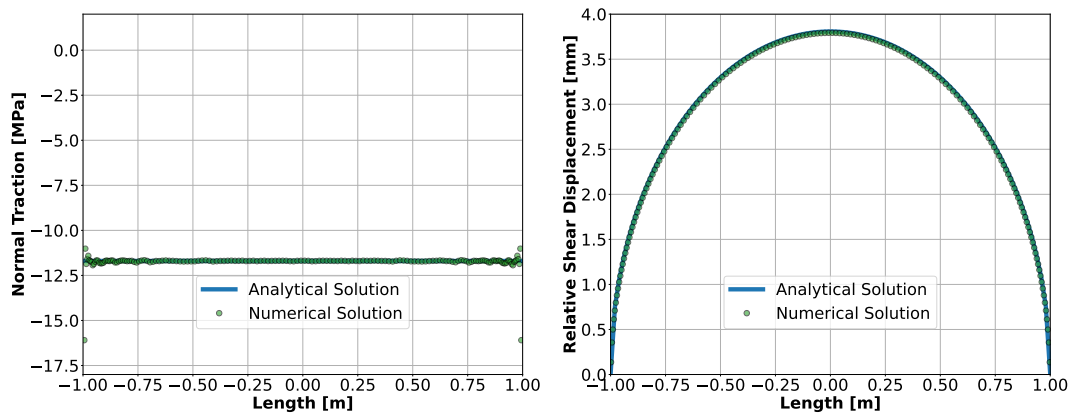
Fig. 4.11: Simulation result of fracture slip

The figure below shows a comparison between the numerical predictions (marks) and the corresponding analytical solutions (solid curves) for the normal traction ( $t_N$ ) and slip ( $g_T$ ) distributions on the fracture surface. One can observe that the numerical results obtained by GEOSX and the analytical solutions are nearly identical.

### To go further

#### Feedback on this example

For any feedback on this example, please submit a [GitHub issue on the project's GitHub page](#).



## Fracture Intersection Problem

### Context

In this example, two fractures intersecting at a right angle are simulated using a Lagrange contact model in a 2D infinite domain and subjected to a constant uniaxial compressive remote stress. Numerical solutions based on the symmetric-Galerkin boundary element method (Phan et al., 2003) is used to verify the accuracy of the GEOSX results for the normal traction, normal opening, and shear slippage on the fracture surfaces, considering frictional contact and fracture-fracture interaction. In this example, the `TimeHistory` function and a Python script are used to output and post-process multi-dimensional data (traction and displacement on the fracture surfaces).

### Input file

Everything required is contained within two xml files located at:

```
inputFiles/lagrangianContactMechanics/ContactMechanics_TFrac_base.xml
```

```
inputFiles/lagrangianContactMechanics/ContactMechanics_TFrac_benchmark.xml
```

## Description of the case

We simulate two intersecting fractures under a remote compressive stress constraint, as shown below. The two fractures sit in an infinite, homogeneous, isotropic, and elastic medium. The vertical fracture is internally pressurized and perpendicularly intersects the middle of the horizontal fracture. A combination of uniaxial compression, frictional contact, and opening of the vertical fracture causes mechanical deformations of the surrounding rock, thus leads to sliding of the horizontal fracture. For verification purposes, a plane strain deformation and Coulomb failure criterion are considered in this numerical model.

To simulate this problem, we use a Lagrange contact model. Displacement and stress fields on the fracture plane are calculated numerically. Predictions of the normal traction and slip along the sliding fracture and mechanical aperture of the pressurized fracture are compared with the corresponding literature work (Phan et al., 2003).

For this example, we focus on the `Mesh`, the `Constitutive`, and the `FieldSpecifications` tags.

### Mesh

The following figure shows the mesh used in this problem.

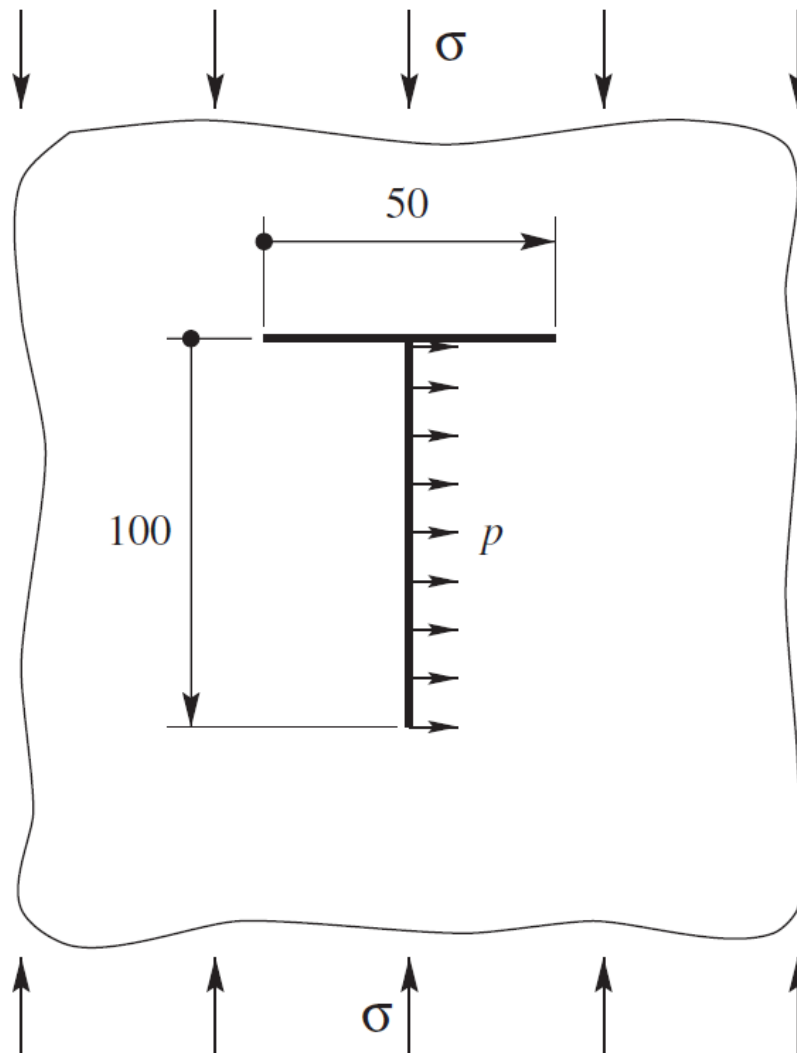


Fig. 4.12: Sketch of the problem (Phan et al., 2003)

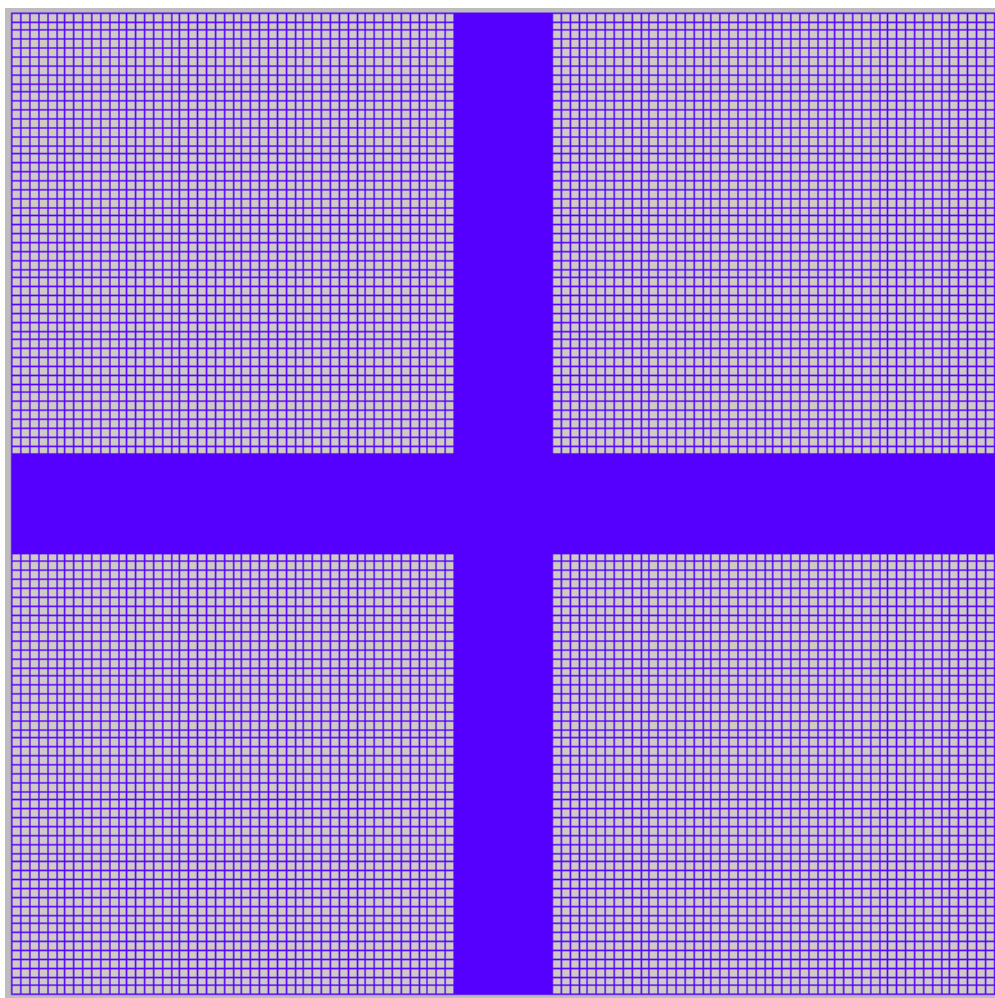


Fig. 4.13: Generated mesh

This mesh was created using the internal mesh generator as parametrized in the `InternalMesh` XML tag. The mesh contains 300 x 300 x 1 eight-node brick elements in the x, y, and z directions respectively. Such eight-node hexahedral elements are defined as `C3D8` `elementTypes`, and their collection forms a mesh with one group of cell blocks named here `cb1`.

```
<Mesh>
  <InternalMesh
    name="mesh1"
    elementTypes="{ C3D8 }"
    xCoords="{ -1000, -100, 100, 1000 }"
    yCoords="{ -1000, -100, 100, 1000 }"
    zCoords="{ 0, 1 }"
    nx="{ 50, 200, 50 }"
    ny="{ 50, 200, 50 }"
    nz="{ 1 }"
    cellBlockNames="{ cb1 }"/>
  </InternalMesh>
</Mesh>
```

Refinement is necessary to conform with the fracture geometry specified in the `Geometry` section.

```
<Geometry>
  <BoundedPlane
    name="fracture1"
    normal="{1.0, 0.0, 0.0}"
    origin="{0.0, 0.0, 0.0}"
    lengthVector="{0.0, 1.0, 0.0}"
    widthVector="{0.0, 0.0, 1.0}"
    dimensions="{ 100, 10 }"/>

  <BoundedPlane
    name="core1"
    normal="{1.0, 0.0, 0.0}"
    origin="{0.0, 0.0, 0.0}"
    lengthVector="{0.0, 1.0, 0.0}"
    widthVector="{0.0, 0.0, 1.0}"
    dimensions="{ 100, 10 }"/>

  <BoundedPlane
    name="fracture2"
    normal="{0.0, 1.0, 0.0}"
    origin="{0.0, 50.0, 0.0}"
    lengthVector="{1.0, 0.0, 0.0}"
    widthVector="{0.0, 0.0, 1.0}"
    dimensions="{ 50, 10 }"/>

  <BoundedPlane
    name="core2"
    normal="{0.0, 1.0, 0.0}"
    origin="{0.0, 50.0, 0.0}"
    lengthVector="{1.0, 0.0, 0.0}"
    widthVector="{0.0, 0.0, 1.0}"
    dimensions="{ 50, 10 }"/>
</BoundedPlane>
</Geometry>
```



## Solid mechanics solver

GEOSX is a multiphysics simulation platform. Different combinations of physics solvers can be applied in different regions of the domain at different stages of the simulation. The `Solvers` tag in the XML file is used to list and parameterize these solvers.

To specify a coupling between two different solvers, we define and characterize each single-physics solver separately. Then, we customize a *coupling solver* between these single-physics solvers as an additional solver. This approach allows for generality and flexibility in constructing multiphysics solvers. Each single-physics solver should be given a meaningful and distinct name, because GEOSX recognizes these single-physics solvers by their given names to create the coupling.

To setup a coupling between rock and fracture deformations, we define three different solvers:

- For solving the frictional contact, we define a Lagrangian contact solver, called here `lagrangiancontact`. In this solver, we specify `targetRegions` that include both the continuum region `Region` and the discontinuum region `Fracture` where the solver is applied to couple rock and fracture deformations. The contact constitutive law used for the fracture elements is named `fractureMaterial`, and is defined later in the `Constitutive` section.
- Rock deformations are handled by a solid mechanics solver `SolidMechanics_LagrangianFEM`. This solid mechanics solver (see *SolidMechanicsLagrangianFEM*) is based on the Lagrangian finite element formulation. The problem runs in `QuasiStatic` mode without inertial effects. The computational domain is discretized by `FE1`, which is defined in the `NumericalMethods` section. The solid material is named `rock` and its mechanical properties are specified later in the `Constitutive` section.
- The solver `SurfaceGenerator` defines the fracture region and rock toughness.

```
<LagrangianContact
  name="lagrangiancontact"
  solidSolverName="lagsolve"
  stabilizationName="TPFAstabilization"
  logLevel="1"
  discretization="FE1"
  targetRegions="{ Region, Fracture }"
  contactRelationName="fractureMaterial"
  fractureRegionName="Fracture">
  <NonlinearSolverParameters
    newtonTol="1.0e-8"
    logLevel="2"
    maxNumConfigurationAttempts="10"
    newtonMaxIter="10"
    lineSearchAction="Require"
    lineSearchMaxCuts="2"
    maxTimeStepCuts="2"/>
  <LinearSolverParameters
    solverType="direct"
    directParallel="0"
    logLevel="0"/>
</LagrangianContact>

<SolidMechanics_LagrangianFEM
  name="lagsolve"
  timeIntegrationOption="QuasiStatic"
  logLevel="0"
  discretization="FE1"
  targetRegions="{ Region, Fracture }">
  <NonlinearSolverParameters
```

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```

    newtonTol="1.0e-6"
    newtonMaxIter="5"/>
  <LinearSolverParameters
    krylovTol="1.0e-10"
    logLevel="0"/>
</SolidMechanics_LagrangianFEM>

<SurfaceGenerator
  name="SurfaceGen"
  logLevel="0"
  fractureRegion="Fracture"
  targetRegions="{ Region }"
  rockToughness="1.0e6"
  mpiCommOrder="1"/>

```

## Constitutive laws

For this problem, we simulate the elastic deformation and fracture slippage caused by the uniaxial compression. A homogeneous and isotropic domain with one solid material is assumed, and its mechanical properties are specified in the Constitutive section.

Fracture surface slippage is assumed to be governed by the Coulomb failure criterion. The contact constitutive behavior is named `fractureMaterial` in the Coulomb block, where cohesion `cohesion="0.0"` and friction coefficient `frictionCoefficient="0.577350269"` are specified.

```

<Constitutive>
  <ElasticIsotropic
    name="rock"
    defaultDensity="2700"
    defaultBulkModulus="38.89e9"
    defaultShearModulus="29.17e9"/>

  <Coulomb
    name="fractureMaterial"
    cohesion="0.0"
    frictionCoefficient="0.577350269"
    apertureTableName="apertureTable"/>
</Constitutive>

```

Recall that in the `SolidMechanics_LagrangianFEM` section, `rock` is the material of the computational domain. Here, the isotropic elastic model `ElasticIsotropic` is used to simulate the mechanical behavior of `rock`.

All constitutive parameters such as density, bulk modulus, and shear modulus are specified in the International System of Units.

## Time history function

In the Tasks section, `PackCollection` tasks are defined to collect time history information from fields. Either the entire field or specified named sets of indices in the field can be collected. In this example, `tractionCollection` and `displacementJumpCollection` tasks are specified to output the local traction `fieldName="traction"` and relative displacement `fieldName="displacementJump"` on the fracture surface.

```

<Tasks>
  <PackCollection
    name="tractionCollection"
    objectPath="ElementRegions/Fracture/faceElementSubRegion"
    fieldName="traction"/>

  <PackCollection
    name="displacementJumpCollection"
    objectPath="ElementRegions/Fracture/faceElementSubRegion"
    fieldName="displacementJump"/>
</Tasks>

```

These two tasks are triggered using the Event manager with a `PeriodicEvent` defined for these recurring tasks. GEOSX writes two files named after the string defined in the `filename` keyword and formatted as HDF5 files (`displacementJump_history.hdf5` and `traction_history.hdf5`). The TimeHistory file contains the collected time history information from each specified time history collector. This information includes datasets for the simulation time, element center defined in the local coordinate system, and the time history information. A Python script is used to read and plot any specified subset of the time history data for verification and visualization.

## Initial and boundary conditions

The next step is to specify fields, including:

- The initial value (the remote compressive stress needs to be initialized),
- The boundary conditions (traction loaded on the vertical fracture and the constraints of the outer boundaries have to be set).

In this tutorial, we specify an uniaxial vertical stress  $\text{SigmaY}$  ( $\sigma_y = -1.0\text{e}8$  Pa). A compressive traction `NormalTraction` ( $P_{in} = -1.0\text{e}8$  Pa) is applied at the surface of vertical fracture. The remaining parts of the outer boundaries are subjected to roller constraints. These boundary conditions are set up through the `FieldSpecifications` section.

```

<FieldSpecifications>
  <FieldSpecification
    name="frac"
    initialCondition="1"
    setNames="{ fracture1, fracture2 }"
    objectPath="faceManager"
    fieldName="ruptureState"
    scale="1"/>

  <FieldSpecification
    name="separableFace"
    initialCondition="1"
    setNames="{ core1, core2 }"
    objectPath="faceManager"
    fieldName="isFaceSeparable"
    scale="1"/>

  <FieldSpecification
    name="xconstraint"
    objectPath="nodeManager"
    fieldName="totalDisplacement"
    component="0"
    scale="0.0"

```

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```

    setNames="{ xpos, xneg }"/>

<FieldSpecification
  name="yconstraint"
  objectPath="nodeManager"
  fieldName="totalDisplacement"
  component="1"
  scale="0.0"
  setNames="{ ypos, yneg }"/>

<FieldSpecification
  name="zconstraint"
  objectPath="nodeManager"
  fieldName="totalDisplacement"
  component="2"
  scale="0.0"
  setNames="{ zpos, zneg }"/>

<Traction
  name="NormalTraction"
  objectPath="faceManager"
  tractionType="normal"
  scale="-1.0e8"
  functionName="ForceTimeFunction"
  setNames="{ core1 }"/>

<FieldSpecification
  name="SigmaY"
  initialCondition="1"
  setNames="{ all }"
  objectPath="ElementRegions/Region"
  fieldName="rock_stress"
  component="1"
  scale="-1.0e8"/>
</FieldSpecifications>

```

Note that the remote stress and internal fracture pressure has a negative value, due to the negative sign convention for compressive stresses in GEOSX.

The parameters used in the simulation are summarized in the following table.

Symbol	Parameter	Unit	Value
$K$	Bulk Modulus	[GPa]	38.89
$G$	Shear Modulus	[GPa]	29.17
$\sigma_y$	Remote Stress	[MPa]	-100.0
$P_{in}$	Internal Pressure	[MPa]	-100.0
$\theta$	Friction Angle	[Degree]	30.0
$L_h$	Horizontal Frac Length	[m]	50.0
$L_v$	Vertical Frac Length	[m]	100.0

## Inspecting results

We request VTK-format output files and use Paraview to visualize the results. The following figure shows the distribution of  $\sigma_{xx}$  in the computational domain.

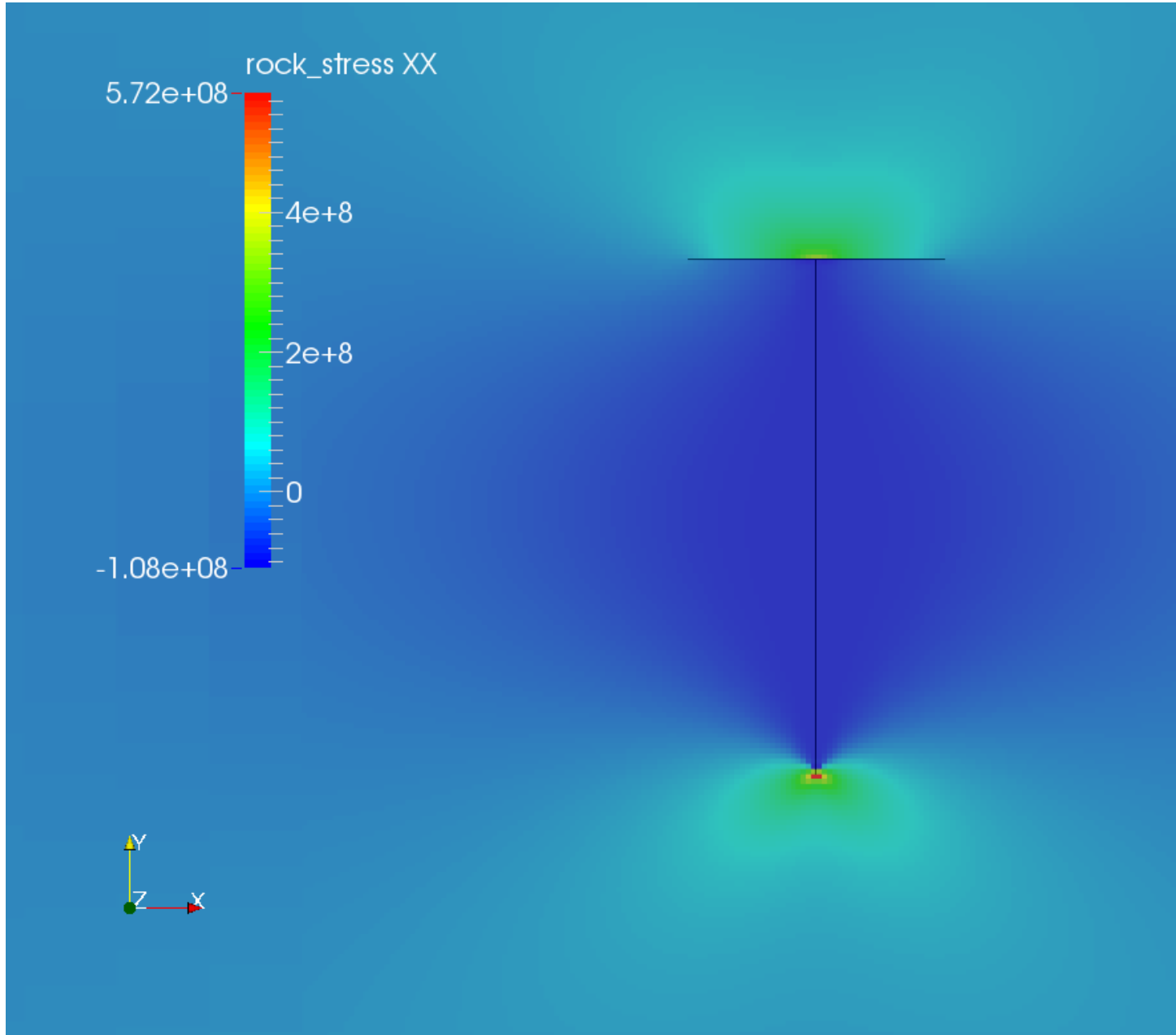


Fig. 4.14: Simulation result of  $\sigma_{xx}$

The next figure shows the distribution of relative shear displacement values along the surface of two intersected fractures.

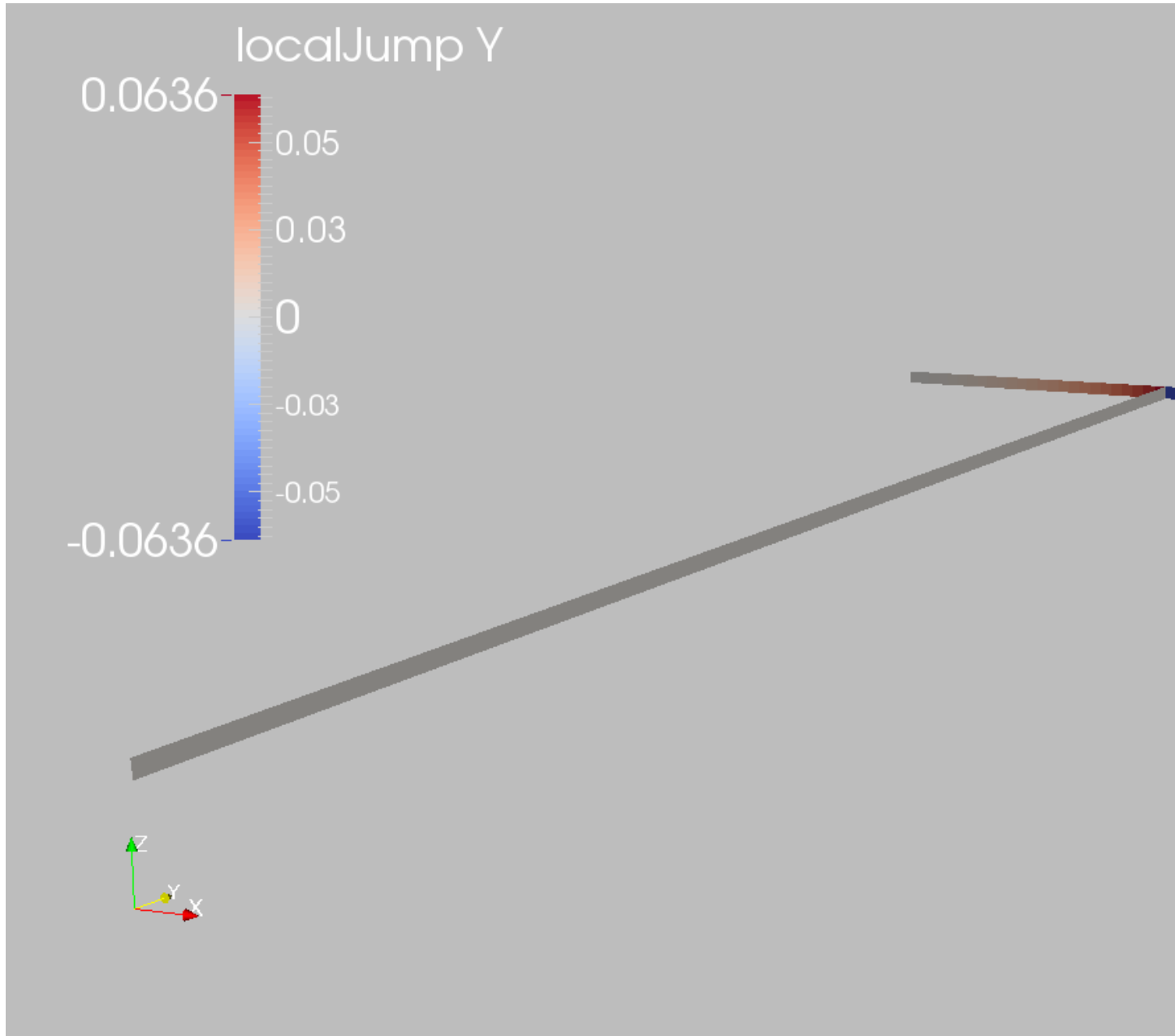
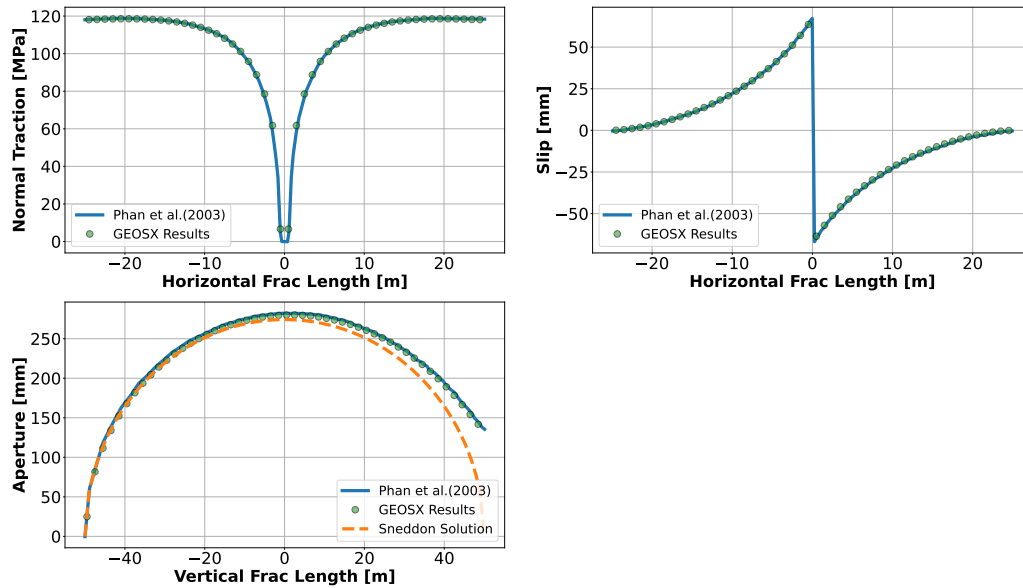


Fig. 4.15: Simulation result of fracture slip

The figure below compares the results from GEOSX (marks) and the corresponding literature reference solution (solid curves) for the normal traction and slip distributions along the horizontal fracture and opening of the vertical fracture. GEOSX reliably captures the mechanical interactions between two intersected fractures and shows excellent agreement with the reference solution. Due to sliding of the horizontal fracture, GEOSX prediction as well as the reference solution on the normal opening of pressurized vertical fracture deviates away from Sneddon's analytical solution, especially near the intersection point.



## To go further

### Feedback on this example

For any feedback on this example, please submit a [GitHub issue](#) on the project's GitHub page.

## Sneddon's Problem

### Objectives

At the end of this example you will know:

- how to define fractures in a porous medium,
- how to use various solvers (EmbeddedFractures, LagrangianContact and HydroFracture) to solve the mechanics problems with fractures.

### Input file

This example uses no external input files and everything required is contained within GEOSX input files.

The xml input files for the case with EmbeddedFractures solver are located at:

```
inputFiles/efemFractureMechanics/Sneddon_embeddedFrac_base.xml
inputFiles/efemFractureMechanics/Sneddon_embeddedFrac_benchmark.xml
```

The xml input files for the case with LagrangianContact solver are located at:

```
inputFiles/lagrangianContactMechanics/Sneddon_contactMechanics_base.xml
inputFiles/lagrangianContactMechanics/Sneddon_contactMechanics_benchmark.xml
```

The xml input files for the case with HydroFracture solver are located at:

```
inputFiles/hydraulicFracturing/Sneddon_hydroFrac_base.xml
inputFiles/hydraulicFracturing/Sneddon_hydroFrac_benchmark.xml
```

## Description of the case

We compute the displacement field induced by the presence of a pressurized fracture, of length  $L_f$ , in a porous medium.

GEOSX will calculate the displacement field in the porous matrix and the displacement jump at the fracture surface. We will use the analytical solution for the fracture aperture,  $w_n$  (normal component of the jump), to verify the numerical results

$$w_n(s) = \frac{4(1 - \nu^2)p_f}{E} \sqrt{\frac{L_f^2}{4} - s^2}$$

where -  $E$  is the Young's modulus -  $\nu$  is the Poisson's ratio -  $p_f$  is the fracture pressure -  $s$  is the local fracture coordinate in  $[-\frac{L_f}{2}, \frac{L_f}{2}]$

In this example, we focus our attention on the Solvers, the ElementRegions, and the Geometry tags.

## Mechanics solver

To define a mechanics solver capable of including embedded fractures, we will define two solvers:

- a `SolidMechanicsEmbeddedFractures` solver, called `mechSolve`
- a small-strain Lagrangian mechanics solver, of type `SolidMechanicsLagrangianSSLE` called here `matrixSolver` (see: [Solid Mechanics Solver](#))

Note that the `name` attribute of these solvers is chosen by the user and is not imposed by GEOSX. It is important to make sure that the `solidSolverName` specified in the embedded fractures solver corresponds to the small-strain Lagrangian solver used in the matrix.

The two single-physics solvers are parameterized as explained in their respective documentation, each with their own tolerances, verbosity levels, target regions, and other solver-specific attributes.

Additionally, we need to specify another solver of type, `EmbeddedSurfaceGenerator`, which is used to discretize the fracture planes.

```
<SolidMechanicsEmbeddedFractures
  name="mechSolve"
  targetRegions="{ Domain, Fracture }"
  fractureRegionName="Fracture"
  initialDt="10"
  solidSolverName="matrixSolver"
  contactRelationName="fractureContact"
  logLevel="1">
  <NonlinearSolverParameters
    newtonTol="1.0e-6"
    newtonMaxIter="2"
    maxTimeStepCuts="1"/>
  <LinearSolverParameters
    solverType="direct"
    directParallel="0"
    logLevel="0"/>
```

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```

</SolidMechanicsEmbeddedFractures>

<SolidMechanicsLagrangianSSLE
  name="matrixSolver"
  timeIntegrationOption="QuasiStatic"
  logLevel="1"
  discretization="FE1"
  targetRegions="{ Domain }"
/>

<EmbeddedSurfaceGenerator
  name="SurfaceGenerator"
  discretization="FE1"
  targetRegions="{ Domain, Fracture }"
  fractureRegion="Fracture"
  logLevel="1"
  mpiCommOrder="1"/>
</Solvers>

```

To setup a coupling between rock and fracture deformations in LagrangianContact solver, we define three different solvers:

- For solving the frictional contact, we define a Lagrangian contact solver, called here `lagrangiancontact`. In this solver, we specify `targetRegions` that include both the continuum region `Region` and the discontinuum region `Fracture` where the solver is applied to couple rock and fracture deformations. The contact constitutive law used for the fracture elements is named `fractureMaterial`, and is defined later in the Constitutive section.
- Rock deformations are handled by a solid mechanics solver `SolidMechanicsLagrangianSSLE`. The problem runs in `QuasiStatic` mode without inertial effects. The computational domain is discretized by `FE1`, which is defined in the NumericalMethods section. The solid material is named `rock` and its mechanical properties are specified later in the Constitutive section.
- The solver `SurfaceGenerator` defines the fracture region and rock toughness.

```

<LagrangianContact
  name="lagrangiancontact"
  solidSolverName="lagsolve"
  stabilizationName="TPFAstabilization"
  logLevel="1"
  discretization="FE1"
  targetRegions="{ Region, Fracture }"
  contactRelationName="fractureMaterial"
  fractureRegionName="Fracture">
  <NonlinearSolverParameters
    newtonTol="1.0e-8"
    logLevel="2"
    newtonMaxIter="10"
    maxNumConfigurationAttempts="10"
    lineSearchAction="Require"
    lineSearchMaxCuts="2"
    maxTimeStepCuts="2"/>
  <LinearSolverParameters
    solverType="direct"
    directParallel="0"
    logLevel="0"/>
</LagrangianContact>

```

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```

<SolidMechanicsLagrangianSSLE
  name="lagsolve"
  timeIntegrationOption="QuasiStatic"
  logLevel="0"
  discretization="FE1"
  targetRegions="{ Region, Fracture }"
>
  <NonlinearSolverParameters
    newtonTol="1.0e-6"
    newtonMaxIter="5"/>
    <LinearSolverParameters
      krylovTol="1.0e-10"
      logLevel="0"/>
  </SolidMechanicsLagrangianSSLE>

<SurfaceGenerator
  name="SurfaceGen"
  logLevel="0"
  fractureRegion="Fracture"
  targetRegions="{ Region }"
  rockToughness="1.0e6"
  mpiCommOrder="1"/>
</Solvers>

```

Three elementary solvers are combined in the solver `Hydrofracture` to model the coupling between fluid flow within the fracture, rock deformation, fracture opening/closure and propagation:

- Rock and fracture deformation are modeled by the solid mechanics solver `SolidMechanicsLagrangianSSLE`. In this solver, we define `targetRegions` that includes both the continuum region and the fracture region. The name of the contact constitutive behavior is also specified in this solver by the `contactRelationName`, besides the `solidMaterialNames`.
- The single phase fluid flow inside the fracture is solved by the finite volume method in the solver `SinglePhaseFVM`.
- The solver `SurfaceGenerator` defines the fracture region and rock toughness.

```

<Hydrofracture
  name="hydrofracture"
  solidSolverName="lagsolve"
  flowSolverName="SinglePhaseFlow"
  surfaceGeneratorName="SurfaceGen"
  couplingTypeOption="FIM"
  logLevel="1"
  targetRegions="{ Fracture }"
  contactRelationName="fractureContact"
  maxNumResolves="2">
    <NonlinearSolverParameters
      newtonTol="1.0e-5"
      newtonMaxIter="20"
      lineSearchMaxCuts="3"/>
    <LinearSolverParameters
      directParallel="0"/>
  </Hydrofracture>

  <SolidMechanicsLagrangianSSLE

```

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```

name="lagsolve"
timeIntegrationOption="QuasiStatic"
discretization="FE1"
targetRegions="{ Domain, Fracture }"
contactRelationName="fractureContact"/>

<SinglePhaseFVM
name="SinglePhaseFlow"
discretization="singlePhaseTPFA"
targetRegions="{ Fracture }"/>

<SurfaceGenerator
name="SurfaceGen"
targetRegions="{ Domain }"
nodeBasedSIF="1"
rockToughness="10.0e6"
mpiCommOrder="1"/>

```

## Events

For the case with EmbeddedFractures solver, we add multiple events defining solver applications:

- an event specifying the execution of the EmbeddedSurfaceGenerator to generate the fracture elements.
- a periodic event specifying the execution of the embedded fractures solver.
- three periodic events specifying the output of simulations results.

```

<Events
maxTime="1.0">
  <SoloEvent
name="preFracture"
target="/Solvers/SurfaceGenerator"/>

  <PeriodicEvent
name="solverApplications"
beginTime="0.0"
endTime="1.0"
forceDt="1.0"
target="/Solvers/mechSolve"/>

  <PeriodicEvent
name="outputs"
targetExactTimestep="0"
target="/Outputs/vtkOutput"/>

  <PeriodicEvent
name="timeHistoryCollection"
timeFrequency="1.0"
targetExactTimestep="0"
target="/Tasks/displacementJumpCollection" />

  <PeriodicEvent
name="timeHistoryOutput"
timeFrequency="1.0"
targetExactTimestep="0"

```

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```
target="/Outputs/timeHistoryOutput"/>
</Events>
```

Similar settings are applied for the other two cases.

## Mesh, material properties, and boundary conditions

Last, let us take a closer look at the geometry of this simple problem, if using EmbeddedFractures solver. We use the internal mesh generator to create a large domain ( $40\text{ m} \times 40\text{ m} \times 1\text{ m}$ ), with one single element along the Z axes, 121 elements along the X axis and 921 elements along the Y axis.

```
<Mesh>
  <InternalMesh
    name="mesh1"
    elementTypes="{ C3D8 }"
    xCoords="{ -20, -4, 4, 20 }"
    yCoords="{ -20, -4, 4, 20 }"
    zCoords="{ 0, 1 }"
    nx="{ 10, 101, 10 }"
    ny="{ 10, 901, 10 }"
    nz="{ 1 }"
    cellBlockNames="{ cb1 }"/>
  </InternalMesh>
</Mesh>
```

The mesh for the case with LagrangianContact solver was also created using the internal mesh generator, as parametrized in the InternalMesh XML tag. The mesh discretizes the same computational domain ( $40\text{ m} \times 40\text{ m} \times 1\text{ m}$ ) with  $300 \times 300 \times 1$  eight-node brick elements in the x, y, and z directions respectively.

```
<Mesh>
  <InternalMesh
    name="mesh1"
    elementTypes="{ C3D8 }"
    xCoords="{ -20, -2, 2, 20 }"
    yCoords="{ -20, -2, 2, 20 }"
    zCoords="{ 0, 1 }"
    nx="{ 40, 220, 40 }"
    ny="{ 40, 220, 40 }"
    nz="{ 1 }"
    cellBlockNames="{ cb1 }"/>
  </InternalMesh>
</Mesh>
```

Similarly, the internal mesh generator was used to discretize the same domain ( $40\text{ m} \times 40\text{ m} \times 1\text{ m}$ ) and generate the mesh for the case with Hydrofracture solver, which contains  $280 \times 280 \times 1$  eight-node brick elements in the x, y, and z directions.

```
<Mesh>
  <InternalMesh
    name="mesh1"
    elementTypes="{ C3D8 }"
    xCoords="{ -20, -2, 2, 20 }"
    yCoords="{ -20, -2, 2, 20 }"
    zCoords="{ 0, 1 }"
    nx="{ 40, 200, 40 }"
    ny="{ 40, 200, 40 }"
    nz="{ 1 }"
    cellBlockNames="{ cb1 }"/>
  </InternalMesh>
</Mesh>
```

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```

    nz="{ 1 }"
    cellBlockNames="{ cb1 }"/>
</Mesh>

```

In all the three cases, eight-node hexahedral elements are defined as C3D8 elementTypes, and their collection forms a mesh with one group of cell blocks named here cb1. Refinement is necessary to conform with the fracture geometry specified in the Geometry section.

The parameters used in the simulation are summarized in the following table.

Symbol	Parameter	Units	Value
$K$	Bulk modulus	[GPa]	16.7
$G$	Shear modulus	[GPa]	10.0
$L_f$	Fracture length	[m]	2.0
$p_f$	Fracture pressure	[MPa]	-2.0

Note that the internal fracture pressure has a negative value, due to the negative sign convention for compressive stresses in GEOSX.

Material properties and boundary conditions are specified in the Constitutive and FieldSpecifications sections.

## Adding a fracture

The static fracture is defined by a nodeset occupying a small region within the computation domain, where the fracture tends to open upon internal pressurization:

- The test case with EmbeddedFractures solver:

```

<Geometry>
  <BoundedPlane
    name="FracturePlane"
    normal="{1.0, 0.0, 0.0}"
    origin="{0.0, 0.0, 0.0}"
    lengthVector="{0.0, 1.0, 0.0}"
    widthVector="{0.0, 0.0, 1.0}"
    dimensions="{ 2, 10 }"/>
  </BoundedPlane>
</Geometry>

```

- The test case with LagrangianContact solver:

```

<Geometry>
  <BoundedPlane
    name="fracture"
    normal="{1.0, 0.0, 0.0}"
    origin="{0.0, 0.0, 0.0}"
    lengthVector="{0.0, 1.0, 0.0}"
    widthVector="{0.0, 0.0, 1.0}"
    dimensions="{ 2, 10 }"/>

  <BoundedPlane
    name="core"
    normal="{1.0, 0.0, 0.0}"
    origin="{0.0, 0.0, 0.0}"

```

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```
lengthVector="{0.0, 1.0, 0.0}"
widthVector="{0.0, 0.0, 1.0}"
dimensions="{ 2, 10 }"/>
</Geometry>
```

- The test case with HydroFracture solver:

```
<Geometry>
  <Box
    name="fracture"
    xMin="{ -0.01, -1.01, -0.01 }"
    xMax="{ 0.01, 1.01, 1.01 }"/>

    <Box
      name="source"
      xMin="{ -0.01, -0.11, -0.01 }"
      xMax="{ 0.01, 0.11, 1.01 }"/>

    <Box
      name="core"
      xMin="{ -0.01, -10.01, -0.01 }"
      xMax="{ 0.01, 10.01, 1.01 }"/>
  </Geometry>
```

To make these cases identical to the analytical example, fracture propagation is not allowed in this example.

## Time history function

In the Tasks section, PackCollection tasks are defined to collect time history information from fields. Either the entire field or specified named sets of indices in the field can be collected. In this example, a task is specified to output fracture aperture (normal opening); however, for different solvers, different `fieldName` and `objectPath` should be called:

- The test case with EmbeddedFractures solver:

```
<Tasks>
  <PackCollection
    name="displacementJumpCollection"
    objectPath="ElementRegions/Fracture/embeddedSurfaceSubRegion"
    fieldName="displacementJump"/>
</Tasks>
```

- The test case with LagrangianContact solver:

```
<Tasks>
  <PackCollection
    name="displacementJumpCollection"
    objectPath="ElementRegions/Fracture/faceElementSubRegion"
    fieldName="displacementJump"/>
</Tasks>
```

- The test case with Hydrofracture solver:

```
<Tasks>
  <PackCollection
```

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```

name="apertureCollection"
objectPath="ElementRegions/Fracture/faceElementSubRegion"
fieldName="elementAperture"/>
</Tasks>

```

These tasks are triggered using the Event manager with a `PeriodicEvent` defined for these recurring tasks. GEOSX writes output files named after the string defined in the `filename` keyword and formatted as HDF5 files. The `TimeHistory` file contains the collected time history information from each specified time history collector. This information includes datasets for the simulation time, element center defined in the local coordinate system, and the time history information. A Python script is used to read and plot any specified subset of the time history data for verification and visualization.

## Running GEOSX

To run these three cases, use the following commands:

```

path/to/geosx -i inputFiles/efemFractureMechanics/Sneddon_embeddedFrac_benchmark.xml
path/to/geosx -i inputFiles/lagrangianContactMechanics/Sneddon_contactMechanics_benchmark.xml
path/to/geosx -i inputFiles/hydraulicFracturing/Sneddon_hydroFrac_benchmark.xml

```

## Inspecting results

This plot compares the analytical solution (continuous lines) with the numerical solutions (markers) for the normal opening of the pressurized fracture. As shown below, consistently, numerical solutions with different solvers correlate very well with the analytical solution.

## To go further

### Feedback on this example

This concludes the Sneddon example. For any feedback on this example, please submit a [GitHub issue on the project's GitHub page](#).

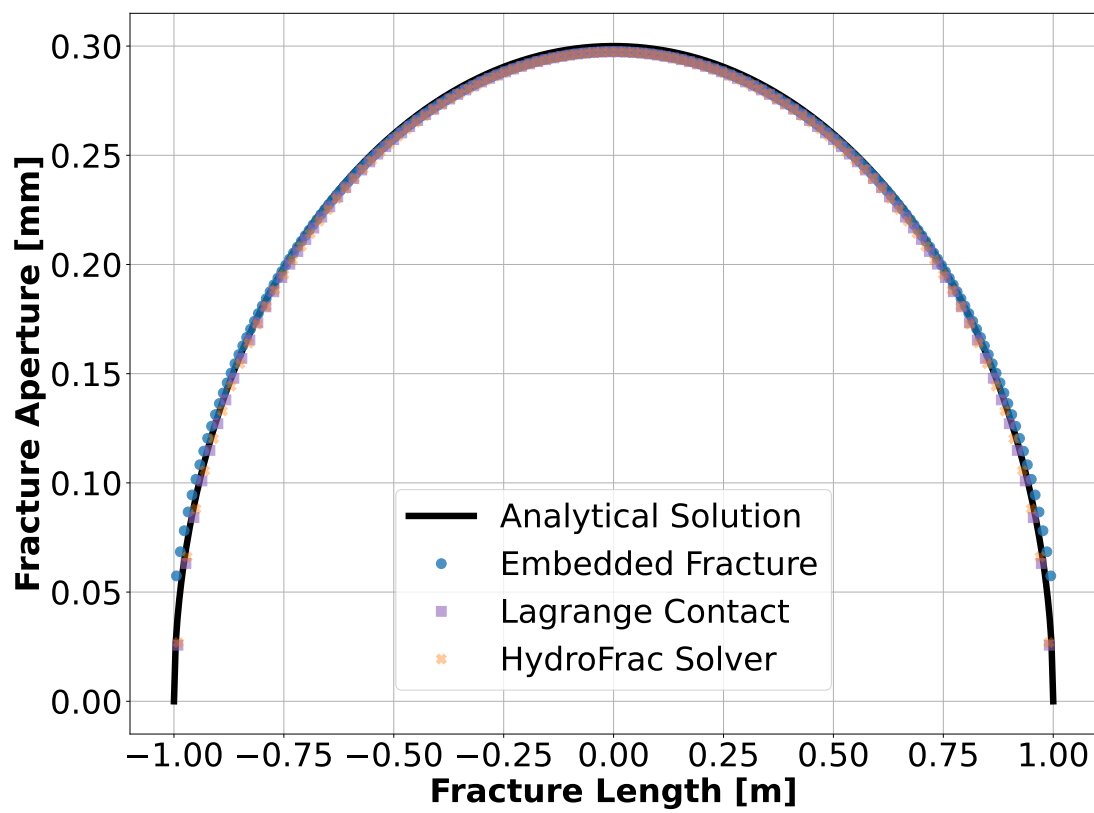
## Mandel's Problem

### Context

In this example, we use the coupled solvers in GEOSX to solve Mandel's 2D consolidation problem, a classic benchmark in poroelasticity. The analytical solution ([Cheng and Detournay, 1988](#)) is employed to verify the accuracy of the modeling predictions on induced pore pressure and the corresponding settlement. In this example, the `TimeHistory` function and a Python script are used to output and post-process multi-dimensional data (pore pressure and displacement field).

### Input file

This example uses no external input files and everything required is contained within two GEOSX input files located at:





```
inputFiles/poromechanics/PoroElastic_Mandel_base.xml
```

```
inputFiles/poromechanics/PoroElastic_Mandel_benchmark.xml
```

### Description of the case

We simulate the consolidation of a poroelastic slab between two rigid and impermeable plates subjected to a constant normal force. The slab is assumed to be fully saturated, homogeneous, isotropic, and infinitely long in the y-direction. We apply a uniform compressive load in the vertical direction. This force leads to a change of pore pressure and mechanical deformations of the sample, evolving with time due to fluid diffusion and coupling effects. The numerical model represents a plane strain deformation and lateral drainage without confinement, showing only a quarter of the computational domain in the x-z plane (the rest follows by symmetry).

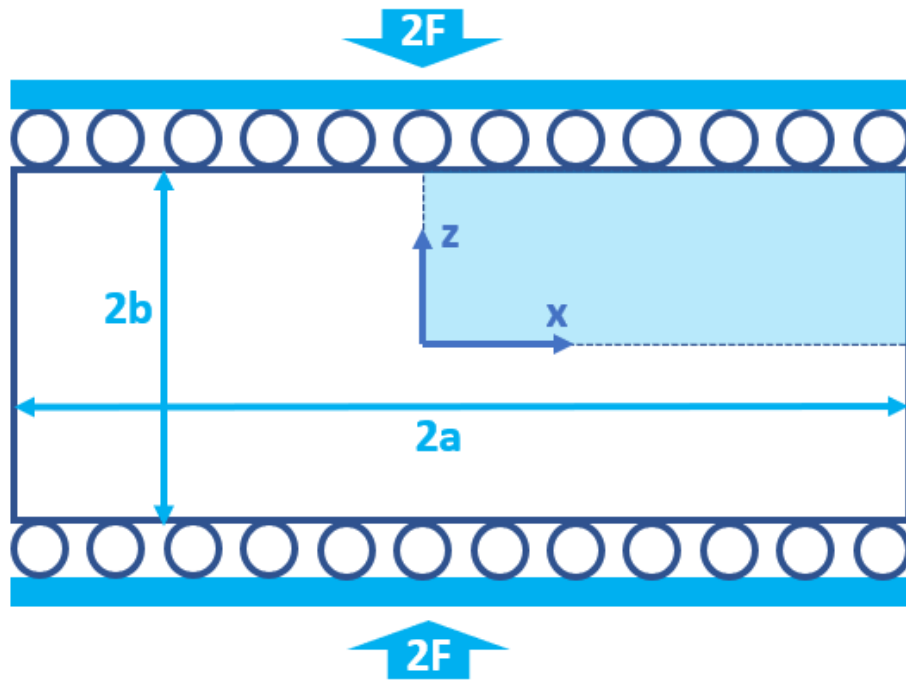


Fig. 4.16: Sketch of the problem

In this example, we set up and solve a poroelastic model to obtain the temporal and spatial solutions of pore pressure ( $p(x, z, t)$ ) and displacement field ( $u_z(x, z, t)$ ) for Mandel's problem. These modeling predictions are validated against corresponding analytical solution (Cheng and Detournay, 1988).

$$p(x, z, t) = 2p_0 \sum_{n=1}^{\infty} \frac{\sin \alpha_n}{\alpha_n - \sin \alpha_n \cos \alpha_n} \left( \cos \frac{\alpha_n x}{a} - \cos \alpha_n \right) \exp \left( -\frac{\alpha_n^2 ct}{a^2} \right)$$

$$u_z(x, z, t) = \left[ -\frac{F(1-\nu)}{2Ga} + \frac{F(1-\nu_u)}{Ga} \sum_{n=1}^{\infty} \frac{\sin \alpha_n \cos \alpha_n}{\alpha_n - \sin \alpha_n \cos \alpha_n} \exp \left( -\frac{\alpha_n^2 ct}{a^2} \right) \right] z$$

with  $\alpha_n$  denoting the positive roots of the following equation:

$$\tan \alpha_n = \frac{1-\nu}{\nu_u - \nu} \alpha_n$$

Upon sudden application of the vertical load, the instantaneous overpressure ( $p_0(x, z)$ ) and settlement ( $u_{z,0}(x, z)$  and  $u_{x,0}(x, z)$ ) across the sample are derived from the Skempton effect:

$$p_0(x, z) = \frac{1}{3a} B (1 + \nu_u) F$$

$$u_{z,0}(x, z) = -\frac{F (1 - \nu_u) z}{2G a}$$

$$u_{x,0}(x, z) = \frac{F \nu_u x}{2G a}$$

where  $\nu$  and  $\nu_u$  are the drained and undrained Poisson's ratio respectively,  $c$  is the consolidation coefficient,  $B$  is Skempton's coefficient,  $G$  is the shear modulus, and  $F$  is the applied force.

For this example, we focus on the `Mesh`, the `Constitutive`, and the `FieldSpecifications` tags.

## Mesh

The following figure shows the mesh used in this problem.

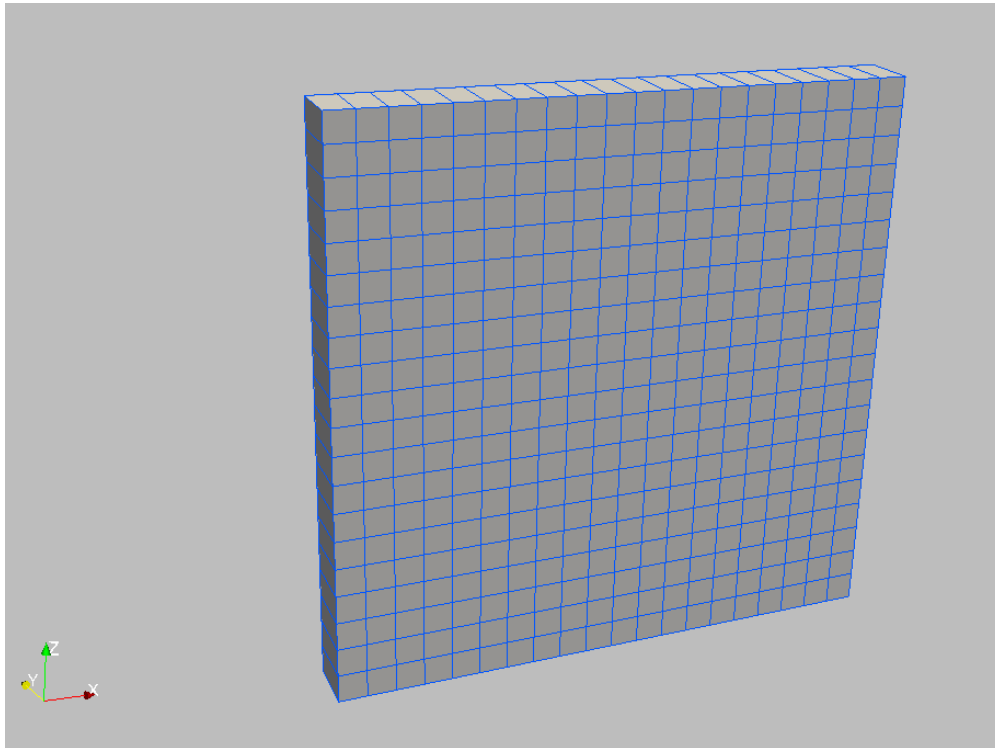


Fig. 4.17: Generated mesh

This mesh was created using the internal mesh generator as parametrized in the `InternalMesh` XML tag. The structured mesh contains 20 x 1 x 20 eight-node brick elements in the x, y, and z directions respectively. Such eight-node hexahedral elements are defined as `C3D8` elementTypes, and their collection forms a mesh with one group of cell blocks named here `cb1`.

```
<Mesh>
  <InternalMesh
    name="mesh1"
```

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```

    elementTypes="{ C3D8 }"
    xCoords="{ 0.0, 1.0 }"
    yCoords="{ 0.0, 0.1 }"
    zCoords="{ 0.0, 1.0 }"
    nx="{ 20 }"
    ny="{ 1 }"
    nz="{ 20 }"
    cellBlockNames="{ cb1 }"/>
</Mesh>

```

## Solid mechanics solver

GEOSX is a multi-physics platform. Different combinations of physics solvers available in the code can be applied in different regions of the domain and be functional at different stages of the simulation. The `Solvers` tag in the XML file is used to list and parameterize these solvers.

To specify a coupling between two different solvers, we define and characterize each single-physics solver separately. Then, we customize a *coupling solver* between these single-physics solvers as an additional solver. This approach allows for generality and flexibility in constructing multi-physics solvers. The order of specifying these solvers is not restricted in GEOSX. Note that end-users should give each single-physics solver a meaningful and distinct name, as GEOSX will recognize these single-physics solvers based on their customized names and create user-expected coupling.

As demonstrated in this example, to setup a poromechanical coupling, we need to define three different solvers in the XML file:

- the mechanics solver, a solver of type `SolidMechanicsLagrangianSSLE` called here `lagsolve` (more information here: [Solid Mechanics Solver](#)),

```

<SolidMechanicsLagrangianSSLE
  name="lagsolve"
  timeIntegrationOption="QuasiStatic"
  logLevel="1"
  discretization="FE1"
  targetRegions="{ Domain }"/>

```

- the single-phase flow solver, a solver of type `SinglePhaseFVM` called here `SinglePhaseFlow` (more information on these solvers at [Singlephase Flow Solver](#)),

```

<SinglePhaseFVM
  name="SinglePhaseFlow"
  logLevel="1"
  discretization="singlePhaseTPFA"
  targetRegions="{ Domain }"/>

```

- the coupling solver (`SinglePhasePoromechanics`) that will bind the two single-physics solvers above, which is named as `poroSolve` (more information at [Poromechanics Solver](#)).

```

<SinglePhasePoromechanics
  name="poroSolve"
  solidSolverName="lagsolve"
  flowSolverName="SinglePhaseFlow"
  logLevel="1"
  targetRegions="{ Domain }">
  <NonlinearSolverParameters

```

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```

    newtonMaxIter="2"
    newtonTol="1.0e-2"
    maxTimeStepCuts="1"
    lineSearchMaxCuts="0"/>
    <LinearSolverParameters
      directParallel="0"/>
  </SinglePhasePoromechanics>

```

The two single-physics solvers are parameterized as explained in their corresponding documentataion pages. We focus on the coupling solver in this example. The solver `poroSolve` uses a set of attributes that specifically describe the coupling process within a poromechanical framework. For instance, we must point this solver to the designated fluid solver (here: `SinglePhaseFlow`) and solid solver (here: `lagsolve`). These solvers interact through the `porousMaterialNames="{ shale }"` with all the constitutive models. We specify the discretization method (FE1, defined in the `NumericalMethods` section), and the target regions (here, we only have one, `Domain`). More parameters are required to characterize a coupling procedure (more information at [Poromechanics Solver](#)). In this way, the two single-physics solvers will be simultaneously called and executed for solving Mandel's problem here.

## Constitutive laws

For this problem, we simulate the poroelastic deformation of a slab under uniaxial compression. A homogeneous and isotropic domain with one solid material is assumed, and its mechanical properties and associated fluid rheology are specified in the `Constitutive` section. `PorousElasticIsotropic` model is used to describe the mechanical behavior of `shaleSolid` when subjected to loading. The single-phase fluid model `CompressibleSinglePhaseFluid` is selected to simulate the response of water upon consolidation.

```

<Constitutive>
  <PorousElasticIsotropic
    name="shale"
    solidModelName="shaleSolid"
    porosityModelName="shalePorosity"
    permeabilityModelName="shalePerm"/>

  <ElasticIsotropic
    name="shaleSolid"
    defaultDensity="0"
    defaultBulkModulus="6.6667e7"
    defaultShearModulus="4.0e7"/>

  <BiotPorosity
    name="shalePorosity"
    grainBulkModulus="1.0e27"
    defaultReferencePorosity="0.375"/>

  <ConstantPermeability
    name="shalePerm"
    permeabilityComponents="{ 1.0e-12, 0.0, 1.0e-12 }"/>

  <CompressibleSinglePhaseFluid
    name="water"
    defaultDensity="1000"
    defaultViscosity="0.001"
    referencePressure="0.000"
    referenceDensity="1"
    compressibility="4.4e-10"

```

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```
referenceViscosity="0.001"
viscosibility="0.0"/>
</Constitutive>
```

All constitutive parameters such as density, viscosity, bulk modulus, and shear modulus are specified in the International System of Units.

## Time history function

In the Tasks section, PackCollection tasks are defined to collect time history information from fields. Either the entire field or specified named sets of indices in the field can be collected. In this example, pressureCollection and displacementCollection tasks are specified to output the time history of pore pressure fieldName="pressure" and displacement field fieldName="TotalDisplacement" across the computational domain.

```
<Tasks>
  <PackCollection
    name="pressureCollection"
    objectPath="ElementRegions/Domain/cb1"
    fieldName="pressure"/>

  <PackCollection
    name="displacementCollection"
    objectPath="nodeManager"
    fieldName="totalDisplacement"/>
</Tasks>
```

These two tasks are triggered using the Event manager with a PeriodicEvent defined for these recurring tasks. GEOSX writes two files named after the string defined in the filename keyword and formatted as HDF5 files (displacement\_history.hdf5 and pressure\_history.hdf5). The TimeHistory file contains the collected time history information from each specified time history collector. This information includes datasets for the simulation time, element center, and the time history information. A Python script is prepared to read and plot any specified subset of the time history data for verification and visualization.

## Initial and boundary conditions

Next, we specify two fields:

- The initial value (the displacements and pore pressure have to be initialized, corresponding to the undrained response),
- The boundary conditions (the vertical displacement applied at the loaded boundary and the constraints of the outer boundaries have to be set).

In this example, the analytical z-displacement is applied at the top surface (zpos) of computational domain to enforce the rigid plate condition. The lateral surface (xpos) is traction-free and allows drainage. The remaining parts of the outer boundaries are subjected to roller constraints. These boundary conditions are set up through the FieldSpecifications section.

```
<FieldSpecifications>
  <FieldSpecification
    name="initialPressure"
    initialCondition="1"
```

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```
setNames="{ all }"
objectPath="ElementRegions/Domain/cb1"
fieldName="pressure"
scale="4934.86"/>

<FieldSpecification
  name="xInitialDisplacement"
  initialCondition="1"
  setNames="{ all }"
  objectPath="nodeManager"
  fieldName="totalDisplacement"
  component="0"
  scale="1.0"
  functionName="initialUxFunc"/>

<FieldSpecification
  name="yInitialDisplacement"
  initialCondition="1"
  setNames="{ all }"
  objectPath="nodeManager"
  fieldName="totalDisplacement"
  component="1"
  scale="0.0"/>

<FieldSpecification
  name="zInitialDisplacement"
  initialCondition="1"
  setNames="{ all }"
  objectPath="nodeManager"
  fieldName="totalDisplacement"
  component="2"
  scale="1.0"
  functionName="initialUzFunc"/>

<FieldSpecification
  name="xnegconstraint"
  objectPath="nodeManager"
  fieldName="totalDisplacement"
  component="0"
  scale="0.0"
  setNames="{ xneg }"/>

<FieldSpecification
  name="yconstraint"
  objectPath="nodeManager"
  fieldName="totalDisplacement"
  component="1"
  scale="0.0"
  setNames="{ yneg, ypos }"/>

<FieldSpecification
  name="zconstraint"
  objectPath="nodeManager"
  fieldName="totalDisplacement"
  component="2"
  scale="0.0"
  setNames="{ zneg }"/>
```

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```

<FieldSpecification
  name="NormalDisplacement"
  objectPath="nodeManager"
  fieldName="totalDisplacement"
  component="2"
  scale="-1.0e-5"
  setNames="{ zpos }"
  functionName="loadFunction"/>

<FieldSpecification
  name="boundaryPressure"
  objectPath="faceManager"
  fieldName="pressure"
  scale="0.0"
  setNames="{ xpos }"/>
</FieldSpecifications>

```

The parameters used in the simulation are summarized in the following table. Note that traction has a negative value, due to the negative sign convention for compressive stresses in GEOSX.

Symbol	Parameter	Unit	Value
$K$	Bulk Modulus	[MPa]	66.667
$G$	Shear Modulus	[MPa]	40.0
$F$	Force per Unit Length	[N/m]	$-10^4$
$\phi$	Porosity	[-]	0.375
$K_s$	Grain Bulk Modulus	[Pa]	$10^{27}$
$\rho_f$	Fluid density	[kg/m <sup>3</sup> ]	$10^3$
$c_f$	Fluid compressibility	[Pa <sup>-1</sup> ]	$4.4 \times 10^{-10}$
$\kappa$	Permeability	[m <sup>2</sup> ]	$10^{-12}$
$\mu$	Fluid viscosity	[Pa s]	$10^{-3}$
$2a$	Slab Length	[m]	2.0
$2b$	Slab Height	[m]	2.0

## Inspecting results

We request VTK-format output files and use Paraview to visualize the results. The following figure shows the distribution of pore pressure ( $p(x, z, t)$ ) at  $t = 10s$  within the computational domain.

The next figure shows the distribution of vertical displacement ( $u_z(x, z, t)$ ) at  $t = 10s$ .

The figure below compares the results from GEOSX (marks) and the corresponding analytical solution (lines) for the pore pressure along the x-direction and vertical displacement along the z-direction. GEOSX reliably captures the short-term Mandel-Cryer effect and shows excellent agreement with the analytical solution at various times.

## To go further

### Feedback on this example

For any feedback on this example, please submit a [GitHub issue](#) on the project's [GitHub](#) page.

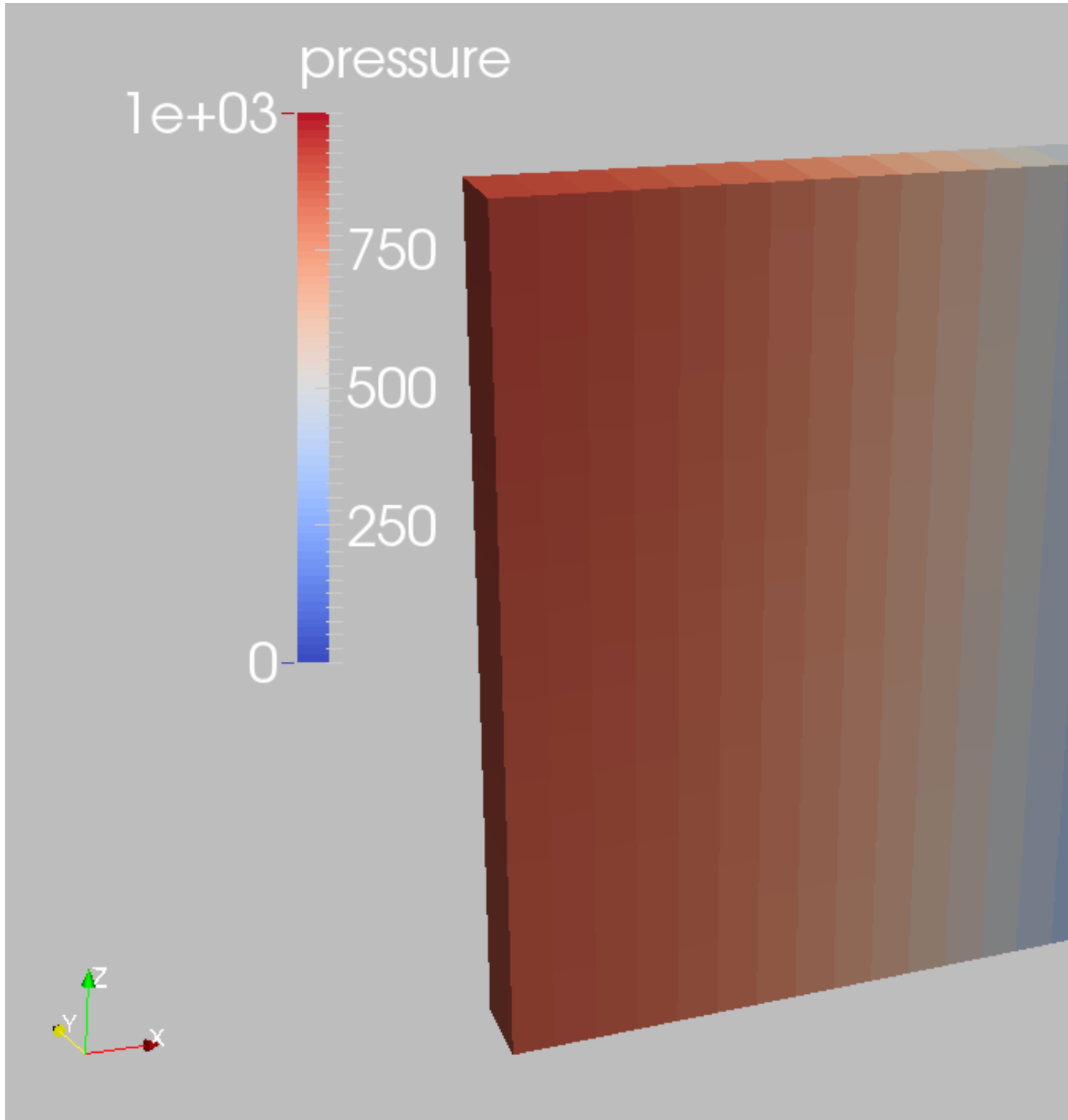


Fig. 4.18: Simulation result of pore pressure at  $t = 10s$



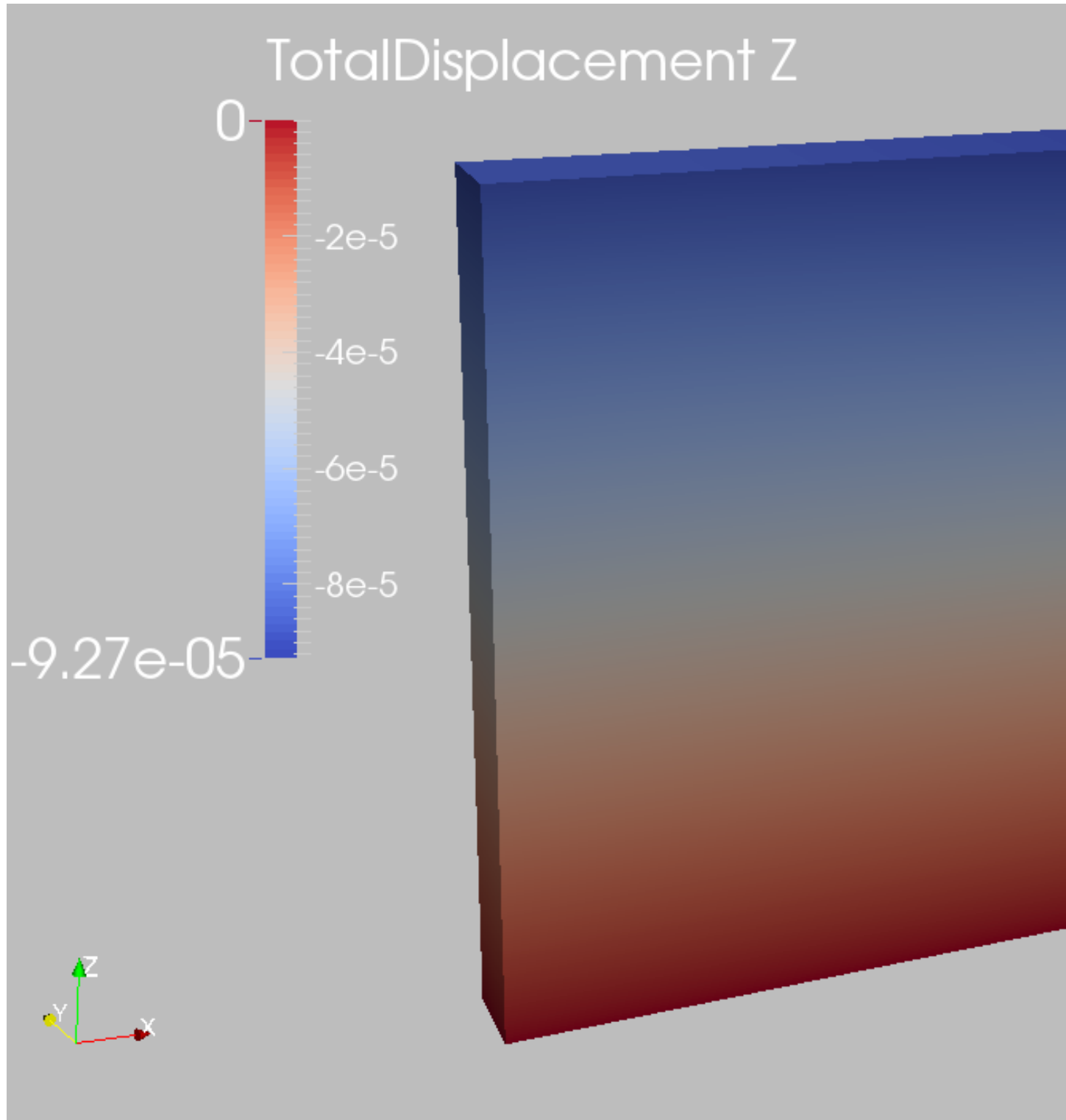
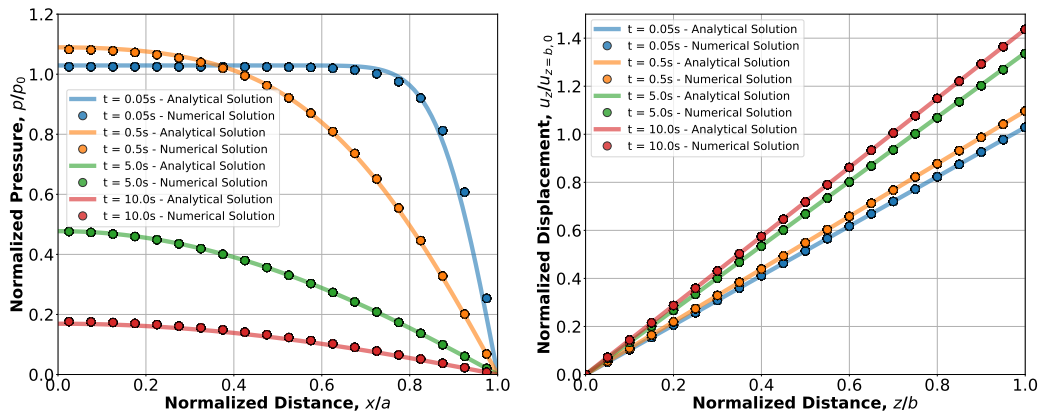


Fig. 4.19: Simulation result of vertical displacement at  $t = 10s$



## Verification of Induced Stresses Along a Fault

### Context

In this example, we evaluate the induced stresses in a pressurized reservoir displaced by a normal fault (permeable or impermeable). This problem is solved using the poroelastic solver in GEOSX to obtain the stress perturbations along the fault plane, which are verified against the corresponding analytical solution (Wu et al., 2020).

### Input file

The xml input files for the test case with impermeable fault are located at:

```
inputFiles/poromechanics/faultPoroelastic_base.xml
inputFiles/poromechanics/impermeableFault_benchmark.xml
```

The xml input files for the test case with permeable fault are located at:

```
inputFiles/poromechanics/faultPoroelastic_base.xml
inputFiles/poromechanics/permeableFault_benchmark.xml
```

A mesh file and a python script for post-processing the simulation results are also provided:

```
inputFiles/poromechanics/faultMesh.vtu
```

```
src/docs/sphinx/advancedExamples/validationStudies/faultMechanics/faultVerification/
↪ faultVerificationFigure.py
```

### Description of the case

We simulate induced stresses along a normal fault in a pressurized reservoir and compare our results against an analytical solution. In conformity to the analytical set-up, the reservoir is divided into two parts by an inclined fault. The fault crosses the entire domain, extending into the overburden and the underburden. The domain is horizontal, infinite, homogeneous, isotropic, and elastic. The reservoir is pressurized uniformly upon injection, and we neglect the transient effect of fluid flow. A pressure buildup is applied to: (i) the whole reservoir in the case of a permeable fault; (ii) the left compartment in the case of an impermeable fault. The overburden and underburden are impermeable (no pressure changes). Due to poromechanical effects, pore pressure changes in the reservoir cause a mechanical deformation of the entire domain. This deformation leads to a stress perturbation on the fault plane that could potentially trigger sliding of the fault. Here, the fault serves only as a flow boundary, and the mechanical separation of the fault plane

(either by shear slippage or normal opening) is prohibited, like in the analytical example. For verification purposes, a plane strain deformation is considered in the numerical model.

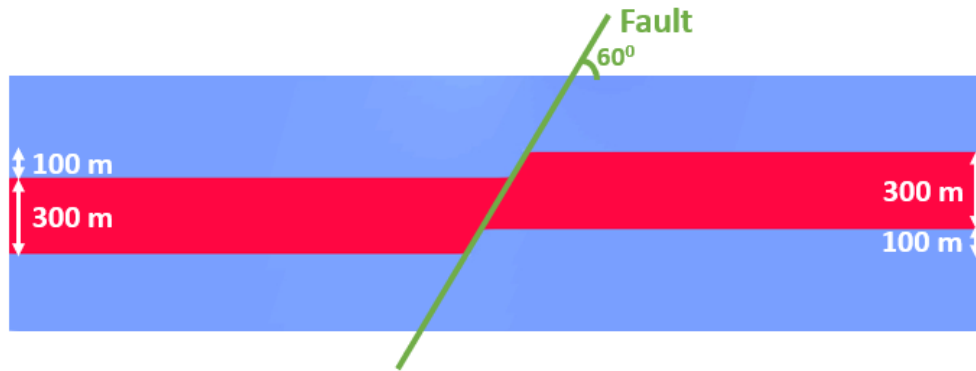


Fig. 4.20: Sketch of the problem

In this example, we set up and solve a poroelastic model to obtain the spatial solutions of displacement and stress fields across the domain upon pressurization. Changes of total stresses along the fault plane are evaluated and compared with the corresponding published work (Wu et al., 2020).

For this example, we focus on the `Mesh`, the `Constitutive`, and the `FieldSpecifications` tags.

## Mesh

The following figure shows the mesh used in this problem.

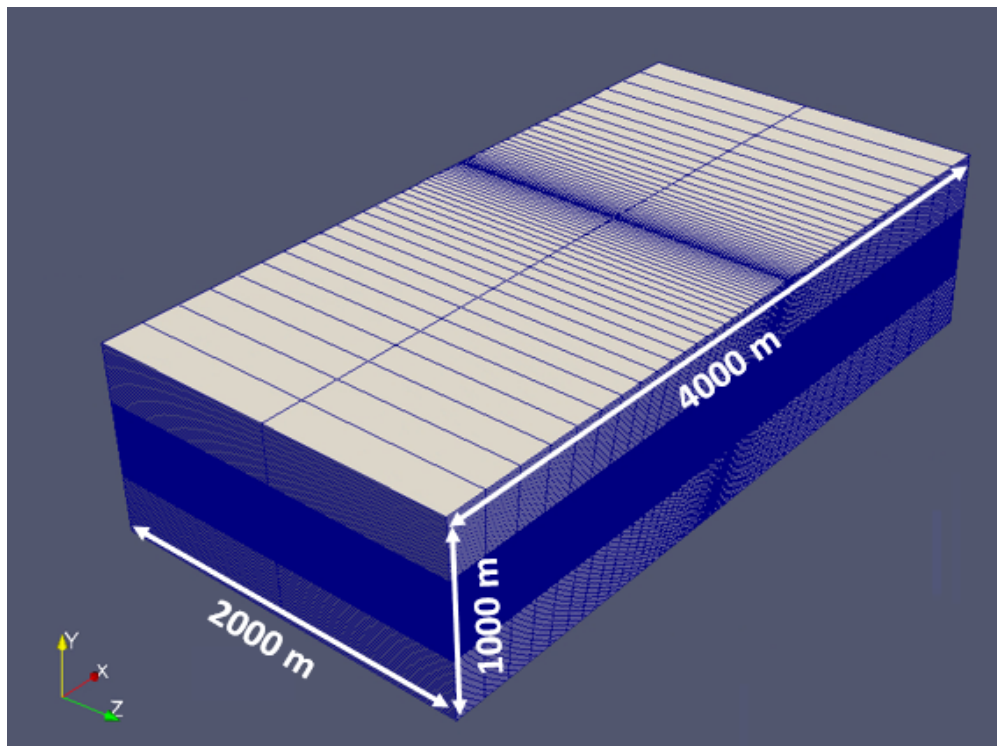


Fig. 4.21: Imported mesh

Here, we load the mesh with `VTKMesh`. The syntax to import external meshes is simple: in the XML file, the mesh file `faultMesh.vtu` is included with its relative or absolute path to the location of the GEOSX XML file and a user-specified label (here `FaultModel`) is given to the mesh object. This mesh contains quadrilateral elements and local refinement to conform with the fault geometry, and two reservoir compartments displaced by the fault. The size of the reservoir should be large enough to avoid boundary effects.

```
<Mesh>
  <VTKMesh
    name="FaultModel"
    file="faultMesh.vtu"
    regionAttribute="CellEntityIds"/>
  </Mesh>
```

## Solid mechanics solver

GEOSX is a multi-physics platform. Different combinations of physics solvers available in the code can be applied in different regions of the domain and be functional at different stages of the simulation. The `Solvers` tag in the XML file is used to list and parameterize these solvers.

To specify a coupling between two different solvers, we define and characterize each single-physics solver separately. Then, we customize a *coupling solver* between these single-physics solvers as an additional solver. This approach allows for generality and flexibility in constructing multi-physics solvers. The order in which solvers are specified is not important in GEOSX. Note that end-users should give each single-physics solver a meaningful and distinct name, as GEOSX will recognize these single-physics solvers based on their customized names to create the expected couplings.

As demonstrated in this example, to setup a poromechanical coupling, we need to define three different solvers in the XML file:

- the mechanics solver, a solver of type `SolidMechanics_LagrangianFEM` called here `mechanicsSolver` (more information here: [Solid Mechanics Solver](#)),

```
<SolidMechanics_LagrangianFEM
  name="mechanicsSolver"
  timeIntegrationOption="QuasiStatic"
  logLevel="1"
  discretization="FE1"
  targetRegions="{ Domain }">
  <NonlinearSolverParameters
    newtonTol = "1.0e-5"
    newtonMaxIter = "15"/>
  <LinearSolverParameters
    solverType="gmres"
    krylovTol="1.0e-10"/>
</SolidMechanics_LagrangianFEM>
```

- the single-phase flow solver, a solver of type `SinglePhaseFVM` called here `singlePhaseFlowSolver` (more information on these solvers at [Singlephase Flow Solver](#)),

```
<SinglePhaseFVM
  name="singlePhaseFlowSolver"
  logLevel="1"
  discretization="singlePhaseTPFA"
  targetRegions="{ Domain }">
  <NonlinearSolverParameters
    newtonTol = "1.0e-6"
```

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```

        newtonMaxIter = "8"
    />
    <LinearSolverParameters
        solverType="gmres"
        krylovTol="1.0e-12"/>
    </SinglePhaseFVM>
</Solvers>

```

- the coupling solver (`SinglePhasePoromechanics`) that will bind the two single-physics solvers above, named `poromechanicsSolver` (more information at [Poromechanics Solver](#)).

```

<Solvers gravityVector="{0.0, 0.0, 0.0}">
    <SinglePhasePoromechanics
        name="poromechanicsSolver"
        solidSolverName="mechanicsSolver"
        flowSolverName="singlePhaseFlowSolver"
        logLevel="1"
        targetRegions="{ Domain }">
        <LinearSolverParameters
            solverType="gmres"
            preconditionerType="mgr"
            logLevel="1"
            krylovAdaptiveTol="1"
        />
        <NonlinearSolverParameters
            newtonMaxIter = "40"
        />
    </SinglePhasePoromechanics>

```

The two single-physics solvers are parameterized as explained in their corresponding documents.

In this example, let us focus on the coupling solver. This solver (`poromechanicsSolver`) uses a set of attributes that specifically describe the coupling process within a poromechanical framework. For instance, we must point this solver to the designated fluid solver (here: `singlePhaseFlowSolver`) and solid solver (here: `mechanicsSolver`). These solvers are forced to interact with all the constitutive models in the target regions (here, we only have one, `Domain`). More parameters are required to characterize a coupling procedure (more information at [Poromechanics Solver](#)). This way, the two single-physics solvers will be simultaneously called and executed for solving the problem.

## Discretization methods for multiphysics solvers

Numerical methods in multiphysics settings are similar to single physics numerical methods. In this problem, we use finite volume for flow and finite elements for solid mechanics. All necessary parameters for these methods are defined in the `NumericalMethods` section.

As mentioned before, the coupling solver and the solid mechanics solver require the specification of a discretization method called `FE1`. In GEOSX, this discretization method represents a finite element method using linear basis functions and Gaussian quadrature rules. For more information on defining finite elements numerical schemes, please see the dedicated [Finite Element Discretization](#) section.

The finite volume method requires the specification of a discretization scheme. Here, we use a two-point flux approximation scheme (`singlePhaseTPFA`), as described in the dedicated documentation (found here: [Finite Volume Discretization](#)).

```
<NumericalMethods>
  <FiniteElements>
    <FiniteElementSpace
      name="FE1"
      order="1"/>
    </FiniteElements>

    <FiniteVolume>
      <TwoPointFluxApproximation
        name="singlePhaseTPFA"
      />
    </FiniteVolume>
  </NumericalMethods>
```

## Constitutive laws

For this problem, a homogeneous and isotropic domain with one solid material is assumed for both the reservoir and its surroundings. The solid and fluid materials are named as `rock` and `water` respectively, and their mechanical properties are specified in the Constitutive section. `PorousElasticIsotropic` model is used to describe the linear elastic isotropic response of `rock` when subjected to fluid injection. And the single-phase fluid model `CompressibleSinglePhaseFluid` is selected to simulate the flow of water.

```
<Constitutive>
  <PorousElasticIsotropic
    name="porousRock"
    solidModelName="rock"
    porosityModelName="rockPorosity"
    permeabilityModelName="rockPerm"
  />

  <ElasticIsotropic
    name="rock"
    defaultDensity="2700"
    defaultYoungModulus="14.95e9"
    defaultPoissonRatio="0.15"
  />

  <CompressibleSinglePhaseFluid
    name="water"
    defaultDensity="1000"
    defaultViscosity="0.001"
    referencePressure="0e6"
    referenceDensity="1000"
    compressibility="2.09028227021e-10"
    referenceViscosity="0.001"
    viscosibility="0.0"
  />

  <BiotPorosity
    name="rockPorosity"
    grainBulkModulus="7.12e10"
    defaultReferencePorosity="0.3"
  />

  <ConstantPermeability
```

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```

    name="rockPerm"
    permeabilityComponents="{1.0e-18, 1.0e-18, 1.0e-18}"
  />
</Constitutive>

```

All constitutive parameters such as density, viscosity, and Young's modulus are specified in the International System of Units.

## Initial and boundary conditions

The next step is to specify fields, including:

- The initial value (the in-situ stresses and pore pressure have to be initialized),
- The boundary conditions (pressure buildup within the reservoir and constraints of the outer boundaries have to be set).

In this example, we need to specify isotropic horizontal stress ( $\sigma_h = -60.0$  MPa and  $\sigma_H = -60.0$  MPa), vertical stress ( $\sigma_v = -70.0$  MPa), and initial reservoir pressure ( $P_0 = 35.0$  MPa). When initializing the model, a normal traction (name="NormalTraction") of -70.0 MPa is imposed on the upper boundary (setNames="{ 91 }") to reach mechanical equilibrium. The lateral and lower boundaries are subjected to roller constraints. These boundary conditions are set up through the FieldSpecifications section.

```

<FieldSpecifications>
  <FieldSpecification
    name="initialPressure"
    initialCondition="1"
    setNames="{all}"
    objectPath="ElementRegions/Domain"
    fieldName="pressure"
    scale="35.0e6"
  />

  <FieldSpecification
    name="stressXX"
    initialCondition="1"
    setNames="{all}"
    objectPath="ElementRegions/Domain"
    fieldName="rock_stress"
    component="0"
    scale="-60.0e6"
  />

  <FieldSpecification
    name="stressYY"
    initialCondition="1"
    setNames="{all}"
    objectPath="ElementRegions/Domain"
    fieldName="rock_stress"
    component="1"
    scale="-70.0e6"
  />

  <FieldSpecification
    name="stressZZ"
    initialCondition="1"

```

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```

    setNames="{all}"
    objectPath="ElementRegions/Domain"
    fieldName="rock_stress"
    component="2"
    scale="-60.0e6"
  />

  <FieldSpecification
    name="xconstraint"
    objectPath="nodeManager"
    fieldName="totalDisplacement"
    component="0"
    scale="0.0"
    setNames="{ 89, 88 }"/>

  <FieldSpecification
    name="yconstraint"
    objectPath="nodeManager"
    fieldName="totalDisplacement"
    component="1"
    scale="0.0"
    setNames="{ 90 }"/>

  <FieldSpecification
    name="zconstraintFront"
    objectPath="nodeManager"
    fieldName="totalDisplacement"
    component="2"
    scale="0.0"
    setNames="{ 92, 93 }"/>

  <Traction
    name="NormalTraction"
    objectPath="faceManager"
    tractionType="normal"
    scale="-70.0e6"
    setNames="{ 91 }"/>
</FieldSpecifications>

```

In this example, the only difference between the impermeable fault and permeable fault cases is how to apply pressure buildup. For the impermeable fault case, a constant pressure buildup is imposed to the left compartment of the reservoir (objectPath="ElementRegions/Domain/97\_hexahedra"):

```

<FieldSpecifications>
  <FieldSpecification
    name="injection"
    initialCondition="0"
    setNames="{all}"
    objectPath="ElementRegions/Domain/97_hexahedra"
    fieldName="pressure"
    scale="55.0e6"/>
</FieldSpecifications>

```

For the permeable fault case, a constant pressure buildup is imposed to both compartments of the reservoir: (objectPath="ElementRegions/Domain/97\_hexahedra" and objectPath="ElementRegions/Domain/96\_hexahedra"):



```

<FieldSpecifications>
  <FieldSpecification
    name="injection"
    initialCondition="0"
    setNames="{all}"
    objectPath="ElementRegions/Domain/97_hexahedra"
    fieldName="pressure"
    scale="55.0e6"/>

  <FieldSpecification
    name="injection2"
    initialCondition="0"
    setNames="{all}"
    objectPath="ElementRegions/Domain/96_hexahedra"
    fieldName="pressure"
    scale="55.0e6"/>
</FieldSpecifications>

```

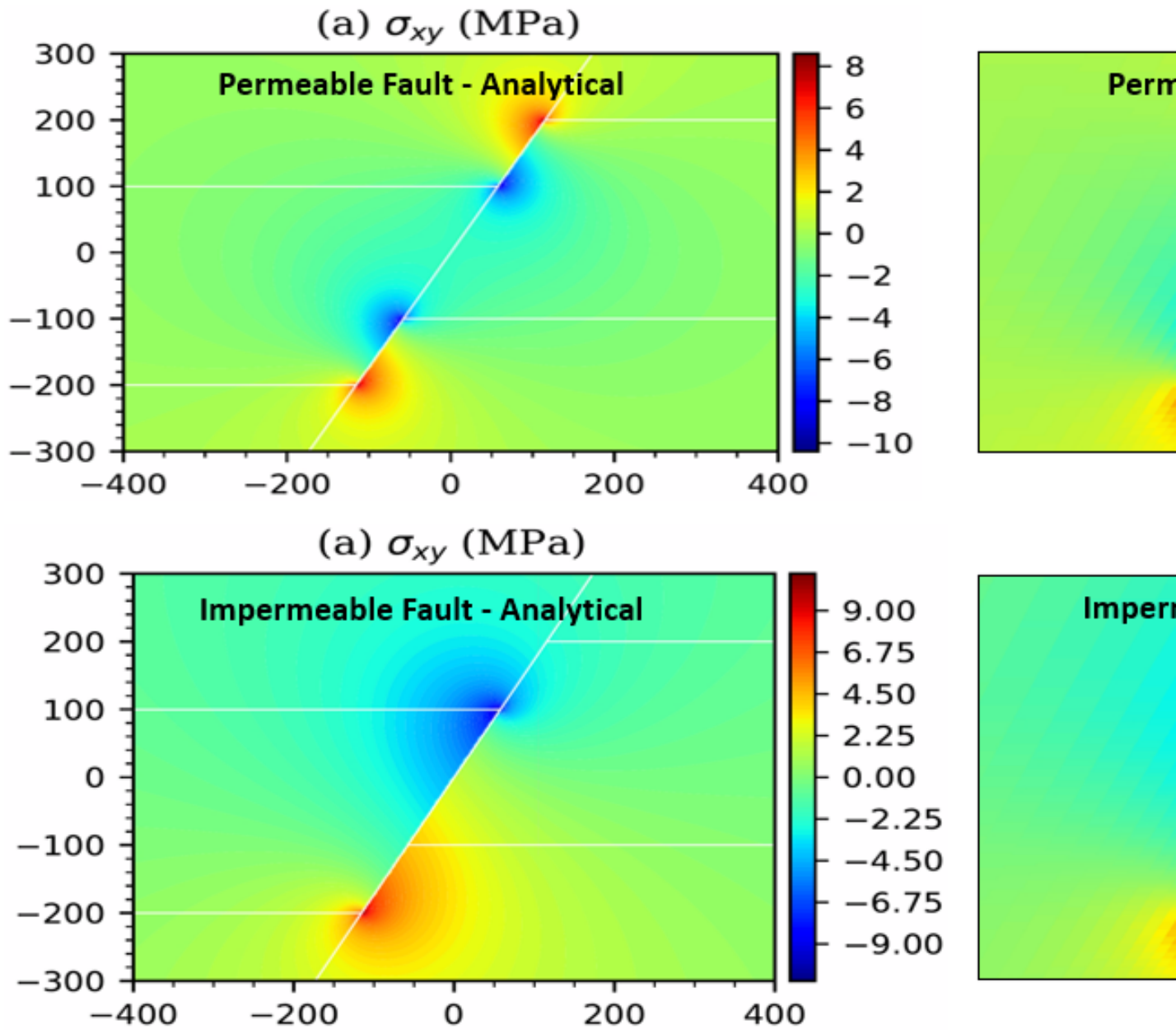
The parameters used in the simulation are summarized in the following table, which are specified in the Constitutive and FieldSpecifications sections. Note that stresses and traction have negative values, due to the negative sign convention for compressive stresses in GEOSX.

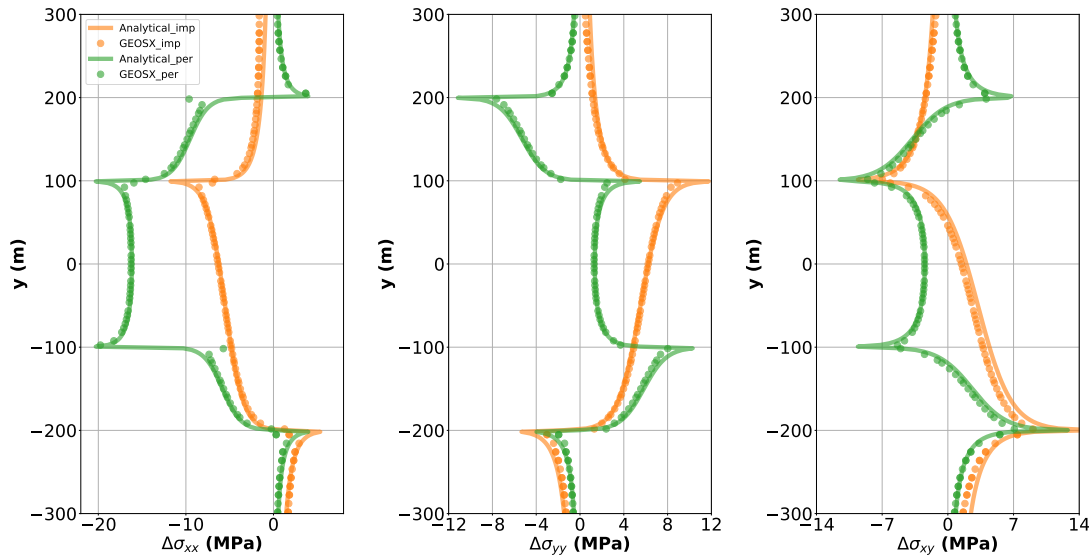
Symbol	Parameter	Unit	Value
$E$	Young's Modulus	[GPa]	14.95
$\nu$	Poisson's Ratio	[-]	0.15
$\sigma_h$	Min Horizontal Stress	[MPa]	-60.0
$\sigma_H$	Max Horizontal Stress	[MPa]	-60.0
$\sigma_v$	Vertical Stress	[MPa]	-70.0
$p_0$	Initial Reservoir Pressure	[MPa]	35.0
$\Delta p$	Pressure Buildup	[MPa]	20.0
$K_s$	Grain Bulk Modulus	[GPa]	71.2
$\theta$	Fault Dip	[Degree]	60.0
$\kappa$	Matrix Permeability	[m <sup>2</sup> ]	1.0*10 <sup>-18</sup>
$\phi$	Porosity	[-]	0.3
$D_L$	Domain Length	[m]	4000.0
$D_W$	Domain Width	[m]	2000.0
$D_T$	Domain Thickness	[m]	1000.0
$Res_T$	Reservoir Thickness	[m]	300.0
$F_{off}$	Fault Vertical Offset	[m]	100.0

## Inspecting results

We request VTK-format output files and use Paraview to visualize the results. The following figure shows the distribution of resulting shear stress ( $\sigma_{xy}$ ) in the computational domain for two different cases (a permeable vs. an impermeable fault). Numerical solutions for both cases are also compared with the corresponding analytical solutions.

The figure below compares the results from GEOSX (marks) and the corresponding analytical solution (solid curves) for the change of total stresses ( $\sigma_{xx}$ ,  $\sigma_{yy}$  and  $\sigma_{xy}$ ) along the fault plane. As shown, GEOSX reliably captures the mechanical deformation of the faulted reservoir and shows excellent agreement with the analytical solutions for two different scenarios. Differences in the stress perturbations between the cases with permeable and impermeable fault are also noticeable, which suggests that fault permeability plays a crucial role in governing reservoir deformation for the problems with reservoir pressurization or depletion.

Fig. 4.22: Simulation results of  $\sigma_{xy}$



## To go further

### Feedback on this example

For any feedback on this example, please submit a [GitHub issue](#) on the project's [GitHub](#) page.

## 4.1.3 Hydraulic Fracture

### Toughness dominated KGD hydraulic fracture

#### Description of the case

In this example, we consider a plane-strain hydraulic fracture propagating in an infinite, homogeneous and elastic medium, due to fluid injection at a rate  $Q_0$  during a period from 0 to  $t_{max}$ . Two dimensional KGD fracture is characterized as a vertical fracture with a rectangle-shaped cross section. For verification purpose, the presented numerical model is restricted to the assumptions used to analytically solve this problem (Bunger et al., 2005). Vertical and impermeable fracture surface is assumed, which eliminate the effect of fracture plane inclination and fluid leakoff. The injected fluid flows within the fracture, which is assumed to be governed by the lubrication equation resulting from the mass conservation and the Poiseuille law. Fracture profile is related to fluid pressure distribution, which is mainly dictated by fluid viscosity  $\mu$ . In addition, fluid pressure contributes to the fracture development through the mechanical deformation of the solid matrix, which is characterized by rock elastic properties, including the Young modulus  $E$ , and the Poisson ratio  $\nu$ .

For toughness-dominated fractures, more work is spent to split the intact rock than that applied to move the fracturing fluid. To make the case identical to the toughness dominated asymptotic solution, incompressible fluid with an ultra-low viscosity of 0.001 cp and medium rock toughness should be defined. Fracture is propagating with the creation of new surface if the stress intensity factor exceeds rock toughness  $K_{IC}$ .

In toughness-storage dominated regime, asymptotic solutions of the fracture length  $\ell$ , the net pressure  $p_0$  and the

fracture aperture  $w_0$  at the injection point for the KGD fracture are provided by (Bunger et al., 2005):

$$\begin{aligned}\ell &= 0.9324X^{-1/6}\left(\frac{E_p Q_0^3}{12\mu}\right)^{1/6}t^{2/3} \\ w_0^2 &= 0.5X^{1/2}\left(\frac{12\mu Q_0}{E_p}\right)^{1/2}\ell \\ w_0 p_0 &= 0.125X^{1/2}(12\mu Q_0 E_p)^{1/2}\end{aligned}$$

where the plane modulus  $E_p$  is defined by

$$E_p = \frac{E}{1 - \nu^2}$$

and the term  $X$  is given as:

$$X = \frac{256}{3\pi^2} \frac{K_{Ic}^4}{\mu Q_0 E_p^3}$$

### Input file

The input xml files for this test case are located at:

```
inputFiles/hydraulicFracturing/kgdToughnessDominated_base.xml
```

and

```
inputFiles/hydraulicFracturing/kgdToughnessDominated_benchmark.xml
```

The corresponding integrated test with coarser mesh and smaller injection duration is also prepared:

```
inputFiles/hydraulicFracturing/kgdToughnessDominated_Smoke.xml
```

Python scripts for post-processing and visualizing the simulation results are also prepared:

```
inputFiles/hydraulicFracturing/scripts/hydrofractureQueries.py
```

```
inputFiles/hydraulicFracturing/scripts/hydrofractureFigure.py
```

### Mechanics solvers

The solver `SurfaceGenerator` defines rock toughness  $K_{Ic}$  as:

```
<SurfaceGenerator
  name="SurfaceGen"
  targetRegions="{ Domain }"
  nodeBasedSIF="1"
  rockToughness="1e6"
  mpiCommOrder="1"/>
```

Rock and fracture deformation are modeled by the solid mechanics solver `SolidMechanicsLagrangianSSLE`. In this solver, we define `targetRegions` that includes both the continuum region and the fracture region. The name of the contact constitutive behavior is also specified in this solver by the `contactRelationName`, besides the `solidMaterialNames`.

```
<SolidMechanicsLagrangianSSLE
  name="lagsolve"
  timeIntegrationOption="QuasiStatic"
  discretization="FE1"
  targetRegions="{ Domain, Fracture }"
  contactRelationName="fractureContact"/>
```

The single phase fluid flow inside the fracture is solved by the finite volume method in the solver `SinglePhaseFVM` as:

```
<SinglePhaseFVM
  name="SinglePhaseFlow"
  discretization="singlePhaseTPFA"
  targetRegions="{ Fracture }"/>
```

All these elementary solvers are combined in the solver `Hydrofracture` to model the coupling between fluid flow within the fracture, rock deformation, fracture opening/closure and propagation. A fully coupled scheme is defined by setting a flag `FIM` for `couplingTypeOption`.

```
<Hydrofracture
  name="hydrofracture"
  solidSolverName="lagsolve"
  flowSolverName="SinglePhaseFlow"
  surfaceGeneratorName="SurfaceGen"
  couplingTypeOption="FIM"
  logLevel="1"
  targetRegions="{ Fracture }"
  contactRelationName="fractureContact"
  maxNumResolves="2">
```

## The constitutive laws

The constitutive law `CompressibleSinglePhaseFluid` defines the default and reference fluid viscosity, compressibility and density. For this toughness dominated example, ultra low fluid viscosity is used:

```
<CompressibleSinglePhaseFluid
  name="water"
  defaultDensity="1000"
  defaultViscosity="1.0e-6"
  referencePressure="0.0"
  compressibility="5e-10"
  referenceViscosity="1.0e-6"
  viscosibility="0.0"/>
```

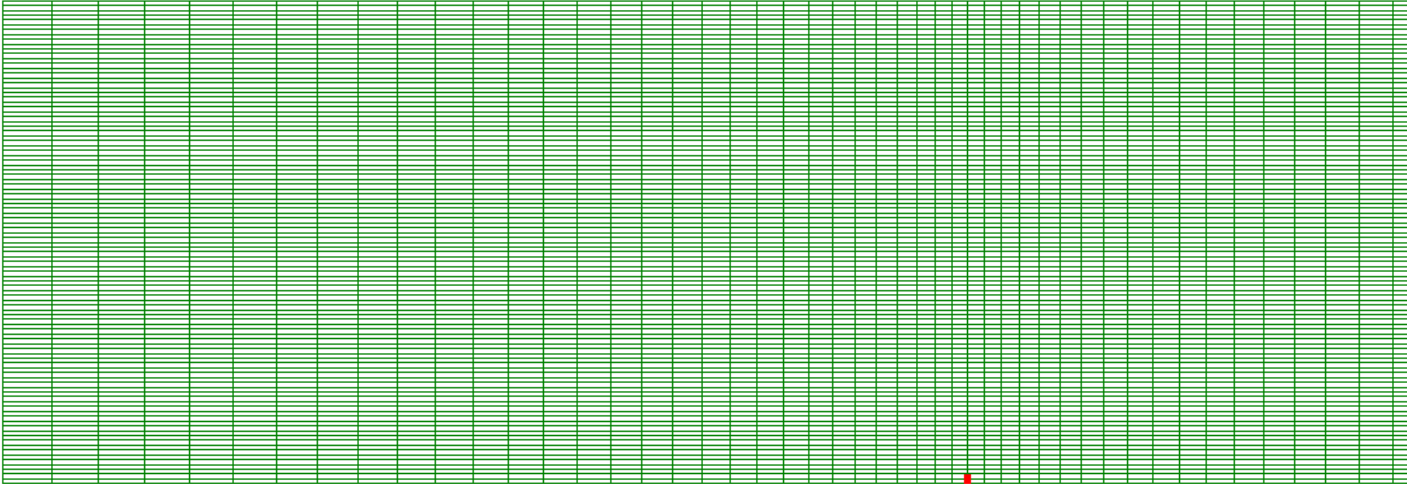
The isotropic elastic Young modulus and Poisson ratio are defined in the `ElasticIsotropic` block. The density of rock defined in this block is useless, as gravity effect is ignored in this example.

```
<ElasticIsotropic
  name="rock"
  defaultDensity="2700"
  defaultYoungModulus="30.0e9"
  defaultPoissonRatio="0.25"/>
```

## Mesh

Internal mesh generator is used to generate the geometry of this example. The domain size is large enough comparing to the final size of the fracture. A sensitivity analysis has shown that the domain size in the direction perpendicular to the fracture plane, i.e. x-axis, must be at least ten times of the final fracture half-length to minimize the boundary effect. However, smaller size along the fracture plane, i.e. y-axis, of only two times the fracture half-length is good enough. It is also important to note that at least two layers are required in z-axis to ensure a good match between the numerical results and analytical solutions, due to the node based fracture propagation criterion. Also in x-axis, bias parameter `xBias` is added for optimizing the mesh by refining the elements near the fracture plane.

```
<InternalMesh
  name="mesh1"
  elementTypes="{C3D8}"
  xCoords="{ -100, 0, 100 }"
  yCoords="{ 0, 50 }"
  zCoords="{ 0, 1 }"
  nx="{ 30, 30 }"
  ny="{ 100 }"
  nz="{ 2 }"
  xBias="{ 0.5, -0.5 }"
  cellBlockNames="{cb1}"/>
```



## Defining the initial fracture

The initial fracture is defined by a nodeset occupying a small area where the KGD fracture starts to propagate:

```
<Box
  name="fracture"
  xMin="{ -0.01, -0.01, -0.01 }"
  xMax="{ 0.01, 1.01, 1.01 }"/>
```

This initial `ruptureState` condition must be specified for this area in the following `FieldSpecification` block:

```
<FieldSpecification
  name="frac"
  initialCondition="1"
  setNames="{ fracture }"
  objectPath="faceManager"
  fieldName="ruptureState"
  scale="1"/>
```

## Defining the fracture plane

The plane within which the KGD fracture propagates is predefined to reduce the computational cost. The fracture plane is outlined by a separable nodeset by the following initial `FieldSpecification` condition:

```
<Box
  name="core"
  xMin="{ -0.01, -0.01, -0.01 }"
  xMax="{ 0.01, 50.01, 1.01 }"/>
```

```
<FieldSpecification
  name="separableFace"
  initialCondition="1"
  setNames="{ core }"
  objectPath="faceManager"
  fieldName="isFaceSeparable"
  scale="1"/>
```

## Defining the injection rate

Fluid is injected into a sub-area of the initial fracture. Only half of the injection rate is defined in this boundary condition because only half-wing of the KGD fracture is modeled regarding its symmetry. Hereby, the mass injection rate is actually defined, instead of the volume injection rate. More precisely, the value given for `scale` is  $Q_0\rho_f/2$  (not  $Q_0/2$ ).

```
<SourceFlux
  name="sourceTerm"
  objectPath="ElementRegions/Fracture"
  scale="-5e-2"
  setNames="{ source }"/>
```

## Time history function

In the `Tasks` section, `PackCollection` tasks are defined to collect time history information from fields. Either the entire field or specified named sets of indices in the field can be collected. In this example, `pressureCollection`, `apertureCollection`, `hydraulicApertureCollection` and `areaCollection` are specified to output the time history of fracture characteristics (pressure, width and area). `objectPath="ElementRegions/Fracture/FractureSubRegion"` indicates that these `PackCollection` tasks are applied to the fracture element subregion.

```
<Tasks>
  <PackCollection
    name="pressureCollection"
```

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```

objectPath="ElementRegions/Fracture/FractureSubRegion"
fieldName="pressure"/>

<PackCollection
  name="apertureCollection"
  objectPath="ElementRegions/Fracture/FractureSubRegion"
  fieldName="elementAperture"/>

<PackCollection
  name="hydraulicApertureCollection"
  objectPath="ElementRegions/Fracture/FractureSubRegion"
  fieldName="hydraulicAperture"/>

<PackCollection
  name="areaCollection"
  objectPath="ElementRegions/Fracture/FractureSubRegion"
  fieldName="elementArea"/>
</Tasks>

```

These tasks are triggered using the Event manager with a `PeriodicEvent` defined for the recurring tasks. GEOSX writes one file named after the string defined in the `filename` keyword and formatted as a HDF5 file (`kgdToughnessDominated_output.hdf5`). This TimeHistory file contains the collected time history information from specified time history collector. This file includes datasets for the simulation time, fluid pressure, element aperture, hydraulic aperture and element area for the propagating hydraulic fracture. A Python script is prepared to read and query any specified subset of the time history data for verification and visualization.

The parameters used in the simulation are summarized in the following table.

Symbol	Parameter	Units	Value
$Q_0$	Injection rate	[m <sup>3</sup> /s]	10 <sup>-4</sup>
$E$	Young's modulus	[GPa]	30
$\nu$	Poisson's ratio	[ - ]	0.25
$\mu$	Fluid viscosity	[Pa.s]	10 <sup>-6</sup>
$K_{Ic}$	Rock toughness	[MPa.m <sup>1/2</sup> ]	1

## Inspecting results

Fracture propagation during the fluid injection period is shown in the figure below.

First, by running the query script

```
python ./hydrofractureQueries.py kgdToughnessDominated
```

the HDF5 output is postprocessed and temporal evolution of fracture characteristics (fluid pressure and fracture width at fluid inlet and fracture half length) are saved into a txt file `model-results.txt`, which can be used for verification and visualization:

```

[['time', 'pressure', 'aperture', 'length']]
2 4.086e+05 8.425e-05 2
4 3.063e+05 0.0001021 3
6 3.121e+05 0.0001238 3.5
8 2.446e+05 0.0001277 4.5
10 2.411e+05 0.0001409 5

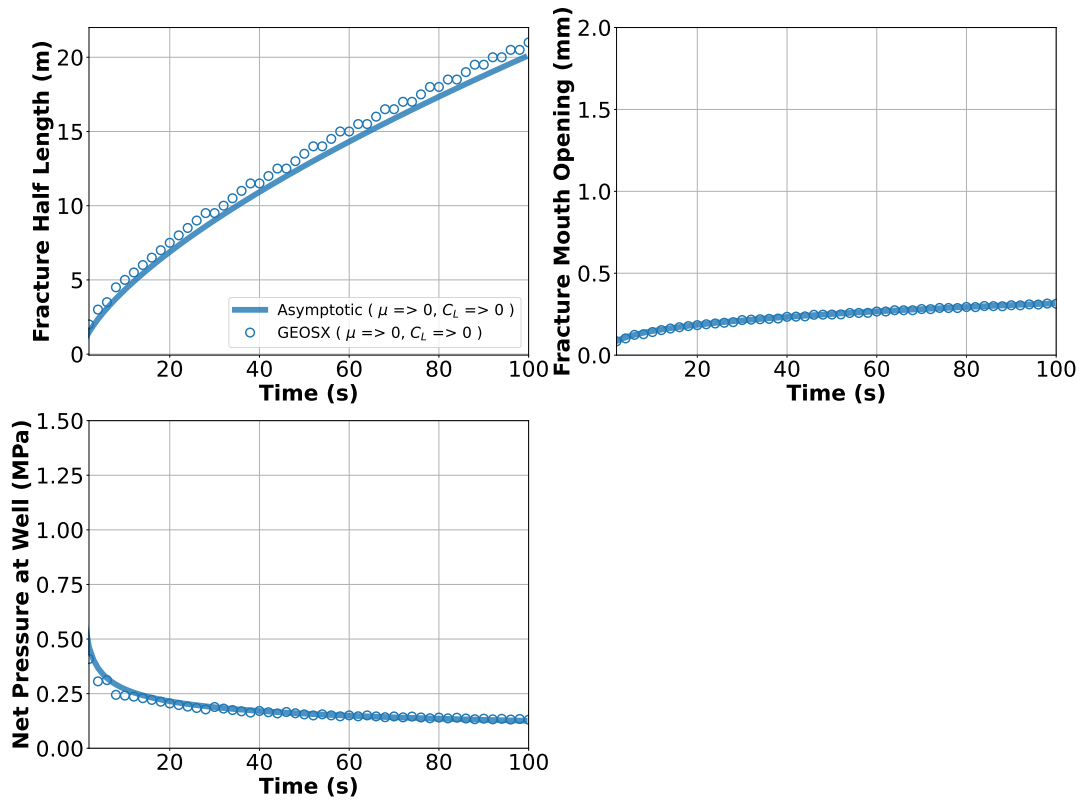
```



Note: GEOSX python tools `geosx_xml_tools` should be installed to run the query script (See [Python Tools Setup](#) for details).

A good agreement between GEOSX results and analytical solutions is shown in the comparison below, which is generated using the visualization script:

```
python ./kgdToughnessDominatedFigure.py
```



## To go further

### Feedback on this example

This concludes the toughness dominated KGD example. For any feedback on this example, please submit a [GitHub issue on the project's GitHub page](#).

## Viscosity dominated KGD hydraulic fracture

### Description of the case

The KGD problem addresses a single plane strain fracture growing in an infinite elastic domain. Basic assumptions and characteristic shape for this example is similar to those of another case ([Viscosity dominated KGD hydraulic fracture](#)) except that the viscosity dominated regime is now considered. In this regime, more work is spent to move

the fracturing fluid than to split the intact rock. In this test, slickwater with a constant viscosity of 1 cp is chosen as fracturing fluid, whose compressibility is neglected. To make the case identical to the viscosity dominated asymptotic solution, an ultra-low rock toughness  $K_{Ic}$  is defined and fracture is assumed to be always propagating following fluid front. Asymptotic solutions of the fracture length  $\ell$ , the net pressure  $p_0$  and the fracture aperture  $w_0$  at the injection point for the KGD fracture with a viscosity dominated regime are provided by (Adachi and Detournay, 2002):

$$\begin{aligned}\ell &= 0.6152 \left( \frac{E_p Q_0^3}{12\mu} \right)^{1/6} t^{2/3} \\ w_0^2 &= 2.1 \left( \frac{12\mu Q_0}{E_p} \right)^{1/2} \ell \\ w_0 p_0 &= 0.62 (12\mu Q_0 E_p)^{1/2}\end{aligned}$$

where the plane modulus  $E_p$  is defined by

$$E_p = \frac{E}{1 - \nu^2}$$

and the term  $X$  is given as:

$$X = \frac{256}{3\pi^2} \frac{K_{Ic}^4}{\mu Q_0 E_p^3}$$

### Input file

The input xml files for this test case are located at:

```
inputFiles/hydraulicFracturing/kgdViscosityDominated_base.xml
```

and

```
inputFiles/hydraulicFracturing/kgdViscosityDominated_benchmark.xml
```

The corresponding integrated test with coarser mesh and smaller injection duration is also prepared:

```
inputFiles/hydraulicFracturing/kgdViscosityDominated_smoke.xml
```

Python scripts for post-processing and visualizing the simulation results are also prepared:

```
inputFiles/hydraulicFracturing/scripts/hydrofractureQueries.py
```

```
inputFiles/hydraulicFracturing/scripts/hydrofractureFigure.py
```

Fluid rheology and rock toughness are defined in the xml blocks below. Please note that setting an absolute zero value for the rock toughness could lead to instability issue. Therefore, a low value of  $K_{Ic}$  is used in this example.

```
<SurfaceGenerator
  name="SurfaceGen"
  targetRegions="{ Domain }"
  nodeBasedSIF="1"
  rockToughness="1e4"
  mpiCommOrder="1"/>
```

```
<CompressibleSinglePhaseFluid
  name="water"
  defaultDensity="1000"
  defaultViscosity="1.0e-3"
```

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```
referencePressure="0.0"
compressibility="5e-10"
referenceViscosity="1.0e-3"
viscosibility="0.0"/>
```

First, by running the query script

```
python ./hydrofractureQueries.py kgdViscosityDominated
```

the HDF5 output is postprocessed and temporal evolution of fracture characteristics (fluid pressure and fracture width at fluid inlet and fracure half length) are saved into a txt file `model-results.txt`, which can be used for verification and visualization:

```
[['      time', ' pressure', ' aperture', '      length']]
 2 1.075e+06 0.0001176      1.5
 4 9.636e+05 0.0001645        2
 6 8.372e+05 0.0001917      2.5
 8 7.28e+05 0.000209        3
10 6.512e+05 0.000222      3.5
```

Note: GEOSX python tools `geosx_xml_tools` should be installed to run the query script (See [Python Tools Setup](#) for details).

A good agreement between GEOSX results and analytical solutions is shown in the comparison below, which is generated using the visualization script:

```
python ./kgdViscosityDominatedFigure.py
```

## To go further

### Feedback on this example

This concludes the viscosity dominated KGD example. For any feedback on this example, please submit a [GitHub issue](#) on the project's [GitHub](#) page.

## Validating KGD Hydraulic Fracture with Experiment

### Context

In this example, we use GEOSX to model a planar hydraulic fracture propagating in a finite domain subject to traction-free external boundaries. Contrary to the classic KGD problems, we do not assume an infinite rock domain. Existing analytical solutions cannot model fracture behavior in this scenario, so this problem is solved using the hydrofracture solver in GEOSX. We validate the simulation results against a benchmark experiment ([Rubin, 1983](#)).

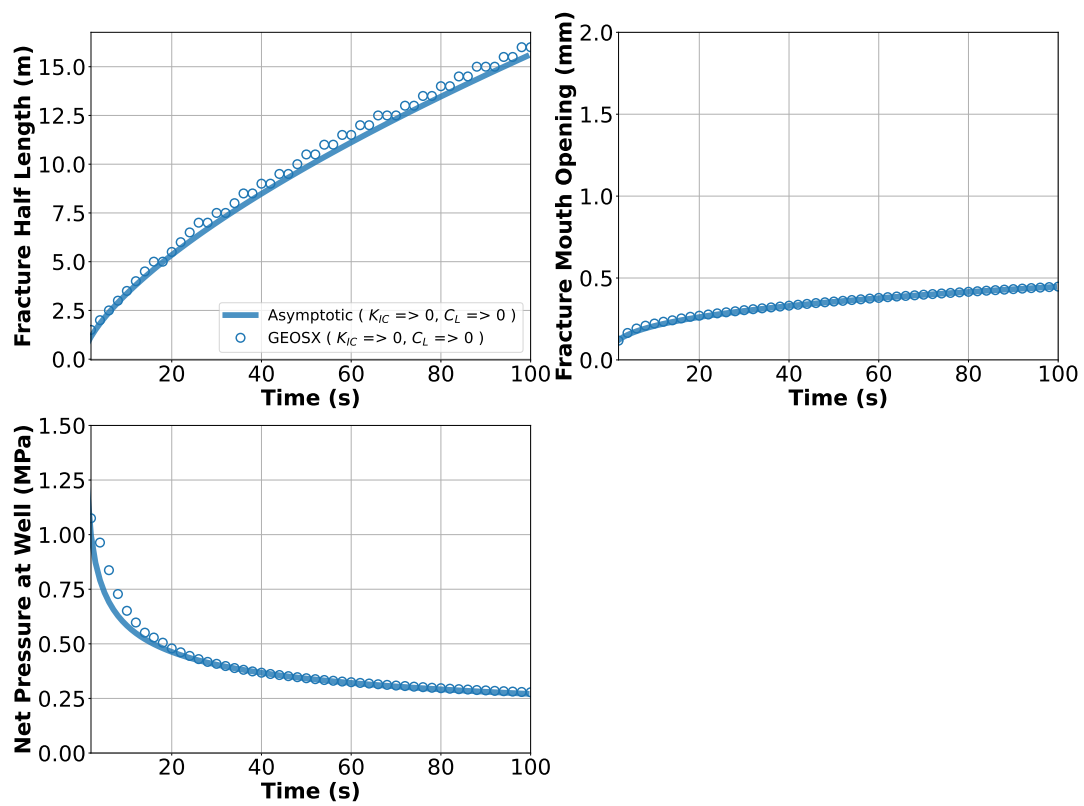
### Input file

This example uses no external input files. Everything we need is contained within two GEOSX input files:

```
inputFiles/hydraulicFracturing/kgdValidation_base.xml
```

```
inputFiles/hydraulicFracturing/kgdValidation_benchmark.xml
```

Python scripts for post-processing and visualizing the simulation results are also prepared:



```
src/docs/sphinx/advancedExamples/validationStudies/hydraulicFracture/kgdValidation/
↳ kgdValidationQueries.py
```

```
src/docs/sphinx/advancedExamples/validationStudies/hydraulicFracture/kgdValidation/
↳ kgdValidationFigure.py
```

## Description of the case

We simulate a hydraulic fracturing experiment within a finite domain made of three layers of polymethylmethacrylate (PMMA). As shown below, we inject viscous fluid to create a single planar fracture in the middle layer. The target layer is bonded weakly to the adjacent layers, so a vertical fracture develops inside the middle layer. Four pressure gages are placed to monitor wellbore pressure (gage 56) and fluid pressure along the fracture length (gage 57, 58, and 59). A linear variable differential transducer (LVDT) measures the fracture aperture at 28.5 mm away from the wellbore. Images are taken at regular time intervals to show the temporal evolution of the fracture extent. All experimental measurements for the time history of pressure, aperture, and length are reported in [Rubin \(1983\)](#). We use GEOSX to reproduce the conditions of this test, including material properties and pumping parameters. In the experiment, the upper and lower layers are used only to restrict the fracture height growth, they are therefore not simulated in GEOSX but are present as boundary conditions. Given the vertical plane of symmetry, only half of the middle layer is modeled. For verification purposes, a plane strain deformation and zero fluid leak-off are considered in the numerical model.

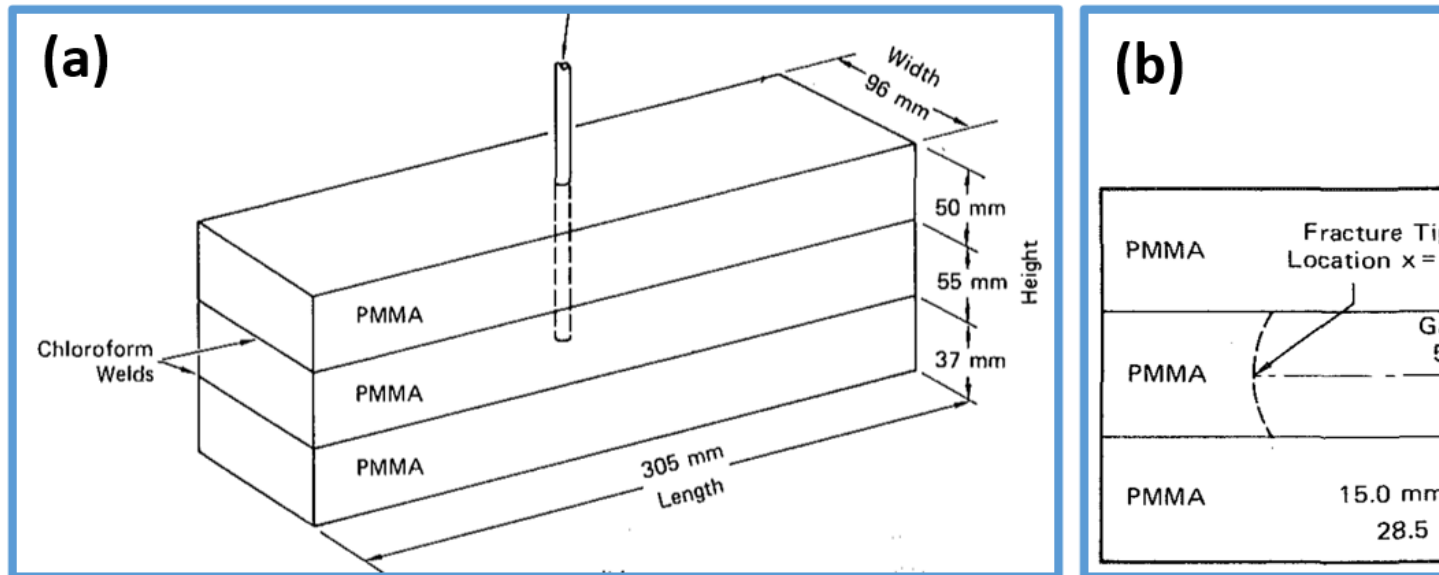


Fig. 4.23: Sketch of the problem

In this example, we solve the hydraulic fracturing problem with the `hydrofrac` solver to obtain the temporal solution of the fracture characteristics (length, aperture and pressure). These modeling predictions are compared with the corresponding experimental results ([Rubin, 1983](#)).

For this example, we focus on the `Mesh`, the `Constitutive`, and the `FieldSpecifications` tags.

## Mesh

The following figure shows the mesh used in this problem.

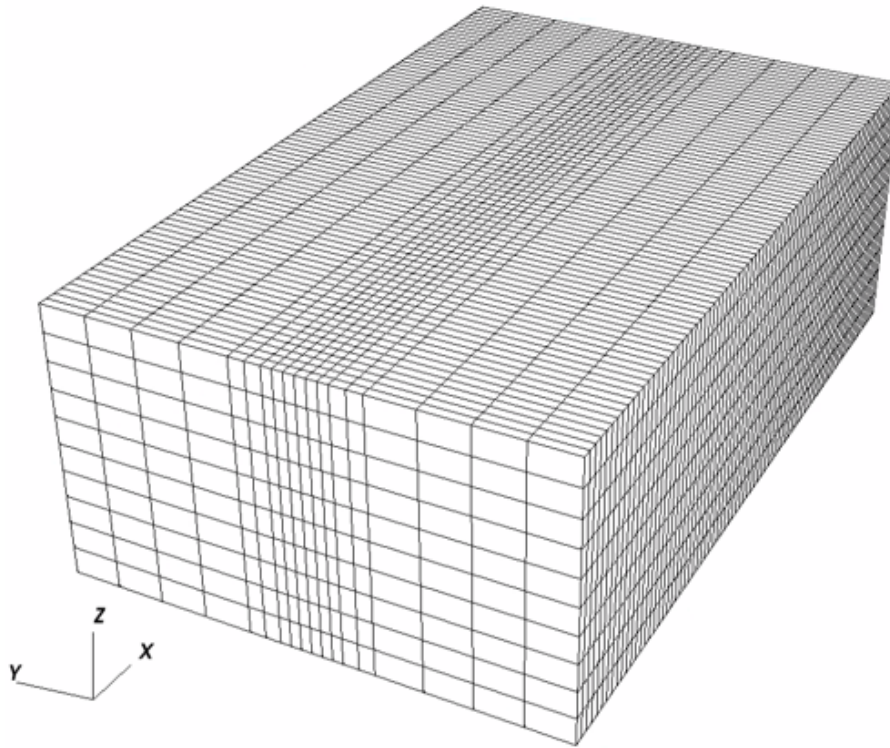


Fig. 4.24: Generated mesh

We use the internal mesh generator to create a computational domain ( $0.1525\text{ m} \times 0.096\text{ m} \times 0.055\text{ m}$ ), as parametrized in the `InternalMesh` XML tag. The structured mesh contains  $80 \times 18 \times 10$  eight-node brick elements in the  $x$ ,  $y$ , and  $z$  directions respectively. Such eight-node hexahedral elements are defined as `C3D8` elementTypes, and their collection forms a mesh with one group of cell blocks named here `cb1`. Along the  $y$ -axis, refinement is performed for the elements in the vicinity of the fracture plane.

```
<InternalMesh
  name="mesh1"
  elementTypes="{C3D8}"
  xCoords="{ 0, 0.1525 }"
  yCoords="{ -0.048, -0.012, -0.006, 0.006, 0.012, 0.048 }"
  zCoords="{ 0.037, 0.092 }"
  nx="{ 80 }"
  ny="{ 4, 2, 6, 2, 4 }"
  nz="{ 10 }"
  cellBlockNames="{cb1}"/>
```

The fracture plane is defined by a nodeset occupying a small region within the computation domain, where the fracture tends to open and propagate upon fluid injection:

```
<Box
  name="core"
  xMin="{ -0.1, -0.001, 0.036 }"
  xMax="{ 0.2, 0.001, 0.093 }"/>
```

## Solid mechanics solver

GEOSX is a multi-physics platform. Different combinations of physics solvers available in the code can be applied in different regions of the domain and be functional at different stages of the simulation. The `Solvers` tag in the XML file is used to list and parameterize these solvers.

Three elementary solvers are combined in the solver `Hydrofracture` to model the coupling between fluid flow within the fracture, rock deformation, fracture deformation and propagation:

```
<Hydrofracture
  name="hydrofracture"
  solidSolverName="lagsolve"
  flowSolverName="SinglePhaseFlow"
  surfaceGeneratorName="SurfaceGen"
  couplingTypeOption="FIM"
  logLevel="1"
  targetRegions="{ Fracture }"
  contactRelationName="fractureContact"
  maxNumResolves="2">
  <NonlinearSolverParameters
    newtonTol="1.0e-5"
    newtonMaxIter="20"
    lineSearchMaxCuts="3"/>
  <LinearSolverParameters
    directParallel="0"/>
</Hydrofracture>
```

- Rock and fracture deformation are modeled by the solid mechanics solver `SolidMechanicsLagrangianSSLE`. In this solver, we define `targetRegions` that includes both the continuum region and the fracture region. The name of the contact constitutive behavior is specified in this solver by the `contactRelationName`.

```
<SolidMechanicsLagrangianSSLE
  name="lagsolve"
  timeIntegrationOption="QuasiStatic"
  discretization="FE1"
  targetRegions="{ Domain, Fracture }"
  contactRelationName="fractureContact"/>
```

- The single-phase fluid flow inside the fracture is solved by the finite volume method in the solver `SinglePhaseFVM`.

```
<SinglePhaseFVM
  name="SinglePhaseFlow"
  discretization="singlePhaseTPFA"
  targetRegions="{ Fracture }"
  inputFluxEstimate="0.00000366"/>
```

- The solver `SurfaceGenerator` defines the fracture region and rock toughness. With `nodeBasedSIF="0"`, edge-based Stress Intensity Factor (SIF) calculation is chosen for the fracture propagation criterion.

```
<SurfaceGenerator
  name="SurfaceGen"
  logLevel="1"
  targetRegions="{ Domain }"
  nodeBasedSIF="0"
```

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```
rockToughness="1.2e6"
mpiCommOrder="1"/>
```

## Constitutive laws

For this problem, a homogeneous and isotropic domain with one solid material is assumed, and its mechanical properties and associated fluid rheology are specified in the Constitutive section. ElasticIsotropic model is used to describe the mechanical behavior of rock, when subjected to fluid injection. The single-phase fluid model CompressibleSinglePhaseFluid is selected to simulate the response of water upon fracture propagation.

```
<Constitutive>
  <CompressibleSinglePhaseFluid
    name="water"
    defaultDensity="1000"
    defaultViscosity="97.7"
    referencePressure="0.0"
    compressibility="5e-12"
    referenceViscosity="97.7"
    viscosibility="0.0"/>

  <ElasticIsotropic
    name="rock"
    defaultDensity="2700"
    defaultBulkModulus="4.110276e9"
    defaultShearModulus="1.19971e9"/>

  <CompressibleSolidParallelPlatesPermeability
    name="fractureFilling"
    solidModelName="nullSolid"
    porosityModelName="fracturePorosity"
    permeabilityModelName="fracturePerm"/>

  <NullModel
    name="nullSolid"/>

  <PressurePorosity
    name="fracturePorosity"
    defaultReferencePorosity="1.00"
    referencePressure="0.0"
    compressibility="0.0"/>

  <ParallelPlatesPermeability
    name="fracturePerm"/>

  <FrictionlessContact
    name="fractureContact"
    penaltyStiffness="1.0"
    apertureTableName="apertureTable" />
</Constitutive>
```

All constitutive parameters such as density, viscosity, bulk modulus, and shear modulus are specified in the International System of Units.



## Time history function

In the `Tasks` section, `PackCollection` tasks are defined to collect time history information from fields. Either the entire field or specified named sets of indices in the field can be collected. In this example, `pressureCollection`, `apertureCollection`, `hydraulicApertureCollection` and `areaCollection` are specified to output the time history of fracture characteristics (pressure, width and area). `objectPath="ElementRegions/Fracture/FractureSubRegion"` indicates that these `PackCollection` tasks are applied to the fracture element subregion.

```
<Tasks>
  <PackCollection
    name="pressureCollection"
    objectPath="ElementRegions/Fracture/FractureSubRegion"
    fieldName="pressure"/>

  <PackCollection
    name="apertureCollection"
    objectPath="ElementRegions/Fracture/FractureSubRegion"
    fieldName="elementAperture"/>

  <PackCollection
    name="hydraulicApertureCollection"
    objectPath="ElementRegions/Fracture/FractureSubRegion"
    fieldName="hydraulicAperture"/>

  <PackCollection
    name="areaCollection"
    objectPath="ElementRegions/Fracture/FractureSubRegion"
    fieldName="elementArea"/>
</Tasks>
```

These tasks are triggered using the Event manager with a `PeriodicEvent` defined for the recurring tasks. GEOSX writes one file named after the string defined in the `filename` keyword and formatted as a HDF5 file (`KGD_validation_output.hdf5`). This `TimeHistory` file contains the collected time history information from specified time history collector. This file includes datasets for the simulation time, fluid pressure, element aperture, hydraulic aperture and element area for the propagating hydraulic fracture. A Python script is prepared to read and query any specified subset of the time history data for verification and visualization.

## Initial and boundary conditions

The next step is to specify fields, including:

- The initial values: the `waterDensity`, `separableFace` and the `ruptureState` of the propagating fracture have to be initialized,
- The boundary conditions: fluid injection rates and the constraints of the outer boundaries have to be set.

In this example, a mass injection rate `SourceFlux` (`scale="-0.0000366"`) is applied at the surfaces of the initial fracture. Only half of the injection rate is defined in this boundary condition because only a half-wing of the fracture is modeled (the problem is symmetric). The value given for `scale` is  $Q_0 \rho_f / 2$  (not  $Q_0 / 2$ ). The lateral surfaces (`xpos`, `ypos` and `yneg`) are traction free. The remaining parts of the outer boundaries are subjected to roller constraints. These boundary conditions are set up through the `FieldSpecifications` section.

```
<FieldSpecifications>
  <FieldSpecification
    name="waterDensity"
```

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```

        initialCondition="1"
        setNames="{ fracture }"
        objectPath="ElementRegions"
        fieldName="water_density"
        scale="1000"/>

<FieldSpecification
  name="separableFace"
  initialCondition="1"
  setNames="{ core }"
  objectPath="faceManager"
  fieldName="isFaceSeparable"
  scale="1"/>

<FieldSpecification
  name="frac"
  initialCondition="1"
  setNames="{ fracture }"
  objectPath="faceManager"
  fieldName="ruptureState"
  scale="1"/>

<FieldSpecification
  name="zconstraint"
  objectPath="nodeManager"
  fieldName="totalDisplacement"
  component="2"
  scale="0.0"
  setNames="{ zneg, zpos }"/>

<FieldSpecification
  name="xConstraint"
  objectPath="nodeManager"
  fieldName="totalDisplacement"
  component="0"
  scale="0.0"
  setNames="{ xneg }"/>

<SourceFlux
  name="sourceTerm"
  objectPath="ElementRegions/Fracture"
  scale="-0.0000366"
  setNames="{ source }"/>
</FieldSpecifications>

```

Note that the applied traction has a negative value, due to the negative sign convention for compressive stresses in GEOSX.

The parameters used in the simulation are summarized in the following table.

Symbol	Parameter	Unit	Value
$K$	Bulk Modulus	[GPa]	4.11
$G$	Shear Modulus	[GPa]	1.2
$K_{Ic}$	Rock Toughness	[MPa.m <sup>1/2</sup> ]	1.2
$\mu$	Fluid Viscosity	[Pa.s]	97.7
$Q_0$	Injection Rate	[m <sup>3</sup> /s]	73.2x10 <sup>-9</sup>
$t_{inj}$	Injection Time	[s]	100
$h_f$	Fracture Height	[mm]	55

## Inspecting results

The following figure shows the distribution of  $\sigma_{yy}$  at  $t = 100s$  within the computational domain..

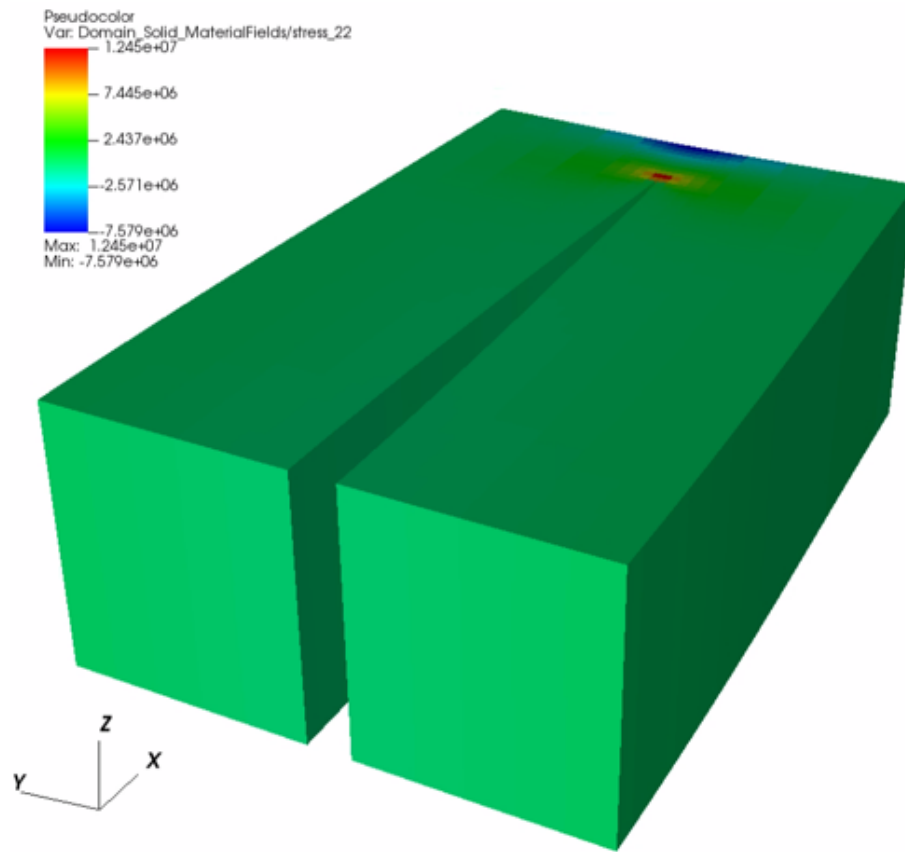


Fig. 4.25: Simulation result of  $\sigma_{xx}$  at  $t = 100s$

By running the query script `kgdValidationQueries.py`, the HDF5 output is postprocessed and temporal evolution of fracture characteristics (fluid pressure and fracture width at fluid inlet and fracure half length) are saved into a txt file `model-results.txt`, which can be used for verification and visualization:

time	wpressure	58pressure	57pressure	Laperture	area
0	0	0	0	0.0001048	
0.1	1.515e+07	0	0	0.0003145	
0.2	1.451e+07	0	0	0.0003774	
0.3	1.349e+07	0	0	0.0004194	

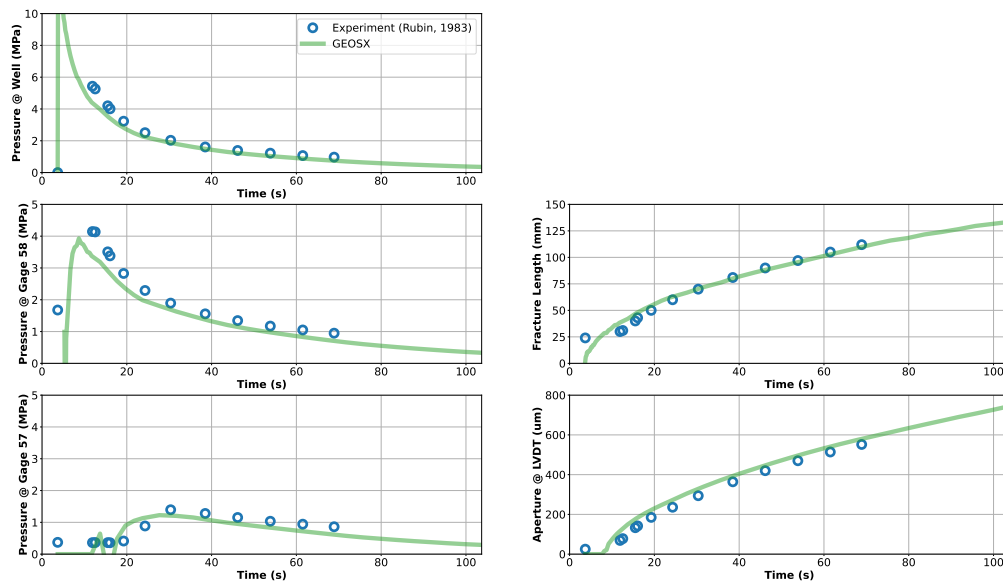
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0.4	1.183e+07	0	0	0	0.0005662
0.5	1.125e+07	0	0	0	0.0005662

Note: GEOSX python tools `geosx_xml_tools` should be installed to run the query script (See [Python Tools Setup](#) for details).

The figure below shows simulation results of the fracture extent at the end of the injection, which is generated using the visualization script `kgdValidationFigure.py`. The temporal evolution of the fracture characteristics (length, aperture and pressure) from the GEOSX simulation are extracted and compared with the experimental data gathered at specific locations. As observed, the time history plots of the modelling predictions (green curves) for the pressure at three gage locations, the fracture length, and the fracture aperture at LVDT location correlate well with the experimental data (blue circles).



## To go further

### Feedback on this example

For any feedback on this example, please submit a [GitHub issue](#) on the project's [GitHub page](#).

## Toughness-Storage-Dominated Penny Shaped Hydraulic Fracture

### Context

In this example, we simulate the growth of a radial hydraulic fracture in toughness-storage-dominated regime, a classic benchmark in hydraulic fracturing ([Settgast et al., 2016](#)). The developed fracture is characterized as a planar fracture with an elliptical cross-section perpendicular to the fracture plane and a circular fracture tip. This problem is solved using the hydrofracture solver in GEOSX. The modeling predictions on the temporal evolutions of the fracture characteristics (length, aperture, and pressure) are verified against the analytical solutions ([Savitski and Detournay, 2002](#)).

## Input file

This example uses no external input files. Everything we need is contained within two GEOSX input files:

```
inputFiles/hydraulicFracturing/pennyShapedToughnessDominated_base.xml
```

```
inputFiles/hydraulicFracturing/pennyShapedToughnessDominated_benchmark.xml
```

Python scripts for post-processing and visualizing the simulation results are also prepared:

```
inputFiles/hydraulicFracturing/scripts/hydrofractureQueries.py
```

```
inputFiles/hydraulicFracturing/scripts/hydrofractureFigure.py
```

## Description of the case

We model a radial fracture emerging from a point source and forming a perfect circular shape in an infinite, isotropic, and homogenous elastic domain. As with the KGD problem, we simplify the model to a radial fracture in a toughness-storage-dominated propagation regime. For toughness-dominated fractures, more work is spent on splitting the intact rock than on moving the fracturing fluid. Storage-dominated propagation occurs if most of the fracturing fluid is contained within the propagating fracture. In this analysis, incompressible fluid with ultra-low viscosity ( $0.001cp$ ) and medium rock toughness ( $3.0MPa\sqrt{m}$ ) are specified. In addition, an impermeable fracture surface is assumed to eliminate the effect of fluid leak-off. This way, the GEOSX simulations represent cases within the valid range of the toughness-storage-dominated assumptions.

In this model, the injected fluid within the fracture follows the lubrication equation resulting from mass conservation and Poiseuille's law. The fracture propagates by creating new surfaces if the stress intensity factor exceeds the local rock toughness  $K_{IC}$ . By symmetry, the simulation is reduced to a quarter-scale to save computational cost. For verification purposes, a plane strain deformation is considered in the numerical model.

In this example, we set up and solve a hydraulic fracture model to obtain the temporal solutions of the fracture radius  $R$ , the net pressure  $p_0$  and the fracture aperture  $w_0$  at the injection point for the penny-shaped fracture developed in this toughness-storage-dominated regime. The numerical predictions from GEOSX are then compared with the corresponding asymptotic solutions (Savitski and Detournay, 2002):

$$R(t) = 0.8546 \left( \frac{E_p^2 Q_0^2 t^2}{K_p^2} \right)^{1/5}$$

$$w_0(t) = 0.6537 \left( \frac{K_p^4 Q_0 t}{E_p^4} \right)^{1/5}$$

$$p_0(t) = 0.3004 \left( \frac{K_p^6}{E_p Q_0 t} \right)^{1/5}$$

where the plane modulus  $E_p$  is related to Young's modulus  $E$  and Poisson's ratio  $\nu$ :

$$E_p = \frac{E}{1 - \nu^2}$$

The term  $K_p$  is proportional to the rock toughness  $K_{IC}$ :

$$K_p = \frac{8}{\sqrt{2\pi}} K_{IC}$$

For this example, we focus on the Mesh, the Constitutive, and the FieldSpecifications tags.

## Mesh

The following figure shows the mesh used in this problem.

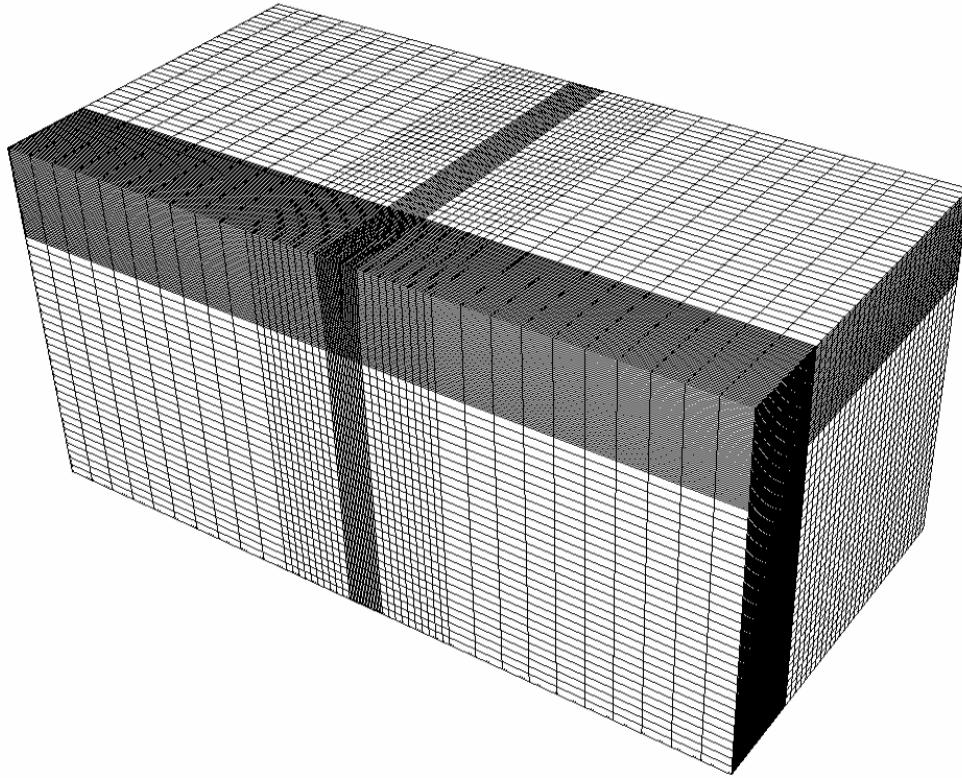


Fig. 4.26: Generated mesh

We use the internal mesh generator to create a computational domain ( $400\text{ m} \times 400\text{ m} \times 800\text{ m}$ ), as parametrized in the `InternalMesh` XML tag. The structured mesh contains  $80 \times 80 \times 60$  eight-node brick elements in the  $x$ ,  $y$ , and  $z$  directions respectively. Such eight-node hexahedral elements are defined as `C3D8` `elementTypes`, and their collection forms a mesh with one group of cell blocks named here `cb1`. Local refinement is performed for the elements in the vicinity of the fracture plane.

Note that the domain size in the direction perpendicular to the fracture plane, i.e.  $z$ -axis, must be at least ten times of the final fracture radius to minimize possible boundary effects.

```
<Mesh>
  <InternalMesh
    name="mesh1"
    elementTypes="{ C3D8 }"
    xCoords="{ 0, 100, 200, 400 }"
    yCoords="{ 0, 100, 200, 400 }"
    zCoords="{ -400, -100, -20, 20, 100, 400 }"
    nx="{ 50, 10, 20 }"
    ny="{ 50, 10, 20 }"
    nz="{ 10, 10, 20, 10, 10 }"
    cellBlockNames="{ cb1 }"/>
  </InternalMesh>
</Mesh>
```

The fracture plane is defined by a nodeset occupying a small region within the computation domain, where the fracture tends to open and propagate upon fluid injection:

```

<Box
  name="core"
  xMin="{ -500.1, -500.1, -0.1 }"
  xMax="{ 500.1, 500.1, 0.1 }"/>

```

## Solid mechanics solver

GEOSX is a multi-physics platform. Different combinations of physics solvers available in the code can be applied in different regions of the domain and be functional at different stages of the simulation. The `Solvers` tag in the XML file is used to list and parameterize these solvers.

Three elementary solvers are combined in the solver `Hydrofracture` to model the coupling between fluid flow within the fracture, rock deformation, fracture deformation and propagation:

```

<Hydrofracture
  name="hydrofracture"
  solidSolverName="lagsolve"
  flowSolverName="SinglePhaseFlow"
  surfaceGeneratorName="SurfaceGen"
  couplingTypeOption="FIM"
  logLevel="1"
  targetRegions="{ Fracture }"
  contactRelationName="fractureContact"
  maxNumResolves="5"
  initialDt="0.1">
  <NonlinearSolverParameters
    newtonTol="1.0e-4"
    newtonMaxIter="50"
    logLevel="1"/>
  <LinearSolverParameters
    solverType="gmres"
    preconditionerType="mgr"
    logLevel="1"
    krylovAdaptiveTol="1"/>
</Hydrofracture>

```

- Rock and fracture deformation are modeled by the solid mechanics solver `SolidMechanicsLagrangianSSLE`. In this solver, we define `targetRegions` that includes both the continuum region and the fracture region. The name of the contact constitutive behavior is specified in this solver by the `contactRelationName`.

```

<SolidMechanicsLagrangianSSLE
  name="lagsolve"
  timeIntegrationOption="QuasiStatic"
  logLevel="1"
  discretization="FE1"
  targetRegions="{ Domain, Fracture }"
  contactRelationName="fractureContact">
  <NonlinearSolverParameters
    newtonTol="1.0e-6"/>
  <LinearSolverParameters
    solverType="gmres"
    krylovTol="1.0e-10"/>
</SolidMechanicsLagrangianSSLE>

```

- The single-phase fluid flow inside the fracture is solved by the finite volume method in the solver

SinglePhaseFVM.

```
<SinglePhaseFVM
  name="SinglePhaseFlow"
  logLevel="1"
  discretization="singlePhaseTPFA"
  targetRegions="{ Fracture }"
  inputFluxEstimate="1.0">
  <NonlinearSolverParameters
    newtonTol="1.0e-5"
    newtonMaxIter="10"/>
  <LinearSolverParameters
    solverType="gmres"
    krylovTol="1.0e-12"/>
</SinglePhaseFVM>
```

- The solver `SurfaceGenerator` defines the fracture region and rock toughness `rockToughness="3.0e6"`. With `nodeBasedSIF="1"`, a node-based Stress Intensity Factor (SIF) calculation is chosen for the fracture propagation criterion.

```
<SurfaceGenerator
  name="SurfaceGen"
  targetRegions="{ Domain }"
  nodeBasedSIF="1"
  rockToughness="3.0e6"
  mpiCommOrder="1"/>
```

## Constitutive laws

For this problem, a homogeneous and isotropic domain with one solid material is assumed. Its mechanical properties and associated fluid rheology are specified in the Constitutive section. `ElasticIsotropic` model is used to describe the mechanical behavior of rock when subjected to fluid injection. The single-phase fluid model `CompressibleSinglePhaseFluid` is selected to simulate the response of water upon fracture propagation.

```
<Constitutive>
  <CompressibleSinglePhaseFluid
    name="water"
    defaultDensity="1000"
    defaultViscosity="1.0e-6"
    referencePressure="0.0"
    compressibility="5e-13"
    referenceViscosity="1.0e-6"
    viscosibility="0.0"/>

  <ElasticIsotropic
    name="rock"
    defaultDensity="2700"
    defaultBulkModulus="20.0e9"
    defaultShearModulus="12.0e9"/>

  <CompressibleSolidParallelPlatesPermeability
    name="fractureFilling"
    solidModelName="nullSolid"
    porosityModelName="fracturePorosity"
    permeabilityModelName="fracturePerm"/>
```

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```

<NullModel
  name="nullSolid"/>

<PressurePorosity
  name="fracturePorosity"
  defaultReferencePorosity="1.00"
  referencePressure="0.0"
  compressibility="0.0"/>

<ParallelPlatesPermeability
  name="fracturePerm"/>

<FrictionlessContact
  name="fractureContact"
  penaltyStiffness="1.0e0"
  apertureTableName="apertureTable"/>
</Constitutive>

```

All constitutive parameters such as density, viscosity, bulk modulus, and shear modulus are specified in the International System of Units.

## Time history function

In the Tasks section, PackCollection tasks are defined to collect time history information from fields. Either the entire field or specified named sets of indices in the field can be collected. In this example, pressureCollection, apertureCollection, hydraulicApertureCollection and areaCollection are specified to output the time history of fracture characteristics (pressure, width and area). objectPath="ElementRegions/Fracture/FractureSubRegion" indicates that these PackCollection tasks are applied to the fracture element subregion.

```

<Tasks>
  <PackCollection
    name="pressureCollection"
    objectPath="ElementRegions/Fracture/FractureSubRegion"
    fieldName="pressure"/>

  <PackCollection
    name="apertureCollection"
    objectPath="ElementRegions/Fracture/FractureSubRegion"
    fieldName="elementAperture"/>

  <PackCollection
    name="hydraulicApertureCollection"
    objectPath="ElementRegions/Fracture/FractureSubRegion"
    fieldName="hydraulicAperture"/>

  <PackCollection
    name="areaCollection"
    objectPath="ElementRegions/Fracture/FractureSubRegion"
    fieldName="elementArea"/>
</Tasks>

```

These tasks are triggered using the Event manager with a PeriodicEvent defined for the recurring tasks. GEOSX writes one file named after the string defined in the filename keyword and formatted as a HDF5 file (pennyShapedToughnessDominated\_output.hdf5). This TimeHistory file contains the collected time his-

tory information from specified time history collector. This file includes datasets for the simulation time, fluid pressure, element aperture, hydraulic aperture and element area for the propagating hydraulic fracture. A Python script is prepared to read and query any specified subset of the time history data for verification and visualization.

## Initial and boundary conditions

The next step is to specify fields, including:

- The initial values: the `waterDensity`, `separableFace` and the `ruptureState` of the propagating fracture have to be initialized,
- The boundary conditions: fluid injection rates and the constraints of the outer boundaries have to be set.

In this example, a mass injection rate `SourceFlux` (`scale="-6.625"`) is applied at the surfaces of the initial fracture. Only one fourth of the total injection rate is defined in this boundary condition because only a quarter of the fracture is modeled (the problem is symmetric). The value given for `scale` is  $Q_0\rho_f/4$  (not  $Q_0/4$ ). All the outer boundaries are subject to roller constraints. These boundary conditions are set through the `FieldSpecifications` section.

```
<FieldSpecifications>
  <FieldSpecification
    name="waterDensity"
    initialCondition="1"
    setNames="{ fracture }"
    objectPath="ElementRegions"
    fieldName="water_density"
    scale="1000"/>

  <FieldSpecification
    name="separableFace"
    initialCondition="1"
    setNames="{ core }"
    objectPath="faceManager"
    fieldName="isFaceSeparable"
    scale="1"/>

  <FieldSpecification
    name="frac"
    initialCondition="1"
    setNames="{ fracture }"
    objectPath="faceManager"
    fieldName="ruptureState"
    scale="1"/>

  <FieldSpecification
    name="yconstraint"
    objectPath="nodeManager"
    fieldName="totalDisplacement"
    component="1"
    scale="0.0"
    setNames="{ yneg, ypos }"/>

  <FieldSpecification
    name="zconstraint"
    objectPath="nodeManager"
    fieldName="totalDisplacement"
    component="2"
```

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```

scale="0.0"
setNames="{ zneg, zpos }"/>

<FieldSpecification
  name="xconstraint"
  objectPath="nodeManager"
  fieldName="totalDisplacement"
  component="0"
  scale="0.0"
  setNames="{ xneg, xpos }"/>

<SourceFlux
  name="sourceTerm"
  objectPath="ElementRegions/Fracture"
  scale="-6.625"
  setNames="{ source }"/>
</FieldSpecifications>

```

The parameters used in the simulation are summarized in the following table.

Symbol	Parameter	Unit	Value
$K$	Bulk Modulus	[GPa]	20.0
$G$	Shear Modulus	[GPa]	12.0
$K_{IC}$	Rock Toughness	[MPa.m <sup>1/2</sup> ]	3.0
$\mu$	Fluid Viscosity	[Pa.s]	1.0x10 <sup>-6</sup>
$Q_0$	Injection Rate	[m <sup>3</sup> /s]	0.0265
$t_{inj}$	Injection Time	[s]	400

## Inspecting results

The following figure shows the distribution of  $\sigma_{zz}$  at  $t = 400s$  within the computational domain..

First, by running the query script

```
python ./hydrofractureQueries.py pennyShapedToughnessDominated
```

the HDF5 output is postprocessed and temporal evolution of fracture characteristics (fluid pressure and fracture width at fluid inlet and fracture radius) are saved into a txt file `model-results.txt`, which can be used for verification and visualization:

```

[['      time', ' pressure', ' aperture', '   length']]
 2 8.207e+05 0.0004661    8.137
 4 6.799e+05 0.0005258   10.59
 6 7.082e+05 0.0006183   11.94
 8 6.07e+05 0.0006163   13.73
10 6.32e+05 0.0006827   14.45

```

Note: GEOSX python tools `geosx_xml_tools` should be installed to run the query script (See [Python Tools Setup](#) for details).

Next, the figure below compares the asymptotic solutions (curves) and the GEOSX simulation results (markers) for this analysis, which is generated using the visualization script:

```
python ./pennyShapedToughnessDominatedFigure.py
```

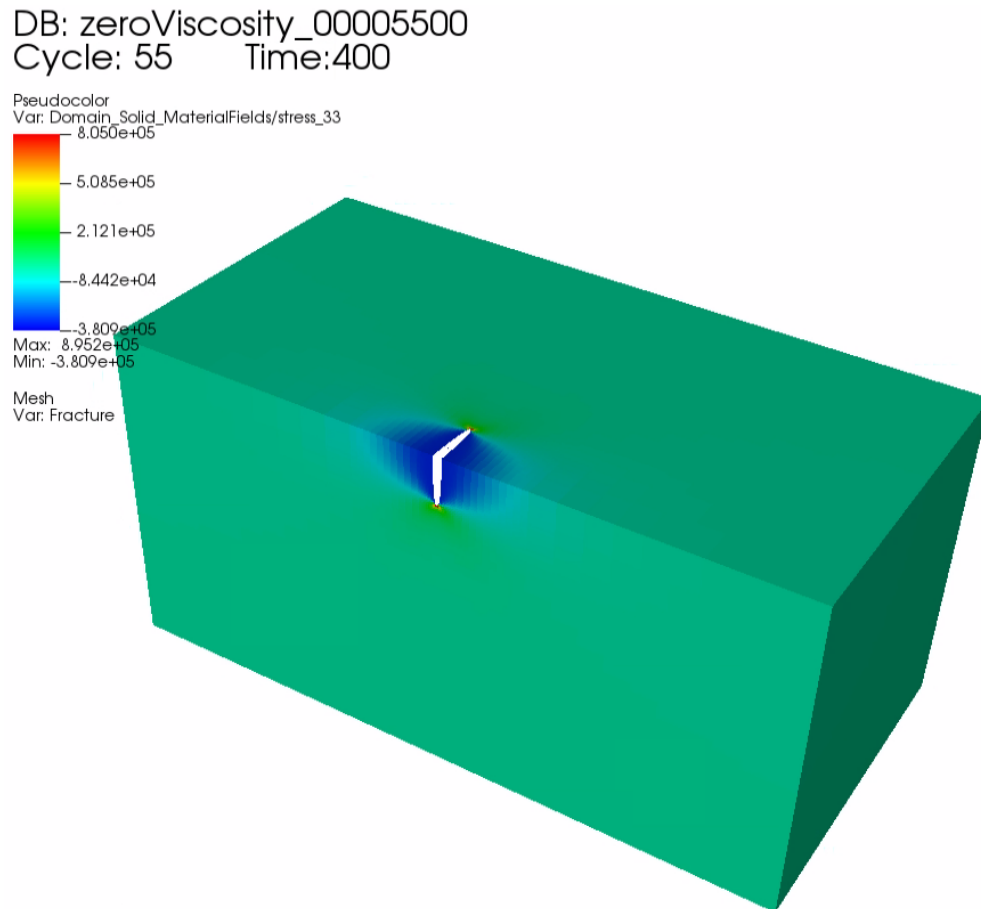
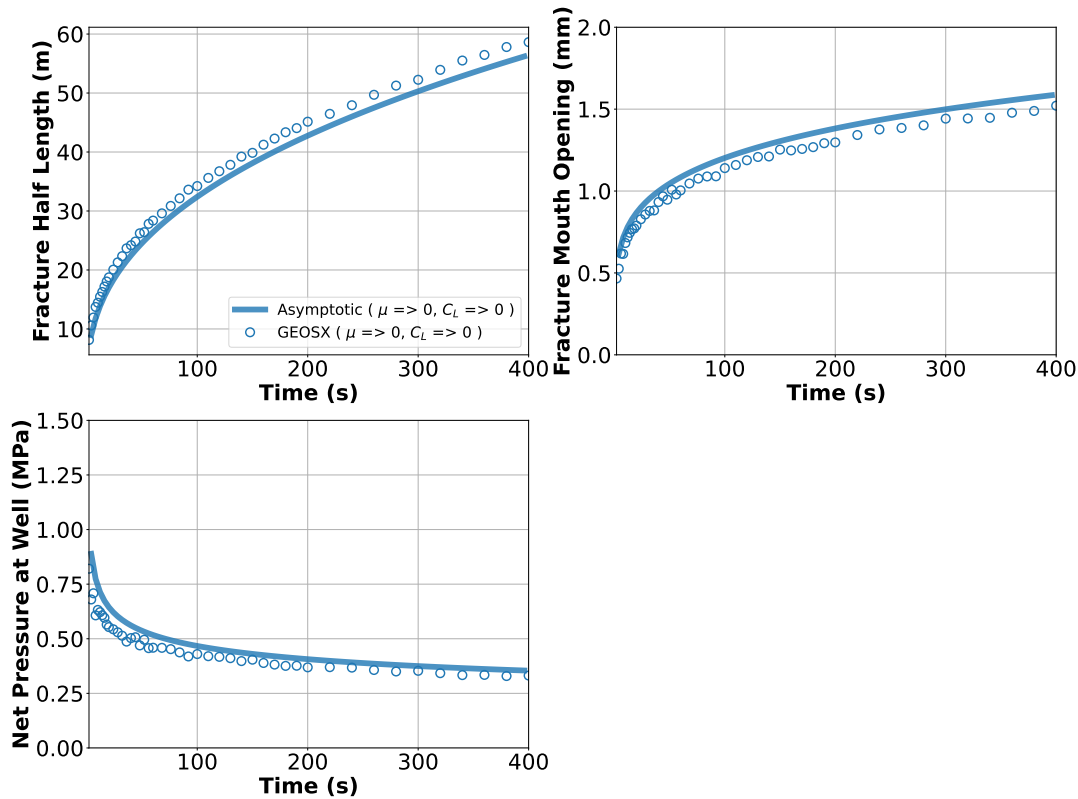


Fig. 4.27: Simulation result of  $\sigma_{zz}$  at  $t = 400s$

The time history plots of fracture radius, fracture aperture and fluid pressure at the point source match the asymptotic solutions, confirming the accuracy of GEOSX simulations.



### To go further

#### Feedback on this example

For any feedback on this example, please submit a [GitHub issue](#) on the project's [GitHub](#) page.

## Viscosity-Storage-Dominated Penny Shaped Hydraulic Fracture

### Context

In this example, we simulate the propagation of a radial hydraulic fracture in viscosity-storage-dominated regime, another classic benchmark in hydraulic fracturing (Settgast et al., 2016). The fracture develops as a planar fracture with an elliptical cross-section perpendicular to the fracture plane and a circular fracture tip. Unlike the toughness-storage-dominated fractures, fluid frictional loss during the transport of viscous fracturing fluids governs the growth of viscosity-storage-dominated fractures. We solve this problem using the hydrofracture solver in GEOSX. We simulate the change in length, aperture, and pressure of the fracture, and compare them against the corresponding analytical solutions (Savitski and Detournay, 2002).

### Input file

This example uses no external input files. Everything we need is contained within two GEOSX input files:

```
inputFiles/hydraulicFracturing/pennyShapedViscosityDominated_base.xml
```

```
inputFiles/hydraulicFracturing/pennyShapedViscosityDominated_benchmark.xml
```

Python scripts for post-processing and visualizing the simulation results are also prepared:

```
inputFiles/hydraulicFracturing/scripts/hydrofractureQueries.py
```

```
inputFiles/hydraulicFracturing/scripts/hydrofractureFigure.py
```

## Description of the case

We model a radial fracture emerging from a point source and forming a perfect circular shape in an infinite, isotropic, and homogenous elastic domain. As with the viscosity-dominated KGD problem, we restrict the model to a radial fracture developed in a viscosity-storage-dominated propagation regime. For viscosity-dominated fractures, more energy is applied to move the fracturing fluid than to split the intact rock. If we neglect fluid leak-off, the storage-dominated propagation occurs from most of the injected fluid confined within the opened surfaces. We use a low rock toughness ( $0.3\text{MPa}\sqrt{\text{m}}$ ), and the slickwater we inject has a constant viscosity value ( $1.0\text{cp}$ ) and zero compressibility. In addition, we assume that the fracture surfaces are impermeable, thus eliminating fluid leak-off. With this configuration, our GEOSX simulations meet the requirements of the viscosity-storage-dominated assumptions.

The fluid injected in the fracture follows the lubrication equation resulting from mass conservation and Poiseuille's law. The fracture propagates by creating new surfaces if the stress intensity factor exceeds the local rock toughness  $K_{IC}$ . By symmetry, the simulation is reduced to a quarter-scale to save computational cost. For verification purposes, a plane strain deformation is considered in the numerical model.

We set up and solve a hydraulic fracture model to obtain the evolution with time of the fracture radius  $R$ , the net pressure  $p_0$  and the fracture aperture  $w_0$  at the injection point for the penny-shaped fracture developed in viscosity-storage-dominated regime. [Savitski and Detournay \(2002\)](#) presented the corresponding asymptotic solutions, used here to validate the results of our GEOSX simulations:

$$\begin{aligned} R(t) &= 0.6955 \left( \frac{E_p Q_0^3 t^4}{M_p} \right)^{1/9} \\ w_0(t) &= 1.1977 \left( \frac{M_p^2 Q_0^3 t}{E_p^2} \right)^{1/9} \\ p_0(\Pi, t) &= \Pi_{mo}(\xi) \left( \frac{E_p^2 M_p}{t} \right)^{1/3} \end{aligned}$$

where the plane modulus  $E_p$  is related to Young's modulus  $E$  and Poisson's ratio  $\nu$ :

$$E_p = \frac{E}{1 - \nu^2}$$

The term  $M_p$  is proportional to the fluid viscosity  $\mu$ :

$$M_p = 12\mu$$

The viscosity scaling function  $\Pi_{mo}$  is given as:

$$\Pi_{mo}(\xi) = A_1 \left[ 2.479 - \frac{2}{3(1 - \xi)^{1/3}} \right] - B \left[ \ln\left(\frac{\xi}{2}\right) + 1 \right]$$

with  $A_1 = 0.3581$ ,  $B = 0.09269$ ,  $c_1 = 0.6846$ ,  $c_2 = 0.07098$ , and  $\xi = r/R(t)$  denoting a dimensionless radial coordinate along the fracture.

For this example, we focus on the Mesh, the Constitutive, and the FieldSpecifications tags.

## Mesh

The following figure shows the mesh used in this problem.

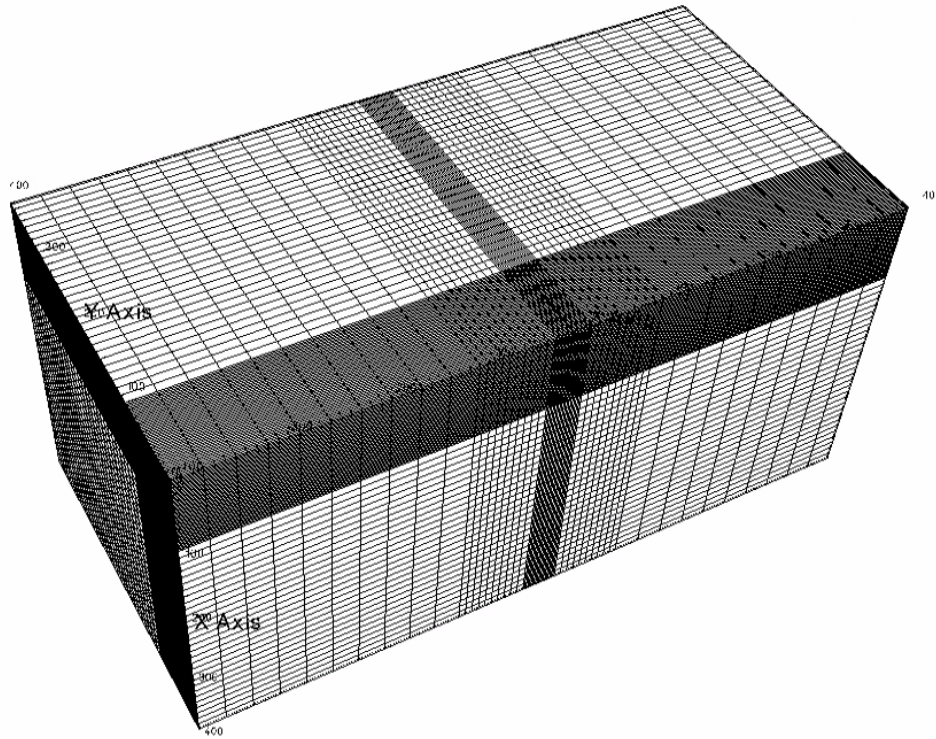


Fig. 4.28: Generated mesh

We use the internal mesh generator to create a computational domain ( $400\text{ m} \times 400\text{ m} \times 800\text{ m}$ ), as parametrized in the `InternalMesh` XML tag. The structured mesh contains  $80 \times 80 \times 60$  eight-node brick elements in the x, y, and z directions respectively. Such eight-node hexahedral elements are defined as `C3D8` `elementTypes`, and their collection forms a mesh with one group of cell blocks named here `cb1`. Local refinement is performed for the elements in the vicinity of the fracture plane.

Note that the domain size in the direction perpendicular to the fracture plane, i.e. z-axis, must be at least ten times of the final fracture radius to minimize possible boundary effects.

```
<Mesh>
  <InternalMesh
    name="mesh1"
    elementTypes="{ C3D8 }"
    xCoords="{ 0, 100, 200, 400 }"
    yCoords="{ 0, 100, 200, 400 }"
    zCoords="{ -400, -100, -20, 20, 100, 400 }"
    nx="{ 50, 10, 20 }"
    ny="{ 50, 10, 20 }"
    nz="{ 10, 10, 20, 10, 10 }"
    cellBlockNames="{ cb1 }"/>
  </InternalMesh>
</Mesh>
```

The fracture plane is defined by a nodeset occupying a small region within the computation domain, where the fracture tends to open and propagate upon fluid injection:

```
<Box
  name="core"
  xMin="{ -500.1, -500.1, -0.1 }"
  xMax="{ 500.1, 500.1, 0.1 }"/>
```

## Solid mechanics solver

GEOSX is a multi-physics platform. Different combinations of physics solvers available in the code can be applied in different regions of the domain and be functional at different stages of the simulation. The `Solvers` tag in the XML file is used to list and parameterize these solvers.

Three elementary solvers are combined in the solver `Hydrofracture` to model the coupling between fluid flow within the fracture, rock deformation, fracture deformation and propagation:

```
<Hydrofracture
  name="hydrofracture"
  solidSolverName="lagsolve"
  flowSolverName="SinglePhaseFlow"
  surfaceGeneratorName="SurfaceGen"
  couplingTypeOption="FIM"
  logLevel="1"
  targetRegions="{ Fracture }"
  contactRelationName="fractureContact"
  maxNumResolves="1"
  initialDt="0.1">
  <NonlinearSolverParameters
    newtonTol="1.0e-4"
    newtonMaxIter="10"
    maxTimeStepCuts="5"
    logLevel="1"/>
  <LinearSolverParameters
    solverType="gmres"
    preconditionerType="mgr"
    logLevel="1"
    krylovAdaptiveTol="1"/>
</Hydrofracture>
```

- Rock and fracture deformation are modeled by the solid mechanics solver `SolidMechanicsLagrangianSSLE`. In this solver, we define `targetRegions` that includes both the continuum region and the fracture region. The name of the contact constitutive behavior is specified in this solver by the `contactRelationName`.

```
<SolidMechanicsLagrangianSSLE
  name="lagsolve"
  timeIntegrationOption="QuasiStatic"
  logLevel="1"
  discretization="FE1"
  targetRegions="{ Domain, Fracture }"
  contactRelationName="fractureContact">
  <NonlinearSolverParameters
    newtonTol="1.0e-6"/>
  <LinearSolverParameters
    solverType="gmres"
    krylovTol="1.0e-10"/>
</SolidMechanicsLagrangianSSLE>
```



- The single-phase fluid flow inside the fracture is solved by the finite volume method in the solver SinglePhaseFVM.

```
<SinglePhaseFVM
  name="SinglePhaseFlow"
  logLevel="1"
  discretization="singlePhaseTPFA"
  targetRegions="{ Fracture }"
  inputFluxEstimate="1.0">
  <NonlinearSolverParameters
    newtonTol="1.0e-5"
    newtonMaxIter="10"/>
  <LinearSolverParameters
    solverType="gmres"
    krylovTol="1.0e-12"/>
</SinglePhaseFVM>
```

- The solver SurfaceGenerator defines the fracture region and rock toughness rockToughness="0.3e6". With nodeBasedSIF="1", a node-based Stress Intensity Factor (SIF) calculation is chosen for the fracture propagation criterion.

```
<SurfaceGenerator
  name="SurfaceGen"
  targetRegions="{ Domain }"
  nodeBasedSIF="1"
  rockToughness="0.3e6"
  mpiCommOrder="1"/>
```

## Constitutive laws

For this problem, a homogeneous and isotropic domain with one solid material is assumed. Its mechanical properties and associated fluid rheology are specified in the Constitutive section. The ElasticIsotropic model is used to describe the mechanical behavior of rock when subjected to fluid injection. The single-phase fluid model CompressibleSinglePhaseFluid is selected to simulate the response of water upon fracture propagation.

```
<Constitutive>
  <CompressibleSinglePhaseFluid
    name="water"
    defaultDensity="1000"
    defaultViscosity="0.001"
    referencePressure="0.0"
    compressibility="5e-12"
    referenceViscosity="1.0e-3"
    viscosibility="0.0"/>

  <ElasticIsotropic
    name="rock"
    defaultDensity="2700"
    defaultBulkModulus="20.0e9"
    defaultShearModulus="12.0e9"/>

  <CompressibleSolidParallelPlatesPermeability
    name="fractureFilling"
    solidModelName="nullSolid"
    porosityModelName="fracturePorosity"
    permeabilityModelName="fracturePerm"/>
```

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```

<NullModel
  name="nullSolid"/>

<PressurePorosity
  name="fracturePorosity"
  defaultReferencePorosity="1.00"
  referencePressure="0.0"
  compressibility="0.0"/>

<ParallelPlatesPermeability
  name="fracturePerm"/>

<FrictionlessContact
  name="fractureContact"
  penaltyStiffness="1.0e0"
  apertureTableName="apertureTable"/>
</Constitutive>

```

All constitutive parameters such as density, viscosity, bulk modulus, and shear modulus are specified in the International System of Units.

### Time history function

In the Tasks section, PackCollection tasks are defined to collect time history information from fields. Either the entire field or specified named sets of indices in the field can be collected. In this example, pressureCollection, apertureCollection, hydraulicApertureCollection and areaCollection are specified to output the time history of fracture characteristics (pressure, width and area). objectPath="ElementRegions/Fracture/FractureSubRegion" indicates that these PackCollection tasks are applied to the fracture element subregion.

```

<Tasks>
  <PackCollection
    name="pressureCollection"
    objectPath="ElementRegions/Fracture/FractureSubRegion"
    fieldName="pressure"/>

  <PackCollection
    name="apertureCollection"
    objectPath="ElementRegions/Fracture/FractureSubRegion"
    fieldName="elementAperture"/>

  <PackCollection
    name="hydraulicApertureCollection"
    objectPath="ElementRegions/Fracture/FractureSubRegion"
    fieldName="hydraulicAperture"/>

  <PackCollection
    name="areaCollection"
    objectPath="ElementRegions/Fracture/FractureSubRegion"
    fieldName="elementArea"/>
</Tasks>

```

These tasks are triggered using the Event manager with a PeriodicEvent defined for the recurring tasks. GEOSX writes one file named after the string defined in the filename keyword and formatted as a HDF5 file

(pennyShapedViscosityDominated\_output.hdf5). This TimeHistory file contains the collected time history information from specified time history collector. This file includes datasets for the simulation time, fluid pressure, element aperture, hydraulic aperture and element area for the propagating hydraulic fracture. A Python script is prepared to read and query any specified subset of the time history data for verification and visualization.

## Initial and boundary conditions

Next, we specify initial and boundary conditions:

- Initial values: the waterDensity, separableFace and the ruptureState of the propagating fracture have to be initialized,
- Boundary conditions: fluid injection rates and the constraints of the outer boundaries have to be set.

In this example, a mass injection rate SourceFlux (scale="-6.625") is applied at the surfaces of the initial fracture. Only one fourth of the total injection rate is defined in this boundary condition because only a quarter of the fracture is modeled (the problem is symmetric). The value given for scale is  $Q_0\rho_f/4$  (not  $Q_0/4$ ). All the outer boundaries are subject to roller constraints. These boundary conditions are set through the FieldSpecifications section.

```
<FieldSpecifications>
  <FieldSpecification
    name="waterDensity"
    initialCondition="1"
    setNames="{ fracture }"
    objectPath="ElementRegions"
    fieldName="water_density"
    scale="1000"/>

  <FieldSpecification
    name="separableFace"
    initialCondition="1"
    setNames="{ core }"
    objectPath="faceManager"
    fieldName="isFaceSeparable"
    scale="1"/>

  <FieldSpecification
    name="frac"
    initialCondition="1"
    setNames="{ fracture }"
    objectPath="faceManager"
    fieldName="ruptureState"
    scale="1"/>

  <FieldSpecification
    name="yconstraint"
    objectPath="nodeManager"
    fieldName="totalDisplacement"
    component="1"
    scale="0.0"
    setNames="{ yneg, ypos }"/>

  <FieldSpecification
    name="zconstraint"
    objectPath="nodeManager"
    fieldName="totalDisplacement"
```

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```

    component="2"
    scale="0.0"
    setNames="{ zneg, zpos }"/>

<FieldSpecification
  name="xconstraint"
  objectPath="nodeManager"
  fieldName="totalDisplacement"
  component="0"
  scale="0.0"
  setNames="{ xneg, xpos }"/>

<SourceFlux
  name="sourceTerm"
  objectPath="ElementRegions/Fracture"
  scale="-6.625"
  setNames="{ source }"/>
</FieldSpecifications>

```

The parameters used in the simulation are summarized in the following table.

Symbol	Parameter	Unit	Value
$K$	Bulk Modulus	[GPa]	20.0
$G$	Shear Modulus	[GPa]	12.0
$K_{IC}$	Rock Toughness	[MPa.m <sup>1/2</sup> ]	0.3
$\mu$	Fluid Viscosity	[Pa.s]	1.0x10 <sup>-3</sup>
$Q_0$	Injection Rate	[m <sup>3</sup> /s]	0.0265
$t_{inj}$	Injection Time	[s]	400

## Inspecting results

The following figure shows the distribution of  $\sigma_{zz}$  at  $t = 400s$  within the computational domain..

First, by running the query script

```
python ./hydrofractureQueries.py pennyShapedViscosityDominated
```

the HDF5 output is postprocessed and temporal evolution of fracture characteristics (fluid pressure and fracture width at fluid inlet and fracture radius) are saved into a txt file `model-results.txt`, which can be used for verification and visualization:

```

[['      time', '      pressure', '      aperture', '      length']]
 2 1.654e+06 0.0006768      8.137
 4 1.297e+06 0.000743      10.59
 6 1.115e+06 0.0007734     12.36
 8 1.005e+06 0.0007918     13.73
10 9.482e+05 0.0008189     15.14

```

Note: GEOSX python tools `geosx_xml_tools` should be installed to run the query script (See [Python Tools Setup](#) for details).

Next, GEOSX simulation results (markers) and asymptotic solutions (curves) for the case with viscosity-storage dominated assumptions are plotted together in the following figure, which is generated using the visualization script:

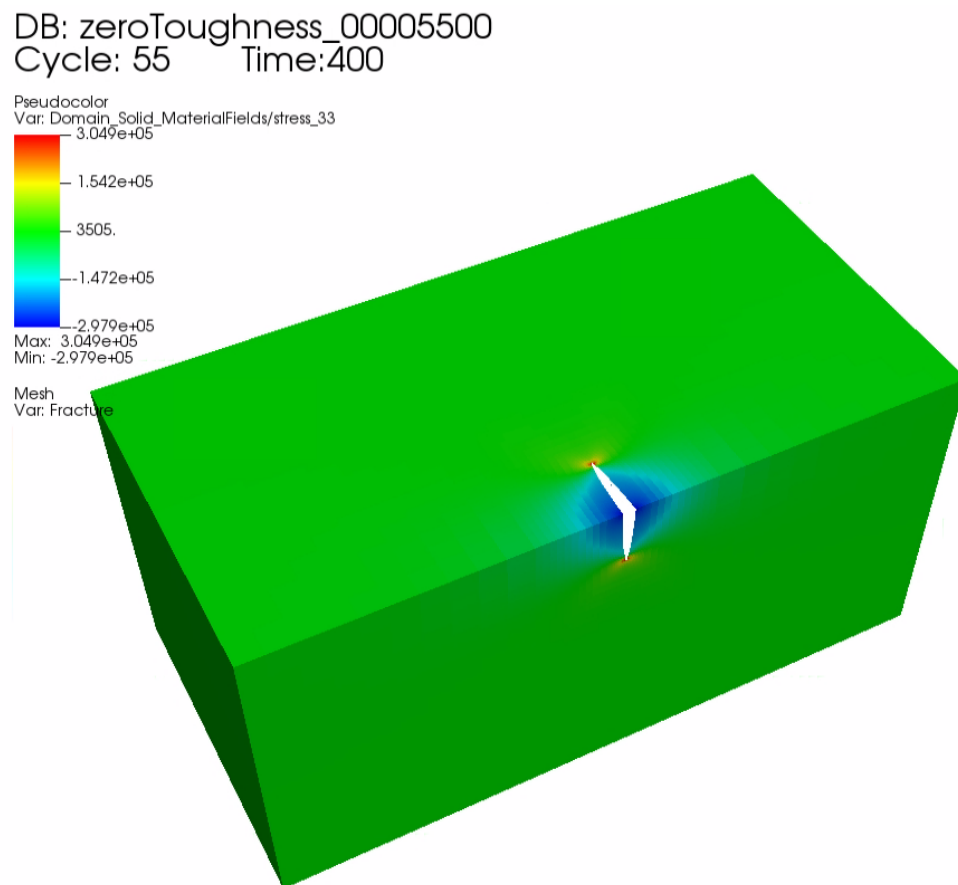
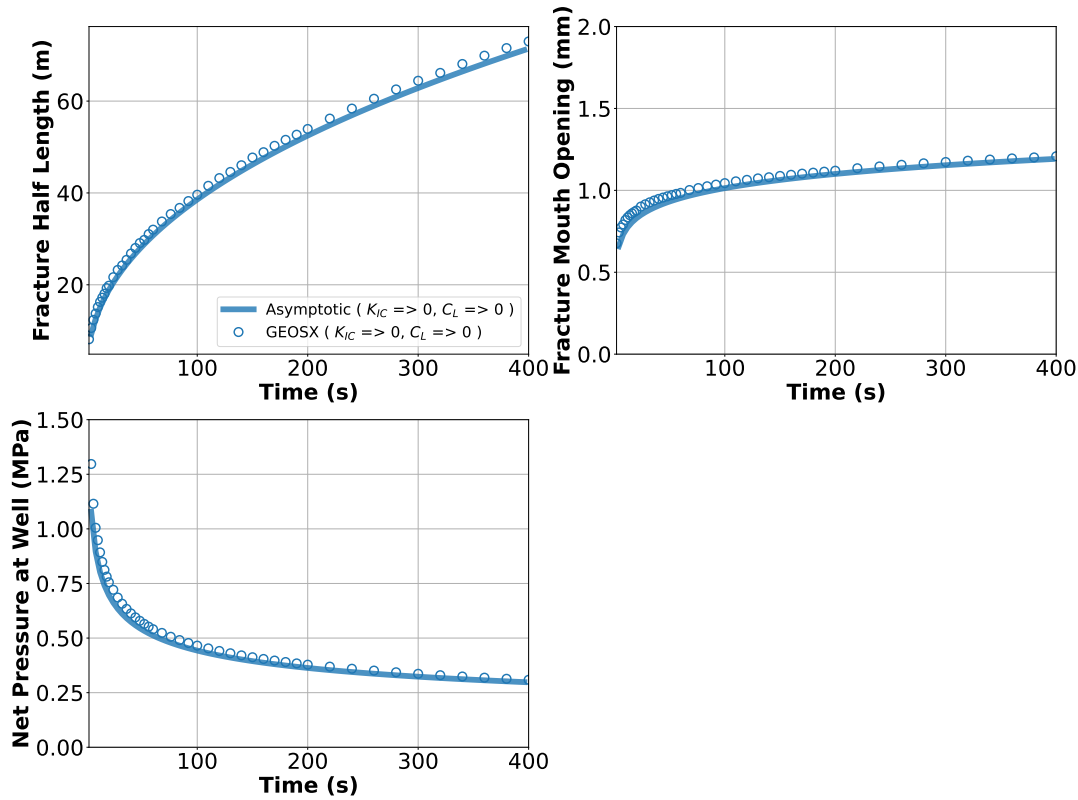


Fig. 4.29: Simulation result of  $\sigma_{zz}$  at  $t = 400s$

```
python ./pennyShapedViscosityDominatedFigure.py
```

As seen, GEOSX predictions of the temporal evolution of fracture radius, wellbore aperture and pressure at fluid inlet are nearly identical to the asymptotic solutions.



## To go further

### Feedback on this example

For any feedback on this example, please submit a [GitHub issue](#) on the project's [GitHub page](#).

## Viscosity-Storage-Dominated PKN Hydraulic Fracture

### Context

In this example, we simulate the propagation of a Perkins–Kern–Nordgren (PKN) fracture in a viscosity-storage-dominated regime, a classic benchmark in hydraulic fracturing. The developed planar fracture displays an elliptical vertical cross-section. Unlike KGD and penny-shaped fractures, the growth height of a PKN fracture is constrained by mechanical barriers (such as bedding layers, sedimentary laminations, or weak interfaces), thus promoting lateral propagation. This problem is solved using the hydrofracture solver in GEOSX to obtain the temporal evolutions of the fracture characteristics (length, aperture, and pressure). We validate these simulated values against existing analytical solutions (Kovalyshen and Detournay, 2010; Economides and Nolte, 2000).

## Input file

This example uses no external input files. Everything we need is contained within two GEOSX input files:

```
inputFiles/hydraulicFracturing/pknViscosityDominated_base.xml
```

```
inputFiles/hydraulicFracturing/pknViscosityDominated_benchmark.xml
```

Python scripts for post-processing and visualizing the simulation results are also prepared:

```
inputFiles/hydraulicFracturing/scripts/hydrofractureQueries.py
```

```
inputFiles/hydraulicFracturing/scripts/hydrofractureFigure.py
```

## Description of the case

In this example, a hydraulic fracture initiates and propagates from the center of a 20m-thick layer. This layer is homogeneous and bounded by neighboring upper and lower layers. For viscosity-dominated fractures, more energy is necessary to move the fracturing fluid than to split the intact rock. If fluid leak-off is neglected, storage-dominated propagation occurs with most of the injected fluid confined within the open surfaces. To meet the requirements of the viscosity-storage-dominated assumptions, impermeable domain (no fluid leak-off), incompressible fluid with constant viscosity (1.0cp) and ultra-low rock toughness ( $0.1MPa\sqrt{m}$ ) are chosen in the GEOSX simulation. With these parameters, the fracture stays within the target layer; it extends horizontally and meets the conditions of the PKN fracture in a viscosity-storage-dominated regime.

We assume that the fluid injected in the fracture follows the lubrication equation resulting from mass conservation and Poiseuille's law. The fracture propagates by creating new surfaces if the stress intensity factor exceeds the local rock toughness  $K_{IC}$ . As the geometry of the PKN fracture exhibits symmetry, the simulation is reduced to a quarter-scale. For verification purposes, a plane strain deformation is considered in the numerical model.

We set up and solve a hydraulic fracture model to obtain the temporal solutions of the fracture half length  $l$ , the net pressure  $p_0$  and the fracture aperture  $w_0$  at the fluid inlet for the PKN fracture propagating in viscosity-storage-dominated regime. [Kovalyshen and Detournay \(2010\)](#) and [Economides and Nolte \(2000\)](#) derived the analytical solutions for this classic hydraulic fracture problem, used here to verify the results of the GEOSX simulations:

$$l(t) = 0.3817 \left( \frac{E_p Q_0^3 t^4}{\mu h^4} \right)^{1/5}$$

$$w_0(t) = 3 \left( \frac{\mu Q_0 l}{E_p} \right)^{1/4}$$

$$p_0(t) = \left( \frac{16 \mu Q_0 E_p^3 l}{\pi h^4} \right)^{1/4}$$

where the plane modulus  $E_p$  is related to Young's modulus  $E$  and Poisson's ratio  $\nu$ :

$$E_p = \frac{E}{1 - \nu^2}$$

For this example, we focus on the `Mesh`, the `Constitutive`, and the `FieldSpecifications` tags.

## Mesh

The following figure shows the mesh used in this problem.

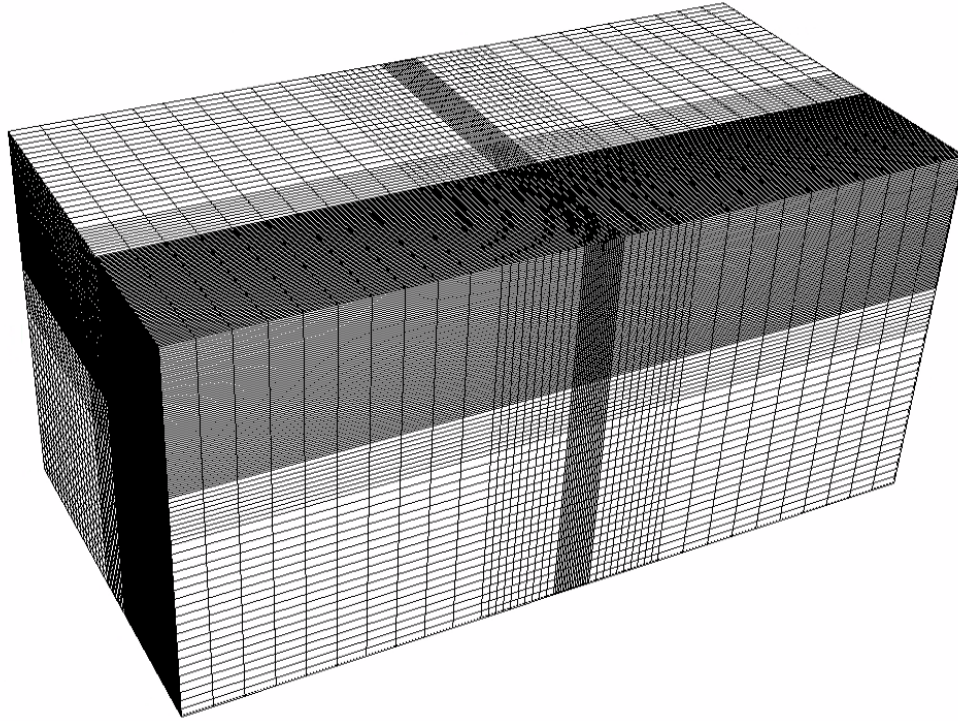


Fig. 4.30: Generated mesh

We use the internal mesh generator to create a computational domain ( $400\text{ m} \times 400\text{ m} \times 800\text{ m}$ ), as parametrized in the `InternalMesh` XML tag. The structured mesh contains  $105 \times 105 \times 60$  eight-node brick elements in the  $x$ ,  $y$ , and  $z$  directions respectively. Such eight-node hexahedral elements are defined as `C3D8` `elementTypes`, and their collection forms a mesh with one group of cell blocks named here `cb1`. Local refinement is performed for the elements in the vicinity of the fracture plane.

```
<Mesh>
  <InternalMesh
    name="mesh1"
    elementTypes="{ C3D8 }"
    xCoords="{ 0, 150, 200, 400 }"
    yCoords="{ 0, 150, 200, 400 }"
    zCoords="{ -400, -100, -20, 20, 100, 400 }"
    nx="{ 75, 10, 20 }"
    ny="{ 75, 10, 20 }"
    nz="{ 10, 10, 20, 10, 10 }"
    cellBlockNames="{ cb1 }"/>
  </InternalMesh>
</Mesh>
```

The fracture plane is defined by a nodeset occupying a small region within the computational domain, where the fracture tends to open and propagate upon fluid injection:

```
<Box
  name="core"
  xMin="{ -500.1, -500.1, -0.1 }"
  xMax="{ 500.1, 10.1, 0.1 }"/>
```



## Solid mechanics solver

GEOSX is a multi-physics platform. Different combinations of physics solvers available in the code can be applied in different regions of the domain and be functional at different stages of the simulation. The `Solvers` tag in the XML file is used to list and parameterize these solvers.

Three elementary solvers are combined in the solver `hydrofracture` to model the coupling between fluid flow within the fracture, rock deformation, fracture deformation and propagation:

```
<Hydrofracture
  name="hydrofracture"
  solidSolverName="lagsolve"
  flowSolverName="SinglePhaseFlow"
  surfaceGeneratorName="SurfaceGen"
  couplingTypeOption="FIM"
  logLevel="1"
  targetRegions="{ Fracture }"
  contactRelationName="fractureContact"
  maxNumResolves="5"
  initialDt="0.1">
  <NonlinearSolverParameters
    newtonTol="1.0e-4"
    newtonMaxIter="10"
    maxTimeStepCuts="5"/>
  <LinearSolverParameters
    solverType="gmres"
    preconditionerType="mgr"
    logLevel="1"
    krylovAdaptiveTol="1"/>
</Hydrofracture>
```

- Rock and fracture deformations are modeled by the solid mechanics solver `SolidMechanicsLagrangianSSLE`. In this solver, we define `targetRegions` that includes both the continuum region and the fracture region. The name of the contact constitutive behavior is specified in this solver by the `contactRelationName`.

```
<SolidMechanicsLagrangianSSLE
  name="lagsolve"
  timeIntegrationOption="QuasiStatic"
  discretization="FE1"
  targetRegions="{ Domain, Fracture }"
  contactRelationName="fractureContact">
  <NonlinearSolverParameters
    newtonTol="1.0e-6"/>
  <LinearSolverParameters
    solverType="gmres"
    krylovTol="1.0e-10"/>
</SolidMechanicsLagrangianSSLE>
```

- The single-phase fluid flow inside the fracture is solved by the finite volume method in the solver `SinglePhaseFVM`.

```
<SinglePhaseFVM
  name="SinglePhaseFlow"
  discretization="singlePhaseTPFA"
  targetRegions="{ Fracture }"
  inputFluxEstimate="1.0">
```

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```

<NonlinearSolverParameters
  newtonTol="1.0e-5"
  newtonMaxIter="10"/>
<LinearSolverParameters
  solverType="gmres"
  krylovTol="1.0e-12"/>
</SinglePhaseFVM>

```

- The solver `SurfaceGenerator` defines the fracture region and rock toughness `rockToughness="0.1e6"`. With `nodeBasedSIF="1"`, a node-based Stress Intensity Factor (SIF) calculation is chosen for the fracture propagation criterion.

```

<SurfaceGenerator
  name="SurfaceGen"
  targetRegions="{ Domain }"
  nodeBasedSIF="1"
  rockToughness="0.1e6"
  mpiCommOrder="1"/>

```

## Constitutive laws

For this problem, a homogeneous and isotropic domain with one solid material is assumed. Its mechanical properties and associated fluid rheology are specified in the Constitutive section. `ElasticIsotropic` model is used to describe the mechanical behavior of rock when subjected to fluid injection. The single-phase fluid model `CompressibleSinglePhaseFluid` is selected to simulate the response of water upon fracture propagation.

```

<Constitutive>
  <CompressibleSinglePhaseFluid
    name="water"
    defaultDensity="1000"
    defaultViscosity="0.001"
    referencePressure="0.0"
    compressibility="5e-12"
    referenceViscosity="1.0e-3"
    viscosibility="0.0"/>

    <ElasticIsotropic
      name="rock"
      defaultDensity="2700"
      defaultBulkModulus="20.0e9"
      defaultShearModulus="12.0e9"/>

    <CompressibleSolidParallelPlatesPermeability
      name="fractureFilling"
      solidModelName="nullSolid"
      porosityModelName="fracturePorosity"
      permeabilityModelName="fracturePerm"/>

    <NullModel
      name="nullSolid"/>

    <PressurePorosity
      name="fracturePorosity"
      defaultReferencePorosity="1.00"

```

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```

referencePressure="0.0"
compressibility="0.0"/>

<ParallelPlatesPermeability
  name="fracturePerm"/>

<FrictionlessContact
  name="fractureContact"
  penaltyStiffness="1.0e0"
  apertureTableName="apertureTable"/>
</Constitutive>

```

All constitutive parameters such as density, viscosity, bulk modulus, and shear modulus are specified in the International System of Units.

## Time history function

In the Tasks section, PackCollection tasks are defined to collect time history information from fields. Either the entire field or specified named sets of indices in the field can be collected. In this example, pressureCollection, apertureCollection, hydraulicApertureCollection and areaCollection are specified to output the time history of fracture characteristics (pressure, width and area). objectPath="ElementRegions/Fracture/FractureSubRegion" indicates that these PackCollection tasks are applied to the fracture element subregion.

```

<Tasks>
  <PackCollection
    name="pressureCollection"
    objectPath="ElementRegions/Fracture/FractureSubRegion"
    fieldName="pressure"/>

  <PackCollection
    name="apertureCollection"
    objectPath="ElementRegions/Fracture/FractureSubRegion"
    fieldName="elementAperture"/>

  <PackCollection
    name="hydraulicApertureCollection"
    objectPath="ElementRegions/Fracture/FractureSubRegion"
    fieldName="hydraulicAperture"/>

  <PackCollection
    name="areaCollection"
    objectPath="ElementRegions/Fracture/FractureSubRegion"
    fieldName="elementArea"/>
</Tasks>

```

These tasks are triggered using the Event manager with a PeriodicEvent defined for the recurring tasks. GEOSX writes one file named after the string defined in the filename keyword and formatted as a HDF5 file (pknViscosityDominated\_output.hdf5). This TimeHistory file contains the collected time history information from specified time history collector. This file includes datasets for the simulation time, fluid pressure, element aperture, hydraulic aperture and element area for the propagating hydraulic fracture. A Python script is prepared to read and query any specified subset of the time history data for verification and visualization.

## Initial and boundary conditions

The next step is to specify:

- The initial values: the `waterDensity`, `separableFace` and the `ruptureState` of the propagating fracture have to be initialized,
- The boundary conditions: fluid injection rates and the constraints of the outer boundaries have to be set.

In this example, a mass injection rate `SourceFlux` (`scale="-6.625"`) is applied at the surfaces of the initial fracture. Only one fourth of the total injection rate is used because only a quarter of the fracture is modeled (the problem is symmetric). The value given for `scale` is  $Q_0\rho_f/4$  (not  $Q_0/4$ ). All the outer boundaries are subject to roller constraints. These boundary conditions are set through the `FieldSpecifications` section.

```
<FieldSpecifications>
  <FieldSpecification
    name="waterDensity"
    initialCondition="1"
    setNames="{ fracture }"
    objectPath="ElementRegions"
    fieldName="water_density"
    scale="1000"/>

  <FieldSpecification
    name="separableFace"
    initialCondition="1"
    setNames="{ core }"
    objectPath="faceManager"
    fieldName="isFaceSeparable"
    scale="1"/>

  <FieldSpecification
    name="frac"
    initialCondition="1"
    setNames="{ fracture }"
    objectPath="faceManager"
    fieldName="ruptureState"
    scale="1"/>

  <FieldSpecification
    name="yconstraint"
    objectPath="nodeManager"
    fieldName="totalDisplacement"
    component="1"
    scale="0.0"
    setNames="{ yneg, ypos }"/>

  <FieldSpecification
    name="zconstraint"
    objectPath="nodeManager"
    fieldName="totalDisplacement"
    component="2"
    scale="0.0"
    setNames="{ zneg, zpos }"/>

  <FieldSpecification
    name="xconstraint"
    objectPath="nodeManager"
```

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```

    fieldName="totalDisplacement"
    component="0"
    scale="0.0"
    setNames="{ xneg, xpos }"/>

<SourceFlux
  name="sourceTerm"
  objectPath="ElementRegions/Fracture"
  scale="-6.625"
  setNames="{ source }"/>
</FieldSpecifications>

```

The parameters used in the simulation are summarized in the following table.

Symbol	Parameter	Unit	Value
$K$	Bulk Modulus	[GPa]	20.0
$G$	Shear Modulus	[GPa]	12.0
$K_{IC}$	Rock Toughness	[MPa.m <sup>1/2</sup> ]	0.1
$\mu$	Fluid Viscosity	[Pa.s]	1.0x10 <sup>-3</sup>
$Q_0$	Injection Rate	[m <sup>3</sup> /s]	0.0265
$t_{inj}$	Injection Time	[s]	200
$h_f$	Fracture Height	[m]	20

## Inspecting results

The following figure shows the distribution of  $\sigma_{zz}$  at  $t = 200s$  within the computational domain..

First, by running the query script

```
python ./hydrofractureQueries.py pknViscosityDominated
```

the HDF5 output is postprocessed and temporal evolution of fracture characteristics (fluid pressure and fracture width at fluid inlet and fracture half length) are saved into a txt file `model-results.txt`, which can be used for verification and visualization:

```

[['      time', ' pressure', ' aperture', '      length']]
 2 1.413e+06 0.0006093      5.6
 4 1.174e+06 0.0007132      8.4
 6 1.077e+06 0.0007849     10.8
 8 1.044e+06 0.0008482     12.8
10 1.047e+06 0.0009098     14.8

```

Note: GEOSX python tools `geosx_xml_tools` should be installed to run the query script (See [Python Tools Setup](#) for details).

Next, figure below shows the comparisons between the results from GEOSX simulations (markers) and the corresponding analytical solutions (curves) for the example with viscosity-storage dominated assumptions, which is generated using the visualization script:

```
python ./pknViscosityDominatedFigure.py
```

The evolution in time of the fracture half-length, the near-wellbore fracture aperture, and the fluid pressure all correlate well with the analytical solutions.

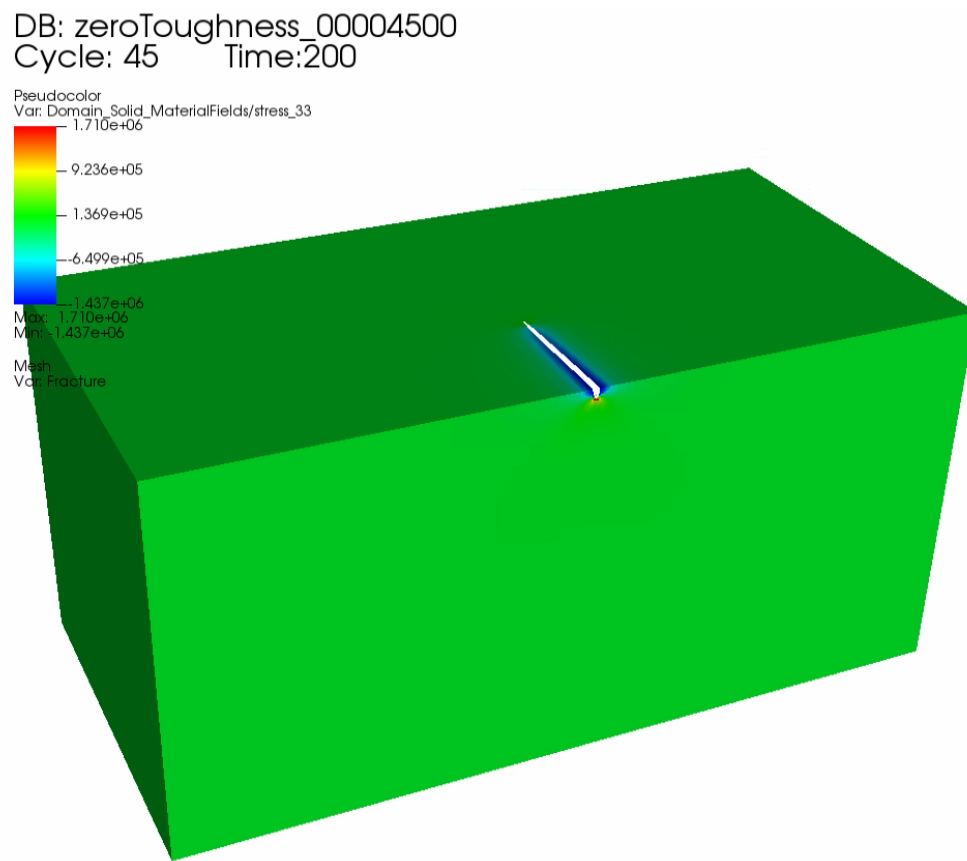
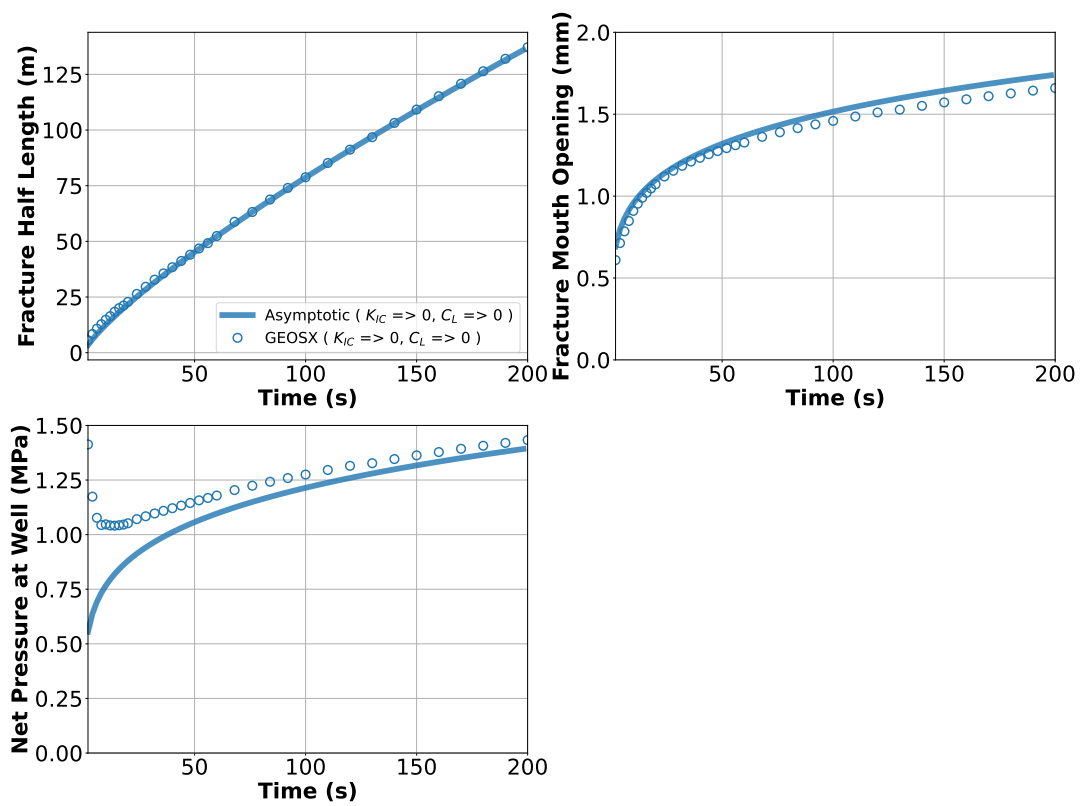


Fig. 4.31: Simulation result of  $\sigma_{zz}$  at  $t = 200s$



## To go further

### Feedback on this example

For any feedback on this example, please submit a [GitHub issue on the project's GitHub page](#).

## Proppant Slot Test

### Context

In this example, a simulation is built up to model a proppant slot test. In this way, the implemented proppant model is validated by comparing numerical results with the corresponding experimental data. Furthermore, this calibrated proppant model can allow field engineers to customize stimulation design and optimize field operations in multiple engineering aspects (Huang et al., 2021).

### Input file

This example uses no external input files and everything is contained within a single xml file that is located at:

```
inputFiles/proppant/ProppantSlotTest_base.xml
```

```
inputFiles/proppant/ProppantSlotTest_benchmark.xml
```

## Description of the case

Chun et al. (2020) conducted slot tests on proppant transport with slickwater. As shown below, a 4 ft X 1 ft slot with 0.3 in gap width was constructed. Three fluid inlets with 0.5 in inner diameter were placed at the right side of the slot, which were three inches away from each other. One outlet was placed on the top side to allow pressure relief. The other one was located on the left side acting as a fluid sink. In their tests, to resemble a slickwater fracturing treatment, the proppant concentration was kept at 1.5 ppg and the viscosity of carrying fluid was approximately 1 cp. The slurry was mixed well and then injected into the flow channel at a constant injection rate of 6 gpm. A simulation case with the same settings is built up to mimic these slot tests. A vertical and impermeable fracture surface is assumed in this case, which eliminates the effect of fracture plane inclination and fluid leak-off. A static fracture with an uniform aperture of 0.3 in is defined and fracture propagation is not involved. 30/50 mesh proppant is injected via the three inlets and is flowed through the slot for 30 seconds.

To simulate proppant transport phenomenon, a proppant solver based on the assumption of multi-component single phase flow is used in this example. Proppant concentration and distribution within the slot are numerically calculated by solving the equations of proppant transport in hydraulic fractures. These numerical predictions are then validated against the corresponding testing results (Chun et al., 2020).

In this example, we focus our attention on the `Solvers`, `Constitutive` and `FieldSpecifications` tags.

### Mesh

The following figure shows the mesh used for solving this problem.

We use the internal mesh generator `InternalMesh` to create a computational domain. This mesh contains 2 x 97 x 24 eight-node brick elements in the x, y and z directions, respectively. Here, a structured three-dimensional mesh is generated with `C3D8` as the `elementTypes` (eight-node hexahedral elements). This mesh is defined as a cell block with the name `cb1`.



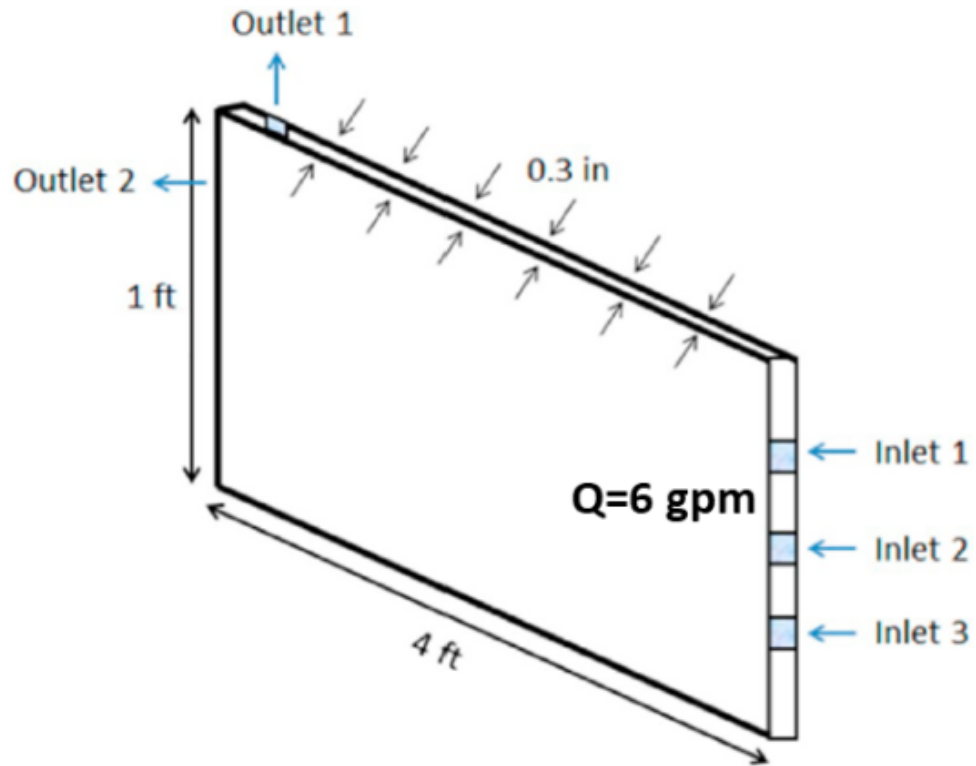


Fig. 4.32: Configuration of the slot for proppant transport experiment (after Chun et al., 2020)

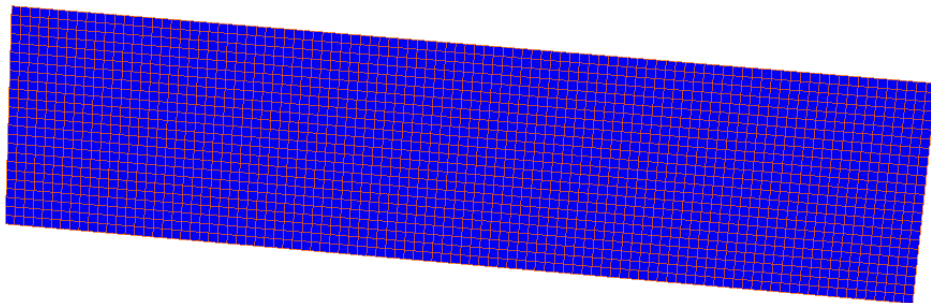


Fig. 4.33: Mesh for simulating the proppant slot tests.

```

<Mesh>
  <InternalMesh
    name="mesh"
    elementTypes="{ C3D8 }"
    xCoords="{ -1, 1 }"
    yCoords="{ 0, 1.2319 }"
    zCoords="{ 0, 0.3048 }"
    nx="{ 2 }"
    ny="{ 97 }"
    nz="{ 24 }"
    cellBlockNames="{ cb1 }"/>
  </InternalMesh>
</Mesh>

```

## Proppant transport solver

GEOSX is a multi-physics platform. Different combinations of physics solvers available in the code can be applied in different regions of the domain and be functional at different stages of the simulation. The `Solvers` tag in the XML file is used to define these solvers.

To specify a coupling between two different solvers, we define and characterize each single-physics solver separately. Then, we customize a *coupling solver* between these single-physics solvers as an additional solver. This approach allows for generality and flexibility in constructing multi-physics solvers. The order of specifying these solvers is not restricted in GEOSX. Note that end-users should give each single-physics solver a meaningful and distinct name, as GEOSX will recognize these single-physics solvers based on their customized names and create user-expected coupling.

As demonstrated in this example, to setup a coupled proppant transport solver, we need to define three different solvers in the XML file:

- the proppant transport solver for the fracture region, a solver of type `ProppantTransport` called here `ProppantTransport` (see [Proppant Transport Solver](#) for more information),

```

<ProppantTransport
  name="ProppantTransport"
  logLevel="1"
  updateProppantPacking="1"
  proppantDiameter="4.5e-4"
  frictionCoefficient="0.04"
  criticalShieldsNumber="0.0"
  maxProppantConcentration="0.62"
  discretization="singlePhaseTPFA"
  targetRegions="{ Fracture }">
  <NonlinearSolverParameters
    newtonTol="1.0e-6"
    newtonMaxIter="8"
    lineSearchAction="None"
    maxTimeStepCuts="5"/>
  <LinearSolverParameters
    solverType="gmres"
    krylovTol="1.0e-7"/>
</ProppantTransport>

```

- the single-phase flow solver, a solver of type `SinglePhaseProppantFVM` called here `SinglePhaseFVM`,

```

<SinglePhaseProppantFVM
  name="SinglePhaseFVM"

```

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```

logLevel="1"
discretization="singlePhaseTPFA"
targetRegions="{ Fracture }">
<NonlinearSolverParameters
  newtonTol="1.0e-6"
  newtonMaxIter="8"
  lineSearchAction="None"
  newtonMinIter="0"/>
<LinearSolverParameters
  solverType="gmres"
  preconditionerType="amg"
  krylovTol="1.0e-7"/>
</SinglePhaseProppantFVM>

```

- the coupling solver (FlowProppantTransport) that binds the two single-physics solvers above, which is named as FlowProppantTransport

```

<FlowProppantTransport
  name="FlowProppantTransport"
  proppantSolverName="ProppantTransport"
  flowSolverName="SinglePhaseFVM"
  targetRegions="{ Fracture }"
  logLevel="1"/>

```

In this example, let us focus on the coupling solver. This solver (FlowProppantTransport) describes the coupling process between proppant and flow transport within the `Fracture` region. In this way, the two single-physics solvers (ProppantTransport and SinglePhaseFVM) are sequentially called to solve the sub-problems (proppant transport and pressure problem, respectively) involved in this test case.

## Constitutive laws

For this slot test, 30/50 mesh proppant is injected via the three inlets and flowing through the slot for 30 seconds. The viscosity of carrying fluid is 0.001 Pa.s to resemble slickwater fracturing. In this example, the solid and fluid materials are named as `sand` and `water` respectively. Proppant characterization and fluid rheology are specified in the Constitutive section:

```

<Constitutive>
  <ProppantSlurryFluid
    name="water"
    referencePressure="1e5"
    referenceDensity="1000"
    compressibility="0.0"
    maxProppantConcentration="0.62"
    referenceViscosity="0.001"
    referenceProppantDensity="2550.0"/>

  <ParticleFluid
    name="sand"
    particleSettlingModel="Stokes"
    hinderedSettlingCoefficient="4.5"
    proppantDensity="2550.0"
    proppantDiameter="4.5e-4"
    maxProppantConcentration="0.62"/>

```

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```

<ProppantSolidProppantPermeability
  name="fractureFilling"
  solidModelName="nullSolid"
  porosityModelName="fracturePorosity"
  permeabilityModelName="fracturePerm" />

<NullModel
  name="nullSolid" />

<ProppantPorosity
  name="fracturePorosity"
  defaultReferencePorosity="1.00"
  maxProppantConcentration="0.62" />

<ProppantPermeability
  name="fracturePerm"
  proppantDiameter="4.5e-4"
  maxProppantConcentration="0.62" />
</Constitutive>

```

The constitutive parameters such as proppant density and proppant diameter are specified in the International System of Units.

## Initial and boundary conditions

The next step is to specify fields, including:

- The initial value (fracture aperture, fluid pressure and proppant concentration within the fracture have to be initialized)
- The boundary conditions (fluid pressure and proppant concentration at fluid inlets and outlets)

These boundary conditions are set up through the `FieldSpecifications` section. At a constant injection rate, the slurry is equally flowing into the open channel through three inlets.

```

<FieldSpecifications>
  <FieldSpecification
    name="frac"
    initialCondition="1"
    setNames="{ fracture }"
    objectPath="faceManager"
    fieldName="ruptureState"
    scale="1" />

  <FieldSpecification
    name="fracAp"
    initialCondition="1"
    objectPath="ElementRegions/Fracture"
    fieldName="elementAperture"
    scale="7.62e-3"
    setNames="{ fracture }" />

  <FieldSpecification
    name="frac1"
    initialCondition="1"
    objectPath="ElementRegions/Fracture"

```

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```

    fieldName="pressure"
    scale="0.0"
    component="0"
    setNames="{ fracture }"/>

<FieldSpecification
  name="frac2"
  initialCondition="1"
  objectPath="ElementRegions/Fracture"
  fieldName="proppantConcentration"
  scale="0.0"
  component="0"
  setNames="{ fracture }"/>

<FieldSpecification
  name="frac3"
  initialCondition="1"
  objectPath="ElementRegions/Fracture"
  fieldName="isProppantBoundary"
  component="0"
  setNames="{ fracture }"/>

<FieldSpecification
  name="frac4"
  initialCondition="1"
  objectPath="ElementRegions/Fracture"
  fieldName="isProppantBoundary"
  scale="1"
  component="0"
  setNames="{ left0 }"/>

<SourceFlux
  name="left1a"
  objectPath="ElementRegions/Fracture"
  fieldName="pressure"
  scale="-0.14"
  component="0"
  setNames="{ left1 }"/>

<FieldSpecification
  name="left1b"
  objectPath="ElementRegions/Fracture"
  fieldName="proppantConcentration"
  scale="0.07"
  component="0"
  setNames="{ left1 }"/>

<SourceFlux
  name="left2a"
  objectPath="ElementRegions/Fracture"
  fieldName="pressure"
  scale="-0.14"
  component="0"
  setNames="{ left2 }"/>

<FieldSpecification
  name="left2b"

```

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```

    objectPath="ElementRegions/Fracture"
    fieldName="proppantConcentration"
    scale="0.07"
    component="0"
    setNames="{ left2 }"/>

<SourceFlux
  name="left3a"
  objectPath="ElementRegions/Fracture"
  fieldName="pressure"
  scale="-0.14"
  component="0"
  setNames="{ left3 }"/>

<FieldSpecification
  name="left3b"
  objectPath="ElementRegions/Fracture"
  fieldName="proppantConcentration"
  scale="0.07"
  component="0"
  setNames="{ left3 }"/>

<FieldSpecification
  name="right1"
  objectPath="ElementRegions/Fracture"
  fieldName="pressure"
  scale="0.0"
  component="0"
  setNames="{ right }"/>

<FieldSpecification
  name="right2"
  objectPath="ElementRegions/Fracture"
  fieldName="proppantConcentration"
  scale="0.0"
  component="0"
  setNames="{ right }"/>
</FieldSpecifications>

```

Note: For static (non-propagating) fracture problems, the fields `ruptureState` and `elementAperture` should be provided in the initial conditions. `fieldName="pressure"` here means that the source flux term is added to the mass balance equation for pressure.

The parameters used in the simulation are summarized in the following table.

Symbol	Parameter	Unit	Value
$d_p$	Proppant Diameter	[m]	0.00045
$f$	Darcy Friction Coefficient	[-]	0.04
$N_{sh}$	Critical Shields Number	[-]	0.0
$c_s$	Max Fraction of Proppant	[-]	0.62
$\rho_f$	Fluid Density	[kg/m <sup>3</sup> ]	1000
$\mu_f$	Fluid Viscosity	[Pa*s]	0.001
$\rho_p$	Proppant Density	[kg/m <sup>3</sup> ]	2550
$\lambda_s$	Hindered Settling Coefficient	[-]	4.5
$c_p$	Proppant Concentration in Slurry	[m <sup>3</sup> /m <sup>3</sup> ]	0.07
$L$	Fracture Length	[m]	1.219
$H$	Fracture Height	[m]	0.3048
$a$	Fracture Aperture	[m]	0.00762
$Q$	Injection Rate	[m <sup>3</sup> /s]	0.0003785

### Inspecting results

The following figure shows the modelling prediction of proppant distribution at 10 s and 30 s, which are compared with the experiments in (Chun et al., 2020). Due to proppant settling in low viscosity fluid, a heterogeneous proppant distribution is obtained, which evolves with injection time. Three different zones (immobile proppant bed, suspended proppant and clean fluid) are visually identified for both the presented experiment and simulation.

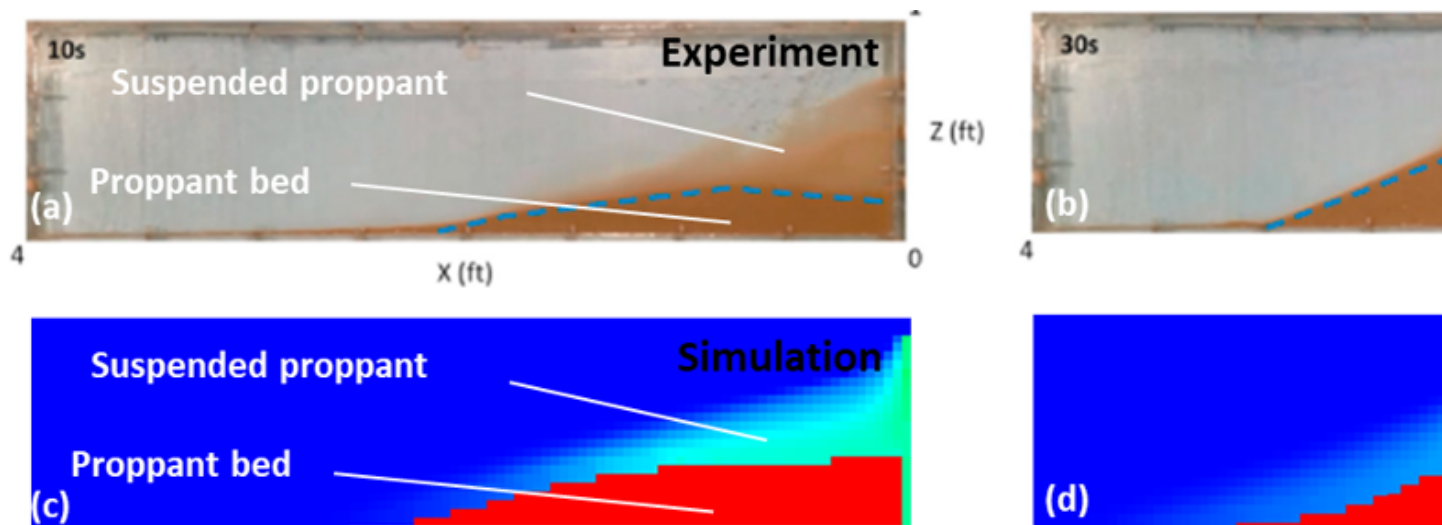


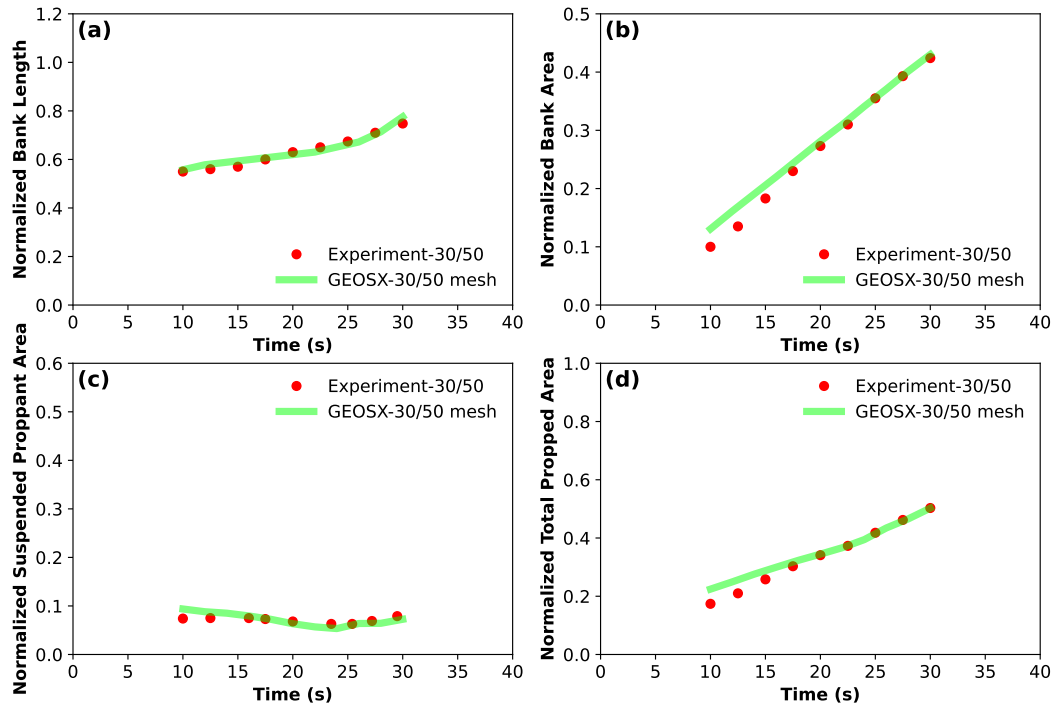
Fig. 4.34: Proppant distribution profile

As shown below, consistently, the modelling predictions (green curve) on proppant transport and distribution show a good agreement with the reported experimental data (red dot) at each time.

### To go further

#### Feedback on this example

This concludes the example on simulating a proppant slot test. For any feedback on this example, please submit a [GitHub issue on the project's GitHub page](#).



For more details

- More on proppant solver, please see [Proppant Transport Solver](#).

## 4.1.4 Wellbore Problems

### Kirsch Wellbore Problem

#### Context

In this example, we simulate a vertical elastic wellbore subjected to in-situ stress and the induced elastic deformation of the reservoir rock. Kirsch's solution to this problem provides the stress and displacement fields developing around a circular cavity, which is hereby employed to verify the accuracy of the numerical results. For this example, the `TimeHistory` function and python scripts are used to output and post-process multi-dimensional data (stress and displacement).

#### Input file

Everything required is contained within two GEOSX input files located at:

```
inputFiles/solidMechanics/KirschProblem_base.xml
```

```
inputFiles/solidMechanics/KirschProblem_benchmark.xml
```



## Description of the case

We solve a drained wellbore problem subjected to anisotropic horizontal stress ( $\sigma_{xx}$  and  $\sigma_{yy}$ ) as shown below. This is a vertical wellbore drilled in an infinite, homogeneous, isotropic, and elastic medium. Far-field in-situ stresses and internal supporting pressure acting at the circular cavity cause a mechanical deformation of the reservoir rock and stress concentration in the near-wellbore region. For verification purpose, a plane strain condition is considered for the numerical model.

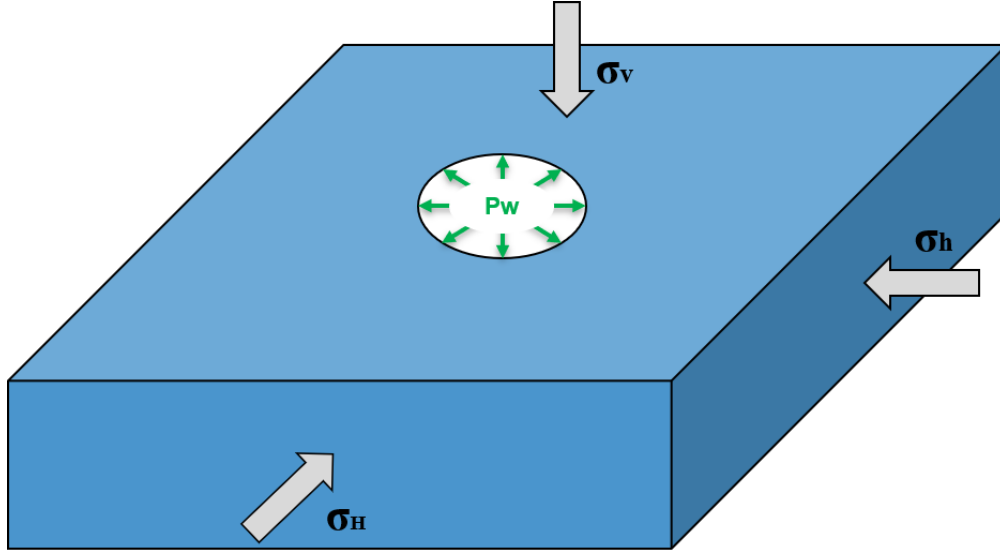


Fig. 4.35: Sketch of the wellbore problem

In this example, stress ( $\sigma_{rr}$ ,  $\sigma_{\theta\theta}$ , and  $\sigma_{r\theta}$ ) and displacement ( $u_r$  and  $u_\theta$ ) fields around the wellbore are calculated numerically. These numerical predictions are compared with the corresponding Kirsch solutions (Poulos and Davis, 1974).

$$\begin{aligned}\sigma_{rr} &= \frac{\sigma_{xx} + \sigma_{yy}}{2} \left[ 1 - \left( \frac{a_0}{r} \right)^2 \right] + \frac{\sigma_{xx} - \sigma_{yy}}{2} \left[ 1 - 4 \left( \frac{a_0}{r} \right)^2 + 3 \left( \frac{a_0}{r} \right)^4 \right] \cos(2\theta) + P_w \left( \frac{a_0}{r} \right)^2 \\ \sigma_{\theta\theta} &= \frac{\sigma_{xx} + \sigma_{yy}}{2} \left[ 1 + \left( \frac{a_0}{r} \right)^2 \right] - \frac{\sigma_{xx} - \sigma_{yy}}{2} \left[ 1 + 3 \left( \frac{a_0}{r} \right)^4 \right] \cos(2\theta) - P_w \left( \frac{a_0}{r} \right)^2 \\ \sigma_{r\theta} &= -\frac{\sigma_{xx} - \sigma_{yy}}{2} \left[ 1 + 2 \left( \frac{a_0}{r} \right)^2 - 3 \left( \frac{a_0}{r} \right)^4 \right] \sin(2\theta) \\ u_r &= -\frac{(a_0)^2}{2Gr} \left[ \frac{\sigma_{xx} + \sigma_{yy}}{2} + \frac{\sigma_{xx} - \sigma_{yy}}{2} (4(1 - \nu) - \left( \frac{a_0}{r} \right)^2) \cos(2\theta) - P_w \right] \\ u_\theta &= \frac{(a_0)^2}{2Gr} \frac{\sigma_{xx} - \sigma_{yy}}{2} [2(1 - 2\nu) + \left( \frac{a_0}{r} \right)^2] \sin(2\theta)\end{aligned}$$

where  $a_0$  is the initial wellbore radius,  $r$  is the radial coordinate,  $\nu$  is the Poisson's ratio,  $G$  is the shear modulus,  $P_w$  is the normal traction acting on the wellbore wall, the angle  $\theta$  is measured with respect to x-z plane and defined as positive in counter-clockwise direction.

In this example, we focus our attention on the `Mesh`, the `Constitutive`, and the `FieldSpecifications` tags.

## Mesh

Following figure shows the generated mesh that is used for solving this wellbore problem.

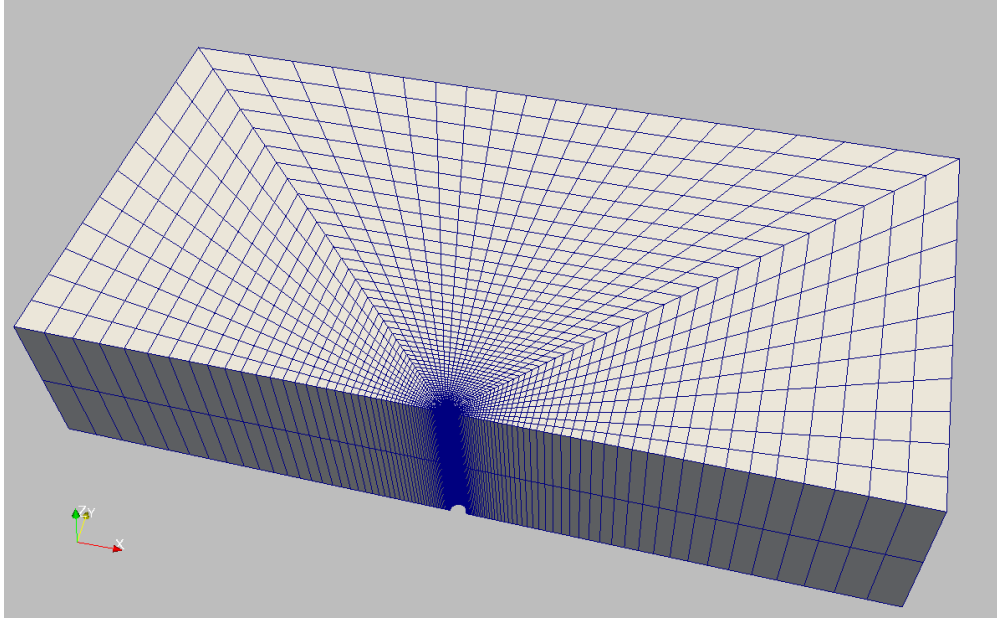


Fig. 4.36: Generated mesh for a vertical wellbore problem

Let us take a closer look at the geometry of this wellbore problem. We use the internal wellbore mesh generator `InternalWellbore` to create a rock domain ( $10\text{ m} \times 5\text{ m} \times 2\text{ m}$ ), with a wellbore of initial radius equal to  $0.1\text{ m}$ . Only half of the domain is modeled by a `theta` angle from  $0$  to  $180$ , assuming symmetry for the rest of the domain. Coordinates of `trajectory` defines the wellbore trajectory, a vertical well in this example. By turning on `autoSpaceRadialElems="{ 1 }"`, the internal mesh generator automatically sets number and spacing of elements in the radial direction, which overrides the values of `nr`. With `useCartesianOuterBoundary="0"`, a Cartesian aligned boundary condition is enforced on the outer blocks. This way, a structured three-dimensional mesh is created with  $50 \times 40 \times 2$  elements in the radial, tangential and  $z$  directions, respectively. All elements are eight-node hexahedral elements (C3D8) and refinement is performed to conform with the wellbore geometry. This mesh is defined as a cell block with the name `cb1`.

```
<Mesh>
  <InternalWellbore
    name="mesh1"
    elementTypes="{ C3D8 }"
    radius="{ 0.1, 5.0 }"
    theta="{ 0, 180 }"
    zCoords="{ -1, 1 }"
    nr="{ 40 }"
    nt="{ 40 }"
    nz="{ 2 }"
    trajectory="{ { 0.0, 0.0, -1.0 },
                  { 0.0, 0.0, 1.0 } }"
    autoSpaceRadialElems="{ 1 }"
    useCartesianOuterBoundary="0"
    cellBlockNames="{ cb1 }"/>
  </InternalWellbore>
</Mesh>
```

## Solid mechanics solver

For a drained wellbore problem, the pore pressure variation is omitted. Therefore, we just need to define a solid mechanics solver, which is called `mechanicsSolver`. This solid mechanics solver (see *Solid Mechanics Solver*) is based on the Lagrangian finite element formulation. The problem is run as `QuasiStatic` without considering inertial effects. The computational domain is discretized by `FE1`, which is defined in the `NumericalMethods` section. The material is named `rock`, whose mechanical properties are specified in the `Constitutive` section.

```
<Solvers gravityVector="{0.0, 0.0, 0.0}">
  <SolidMechanics_LagrangianFEM
    name="mechanicsSolver"
    timeIntegrationOption="QuasiStatic"
    logLevel="1"
    discretization="FE1"
    targetRegions="{Omega}">
    <NonlinearSolverParameters
      newtonTol = "1.0e-5"
      newtonMaxIter = "15"
    />
    <LinearSolverParameters
      directParallel="0"/>
  </SolidMechanics_LagrangianFEM>
</Solvers>
```

## Constitutive laws

For this drained wellbore problem, we simulate a linear elastic deformation around the circular cavity. A homogeneous and isotropic domain with one solid material is assumed, with mechanical properties specified in the `Constitutive` section:

```
<Constitutive>
  <ElasticIsotropic
    name="rock"
    defaultDensity="2700"
    defaultBulkModulus="5.0e8"
    defaultShearModulus="3.0e8"
  />
</Constitutive>
```

Recall that in the `SolidMechanics_LagrangianFEM` section, `rock` is the material in the computational domain. Here, the isotropic elastic model `ElasticIsotropic` simulates the mechanical behavior of `rock`.

The constitutive parameters such as the density, the bulk modulus, and the shear modulus are specified in the International System of Units.

## Time history function

In the `Tasks` section, `PackCollection` tasks are defined to collect time history information from fields. Either the entire field or specified named sets of indices in the field can be collected. In this example, `stressCollection` and `displacementCollection` tasks are specified to output the resultant stresses (tensor stored as an array with Voigt notation) and total displacement field (stored as a 3-component vector) respectively.

```

<Tasks>
  <PackCollection
    name="stressCollection"
    objectPath="ElementRegions/Omega/cb1"
    fieldName="rock_stress"/>

  <PackCollection
    name="displacementCollection"
    objectPath="nodeManager"
    fieldName="totalDisplacement"/>
</Tasks>

```

These two tasks are triggered using the Event management, where `PeriodicEvent` are defined for these recurring tasks. GEOSX writes two files named after the string defined in the `filename` keyword and formatted as HDF5 files (`displacement_history.hdf5` and `stress_history.hdf5`). The `TimeHistory` file contains the collected time history information from each specified time history collector. This information includes datasets for the simulation time, element center or nodal position, and the time history information. Then, a Python script is prepared to access and plot any specified subset of the time history data for verification and visualization.

## Initial and boundary conditions

The next step is to specify fields, including:

- The initial value (the in-situ stresses and traction at the wellbore wall have to be initialized),
- The boundary conditions (constraints of the outer boundaries have to be set).

Here, we specify anisotropic horizontal stress values ( $\sigma_{yy} = -9.0$  MPa and  $\sigma_{xx} = -11.25$  MPa) and a vertical stress ( $\sigma_{zz} = -15.0$  MPa). A compressive traction (`WellLoad`)  $P_w = -2.0$  MPa is loaded at the wellbore wall `rneg`. The remaining parts of the outer boundaries are subjected to roller constraints. These boundary conditions are set in the `FieldSpecifications` section.

```

<FieldSpecifications>
  <FieldSpecification
    name="Sxx"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions"
    fieldName="rock_stress"
    component="0"
    scale="-11.25e6"
  />

  <FieldSpecification
    name="Syy"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions"
    fieldName="rock_stress"
    component="1"
    scale="-9.0e6"
  />

  <FieldSpecification
    name="Szz"
    initialCondition="1"

```

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```

    setNames="{ all }"
    objectPath="ElementRegions"
    fieldName="rock_stress"
    component="2"
    scale="-15.0e6"
  />

  <Traction
    name="WellLoad"
    setNames="{ rneg }"
    objectPath="faceManager"
    scale="-2.0e6"
    tractionType="normal"
  />

  <FieldSpecification
    name="xconstraint"
    objectPath="nodeManager"
    fieldName="totalDisplacement"
    component="0"
    scale="0.0"
    setNames="{ xneg, xpos }"
  />

  <FieldSpecification
    name="yconstraint"
    objectPath="nodeManager"
    fieldName="totalDisplacement"
    component="1"
    scale="0.0"
    setNames="{ tneg, tpos, ypos }"
  />

  <FieldSpecification
    name="zconstraint"
    objectPath="nodeManager"
    fieldName="totalDisplacement"
    component="2"
    scale="0.0"
    setNames="{ zneg, zpos }"
  />
</FieldSpecifications>

```

With `tractionType="normal"`, traction is applied to the wellbore wall `rneg` as a pressure specified as the scalar product of scale `scale="-2.0e6"` and the outward face normal vector. In this case, the loading magnitude of the traction does not change with time.

You may note :

- All initial value fields must have `initialCondition` field set to 1;
- The `setName` field points to the previously defined set to apply the fields;
- `nodeManager` and `faceManager` in the `objectPath` indicate that the boundary conditions are applied to the element nodes and faces, respectively;
- `fieldName` is the name of the field registered in GEOSX;
- Component 0, 1, and 2 refer to the x, y, and z direction, respectively;

- And the non-zero values given by `scale` indicate the magnitude of the loading;
- Some shorthand, such as `xneg` and `xpos`, are used as the locations where the boundary conditions are applied in the computational domain. For instance, `xneg` means the face of the computational domain located at the left-most extent in the x-axis, while `xpos` refers to the face located at the right-most extent in the x-axis. Similar shorthands include `ypos`, `yneg`, `zpos`, and `zneg`;
- The mud pressure loading and in situ stresses have negative values due to the negative sign convention for compressive stress in GEOSX.

The parameters used in the simulation are summarized in the following table.

Symbol	Parameter	Unit	Value
$K$	Bulk Modulus	[MPa]	500.0
$G$	Shear Modulus	[MPa]	300.0
$\sigma_{yy}$	Min Horizontal Stress	[MPa]	-9.0
$\sigma_{xx}$	Max Horizontal Stress	[MPa]	-11.25
$\sigma_{zz}$	Vertical Stress	[MPa]	-15.0
$a_0$	Initial Well Radius	[m]	0.1
$P_w$	Traction at Well	[MPa]	-2.0

## Inspecting results

In the above examples, we request VTK output files that can be imported into Paraview to visualize the outcome. The following figure shows the distribution of  $\sigma_{xx}$  in the near wellbore region.

We use time history function to collect time history information and run a Python script to query and plot the results. The figure below shows the comparisons between the numerical predictions (marks) and the corresponding analytical solutions (solid curves) with respect to the distributions of stress components and displacement at  $\theta = 45$  degrees. Predictions computed by GEOSX match the analytical results.

## To go further

### Feedback on this example

For any feedback on this example, please submit a [GitHub issue on the project's GitHub page](#).

## Cased Elastic Wellbore Problem

### Problem description

This example uses the solid mechanics solver to handle a cased wellbore problem subjected to a pressure test. The completed wellbore is composed of a steel casing, a cement sheath and rock formation. Isotropic linear elastic behavior is assumed for all the three materials. No separation is allowed for the casing-cement and cement-rock contact interfaces.

Analytical results of the radial and hoop stresses,  $\sigma_{rr}$  and  $\sigma_{\theta\theta}$ , in casing, cement sheath and rock are expressed as (Hervé and Zaoui, 1995) :

$$\sigma_{rr} = (2\lambda + 2G)A - \frac{2GB}{r^2}$$

$$\sigma_{\theta\theta} = (2\lambda + 2G)A + \frac{2GB}{r^2}$$

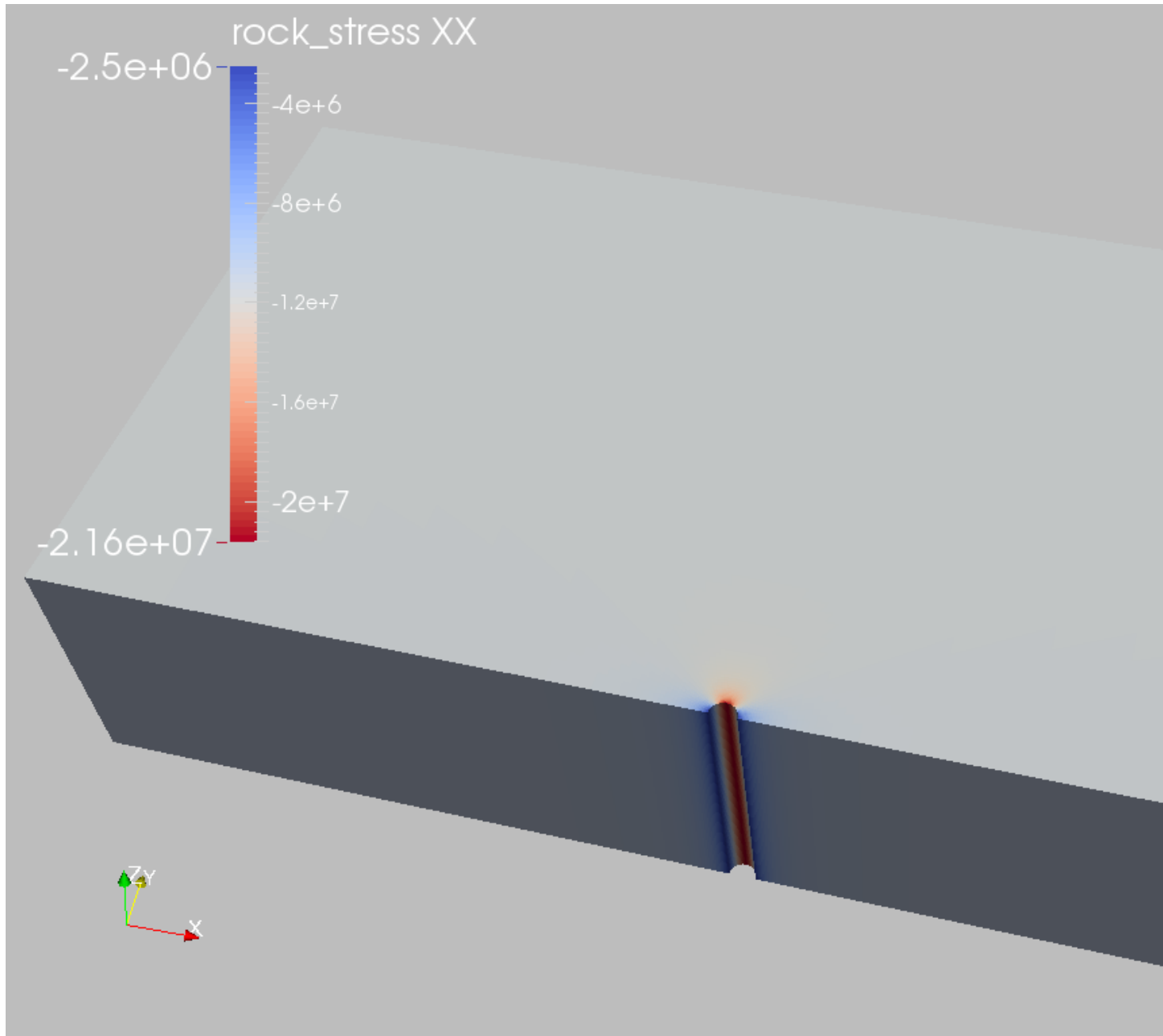
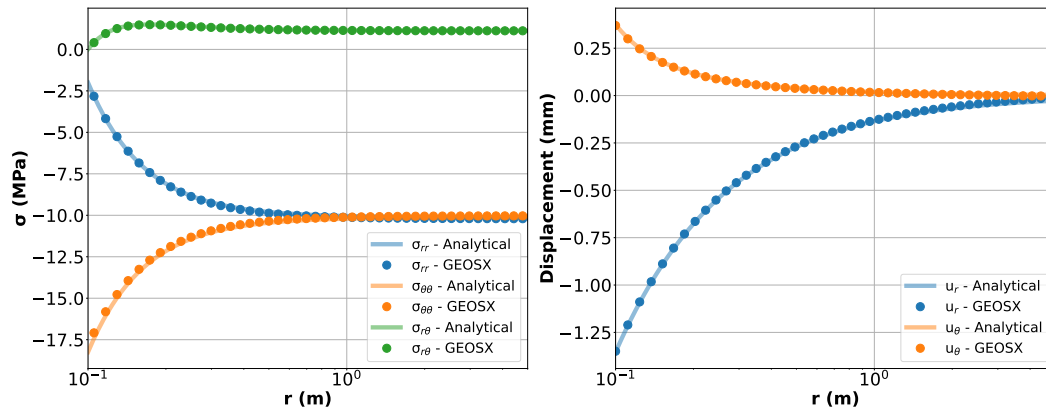


Fig. 4.37: Simulation result of  $\sigma_{xx}$



where  $\lambda$  and  $G$  are the Lamé moduli,  $r$  is the radial coordinate,  $A$  and  $B$  are piecewise constants that are obtained by solving the boundary and interface conditions, as detailed in the post-processing script.

### Input file

This benchmark example uses no external input files and everything required is contained within two GEOSX xml files that are located at:

```
inputFiles/wellbore/CasedElasticWellbore_base.xml
```

and

```
inputFiles/wellbore/CasedElasticWellbore_benchmark.xml
```

The corresponding integrated test is

```
inputFiles/wellbore/CasedElasticWellbore_smoke.xml
```

In this example, we would focus our attention on the `Solvers`, `Mesh` and `Constitutive` tags.

### Solid mechanics solver

As fluid flow is not considered, only the solid mechanics `SolidMechanicsLagrangianSSLE` solver is required for solving this linear elastic problem. In this solver, the three regions and three materials associated to casing, cement sheath and rock are respectively defined by `targetRegions` and `solidMaterialNames`.

```
<SolidMechanicsLagrangianSSLE
  name="lagsolve"
  timeIntegrationOption="QuasiStatic"
  discretization="FE1"
  logLevel="0"
  targetRegions="{ casing, cement, rock }">
```

### Cased wellbore mesh

The internal wellbore mesh generator `InternalWellbore` is employed to create the mesh of this wellbore problem. The radii of the casing cylinder, the cement sheath cylinder and the far-field boundary of the surrounding rock formation are defined by a vector radius. In the tangent direction, `theta` angle is specified from 0 to 360 degree



for a full geometry of the domain. Note that a half or a quarter of the domain can be defined by a `theta` angle from 0 to 180 or 90 degree, respectively. The trajectory of the well is defined by `trajectory`, which is vertical in this case. The `autoSpaceRadialElems` parameters allow optimally increasing the element size from local zone around the wellbore to the far-field zone. In this example, the auto spacing option is only applied for the rock formation. The `useCartesianOuterBoundary` transforms the far-field boundary to a squared shape to enforce a Cartesian aligned outer boundary, which eases the loading of the boundary conditions. The `cellBlockNames` and `elementTypes` define the regions and related element types associated to casing, cement sheath and rock.

```
<Mesh>
  <InternalWellbore
    name="mesh1"
    elementTypes="{ C3D8, C3D8, C3D8 }"
    radius="{ 0.1, 0.106, 0.133, 2.0 }"
    theta="{ 0, 360 }"
    zCoords="{ 0, 1 }"
    nr="{ 10, 20, 10 }"
    nt="{ 320 }"
    nz="{ 1 }"
    trajectory="{ { 0.0, 0.0, 0.0 },
                  { 0.0, 0.0, 1.0 } }"
    autoSpaceRadialElems="{ 0, 0, 1 }"
    useCartesianOuterBoundary="2"
    cellBlockNames="{ casing, cement, rock }"
  />
</Mesh>
```

## Steel, cement, and rock constitutive laws

Isotropic linear elastic constitutive behavior is considered for all the three materials. Note that the default density is useless for this case.

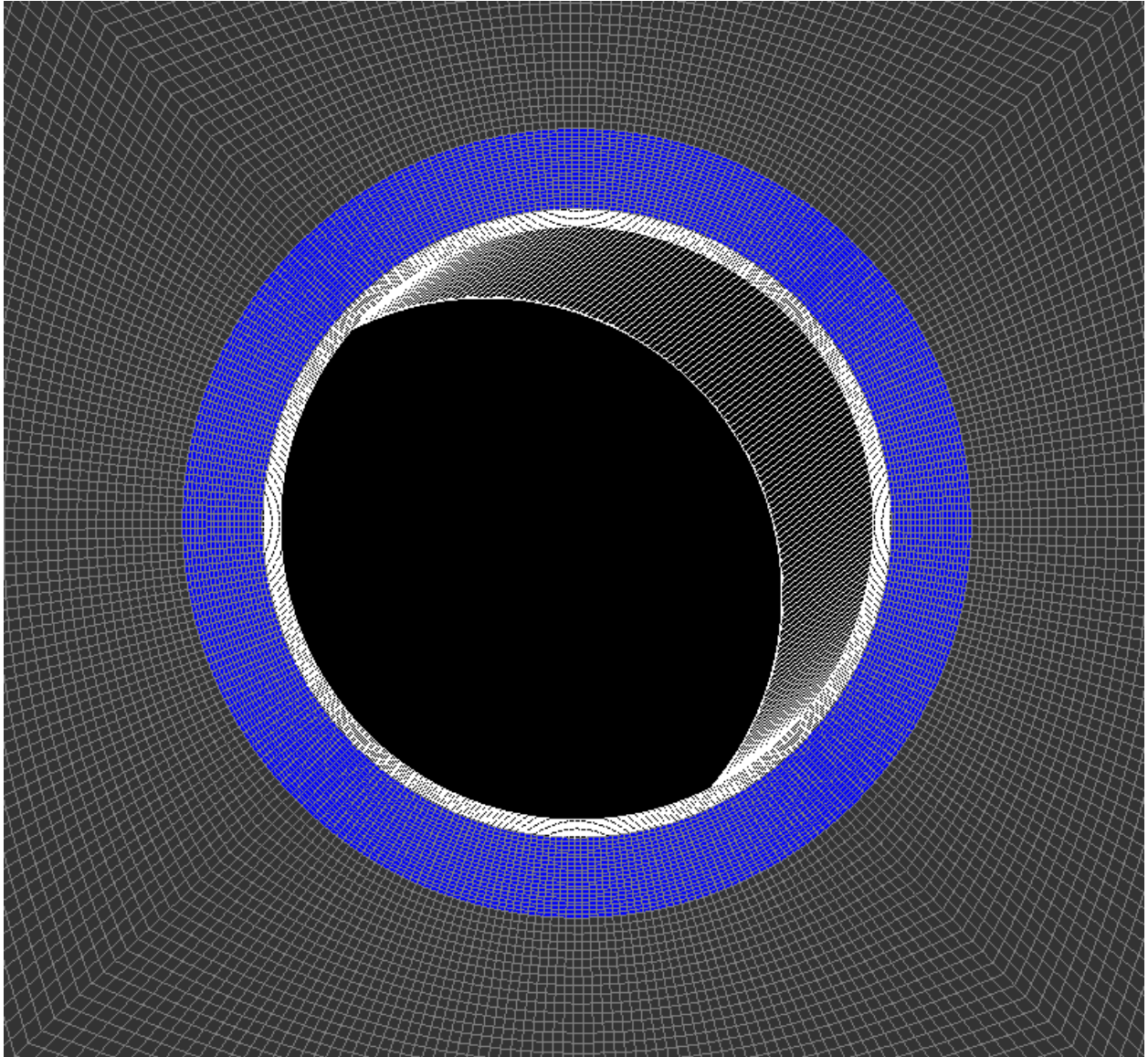
```
<ElasticIsotropic
  name="casing"
  defaultDensity="2700"
  defaultBulkModulus="175e9"
  defaultShearModulus="80.8e9"/>

<ElasticIsotropic
  name="cement"
  defaultDensity="2700"
  defaultBulkModulus="10.3e9"
  defaultShearModulus="6.45e9"/>

<ElasticIsotropic
  name="rock"
  defaultDensity="2700"
  defaultBulkModulus="5.5556e9"
  defaultShearModulus="4.16667e9"/>
```

## Boundary conditions

Far-field boundary are subjected to roller constraints. The normal traction on the inner face of the casing is defined by `Traction` field specification. The nodeset generated by the internal wellbore generator for this face is named as



rneg. The traction type is normal to mimic a casing test pressure that is applied normal to the casing inner face . The negative sign of the scale is attributed to the negative sign convention for compressive stress in GEOSX.

```
<FieldSpecifications>

  <FieldSpecification
    name="xConstraint"
    objectPath="nodeManager"
    fieldName="totalDisplacement"
    component="0"
    scale="0.0"
    setNames="{ xneg, xpos }"/>

  <FieldSpecification
    name="yConstraint"
    objectPath="nodeManager"
    fieldName="totalDisplacement"
    component="1"
    scale="0.0"
    setNames="{ yneg, ypos }"/>

  <FieldSpecification
    name="zconstraint"
    objectPath="nodeManager"
    fieldName="totalDisplacement"
    component="2"
    scale="0.0"
    setNames="{ zneg, zpos }"/>

  <Traction
    name="innerPressure"
    objectPath="faceManager"
    tractionType="normal"
    scale="-10.0e6"
    setNames="{ rneg }"/>
</FieldSpecifications>
```

## Results and benchmark

A good agreement between the GEOSX results and analytical results is shown in the figure below:

## To go further

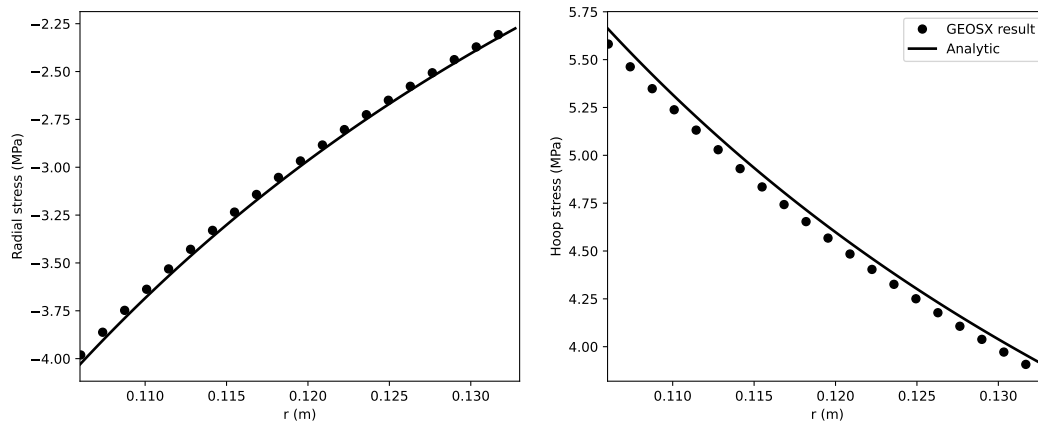
### Feedback on this example

This concludes the cased wellbore example. For any feedback on this example, please submit a [GitHub issue](#) on the project's GitHub page.

## Deviated Elastic Wellbore Problem

### Problem description

This example uses the solid mechanics solver to handle a deviated wellbore problem with open hole completion. This wellbore is subjected to a mud pressure at wellbore wall and undrained condition is assumed (no fluid flow in the



rock formation). A segment of the wellbore with isotropic linear elastic deformation is simulated in this case. Far field stresses and gravity effect are excluded. The main goal of this example is to validate the internal wellbore mesh generator and mechanics solver for the case of an inclined wellbore.

Analytical results of the radial and hoop stresses,  $\sigma_{rr}$  and  $\sigma_{\theta\theta}$ , around the wellbore are expressed as (Detournay and Cheng, 1988) :

$$\sigma_{rr} = p_0 \frac{a^2}{r^2}$$

$$\sigma_{\theta\theta} = -p_0 \frac{a^2}{r^2}$$

where  $p_0$  is the applied mud pressure at wellbore wall,  $a$  is the wellbore radius and  $r$  is the radial coordinate.

### Input file

This benchmark example uses no external input files and everything required is contained within two GEOSX xml files that are located at:

```
inputFiles/wellbore/DeviatedElasticWellbore_base.xml
```

and

```
inputFiles/wellbore/DeviatedElasticWellbore_benchmark.xml
```

The corresponding xml file for the integrated test is

```
inputFiles/wellbore/DeviatedElasticWellbore_smoke.xml
```

In this example, we would focus our attention on the Mesh tag.

### Solid mechanics solver

As fluid flow is not considered, only the solid mechanics solver `SolidMechanicsLagrangianSSLE` is required for solving this wellbore problem.

```
<SolidMechanicsLagrangianSSLE
  name="lagsolve"
  timeIntegrationOption="QuasiStatic"
```

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```

discretization="FE1"
logLevel="0"
targetRegions="{ Omega }"
>

```

## Deviated wellbore mesh

The internal wellbore mesh generator `InternalWellbore` is employed to create the mesh of this wellbore problem. The radius of the wellbore and the size of the surrounding rock formation are defined by a vector `radius`. In the tangent direction, `theta` angle is specified from 0 to 180 degree for a half of the domain regarding its symmetry. Note that the whole domain could be specified with a `theta` angle from 0 to 360 degree, if modeling complicated scenarios. The trajectory of the well is defined by `trajectory`. In this example, the wellbore is inclined in the x-z plane by an angle of 45 degree. The `autoSpaceRadialElems` parameter allows optimally increasing the element size from local zone around the wellbore to the far-field zone, which is set to 1 to activate this option. The `useCartesianOuterBoundary` transforms the far-field boundary to a squared shape to enforce a Cartesian aligned outer boundary, which eases the loading of the far-field boundary conditions. In this example, this value is set to 0 for the single region along the radial direction.

```

<Mesh>
  <InternalWellbore
    name="mesh1"
    elementTypes="{ C3D8 }"
    radius="{ 0.1, 2 }"
    theta="{ 0, 180 }"
    zCoords="{ -0.5, 0.5 }"
    nr="{ 30 }"
    nt="{ 80 }"
    nz="{ 100 }"
    trajectory="{ { -0.5, 0.0, -0.5 },
                  { 0.5, 0.0, 0.5 } }"
    autoSpaceRadialElems="{ 1 }"
    useCartesianOuterBoundary="0"
    cellBlockNames="{ cb1 }"/>
  </InternalWellbore>
</Mesh>

```

## Constitutive law

Isotropic linear elastic constitutive behavior is considered for the rock around the wellbore. Note that the default density is useless in this specific example, as gravity effect is neglected.

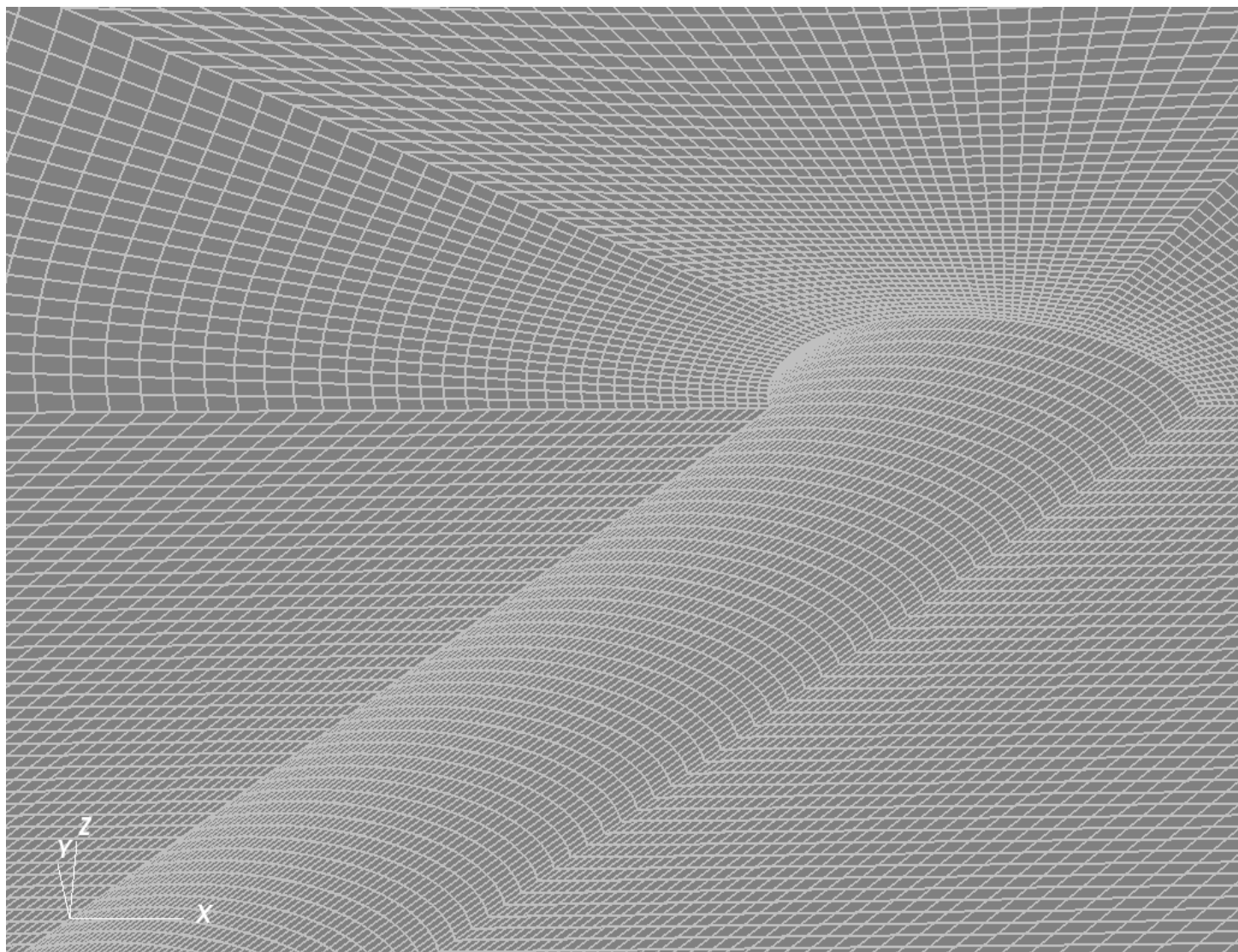
```

<ElasticIsotropic
  name="shale"
  defaultDensity="2700"
  defaultBulkModulus="5.5556e9"
  defaultShearModulus="4.16667e9"/>

```

## Boundary conditions

Far-field boundaries are subjected to roller constraints and in-situ stresses are not considered. The mud pressure on the wellbore wall is defined by `Traction` field specification. The nodeset generated by the internal wellbore generator



for this face is named as `rneg`. The traction type is `normal` to mimic a pressure that is applied normal to the wellbore wall. The negative sign of the scale is attributed to the negative sign convention for compressive stresses in GEOSX.

```
<FieldSpecifications>

  <FieldSpecification
    name="xConstraint"
    objectPath="nodeManager"
    fieldName="totalDisplacement"
    component="0"
    scale="0.0"
    setNames="{ xneg }"/>

  <FieldSpecification
    name="yConstraint"
    objectPath="nodeManager"
    fieldName="totalDisplacement"
    component="1"
    scale="0.0"
    setNames="{ tneg, tpos }"/>

  <FieldSpecification
    name="zconstraint"
    objectPath="nodeManager"
    fieldName="totalDisplacement"
    component="2"
    scale="0.0"
    setNames="{ zneg }"/>

  <Traction
    name="innerPressure"
    objectPath="faceManager"
    tractionType="normal"
    scale="-10.e6"
    setNames="{ rneg }"/>
</FieldSpecifications>
```

## Results and benchmark

A good agreement between the GEOSX results and the corresponding analytical solutions is shown in the figure below:

## To go further

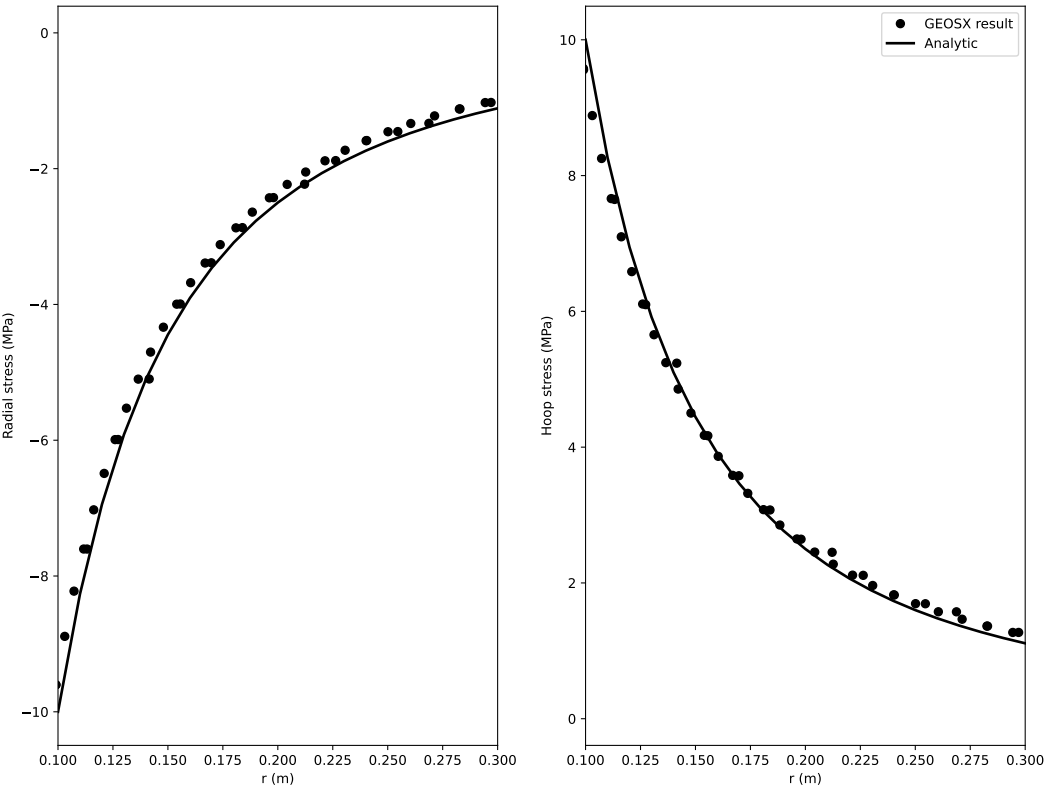
### Feedback on this example

This concludes the deviated elastic wellbore example. For any feedback on this example, please submit a [GitHub issue](#) on the project's [GitHub page](#).

## Elasto-Plastic Near-Well Deformation

### Context

The main goal of this example is to learn how to use the internal wellbore mesh generator and an elasto-plastic model to handle wellbore problems in GEOSX. The Extended Drucker-Prager model (see [Model: Extended Drucker-Prager](#)) is applied to solve for elastoplastic deformation within the vicinity of a vertical wellbore. For the presented example,





an analytical solution is employed to verify the accuracy of the numerical results. The resulting model can be used as a base for more complex analysis (e.g., wellbore drilling, fluid injection and storage scenarios).

## Objectives

At the end of this example you will know:

- how to construct meshes for wellbore problems with the internal mesh generator,
- how to specify initial and boundary conditions, such as in-situ stresses and variation of traction at the wellbore wall,
- how to use a plastic model for mechanical problems in the near wellbore region.

## Input file

This example uses no external input files and everything required is contained within two xml files that are located at:

```
inputFiles/solidMechanics/ExtendedDruckerPragerWellbore_base.xml
```

```
inputFiles/solidMechanics/ExtendedDruckerPragerWellbore_benchmark.xml
```

## Description of the case

We simulate a drained wellbore problem subjected to isotropic horizontal stress ( $\sigma_h$ ) and vertical stress ( $\sigma_v$ ). By lowering the wellbore supporting pressure ( $P_w$ ), the wellbore contracts, and the reservoir rock experiences elastoplastic deformation. A plastic zone develops in the near wellbore region, as shown below.

To simulate this phenomenon, the strain hardening Extended Drucker-Prager model with an associated plastic flow rule in GEOSX is used in this example. Displacement and stress fields around the wellbore are numerically calculated. These numerical predictions are then compared with the corresponding analytical solutions (Chen and Abousleiman, 2017) from the literature.

All inputs for this case are contained inside a single XML file. In this example, we focus our attention on the Mesh tags, the Constitutive tags, and the FieldSpecifications tags.

## Mesh

Following figure shows the generated mesh that is used for solving this 3D wellbore problem

Let us take a closer look at the geometry of this wellbore problem. We use the internal mesh generator `InternalWellbore` to create a rock domain ( $10\text{ m} \times 5\text{ m} \times 2\text{ m}$ ), with a wellbore of initial radius equal to 0.1 m. Coordinates of trajectory defines the wellbore trajectory, which represents a vertical well in this example. By turning on `autoSpaceRadialElems="{ 1 }"`, the internal mesh generator automatically sets number and spacing of elements in the radial direction, which overrides the values of `nr`. With `useCartesianOuterBoundary="0"`, a Cartesian aligned outer boundary on the outer block is enforced. In this way, a structured three-dimensional mesh is created with  $100 \times 80 \times 2$  elements in the radial, tangential and z directions, respectively. All the elements are eight-node hexahedral elements (C3D8) and refinement is performed to conform with the wellbore geometry. This mesh is defined as a cell block with the name `cb1`.

```
<Mesh>
  <InternalWellbore
    name="mesh1"
    elementTypes="{ C3D8 }"
    radius="{ 0.1, 5.0 }"
    theta="{ 0, 180 }"
```

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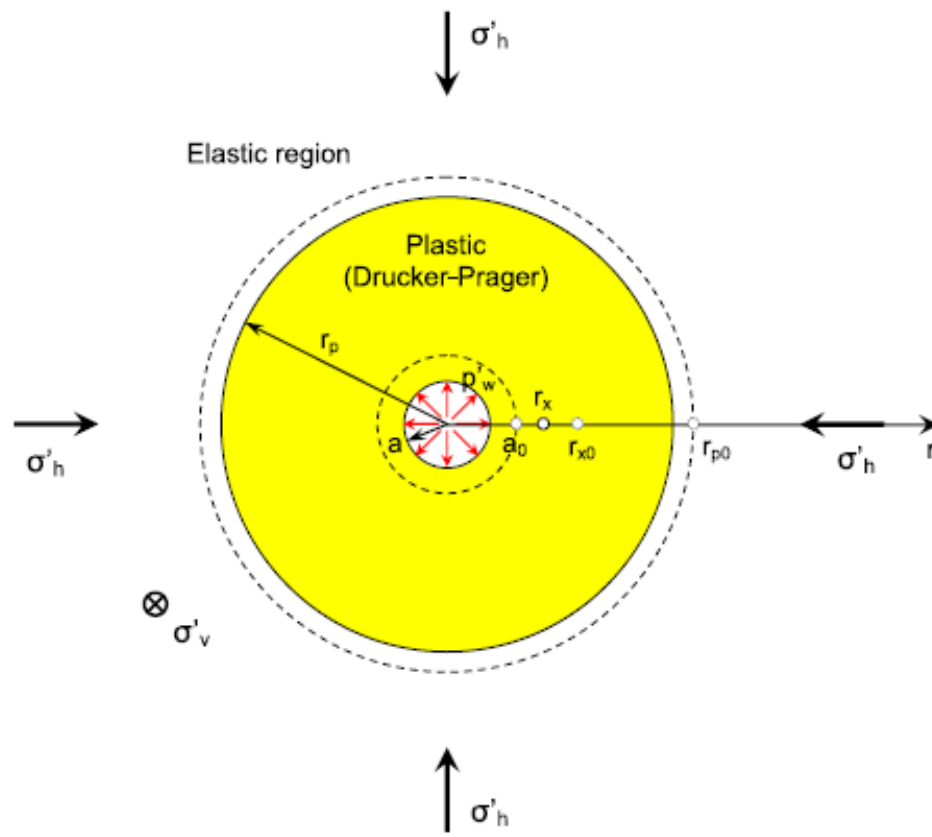


Fig. 4.38: Sketch of the wellbore problem (Chen and Abousleiman, 2017)

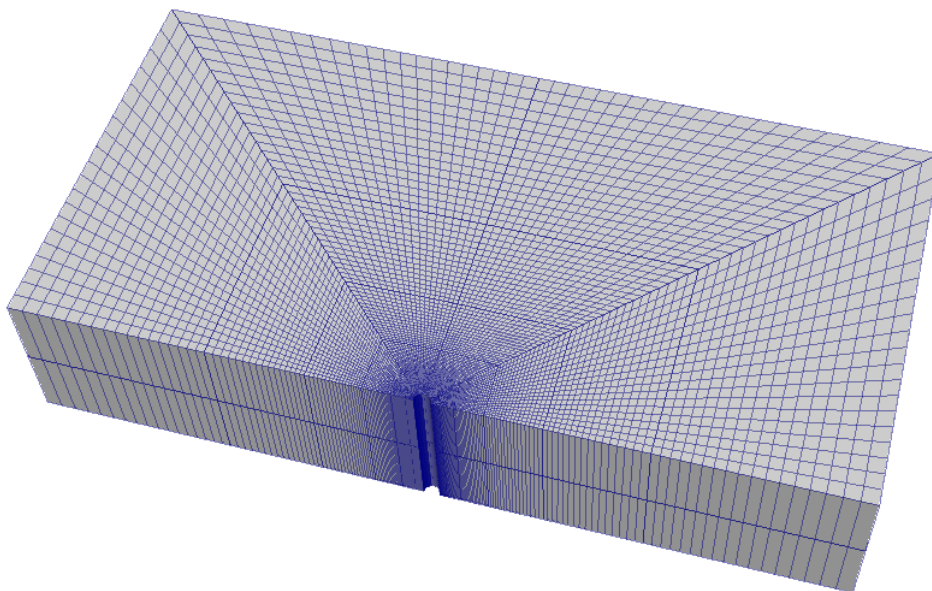


Fig. 4.39: Generated mesh for the wellbore problem

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```

zCoords="{ -1, 1 }"
nr="{ 40 }"
nt="{ 80 }"
nz="{ 2 }"
trajectory="{ { 0.0, 0.0, -1.0 },
               { 0.0, 0.0, 1.0 } }"
autoSpaceRadialElems="{ 1 }"
useCartesianOuterBoundary="0"
cellBlockNames="{ cb1 }"/>
</Mesh>

```

## Solid mechanics solver

For the drained wellbore problem, the pore pressure variation is omitted and can be subtracted from the analysis. Therefore, we just need to define a solid mechanics solver, which is called `mechanicsSolver`. This solid mechanics solver (see *Solid Mechanics Solver*) is based on the Lagrangian finite element formulation. The problem is run as `QuasiStatic` without considering inertial effects. The computational domain is discretized by `FE1`, which is defined in the `NumericalMethods` section. The material is named as `rock`, whose mechanical properties are specified in the `Constitutive` section.

```

<Solvers
  gravityVector="{ 0.0, 0.0, 0.0 }">
  <SolidMechanics_LagrangianFEM
    name="mechanicsSolver"
    timeIntegrationOption="QuasiStatic"
    logLevel="1"
    discretization="FE1"
    targetRegions="{ Omega }"
  >
  <LinearSolverParameters
    directParallel="0"/>
  <NonlinearSolverParameters
    newtonTol="1.0e-5"
    newtonMaxIter="15"/>
  </SolidMechanics_LagrangianFEM>
</Solvers>

```

## Constitutive laws

For this drained wellbore problem, we simulate the elastoplastic deformation caused by wellbore contraction. A homogeneous domain with one solid material is assumed, whose mechanical properties are specified in the `Constitutive` section:

```

<Constitutive>
  <ExtendedDruckerPrager
    name="rock"
    defaultDensity="2700"
    defaultBulkModulus="5.0e8"
    defaultShearModulus="3.0e8"
    defaultCohesion="0.0"
    defaultInitialFrictionAngle="15.27"
    defaultResidualFrictionAngle="23.05"
  >
</Constitutive>

```

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```

    defaultDilationRatio="1.0"
    defaultHardening="0.01"/>
</Constitutive>

```

Recall that in the `SolidMechanics_LagrangianFEM` section, rock is designated as the material in the computational domain. Here, Extended Drucker Prager model `ExtendedDruckerPrager` is used to simulate the elastoplastic behavior of rock. As for the material parameters, `defaultInitialFrictionAngle`, `defaultResidualFrictionAngle` and `defaultCohesion` denote the initial friction angle, the residual friction angle, and cohesion, respectively, as defined by the Mohr-Coulomb failure envelope. As the residual friction angle `defaultResidualFrictionAngle` is larger than the initial one `defaultInitialFrictionAngle`, a strain hardening model is adopted, whose hardening rate is given as `defaultHardening="0.01"`. If the residual friction angle is set to be less than the initial one, strain weakening will take place. Setting `defaultDilationRatio="1.0"` corresponds to an associated flow rule. The constitutive parameters such as the density, the bulk modulus, and the shear modulus are specified in the International System of Units.

## Initial and boundary conditions

The next step is to specify fields, including:

- The initial value (the in-situ stresses and traction at the wellbore wall have to be initialized)
- The boundary conditions (the reduction of wellbore pressure and constraints of the outer boundaries have to be set)

In this example, we need to specify isotropic horizontal stress ( $\sigma_h = -11.25$  MPa) and vertical stress ( $\sigma_v = -15.0$  MPa). To reach equilibrium, a compressive traction  $P_w = -11.25$  MPa is instantaneously applied at the wellbore wall `rneg` at time  $t = 0$  s, which will then be gradually reduced to a lower value (-2.0 MPa) to let wellbore contract. The remaining parts of the outer boundaries are subjected to roller constraints. These boundary conditions are set up through the `FieldSpecifications` section.

```

<FieldSpecifications>
  <FieldSpecification
    name="stressXX"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions/Omega/cb1"
    fieldName="rock_stress"
    component="0"
    scale="-11250000.0"/>

  <FieldSpecification
    name="stressYY"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions/Omega/cb1"
    fieldName="rock_stress"
    component="1"
    scale="-11250000.0"/>

  <FieldSpecification
    name="stressZZ"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions/Omega/cb1"
    fieldName="rock_stress"

```

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```

    component="2"
    scale="-15000000.0"/>

    <Traction
      name="ExternalLoad"
      setNames="{ rneg }"
      objectPath="faceManager"
      scale="-11.25e6"
      tractionType="normal"
      functionName="timeFunction"/>

    <FieldSpecification
      name="xconstraint"
      objectPath="nodeManager"
      fieldName="totalDisplacement"
      component="0"
      scale="0.0"
      setNames="{ xneg, xpos }"/>

    <FieldSpecification
      name="yconstraint"
      objectPath="nodeManager"
      fieldName="totalDisplacement"
      component="1"
      scale="0.0"
      setNames="{ tneg, tpos, ypos }"/>

    <FieldSpecification
      name="zconstraint"
      objectPath="nodeManager"
      fieldName="totalDisplacement"
      component="2"
      scale="0.0"
      setNames="{ zneg, zpos }"/>
  </FieldSpecifications>

```

With `tractionType="normal"`, traction is applied to the wellbore wall `rneg` as a pressure specified from the product of scale `scale="-11.25e6"` and the outward face normal. A table function `timeFunction` is used to define the time-dependent traction `ExternalLoad`. The coordinates and values form a time-magnitude pair for the loading time history. In this case, the loading magnitude decreases linearly as the time evolves.

```

<Functions>
  <TableFunction
    name="timeFunction"
    inputVarNames="{ time }"
    coordinates="{ 0.0, 1.0 }"
    values="{ 1.0, 0.1778 }"/>
</Functions>

```

You may note :

- All initial value fields must have `initialCondition` field set to 1;
- The `setName` field points to the previously defined box to apply the fields;
- `nodeManager` and `faceManager` in the `objectPath` indicate that the boundary conditions are applied to the element nodes and faces, respectively;

- `fieldName` is the name of the field registered in GEOSX;
- Component 0, 1, and 2 refer to the x, y, and z direction, respectively;
- And the non-zero values given by `Scale` indicate the magnitude of the loading;
- Some shorthand, such as `xneg` and `xpos`, are used as the locations where the boundary conditions are applied in the computational domain. For instance, `xneg` means the portion of the computational domain located at the left-most in the x-axis, while `xpos` refers to the portion located at the right-most area in the x-axis. Similar shorthand include `ypos`, `yneg`, `zpos`, and `zneg`;
- The mud pressure loading has a negative value due to the negative sign convention for compressive stress in GEOSX.

The parameters used in the simulation are summarized in the following table.

Symbol	Parameter	Unit	Value
$K$	Bulk modulus	[MPa]	500
$G$	Shear Modulus	[MPa]	300
$C$	Cohesion	[MPa]	0.0
$\phi_i$	Initial Friction Angle	[degree]	15.27
$\phi_r$	Residual Friction Angle	[degree]	23.05
$c_h$	Hardening Rate	[-]	0.01
$\sigma_h$	Horizontal Stress	[MPa]	-11.25
$\sigma_v$	Vertical Stress	[MPa]	-15.0
$a_0$	Initial Well Radius	[m]	0.1
$P_w$	Mud Pressure	[MPa]	-2.0

## Inspecting results

In the above example, we requested silo-format output files. We can therefore import these into VisIt and use python scripts to visualize the outcome. Below figure shows the comparisons between the numerical predictions (marks) and the corresponding analytical solutions (solid curves) with respect to the distributions of normal stress components, stress path, the supporting wellbore pressure and wellbore size. It is clear that the GEOSX predictions are in excellent agreement with the analytical results.

For the same wellbore problem, using different constitutive models (plastic vs. elastic), obviously, distinct differences in rock deformation and distribution of resultant stresses is also observed and highlighted.

## To go further

### Feedback on this example

This concludes the example on Plasticity Model for Wellbore Problems. For any feedback on this example, please submit a [GitHub issue](#) on the project's [GitHub](#) page.

### For more details

- More on plasticity models, please see *Model: Extended Drucker-Prager*.
- More on functions, please see *Functions*.

## Modified Cam-Clay Model for Wellbore Problems

### Context

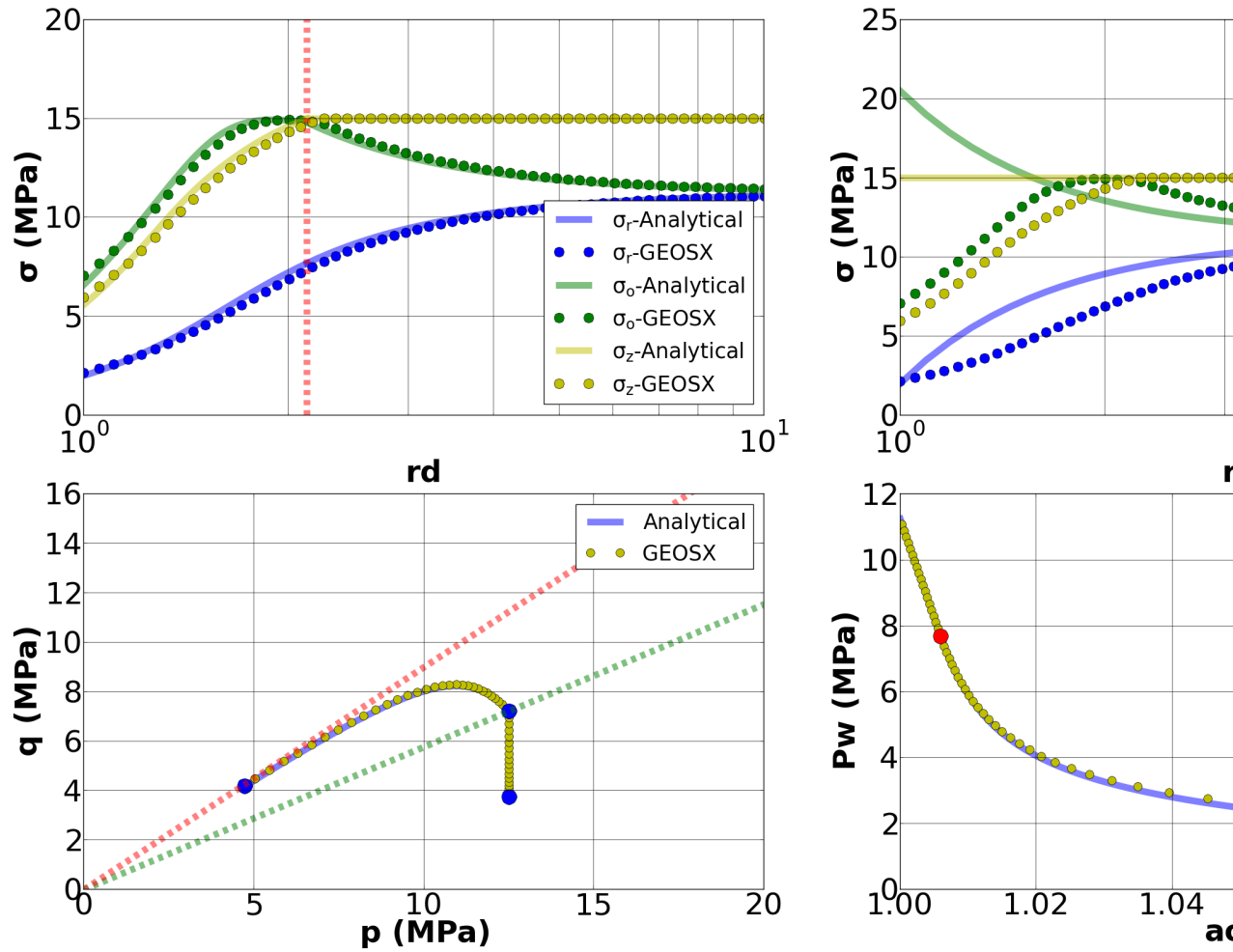


Fig. 4.40: Comparing GEOSX results with analytical solutions

In this benchmark example, the Modified Cam-Clay model (see [Model: Modified Cam-Clay](#)) is applied to solve for elastoplastic deformation within the vicinity of a vertical wellbore. For the presented example, an analytical solution is employed to verify the accuracy of the numerical results. The resulting model can be used as a base for more complex analysis (e.g., wellbore drilling, fluid injection and storage scenarios).

### Input file

Everything required is contained within two GEOSX input files located at:

```
inputFiles/solidMechanics/ModifiedCamClayWellbore_base.xml
```

```
inputFiles/solidMechanics/ModifiedCamClayWellbore_benchmark.xml
```

### Description of the case

We simulate a drained wellbore problem subjected to isotropic horizontal stress ( $\sigma_h$ ) and vertical stress ( $\sigma_v$ ), as shown below. By increasing the wellbore supporting pressure ( $P_w$ ), the wellbore expands, and the formation rock experiences elastoplastic deformation. A plastic zone develops in the near wellbore region.

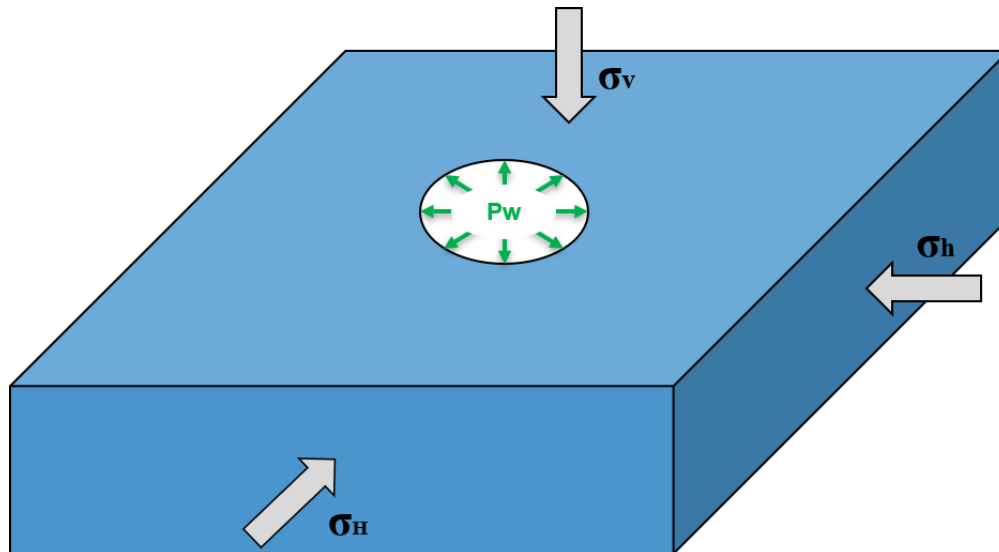


Fig. 4.41: Sketch of the wellbore problem

To simulate this phenomenon, the Modified Cam-Clay model is used in this example. Displacement and stress fields around the wellbore are numerically calculated. These numerical predictions are then compared with the corresponding analytical solutions ([Chen and Abousleiman, 2013](#)) from the literature.

In this example, we focus our attention on the `Mesh` tags, the `Constitutive` tags, and the `FieldSpecifications` tags.

### Mesh

Following figure shows the generated mesh that is used for solving this wellbore problem.

Let us take a closer look at the geometry of this wellbore problem. We use the internal wellbore mesh generator `InternalWellbore` to create a rock domain ( $10\text{ m} \times 5\text{ m} \times 2\text{ m}$ ), with a wellbore of initial radius equal to  $0.1\text{ m}$ . Coordinates of `trajectory` defines the wellbore trajectory, which represents a vertical well



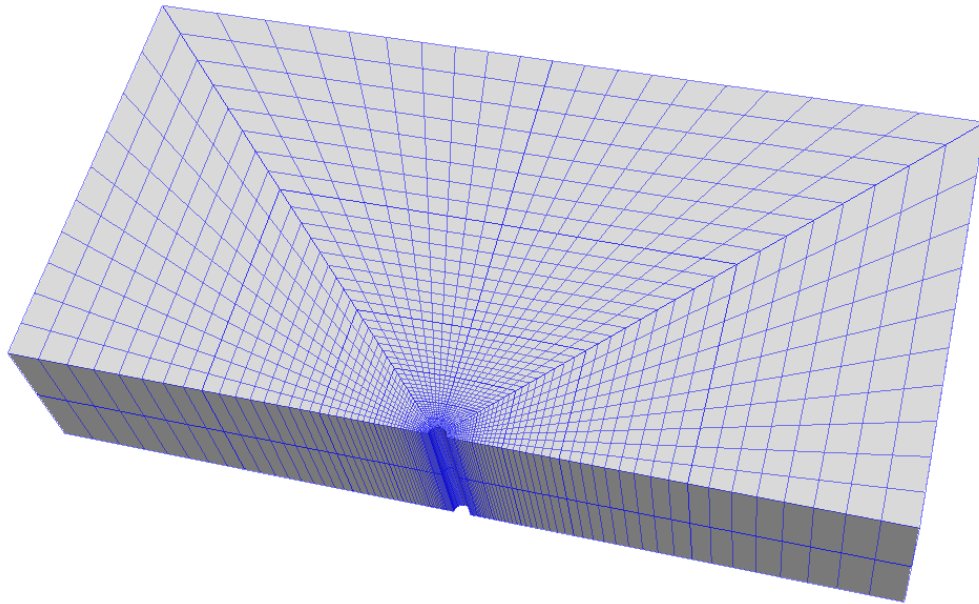


Fig. 4.42: Generated mesh for a vertical wellbore problem

in this example. By turning on `autoSpaceRadialElems="{ 1 }"`, the internal mesh generator automatically sets number and spacing of elements in the radial direction, which overrides the values of `nr`. With `useCartesianOuterBoundary="0"`, a Cartesian aligned outer boundary on the outer block is enforced. In this way, a structured three-dimensional mesh is created with  $50 \times 40 \times 2$  elements in the radial, tangential and  $z$  directions, respectively. All the elements are eight-node hexahedral elements (C3D8) and refinement is performed to conform with the wellbore geometry. This mesh is defined as a cell block with the name `cb1`.

```
<Mesh>
  <InternalWellbore
    name="mesh1"
    elementTypes="{ C3D8 }"
    radius="{ 0.1, 5.0 }"
    theta="{ 0, 180 }"
    zCoords="{ -1, 1 }"
    nr="{ 40 }"
    nt="{ 40 }"
    nz="{ 2 }"
    trajectory="{ { 0.0, 0.0, -1.0 },
                  { 0.0, 0.0, 1.0 } }"
    autoSpaceRadialElems="{ 1 }"
    useCartesianOuterBoundary="0"
    cellBlockNames="{ cb1 }"/>
  </InternalWellbore>
</Mesh>
```

## Solid mechanics solver

For the drained wellbore problem, the pore pressure variation is omitted. Therefore, we just need to define a solid mechanics solver, which is called `mechanicsSolver`. This solid mechanics solver (see [Solid Mechanics Solver](#)) is based on the Lagrangian finite element formulation. The problem is run as `QuasiStatic` without considering inertial effects. The computational domain is discretized by FE1, which is defined in the `NumericalMethods`

section. The material is named as `rock`, whose mechanical properties are specified in the `Constitutive` section.

```
<Solvers
  gravityVector="{ 0.0, 0.0, 0.0 }">
  <SolidMechanics_LagrangianFEM
    name="mechanicsSolver"
    timeIntegrationOption="QuasiStatic"
    logLevel="1"
    discretization="FE1"
    targetRegions="{ Omega }"
  >
  <LinearSolverParameters
    directParallel="0"/>
  <NonlinearSolverParameters
    newtonTol="1.0e-5"
    newtonMaxIter="15"/>
  </SolidMechanics_LagrangianFEM>
</Solvers>
```

## Constitutive laws

For this drained wellbore problem, we simulate the elastoplastic deformation caused by wellbore expansion. A homogeneous domain with one solid material is assumed, whose mechanical properties are specified in the `Constitutive` section:

```
<Constitutive>
  <ModifiedCamClay
    name="rock"
    defaultDensity="2700"
    defaultRefPressure="-1.2e5"
    defaultRefStrainVol="-0.0"
    defaultShearModulus="4.302e6"
    defaultPreConsolidationPressure="-1.69e5"
    defaultCslSlope="1.2"
    defaultVirginCompressionIndex="0.072676"
    defaultRecompressionIndex="0.014535"
  />
</Constitutive>
```

Recall that in the `SolidMechanics_LagrangianFEM` section, `rock` is designated as the material in the computational domain. Here, `ModifiedCamClay` is used to simulate the elastoplastic behavior of rock.

The following material parameters should be defined properly to reproduce the analytical example:

Name	Type	Default	Description
defaultCslSlope	real64	1	Slope of the critical state line
defaultDensity	real64	required	Default Material Density
defaultPreConsolidationPressure	real64	-1.5	Initial preconsolidation pressure
defaultRecompressionIndex	real64	0.002	Recompression Index
defaultRefPressure	real64	-1	Reference Pressure
defaultRefStrainVol	real64	0	Reference Volumetric Strain
defaultShearModulus	real64	-1	Elastic Shear Modulus Parameter
defaultVirginCompressionIndex	real64	0.005	Virgin compression index
name	string	required	A name is required for any non-unique nodes

The constitutive parameters such as the density, the bulk modulus, and the shear modulus are specified in the International System of Units.

## Initial and boundary conditions

The next step is to specify fields, including:

- The initial value (the in-situ stresses and traction at the wellbore wall have to be initialized)
- The boundary conditions (the reduction of wellbore pressure and constraints of the outer boundaries have to be set)

In this tutorial, we need to specify isotropic horizontal stress ( $\sigma_h = -100$  kPa) and vertical stress ( $\sigma_v = -160$  kPa). To reach equilibrium, a compressive traction  $P_w = -100$  kPa is instantaneously applied at the wellbore wall `rneg` at time  $t = 0$  s, which will then be gradually increased to a higher value (-300 kPa) to let wellbore expand. The remaining parts of the outer boundaries are subjected to roller constraints. These boundary conditions are set up through the `FieldSpecifications` section.

```
<FieldSpecifications>
  <FieldSpecification
    name="stressXX"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions"
    fieldName="rock_stress"
    component="0"
    scale="-1.0e5"/>

  <FieldSpecification
    name="stressYY"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions"
    fieldName="rock_stress"
    component="1"
    scale="-1.0e5"/>

  <FieldSpecification
    name="stressZZ"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions"
    fieldName="rock_stress"
    component="2"
    scale="-1.6e5"/>

  <Traction
    name="ExternalLoad"
    setNames="{ rneg }"
    objectPath="faceManager"
    scale="-1.0e5"
    tractionType="normal"
    functionName="timeFunction"/>

  <FieldSpecification
    name="xconstraint"
    objectPath="nodeManager"
```

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```

    fieldName="totalDisplacement"
    component="0"
    scale="0.0"
    setNames="{ xneg, xpos }"/>

<FieldSpecification
  name="yconstraint"
  objectPath="nodeManager"
  fieldName="totalDisplacement"
  component="1"
  scale="0.0"
  setNames="{ tneg, tpos, ypos }"/>

<FieldSpecification
  name="zconstraint"
  objectPath="nodeManager"
  fieldName="totalDisplacement"
  component="2"
  scale="0.0"
  setNames="{ zneg, zpos }"/>
</FieldSpecifications>

```

With `tractionType="normal"`, traction is applied to the wellbore wall `rneg` as a pressure specified from the product of scale `scale="-1.0e5"` and the outward face normal. A table function `timeFunction` is used to define the time-dependent traction `ExternalLoad`. The coordinates and values form a time-magnitude pair for the loading time history. In this case, the loading magnitude increases linearly as the time evolves.

```

<Functions>
  <TableFunction
    name="timeFunction"
    inputVarNames="{ time }"
    coordinates="{ 0.0, 1.0 }"
    values="{ 1.0, 3.0 }"/>
</Functions>

```

You may note :

- All initial value fields must have `initialCondition` field set to 1;
- The `setName` field points to the previously defined set to apply the fields;
- `nodeManager` and `faceManager` in the `objectPath` indicate that the boundary conditions are applied to the element nodes and faces, respectively;
- `fieldName` is the name of the field registered in GEOSX;
- Component 0, 1, and 2 refer to the x, y, and z direction, respectively;
- And the non-zero values given by `scale` indicate the magnitude of the loading;
- Some shorthand, such as `xneg` and `xpos`, are used as the locations where the boundary conditions are applied in the computational domain. For instance, `xneg` means the face of the computational domain located at the left-most extent in the x-axis, while `xpos` refers to the face located at the right-most extent in the x-axis. Similar shorthands include `ypos`, `yneg`, `zpos`, and `zneg`;
- The mud pressure loading has a negative value due to the negative sign convention for compressive stress in GEOSX.

The parameters used in the simulation are summarized in the following table.

Symbol	Parameter	Units	Value
$P_r$	Reference Pressure	[kPa]	120
$G$	Shear Modulus	[kPa]	4302
$P_c$	PreConsolidation Pressure	[kPa]	169
$M$	Slope of CSL	[-]	1.2
$c_c$	Virgin Compression Index	[-]	0.072676
$c_r$	Recompression Index	[-]	0.014535
$\sigma_h$	Horizontal Stress	[kPa]	-100
$\sigma_v$	Vertical Stress	[kPa]	-160
$a_0$	Initial Well Radius	[m]	0.1
$P_w$	Mud Pressure	[kPa]	-300

## Inspecting results

In the above example, we requested silo-format output files. We can therefore import these into VisIt and use python scripts to visualize the outcome. The following figure shows the distribution of  $\sigma_{\theta\theta}$  in the near wellbore region.

The figure below shows the comparisons between the numerical predictions (marks) and the corresponding analytical solutions (solid curves) with respect to the distributions of normal stress components, stress path, the supporting wellbore pressure and wellbore size. It is evident that the predictions well match the analytical results.

## To go further

### Feedback on this example

For any feedback on this example, please submit a [GitHub issue on the project's GitHub page](#).

## Deviated Poro-Elastic Wellbore Subjected to Fluid Injection

### Problem description

This example aims to solve a typical injection problem of a deviated wellbore subjected to a fluid pressure loaded at wellbore wall. The problem geometry is generated with the internal wellbore mesh generator. Open hole completion and poroelastic deformation are assumed. The coupled poroelastic solver, which combines the solid mechanics solver and the single phase flow solver, is hereby employed to solve this specific problem. In-situ stresses and gravity effect are excluded from this example. Please refer to the case *Deviated Poro-Elastic Wellbore Subjected to In-situ Stresses and Pore Pressure* for in-situ stresses and pore pressure effects.

Analytical solutions of the pore pressure, the radial and hoop stresses in the near wellbore region are expressed in the Laplace space as (Detournay and Cheng, 1988) :

$$\begin{aligned}
 p &= p_0 \frac{k_0(R\sqrt{s})}{s k_0(\sqrt{s})} \\
 \sigma_{rr} &= -b \frac{1-2\nu}{1-\nu} p_0 \frac{-R k_1(R\sqrt{s}) + k_1(\sqrt{s})}{R^2 \sqrt{s^3} k_0(\sqrt{s})} \\
 \sigma_{\theta\theta} &= -b \frac{1-2\nu}{1-\nu} p - \sigma_{rr}
 \end{aligned}$$

where  $s$  is the Laplace variable normalized by the fluid diffusion coefficient,  $k_0$  and  $k_1$  are respectively the modified Bessel functions of second kind of order 0 and 1,  $R$  is the dimensionless radial coordinate that is defined by the radial coordinate normalized by the wellbore radius,  $\nu$  is the Poisson ratio and  $b$  is the Biot coefficient. Fluid pressure and

DB: plot\_00020000  
Cycle: 200      Time: 1

Mesh  
Var: Omega

Pseudocolor  
Var: Omega\_Solid\_MaterialFields/principalStressVector0\_magnitude

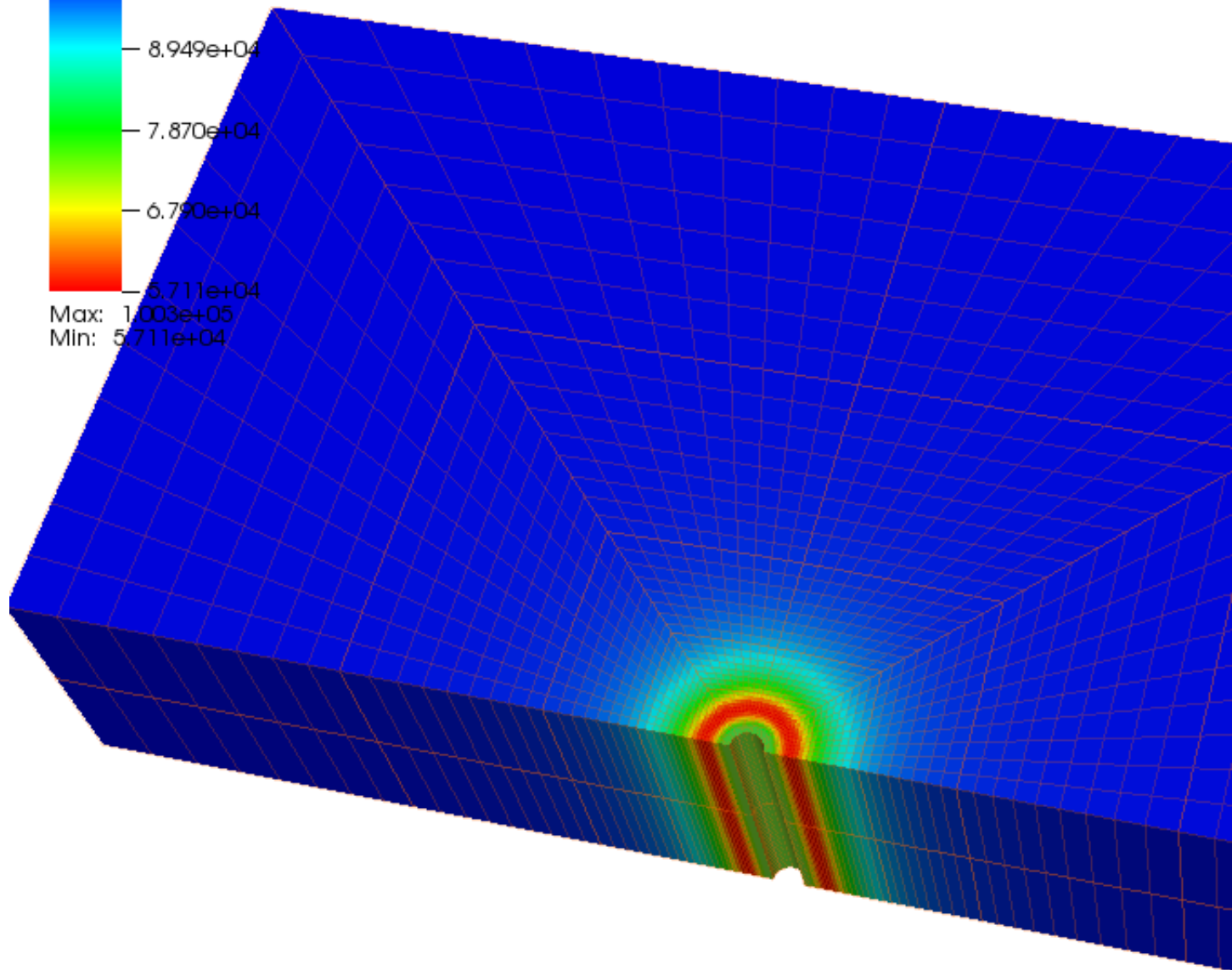
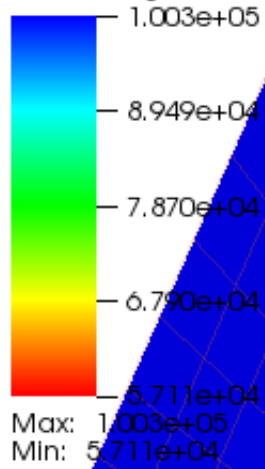
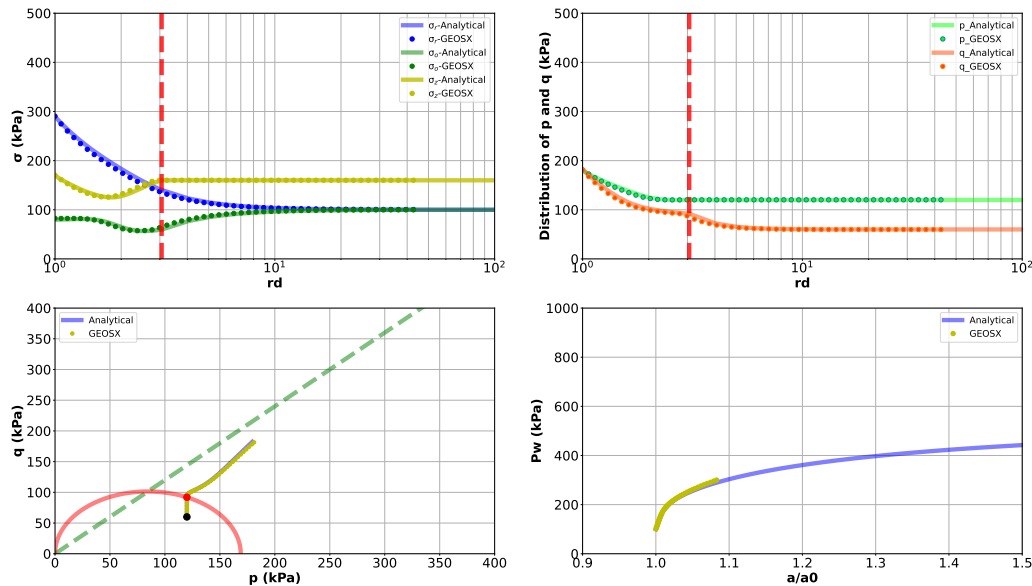


Fig. 4.43: Simulation result of  $\sigma_{\theta\theta}$



stresses in time space are obtained from these analytical expressions by the inverse Laplace transform (see the attached Python script for more details).

### Input file

Everything required is contained within two GEOSX xml files that are located at:

```
inputFiles/wellbore/DeviatedPoroElasticWellbore_Injection_base.xml
```

```
inputFiles/wellbore/DeviatedPoroElasticWellbore_Injection_benchmark.xml
```

In this example, we would focus our attention on the `Solvers` and the `Mesh` tags.

### Poroelectric solver

The coupled Poroelectric solver, that defines a coupling strategy between the solid mechanics solver `SolidMechanicsLagrangianSSLE` and the single phase flow solver `SinglePhaseFVM`, is required for solving this wellbore problem.

```
<SinglePhasePoromechanics
  name="poroSolve"
  solidSolverName="lagsolve"
  flowSolverName="SinglePhaseFlow"
  logLevel="1"
  targetRegions="{ Omega }">
```

```
<SolidMechanicsLagrangianSSLE
  name="lagsolve"
  timeIntegrationOption="QuasiStatic"
  discretization="FE1"
```

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```

logLevel="0"
targetRegions="{ Omega }"
>

```

```

<SinglePhaseFVM
  name="SinglePhaseFlow"
  logLevel="1"
  discretization="singlePhaseTPFA"
  targetRegions="{ Omega }">

```

## Deviated wellbore mesh

The internal wellbore mesh generator `InternalWellbore` is employed to create the mesh of this wellbore problem. The radius of the wellbore and the size of the surrounding rock formation are defined by a vector `radius`. In the tangent direction, `theta` angle is specified from 0 to 180 degree for a half of the domain regarding its symmetry. Note that the whole domain could be specified with a `theta` angle from 0 to 360 degree, if modeling complicated scenarios. The trajectory of the well is defined by `trajectory`. In this example, the wellbore is inclined in the x-z plane by an angle of 45 degree. The `autoSpaceRadialElems` parameter allows optimally increasing the element size from local zone around the wellbore to the far-field zone, which is set to 1 to activate this option. The `useCartesianOuterBoundary` transforms the far-field boundary to a squared shape to enforce a Cartesian aligned outer boundary, which eases the loading of the far-field boundary conditions. In this example, this value is set to 0 for the single region along the radial direction.

```

<Mesh>
  <InternalWellbore
    name="mesh1"
    elementTypes="{ C3D8 }"
    radius="{ 0.1, 4 }"
    theta="{ 0, 180 }"
    zCoords="{ -1, 1 }"
    nr="{ 30 }"
    nt="{ 80 }"
    nz="{ 10 }"
    trajectory="{ { -1.0, 0.0, -1.0 },
                  { 1.0, 0.0, 1.0 } }"
    autoSpaceRadialElems="{ 1 }"
    useCartesianOuterBoundary="0"
    cellBlockNames="{ cb1 }"/>
  </InternalWellbore>
</Mesh>

```

## Constitutive law

Isotropic elastic constitutive block `ElasticIsotropic`, with the specified bulk and shear elastic moduli, is considered for the rock around the wellbore. Fluid properties, such as dynamic viscosity and compressibility, are given in the `CompressibleSinglePhaseFluid` constitutive block. The grain bulk modulus, that is required for computing the Biot coefficient, as well as the default porosity are located in the `BiotPorosity` block. The constant permeability is given in the `ConstantPermeability` block.

```

<PorousElasticIsotropic
  name="porousRock"
  solidModelName="rock"

```

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```

porosityModelName="rockPorosity"
permeabilityModelName="rockPerm"/>

<ElasticIsotropic
  name="rock"
  defaultDensity="0"
  defaultBulkModulus="11039657020.4"
  defaultShearModulus="8662741799.83"/>

<!-- BiotCoefficient="0.771"
      BiotModulus=15.8e9 -->
<CompressibleSinglePhaseFluid
  name="water"
  defaultDensity="1000"
  defaultViscosity="0.001"
  referencePressure="0e6"
  compressibility="1.78403329184e-10"
  viscosibility="0.0"/>

<BiotPorosity
  name="rockPorosity"
  grainBulkModulus="48208109259"
  defaultReferencePorosity="0.3"/>

<ConstantPermeability
  name="rockPerm"
  permeabilityComponents="{ 1.0e-17, 1.0e-17, 1.0e-17 }"/>

```

## Boundary conditions

Far-field boundaries are impermeable and subjected to roller constraints. The pressure on the wellbore wall is defined by face pressure field specification. The nodeset generated by the internal wellbore generator for this face is named as rneg. The negative sign of the scale denotes the fluid injection. Initial fluid pressure and the corresponding initial porosity are also given for the computational domain. In this example, uniform isotropic permeability is assumed.

```

<FieldSpecifications>
  <FieldSpecification
    name="initialPorosity"
    initialCondition="1"
    setNames="{all}"
    objectPath="ElementRegions/Omega/cb1"
    fieldName="rockPorosity_porosity"
    scale="0.3"/>

  <FieldSpecification
    name="initialPressure"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions/Omega/cb1"
    fieldName="pressure"
    scale="0e6"/>

  <FieldSpecification
    name="xConstraint"
    objectPath="nodeManager"

```

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```

    fieldName="totalDisplacement"
    component="0"
    scale="0.0"
    setNames="{ xneg, xpos }"/>

<FieldSpecification
  name="yConstraint"
  objectPath="nodeManager"
  fieldName="totalDisplacement"
  component="1"
  scale="0.0"
  setNames="{ tneg, tpos, ypos }"/>

<FieldSpecification
  name="zconstraint"
  objectPath="nodeManager"
  fieldName="totalDisplacement"
  component="2"
  scale="0.0"
  setNames="{ zneg, zpos }"/>

<FieldSpecification
  name="innerPorePressure"
  objectPath="faceManager"
  fieldName="pressure"
  scale="10e6"
  setNames="{ rneg }"/>
</FieldSpecifications>

```

## Results and benchmark

Result of the fluid pressure distribution after 78 s injection is shown in the figure below:

A good agreement between the GEOSX results and the corresponding analytical solutions is shown in the figure below:

## To go further

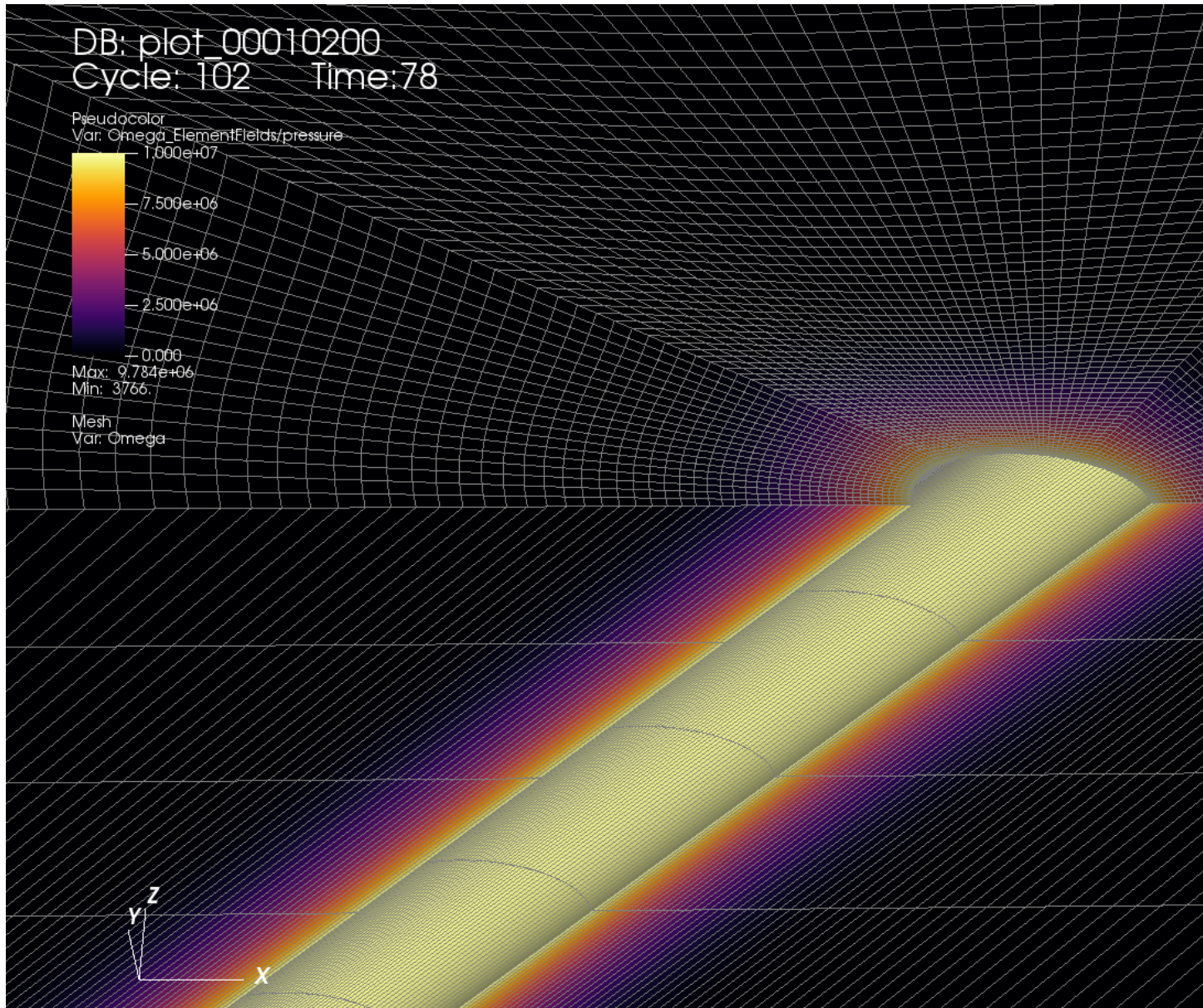
### Feedback on this example

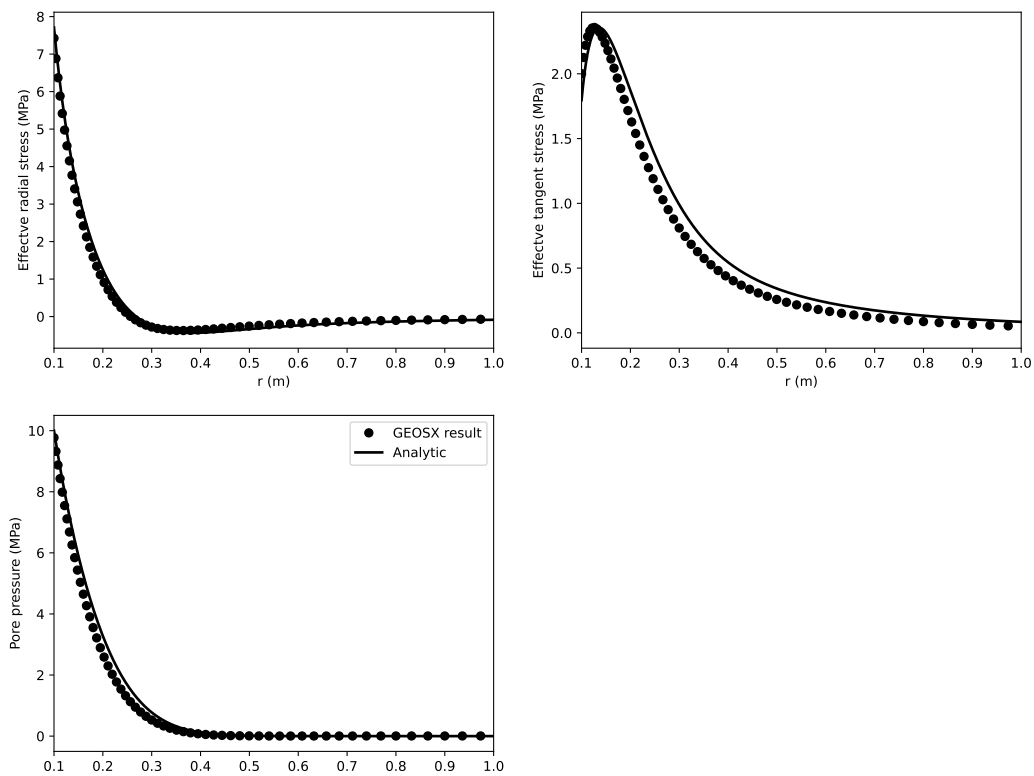
This concludes the deviated poro-elastic wellbore example. For any feedback on this example, please submit a [GitHub issue](#) on the [project's GitHub page](#).

## Deviated Poro-Elastic Wellbore Subjected to In-situ Stresses and Pore Pressure

### Problem description

This example deals with the problem of drilling a deviated poro-elastic wellbore. This is an extension of the poroelastic wellbore example *Deviated Poro-Elastic Wellbore Subjected to Fluid Injection* with the consideration of in-situ stresses and in-situ pore pressure. Both pore pressure and mud pressure are supposed to be nil at the borehole wall following the consideration of (Abousleiman and Cui, 1998). Also, the in-situ horizontal stresses are anisotropic, i.e.  $\sigma_{hmax} > \sigma_{hmin}$ . The wellbore trajectory is deviated from the directions of the in-situ stresses. Analytical solutions of the pore pressure, the radial and hoop stresses in the near wellbore region are given by (Abousleiman and Cui, 1998). They are hereby used to verify the modeling predictions.





## Input file

Everything required is contained within two GEOSX xml files that are located at:

```
inputFiles/wellbore/DeviatedPoroElasticWellbore_Drilling_base.xml
```

```
inputFiles/wellbore/DeviatedPoroElasticWellbore_Drilling_benchmark.xml
```

This case is nearly identical to another example *Deviated Poro-Elastic Wellbore Subjected to Fluid Injection*, except for the FieldSpecifications tag. For this specific case, we need to consider following additional field specifications to define the in-situ stresses, in-situ pore pressure, as well as the zero pore pressure at the borehole wall.

```
<FieldSpecification
  name="initialPorePressure"
  initialCondition="1"
  setNames="{all}"
  objectPath="ElementRegions/Omega/cb1"
  fieldName="pressure"
  scale="10e6"/>

  <FieldSpecification
    name="Sx"
    initialCondition="1"
    setNames="{all}"
    objectPath="ElementRegions/Omega/cb1"
    fieldName="rock_stress"
    component="0"
    scale="-21.9e6"/>

    <FieldSpecification
      name="Sy"
      initialCondition="1"
      setNames="{all}"
      objectPath="ElementRegions/Omega/cb1"
      fieldName="rock_stress"
      component="1"
      scale="-12.9e6"/>

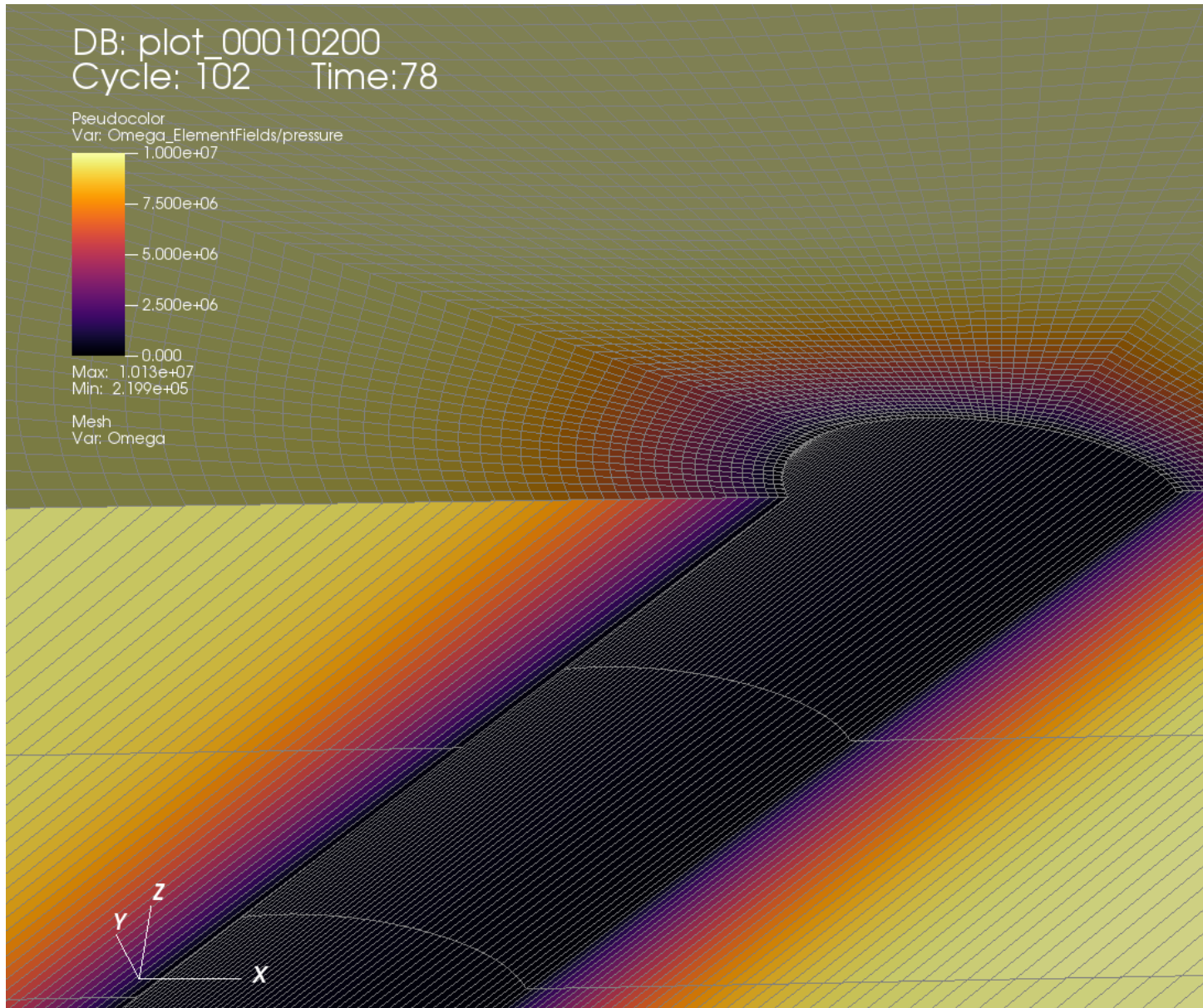
      <FieldSpecification
        name="Sz"
        initialCondition="1"
        setNames="{all}"
        objectPath="ElementRegions/Omega/cb1"
        fieldName="rock_stress"
        component="2"
        scale="-17.9e6"/>

        <FieldSpecification
          name="innerPorePressure"
          objectPath="faceManager"
          fieldName="pressure"
          scale="0e6"
          setNames="{ rneg }"/>
```

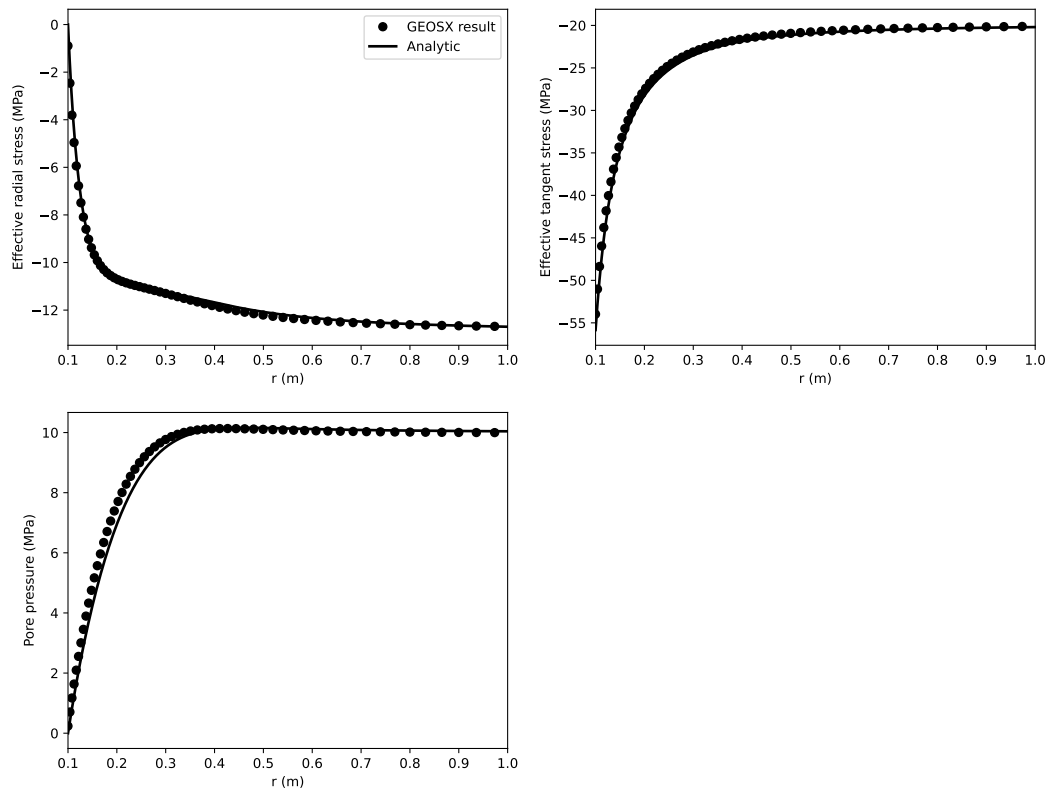
## Results and benchmark

Pore pressure distribution after 78 s injection is shown in the figure below:





A good agreement between the GEOSX results and the corresponding analytical solutions (Abousleiman and Cui, 1998) is shown in the figure below:



## To go further

### Feedback on this example

This concludes the deviated poro-elastic wellbore example with in-situ stresses and pore pressure effects. For any feedback on this example, please submit a [GitHub issue on the project's GitHub page](#).

## Vertical PoroElasto-Plastic Wellbore Problem

### Context

The main objective of this example is to demonstrate how to use the internal wellbore mesh generator and poromechanical solvers in GEOSX to tackle wellbore problems in porous media. In this example, a poroplastic model is applied to find the solution of rock deformation within the vicinity of a vertical wellbore, considering elastoplastic deformation, fluid diffusion and poromechanical coupling effect. To do so, a single phase flow solver is fully coupled with a Lagrangian mechanics solver and the Extended Drucker-Prager model (see [Model: Extended Drucker-Prager](#)) is chosen as the material model for the solid domain. We first solve this problem with a poroelastic model and verify

the modeling results with the corresponding analytical solutions. Then, the verified case is modified to test a poroplastic version, whose results are compared with the ones obtained from the poroelastic case to highlight the impact of plasticity in this specific problem.

### Objectives

At the end of this example you will know:

- how to construct meshes for wellbore problems with the internal wellbore mesh generator,
- how to specify initial and boundary conditions, such as reservoir properties, in-situ stresses, mixed loading (mechanical and fluid) at wellbore wall and far-field constraints,
- how to use multiple solvers in GEOSX for predicting poroplastic deformations in the near wellbore region.

### Input file

This example uses no external input files and everything required is contained within a single GEOSX input file.

The xml input files for the test case with poroelasticity are located at:

```
inputFiles/poromechanics/PoroElasticWellbore_base.xml
inputFiles/poromechanics/PoroElasticWellbore_benchmark.xml
```

The xml input files for the test case with poroplasticity are located at:

```
inputFiles/poromechanics/PoroDruckerPragerWellbore_base.xml
inputFiles/poromechanics/PoroDruckerPragerWellbore_benchmark.xml
```

### Description of the case

We simulate the wellbore problem subjected to anisotropic horizontal stress ( $\sigma_h$  and  $\sigma_H$ ) and vertical stress ( $\sigma_v$ ), as shown below. This is a vertical wellbore, which is drilled in a porous medium. By changing the wellbore supporting pressure, the mechanical deformation of the reservoir rock will be induced and evolve with time, due to fluid diffusion and coupling effect. Considering inelastic constitutive behavior, the reservoir rock in the near wellbore region will experience elastoplastic deformation and a plastic zone will be developed and expand with time. To setup the base case, a poroelastic version is employed to find the poroelastic solutions of this wellbore problem, which are verified with the analytical solution (Detournay and Cheng, 1993) from the literature. Following that, a poroplastic version is built and used to obtain the temporal and spatial solutions of pore pressure, displacement and stress fields around the wellbore, considering induced plastic deformation.

All inputs for this case are contained inside a single XML file. In this example, we focus our attention on the Mesh tags, the Solver tags, the Constitutive tags, and the FieldSpecifications tags.

### Mesh

The following figure shows the generated mesh that is used for solving this wellbore problem

Let us take a closer look at the geometry of this wellbore problem. We use the internal mesh generator `InternalWellbore` to create a rock domain ( $10\text{ m} \times 5\text{ m} \times 2\text{ m}$ ), with a wellbore of initial radius equal to 0.1 m. Coordinates of `trajectory` defines the wellbore trajectory, which represents a perfect vertical well in this example. By turning on `autoSpaceRadialElems="{ 1 }"`, the internal mesh generator automatically sets number and spacing of elements in the radial direction, which overrides the values of `nr`. With `useCartesianOuterBoundary="0"`, a Cartesian aligned outer boundary on the outer block is enforced. In this way, a structured three-dimensional mesh is created with  $100 \times 80 \times 2$  elements in the radial, tangential and z directions, respectively. All the elements are eight-node hexahedral elements (C3D8) and refinement is performed to conform with the wellbore geometry. This mesh is defined as a cell block with the name `cb1`.



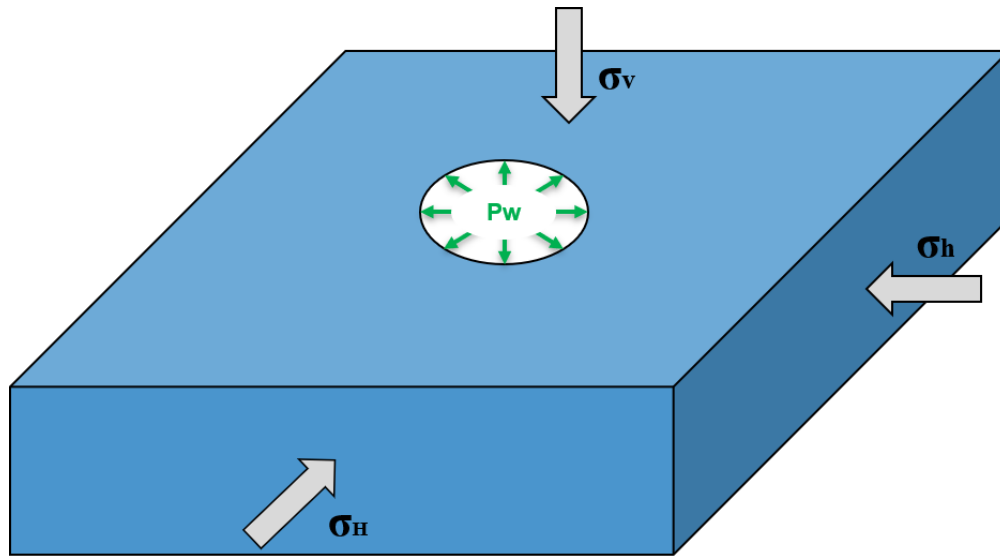


Fig. 4.44: Sketch of the wellbore problem

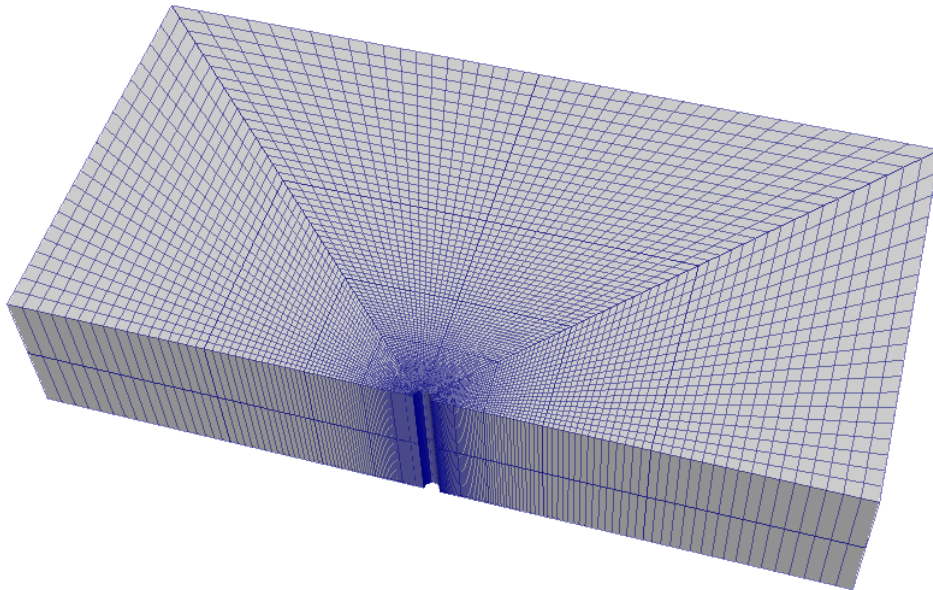


Fig. 4.45: Generated mesh for the wellbore problem

```

<Mesh>
  <InternalWellbore
    name="mesh1"
    elementTypes="{ C3D8 }"
    radius="{ 0.1, 5.0 }"
    theta="{ 0, 180 }"
    zCoords="{ -1, 1 }"
    nr="{ 40 }"
    nt="{ 80 }"
    nz="{ 2 }"
    trajectory="{ { 0.0, 0.0, -1.0 },
                  { 0.0, 0.0, 1.0 } }"
    autoSpaceRadialElems="{ 1 }"
    useCartesianOuterBoundary="0"
    cellBlockNames="{ cb1 }"/>
  </Mesh>

```

## Solid mechanics solver

GEOSX is a multi-physics platform. Different combinations of physics solvers available in the code can be applied in different regions of the domain and be functional at different stages of the simulation. The `Solvers` tag in the XML file is used to list and parameterize these solvers.

To specify a coupling between two different solvers, we define and characterize each single-physics solver separately. Then, we customize a *coupling solver* between these single-physics solvers as an additional solver. This approach allows for generality and flexibility in constructing multi-physics solvers. The order of specifying these solvers is not restricted in GEOSX. Note that end-users should give each single-physics solver a meaningful and distinct name, as GEOSX will recognize these single-physics solvers based on their customized names and create user-expected coupling.

As demonstrated in this example, to setup a poromechanical coupling, we need to define three different solvers in the XML file:

- the mechanics solver, a solver of type `SolidMechanics_LagrangianFEM` called here `mechanicsSolver` (more information here: [Solid Mechanics Solver](#)),

```

<SolidMechanics_LagrangianFEM
  name="mechanicsSolver"
  timeIntegrationOption="QuasiStatic"
  logLevel="1"
  discretization="FE1"
  targetRegions="{ Omega }"
  >
    <NonlinearSolverParameters
      newtonTol = "1.0e-5"
      newtonMaxIter = "15"
    />
  </SolidMechanics_LagrangianFEM>

```

- the single-phase flow solver, a solver of type `SinglePhaseFVM` called here `SinglePhaseFlowSolver` (more information on these solvers at [Singlephase Flow Solver](#)),

```

<SinglePhaseFVM
  name="SinglePhaseFlowSolver"
  logLevel="1"
  discretization="singlePhaseTPFA"

```

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```
targetRegions="{Omega}">
<NonlinearSolverParameters
  newtonTol = "1.0e-6"
  newtonMaxIter = "8"
/>
</SinglePhaseFVM>
</Solvers>
```

- the coupling solver (SinglePhasePoromechanics) that will bind the two single-physics solvers above, which is named as PoromechanicsSolver (more information at [Poromechanics Solver](#)).

```
<Solvers gravityVector="{0.0, 0.0, 0.0}">
<SinglePhasePoromechanics
  name="PoromechanicsSolver"
  solidSolverName="mechanicsSolver"
  flowSolverName="SinglePhaseFlowSolver"
  logLevel="1"
  targetRegions="{Omega}">
  <LinearSolverParameters
    solverType="direct"
    directParallel="0"
    logLevel="0"
  />
  <NonlinearSolverParameters
    newtonMaxIter = "40"
  />
</SinglePhasePoromechanics>
```

The two single-physics solvers are parameterized as explained in their corresponding documents.

In this example, let us focus on the coupling solver. This solver (PoromechanicsSolver) uses a set of attributes that specifically describe the coupling process within a poromechanical framework. For instance, we must point this solver to the designated fluid solver (here: SinglePhaseFlowSolver) and solid solver (here: mechanicsSolver). These solvers are forced to interact through the porousMaterialNames="{porousRock}" with all the constitutive models. We specify the discretization method (FE1, defined in the NumericalMethods section), and the target regions (here, we only have one, Omega). More parameters are required to characterize a coupling procedure (more information at [Poromechanics Solver](#)). In this way, the two single-physics solvers will be simultaneously called and executed for solving the wellbore problem here.

## Discretization methods for multiphysics solvers

Numerical methods in multiphysics settings are similar to single physics numerical methods. In this problem, we use finite volume for flow and finite elements for solid mechanics. All necessary parameters for these methods are defined in the NumericalMethods section.

As mentioned before, the coupling solver and the solid mechanics solver require the specification of a discretization method called FE1. In GEOSX, this discretization method represents a finite element method using linear basis functions and Gaussian quadrature rules. For more information on defining finite elements numerical schemes, please see the dedicated [Finite Element Discretization](#) section.

The finite volume method requires the specification of a discretization scheme. Here, we use a two-point flux approximation scheme (singlePhaseTPFA), as described in the dedicated documentation (found here: [Finite Volume Discretization](#)).

```

<NumericalMethods>
  <FiniteElements>
    <FiniteElementSpace
      name="FE1"
      order="1"/>
    </FiniteElements>
  <FiniteVolume>
    <TwoPointFluxApproximation
      name="singlePhaseTPFA"
    />
  </FiniteVolume>
</NumericalMethods>

```

## Constitutive laws

For this test problem, the solid and fluid materials are named as `rock` and `water` respectively, whose mechanical properties are specified in the `Constitutive` section. In this example, different material models, linear elastic isotropic model (see *Model: Elastic Isotropic*) and Extended Drucker-Prager model (see *Model: Extended Drucker-Prager*), are used to solve the mechanical deformation, which is the only difference between the poroelastic and poroplastic cases in this example.

For the poroelastic case, `PorousElasticIsotropic` model is used to describe the linear elastic isotropic response of rock to loading. And the single-phase fluid model `CompressibleSinglePhaseFluid` is selected to simulate the flow of water upon injection:

```

<Constitutive>
  <PorousElasticIsotropic
    name="porousRock"
    solidModelName="rock"
    porosityModelName="rockPorosity"
    permeabilityModelName="rockPerm"
  />
  <ElasticIsotropic
    name="rock"
    defaultDensity="2700"
    defaultBulkModulus="1.1111e10"
    defaultShearModulus="8.3333e9"
  />
  <CompressibleSinglePhaseFluid
    name="water"
    defaultDensity="1000"
    defaultViscosity="0.001"
    referencePressure="0e6"
    referenceDensity="1000"
    compressibility="2.09028227021e-10"
    referenceViscosity="0.001"
    viscosibility="0.0"
  />
  <BiotPorosity
    name="rockPorosity"
    grainBulkModulus="1.0e27"
    defaultReferencePorosity="0.3"
  />
  <ConstantPermeability
    name="rockPerm"
  />

```

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```

    permeabilityComponents="{1.0e-20, 1.0e-20, 1.0e-20}"
  />
</Constitutive>

```

For the poroplastic case, `PorousExtendedDruckerPrager` model is used to simulate the elastoplastic behavior of rock. And the single-phase fluid model `CompressibleSinglePhaseFluid` is employed to handle the storage and flow of water:

```

<Constitutive>
  <PorousExtendedDruckerPrager
    name="porousRock"
    solidModelName="rock"
    porosityModelName="rockPorosity"
    permeabilityModelName="rockPerm"
  />
  <ExtendedDruckerPrager
    name="rock"
    defaultDensity="2700"
    defaultBulkModulus="1.1111e10"
    defaultShearModulus="8.3333e9"
    defaultCohesion="1.0e6"
    defaultInitialFrictionAngle="15.27"
    defaultResidualFrictionAngle="23.05"
    defaultDilationRatio="1.0"
    defaultHardening="0.01"
  />
  <CompressibleSinglePhaseFluid
    name="water"
    defaultDensity="1000"
    defaultViscosity="0.001"
    referencePressure="0e6"
    referenceDensity="1000"
    compressibility="2.09028227021e-10"
    referenceViscosity="0.001"
    viscosibility="0.0"
  />
  <BiotPorosity
    name="rockPorosity"
    grainBulkModulus="1.0e27"
    defaultReferencePorosity="0.3"
  />
  <ConstantPermeability
    name="rockPerm"
    permeabilityComponents="{1.0e-20, 1.0e-20, 1.0e-20}"
  />
</Constitutive>

```

As for the material parameters, `defaultInitialFrictionAngle`, `defaultResidualFrictionAngle` and `defaultCohesion` denote the initial friction angle, the residual friction angle, and cohesion, respectively, as defined by the Mohr-Coulomb failure envelope. As the residual friction angle `defaultResidualFrictionAngle` is larger than the initial one `defaultInitialFrictionAngle`, a strain hardening model is automatically chosen, whose hardening rate is given as `defaultHardening="0.01"`. If the residual friction angle is set to be less than the initial one, strain weakening will take place. `defaultDilationRatio="1.0"` corresponds to an associated flow rule. If using an incompressible fluid, the user can lower the fluid compressibility `compressibility` to 0. The constitutive parameters such as the density, the bulk modulus, and the shear modulus are specified in the International System of Units. A stress-dependent

porosity model `rockPorosity` and constant permeability `rockPerm` model are defined in this section.

## Initial and boundary conditions

The next step is to specify fields, including:

- The initial value (the in-situ stresses and pore pressure have to be initialized)
- The boundary conditions (traction and fluid loading at the wellbore wall and constraints of the outer boundaries have to be set)

In this example, we need to specify anisotropic horizontal stress ( $\sigma_h = -9.0$  MPa and  $\sigma_H = -11.0$  MPa) and vertical stress ( $\sigma_v = -12.0$  MPa). A compressive traction (InnerMechanicalLoad)  $P_w = -10$  MPa and fluid loading (InnerFluidLoad)  $P_f = 10$  MPa are applied at the wellbore wall `rneg`. The remaining parts of the outer boundaries are subjected to roller constraints. These boundary conditions are set up through the `FieldSpecifications` section.

```
<FieldSpecifications>
  <FieldSpecification
    name="stressXX"
    initialCondition="1"
    setNames="{all}"
    objectPath="ElementRegions/Omega/cb1"
    fieldName="rock_stress"
    component="0"
    scale="-9.0e6"
  />

  <FieldSpecification
    name="stressYY"
    initialCondition="1"
    setNames="{all}"
    objectPath="ElementRegions/Omega/cb1"
    fieldName="rock_stress"
    component="1"
    scale="-11.0e6"
  />

  <FieldSpecification
    name="stressZZ"
    initialCondition="1"
    setNames="{all}"
    objectPath="ElementRegions/Omega/cb1"
    fieldName="rock_stress"
    component="2"
    scale="-12.0e6"
  />

  <FieldSpecification
    name="initialPressure"
    initialCondition="1"
    setNames="{all}"
    objectPath="ElementRegions/Omega/cb1"
    fieldName="pressure"
    scale="0e6"
  />
```

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```

<FieldSpecification
  name="xconstraint"
  objectPath="nodeManager"
  fieldName="totalDisplacement"
  component="0"
  scale="0.0"
  setNames="{xneg, xpos}"
/>

<FieldSpecification
  name="yconstraint"
  objectPath="nodeManager"
  fieldName="totalDisplacement"
  component="1"
  scale="0.0"
  setNames="{tneg, tpos, ypos}"
/>

<FieldSpecification
  name="zconstraint"
  objectPath="nodeManager"
  fieldName="totalDisplacement"
  component="2"
  scale="0.0"
  setNames="{zneg, zpos}"
/>

<Traction
  name="InnerMechanicalLoad"
  setNames="{ rneg }"
  objectPath="faceManager"
  scale="-10.0e6"
  tractionType="normal"
  functionName="timeFunction"
/>

<FieldSpecification
  name="InnerFluidLoad"
  setNames="{ rneg }"
  objectPath="faceManager"
  fieldName="pressure"
  scale="10e6"
  functionName="timeFunction"
/>
</FieldSpecifications>

```

With `tractionType="normal"`, traction is applied to the wellbore wall `rneg` as a pressure specified from the product of scale `scale="-10.0e6"` and the outward face normal. A table function `timeFunction` is used to define the time-dependent loading. The coordinates and values form a time-magnitude pair for the loading time history. In this case, the loading magnitude is given as:

```

<Functions>
  <TableFunction
    name="timeFunction"
    inputVarNames="{time}"
    coordinates="{0.0, 0.1, 1e6}"

```

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```

    values="{0.0, 1.0, 1.0}"
  />
</Functions>

```

You may note :

- All initial value fields must have `initialCondition` field set to 1;
- The `setName` field points to the previously defined box to apply the fields;
- `nodeManager` and `faceManager` in the `objectPath` indicate that the boundary conditions are applied to the element nodes and faces, respectively;
- `fieldName` is the name of the field registered in GEOSX;
- Component 0, 1, and 2 refer to the x, y, and z direction, respectively;
- And the non-zero values given by `scale` indicate the magnitude of the loading;
- Some shorthands, such as `xneg` and `xpos`, are used as the locations where the boundary conditions are applied in the computational domain. For instance, `xneg` means the portion of the computational domain located at the left-most in the x-axis, while `xpos` refers to the portion located at the right-most area in the x-axis. Similar shorthands include `ypos`, `yneg`, `zpos`, and `zneg`;
- The mud pressure loading has a negative value due to the negative sign convention for compressive stress in GEOSX.

The parameters used in the simulation are summarized in the following table, which are specified in the `Constitutive` and `FieldSpecifications` sections.

Symbol	Parameter	Unit	Value
$K$	Bulk Modulus	[GPa]	11.11
$G$	Shear Modulus	[GPa]	8.33
$C$	Cohesion	[MPa]	1.0
$\phi_i$	Initial Friction Angle	[degree]	15.27
$\phi_r$	Residual Friction Angle	[degree]	23.05
$c_h$	Hardening Rate	[-]	0.01
$\sigma_h$	Min Horizontal Stress	[MPa]	-9.0
$\sigma_H$	Max Horizontal Stress	[MPa]	-11.0
$\sigma_v$	Vertical Stress	[MPa]	-12.0
$a_0$	Initial Well Radius	[m]	0.1
$P_w$	Traction at Well	[MPa]	-10.0
$P_f$	Fluid Pressure at Well	[MPa]	10.0
$\rho_f$	Fluid Density	[kg/m <sup>3</sup> ]	1000.0
$\mu$	Fluid Viscosity	[Pa s]	0.001
$c_f$	Fluid Compressibility	[Pa <sup>-1</sup> ]	$2.09 \cdot 10^{-10}$
$\kappa$	Matrix Permeability	[m <sup>2</sup> ]	$1.0 \cdot 10^{-20}$
$\phi$	Porosity	[-]	0.3

## Inspecting results

As defined in the `Events` section, we run this simulation for 497640 seconds. In the above examples, we requested silo-format output files. We can therefore import these into VisIt and use python scripts to visualize the outcome. Please note that a non-dimensional time is used in the analytical solution, and the end time here leads to a non-dimensional end time of  $t^* = 4.62$ .



Using the poroelastic solver, below figure shows the prediction of pore pressure distribution upon fluid injection.

For the above poroelastic example, an analytical solution (Detournay and Cheng, 1993) is hereby employed to verify the accuracy of the numerical results. Following figure shows the comparisons between the numerical predictions (marks) and the corresponding analytical solutions (solid curves) with respect to the distributions of pore pressure, radial displacement, effective radial and tangential stresses along the minimum horizontal stress direction (x-axis). One can observe that GEOSX results correlate very well with the analytical solutions for the poroelastic case.

For the same 3D wellbore problem, the poroplastic case is thereafter tested and compared with the poroelastic one. The figure below shows the distribution of  $\sigma_{yy}$  in the near wellbore region for both cases. As expected, a relaxation of the tangential stress along the direction of minimum horizontal stress is detected, which can be attributed to the plastic response of the rock.

By using python scripts, we can extract the simulation results along any direction and provide detailed comparisons between different cases. Here, the pore pressure, radial displacement, radial and tangential effective stresses along the direction of minimum horizontal stress are obtained at different time steps and plotted against the corresponding ones of the poroelastic case. Because of fluid diffusion and coupling effect, following figure shows that these solutions evolve with time for both cases. As mentioned above, a plastic zone is developed in the vicinity of the wellbore, due to stress concentration. As for the far field region, these two cases become almost identical, with the rock deformation governed by poroelasticity.

## To go further

### Feedback on this example

This concludes the example on PoroPlasticity Model for Wellbore Problems. For any feedback on this example, please submit a [GitHub issue on the project's GitHub page](#).

### For more details

- More on plasticity models, please see *Model: Extended Drucker-Prager*.
- More on multiphysics solvers, please see *Poromechanics Solver*.

## 4.2 Performance Benchmarks

## 4.3 Application Studies

## 4.4 pygeosx Examples

The *pygeosx — GEOSX in Python* enables users to query and/or manipulate the GEOSX datastructure with python in real-time. The following examples show how to use *pygeosx* interface and the supplemental *pygeosx\_tools\_package*.

A quick warning: these examples are recommended for advanced users only. The *pygeosx* interface requires users to be familiar with the GEOSX datastructure, and can trigger the code to crash if an object/variable is inappropriately modified.

### 4.4.1 In Situ Data Monitor

#### Objectives

At the end of this example you will know:

- how to run a problem using the *pygeosx* interface,

DB: plot\_00103200  
Cycle: 1032 Time:497640

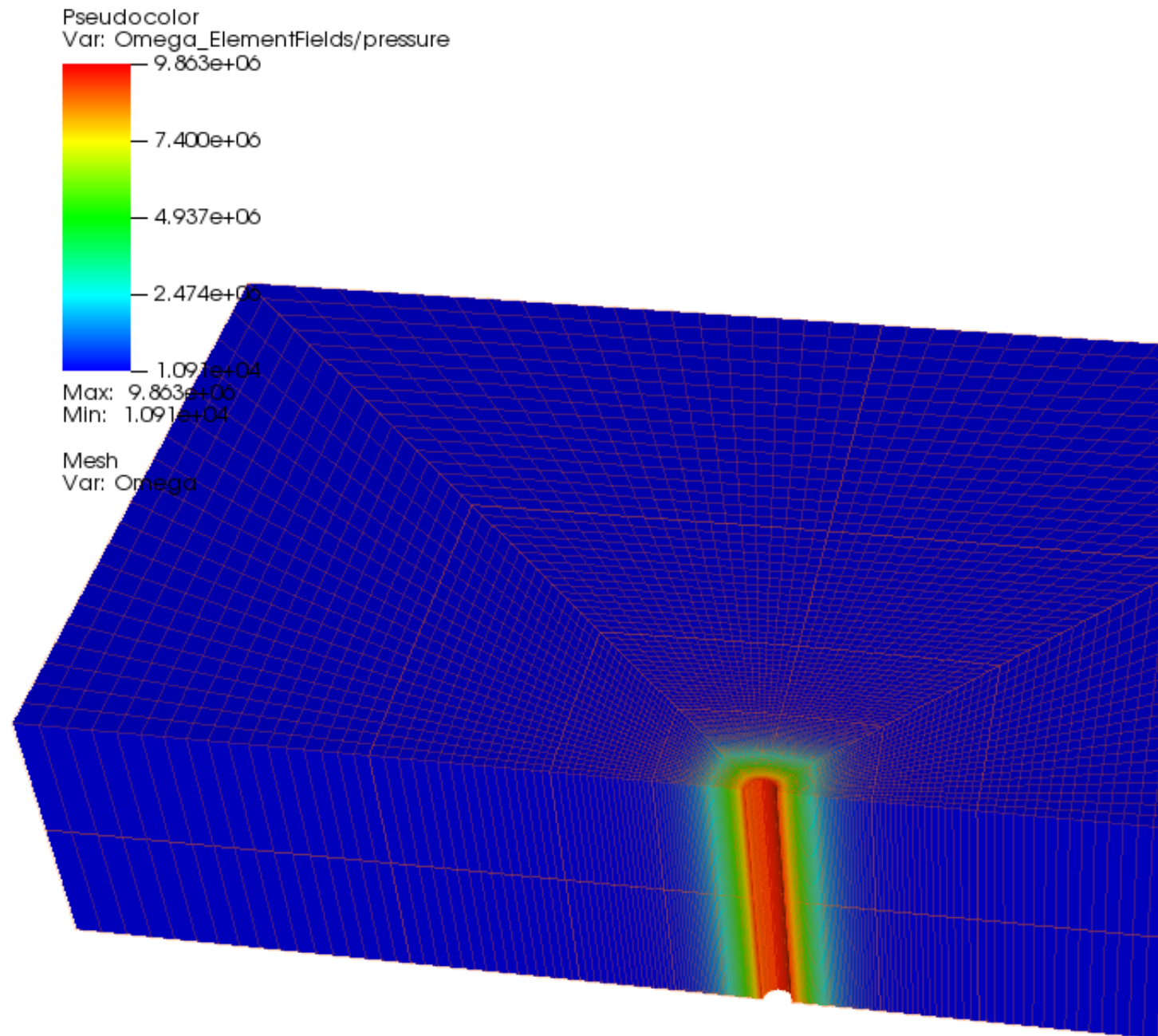


Fig. 4.46: Simulation result of pore pressure distribution

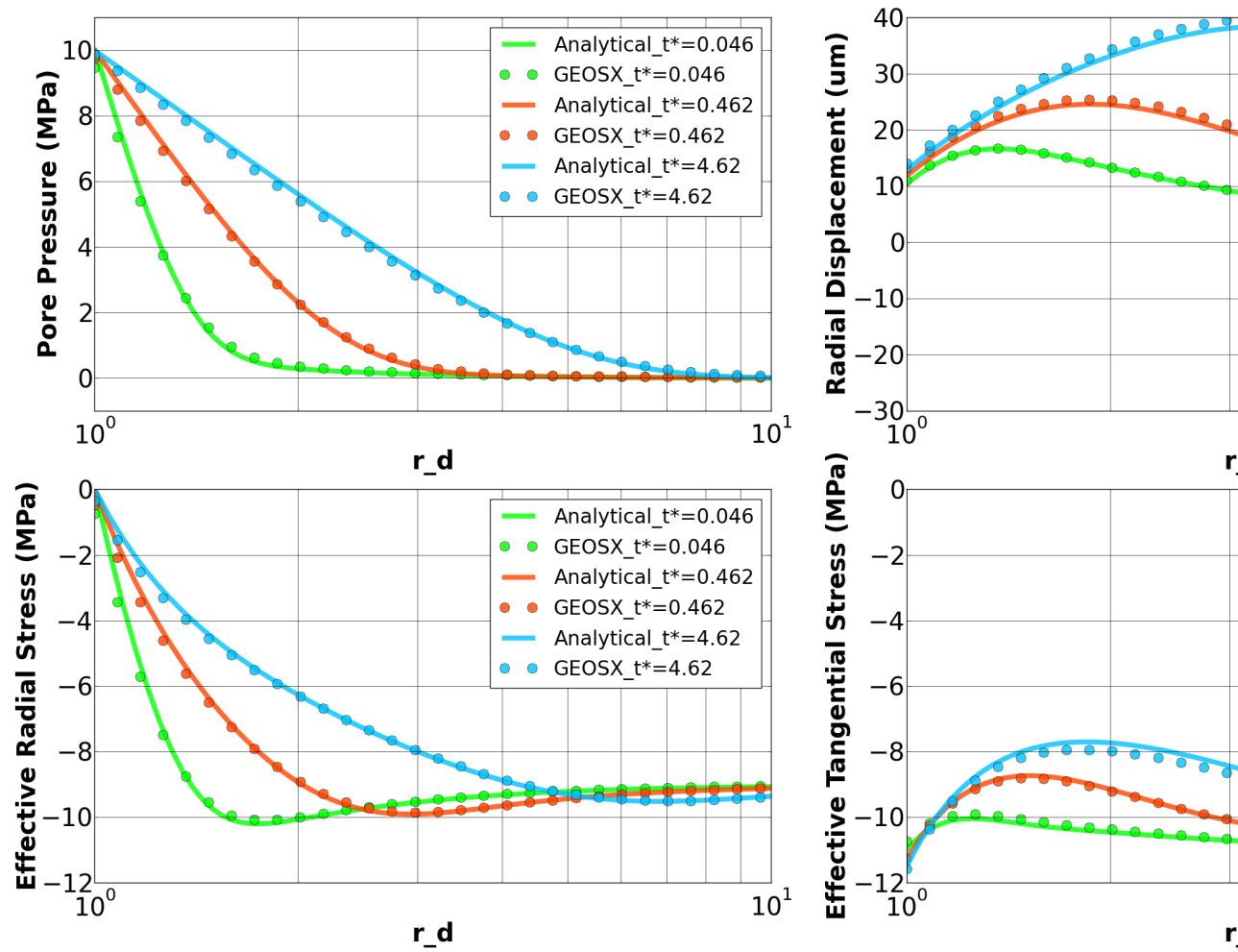


Fig. 4.47: Comparing GEOSX results with analytical solutions

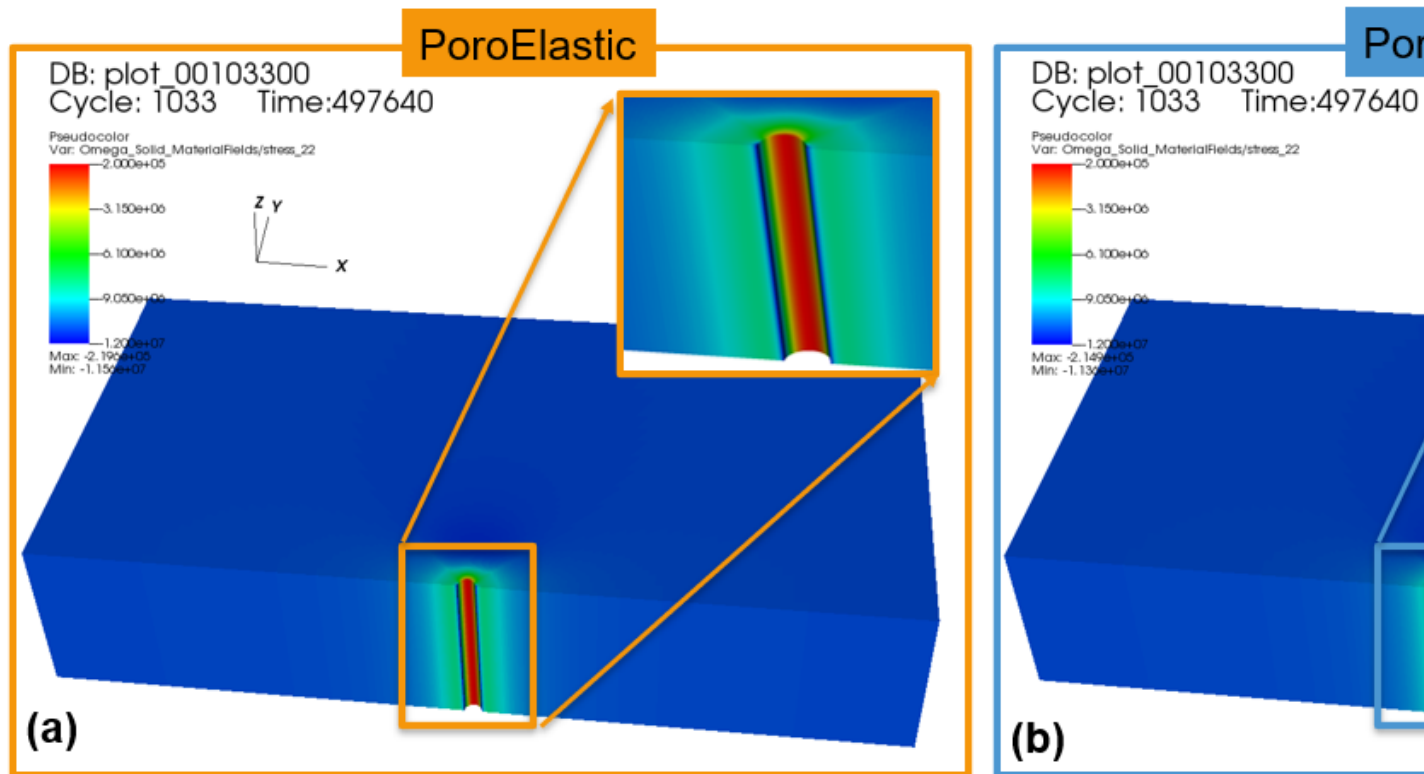


Fig. 4.48: Simulation result of  $S_{yy}$ : PoroElastic vs. PoroPlastic

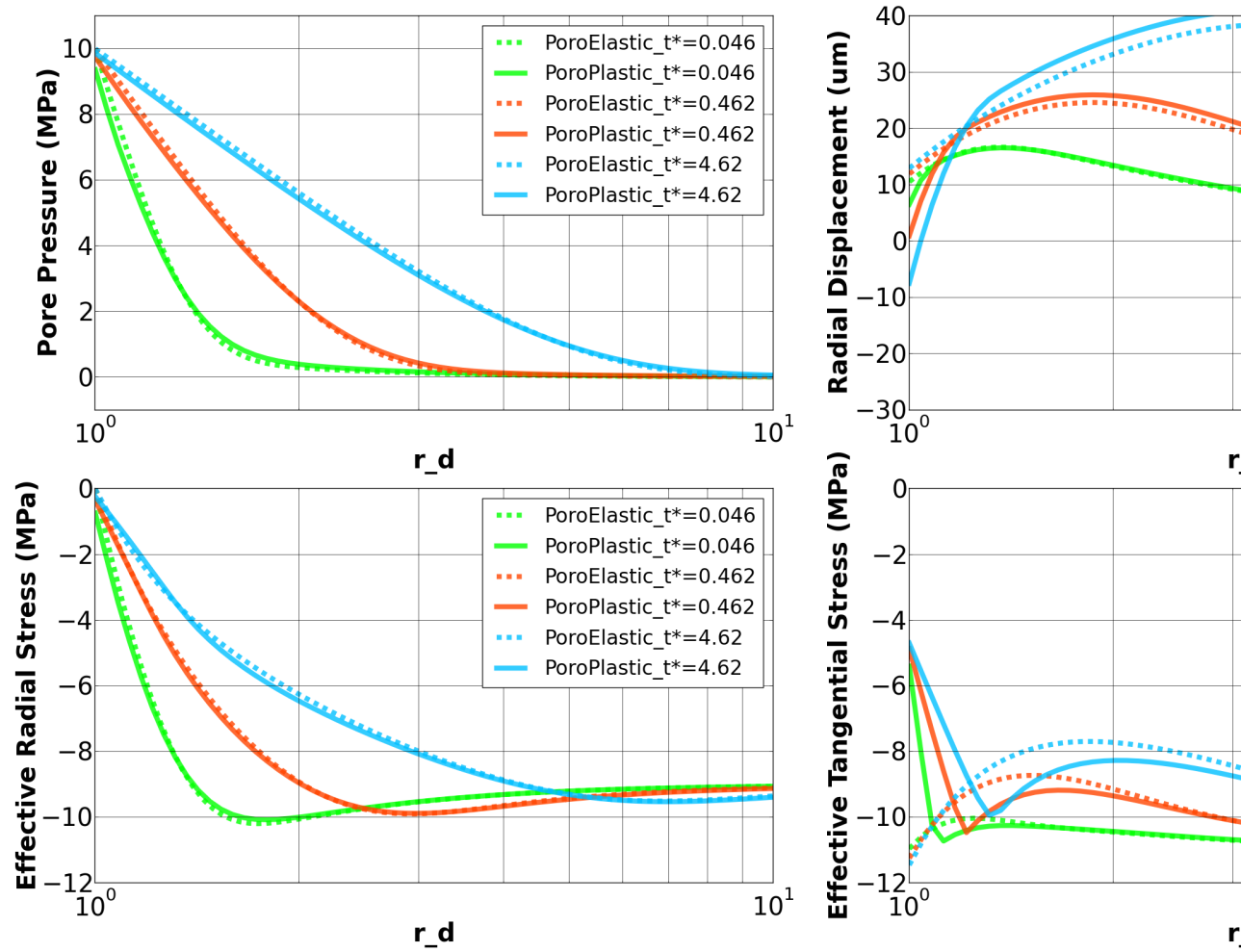


Fig. 4.49: Comparing the PoroPlastic case with the PoroElastic case at different times

- how to process advanced xml features using pygeosx,
- how to extract and monitor values within the GEOSX datastructure in real-time

### Input files

This example requires two input xml files and one python script located at:

```
GEOSX/examples/pygeosxExamples/hydraulicFractureWithMonitor
```

### Description of the case

This example is derived from this basic example: *Hydraulic Fracturing*, which solves for the propagation of a single hydraulic fracture within a heterogeneous reservoir. The pygeosx interface is used to monitor the maximum hydraulic aperture and fracture extents over time.

### XML Configuration

The input xml file for this example requires some modification in order to work with pygeosx. First, we use the advanced xml input features to include the base problem and override the *table\_root* parameter that points to the table files. Note that these paths will need to be updated if you run this problem outside of the example directory.

```
<Included>
  <File
    name="../../hydraulicFracturing/heterogeneousInSituProperties/
    ↪heterogeneousInSitu_singleFracture.xml"/>
  </Included>

  <Parameters>
    <Parameter
      name="table_root"
      value="../../hydraulicFracturing/heterogeneousInSituProperties/tables"/>
    </Parameter>
```

Next, we add a new entry to the output block Python and an entry in the Events block. Whenever the python event is triggered, GEOSX will pause and return to the controlling python script (in this case, every 10 cycles).

```
<Outputs>
  <Python
    name="pythonOutput"/>
</Outputs>

<Events>
  <PeriodicEvent
    name="python"
    cycleFrequency="10"
    target="/Outputs/pythonOutput"/>
</Events>
```

### Python Script

Problems that use the pygeosx interface are driven by a custom python script. To begin, we import a number of packages and check whether this is a parallel run. The custom packages include pygeosx, which provides an interface to GEOSX, and pygeosx\_tools, which provides a number of common tools for working with the datastructure and dealing with parallel communication.

```

from mpi4py import MPI
import pygeosx
from pygeosx_tools import wrapper
from geosx_xml_tools.main import preprocess_parallel
import matplotlib.pyplot as plt

```

In the next step, we apply the xml preprocessor to resolve the advanced xml features. Note that this step will modify the input arguments to reflect the location of the compiled xml file, which is processed directly by GEOSX. The script then initializes GEOSX and receives the *problem* handle, which is the scripts view into the datastructure. There is an opportunity to interact with the GEOSX before the initial conditions are set, which we do not use in this example.

```

# Get the MPI rank
comm = MPI.COMM_WORLD
rank = comm.Get_rank()

# Initialize the code and set initial conditions
args = preprocess_parallel()
problem = pygeosx.initialize(rank, args)
pygeosx.apply_initial_conditions()

```

To extract information from the problem, you need to know the full path (or ‘key’) to the target object. These keys can be quite long, and can change depending on the xml input. In the next step, we use a method from the `pygeosx_tools` package to search for these keys using a list of keywords. If the keys are known beforehand, then this step could be skipped. Note that these functions will throw an error if they do not find a matching key, or if they find multiple matching keys.

```

# Rather than specifying the wrapper paths explicitly,
# search for them using a set of filters
fracture_location_key = wrapper.get_matching_wrapper_path(problem, ['Fracture',
↪ 'elementCenter'])
fracture_aperture_key = wrapper.get_matching_wrapper_path(problem, ['Fracture',
↪ 'effectiveAperture'])

```

Next, we setup a dictionary that will allow us to use `pygeosx_tools` to automatically query the problem. The root level of this dictionary contains the target keys (fracture location and aperture) and the required *time* key. These each point to a sub-dictionary that holds an axis label, a scale factor, and an empty list to hold the time history. The target dictionaries also hold an entry *fhandle*, which contains a matplotlib figure handle that we can use to display the results.

```

# Setup values to record
records = {fracture_location_key: {'label': 'Fracture Extents (m)',
                                   'scale': 1.0,
                                   'history': [],
                                   'fhandle': plt.figure()},
           fracture_aperture_key: {'label': 'Aperture (mm)',
                                   'scale': 1e3,
                                   'history': [],
                                   'fhandle': plt.figure()},
           'time': {'label': 'Time (min)',
                    'scale': 1.0 / 60.0,
                    'history': []}}

```

After setting up the problem, we enter the main problem loop. Upon calling `pygeosx.run()`, the code will execute until a Python event is triggered in the Event loop. At those points, we have the option to interact with the problem before continuing processing. Here, we use `pygeosx_tools` to query the datastructure and occasionally plot the results to the screen.

```
# Setup values to record
records = {fracture_location_key: {'label': 'Fracture Extents (m)',
                                  'scale': 1.0,
                                  'history': [],
                                  'fhandle': plt.figure()},
          fracture_aperture_key: {'label': 'Aperture (mm)',
                                  'scale': 1e3,
                                  'history': [],
                                  'fhandle': plt.figure()},
          'time': {'label': 'Time (min)',
                   'scale': 1.0 / 60.0,
                   'history': []}}
```

## Manual Query

To obtain and manually inspect an object in the problem, you can use the methods in `pygeosx_tools.wrapper`. These are designed to handle any parallel communication that may be required in your analysis. For example, to get the fracture aperture as a numpy array, you could call:

```
from pygeosx_tools import wrapper

# (problem initialization / configuration)

# Grab aperture as a numpy array, using three different approaches

# Local copy (the write flag indicates that we do not plan to modify the result)
aperture_local = wrapper.get_wrapper(problem, fracture_aperture_key, write_flag=False)

# Global copy on the root rank
aperture_global = wrapper.gather_wrapper(problem, fracture_aperture_key)

# Global copy on the all ranks
aperture_global = wrapper.allgather_wrapper(problem, fracture_aperture_key)
```

## Running the Problem

To run the problem, you must use the specific version of python where pygeosx is installed. This is likely located here:

```
GEOSX/[build_dir]/lib/PYGEOSX/bin/python
```

Note that you may need to manually install the `pygeosx_tools` package (and its pre-requisites) into this python distribution. To do so, you can run the following:

```
cd GEOSX/[build_dir]/lib/PYGEOSX/bin
pip install --upgrade ../../../../src/coreComponents/python/modules/pygeosx_tools_
↪package/
```

To run the code, you will call the `pygeosx` run script with python, and supply the typical `geosx` command-line arguments and any parallel arguments. For example:

```
# Load the correct python environment
# If you are not using a bash shell, you may need to target one of
# the other activation scripts
```

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```
source GEOSX/[build_dir]/lib/PYGEOSX/bin/activate

# Move to the correct directory and run
cd /path/to/problem
srun -n 36 -ppdebug python hydraulicFractureWithMonitor.py -i hydraulicFracture.xml -
↪x 6 -y 2 -z 3 -o hf_results
```

## To go further

### Feedback on this example

For any feedback on this example, please submit a [GitHub issue](#) on the project's [GitHub page](#).

### For more details

- More on advanced xml features, please see [Advanced XML Features](#).
- More on the pygeosx interface, please see [pygeosx — GEOSX in Python](#).

## 4.4.2 Initial Condition Modification

### Objectives

At the end of this example you will know:

- how to modify GEOSX arrays using pygeosx
- handle parallel communication with pygeosx\_tools

### Input files

This example requires an input xml and python script located at:

```
GEOSX/examples/pygeosxExamples/sedovWithStressFunction
```

### Description of the case

This example is derived from the sedov integrated test (*GEOSX/src/coreComponents/physicsSolvers/solidMechanics/integratedTests/sedov*) which looks at the propagation of elastic waves due to an initial stress field. The pygeosx interface is used to modify the initial conditions of the problem to something of our choosing.

### XML Configuration

As before, the basic sedov input xml file for this example requires some modification in order to work with pygeosx. First, we use the advanced xml input features to include the base problem (this path may need to be updated, depending on where you run the problem).

```
<Included>
  <File
    name="../../inputFiles/solidMechanics/sedov.xml"/>
  </Included>
```

Next, we add a new entry to the output block Python and an entry in the Events block. Whenever the python event is triggered, GEOSX will pause and return to the controlling python script.

```

<Events>
  <PeriodicEvent
    name="python"
    cycleFrequency="5"
    target="/Outputs/pythonOutput"/>
  </Events>

<Outputs>
  <Python
    name="pythonOutput"/>
  </Outputs>

```

## Python Script

Similar to the previous example, the python script begins by importing the required packages, applying the xml pre-processor, GEOSX initialization, and key search.

```

# Get the MPI rank
comm = MPI.COMM_WORLD
rank = comm.Get_rank()

# Initialize the code and set initial conditions
args = preprocess_parallel()
problem = pygeosx.initialize(rank, args)
pygeosx.apply_initial_conditions()

# Rather than specifying the wrapper paths explicitly,
# search for them using a set of filters
location_key = wrapper.get_matching_wrapper_path(problem, ['Region2',
↪ 'elementCenter'])
stress_key = wrapper.get_matching_wrapper_path(problem, ['Region2', 'shale',
↪ 'stress'])
ghost_key = wrapper.get_matching_wrapper_path(problem, ['Region2', 'cb1',
↪ 'ghostRank'])

```

The next steps rely on a python function that we use to set stress. The argument to this function, *x*, is assumed to be a numpy array of element centers:

```

def stress_fn(x):
    """
    Function to set stress values

    Args:
        x (np.ndarray) the element centers

    Returns:
        np.ndarray: stress values
    """
    R = x[:, 0]**2 + x[:, 1]**2 + x[:, 2]**2
    return np.sin(2.0 * np.pi * R / np.amax(R))

```

In the following section, we zero out the initial stress and then set it based on *stress\_fn*. While doing this, we use *wrapper.print\_global\_value\_range* to check on the process.

```

# Print initial stress
wrapper.print_global_value_range(problem, stress_key, 'stress')

```

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```

# Zero out stress
wrapper.set_wrapper_to_value(problem, stress_key, 0.0)
wrapper.print_global_value_range(problem, stress_key, 'stress')

# Set stress via a function
wrapper.set_wrapper_with_function(problem, stress_key, location_key, stress_fn,
↪target_index=0)
wrapper.set_wrapper_with_function(problem, stress_key, location_key, stress_fn,
↪target_index=1)
wrapper.set_wrapper_with_function(problem, stress_key, location_key, stress_fn,
↪target_index=2)
wrapper.print_global_value_range(problem, stress_key, 'stress')

```

Finally, we run the simulation. As an optional step, we extract numpy arrays from the datastructure using different parallel approaches:

```

# Run the code
while pygeosx.run() != pygeosx.COMPLETED:
    wrapper.print_global_value_range(problem, stress_key, 'stress')

# Gather/allgather tests
tmp = wrapper.gather_wrapper(problem, stress_key)
print(wrapper.rank, 'gather', np.shape(tmp), flush=True)

tmp = wrapper.allgather_wrapper(problem, stress_key)
print(wrapper.rank, 'allgather', np.shape(tmp), flush=True)

tmp = wrapper.allgather_wrapper(problem, stress_key, ghost_key=ghost_key)
print(wrapper.rank, 'allgather_ghost_filtered', np.shape(tmp), flush=True)

```

## Running the Problem

To run the problem, you must use the specific version of python where pygeosx is installed. This is likely located here:

```
GEOSX/[build_dir]/lib/PYGEOSX/bin/python
```

Note that you may need to manually install the pygeosx\_tools package (and its pre-requisites) into this python distribution. To do so, you can run the following:

```

cd GEOSX/[build_dir]/lib/PYGEOSX/bin
pip install --upgrade ../../../../src/coreComponents/python/modules/pygeosx_tools_
↪package/

```

To run the code, you will call the pygeosx run script with python, and supply the typical geosx command-line arguments and any parallel arguments. For example:

```

# Load the correct python environment
# If you are not using a bash shell, you may need to target one of
# the other activation scripts
source GEOSX/[build_dir]/lib/PYGEOSX/bin/activate

# Move to the correct directory and run

```

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```
cd /path/to/problem
python run_sedov_problem.py -i modified_sedov.xml -o results
```

### To go further

#### Feedback on this example

For any feedback on this example, please submit a [GitHub issue](#) on the project's [GitHub page](#).

#### For more details

- More on advanced xml features, please see *Advanced XML Features*.
- More on the pygeosx interface, please see *pygeosx — GEOSX in Python*.

Welcome to the GEOSX user guide.

## 5.1 Input Files

### 5.1.1 XML

GEOSX is configured via one (or more) [Extensible Markup Language \(XML\)](#) files. These files contain a set of elements and attributes that closely follow the internal datastructure of GEOSX. When running GEOSX, these files are specified using the *-i* argument:

```
geosx -i input.xml
```

### XML Components

The following illustrates some of the key features of a GEOSX-format xml file:

```
<?xml version="1.0" ?>

<Problem>
  <BlockA
    someAttribute="1.234">

    <!-- Some comment -->
    <BlockB
      name="firstNamedBlock"
      anotherAttribute="0"/>
    <BlockB
      name="secondNamedBlock"
      anotherAttribute="1"/>
```

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```
</BlockA>
</Problem>
```

The two basic components of an xml file are blocks, which are specified using angle brackets (“<BlockA> </BlockA>”), and attributes that are attached to blocks (attributeName=“attributeValue”). Block and attributes can use any ASCII character aside from <, &, ‘, and “ (if necessary, use &lt;, &amp;, &apos;, or &quot;). Comments are indicated as follows: <!-- Some comment -->.

At the beginning of a GEOSX input file, you will find an optional xml declaration (<?xml version=“1.0” ?>) that is used to indicate the format to certain text editors. You will also find the root *Problem* block, where the GEOSX configuration is placed. Note that, aside from these elements and commented text, the xml format requires that no other objects exist at the first level.

In the example above, there is a single element within the *Problem* block: *BlockA*. *BlockA* has an attribute *someAttribute*, which has a value of 1.234, and has three children: a commented string “Some comment” and two instances of *BlockB*. The *name* attribute is required for blocks that allow multiple instances, and should include a unique string to avoid potential errors. Where applicable these blocks will be executed in the order in which they are specified in input file.

## 5.1.2 Input Validation

The optional *xmlns:xsi* and *xsi:noNamespaceSchemaLocation* attributes in the *Problem* block can be used to indicate the type of document and the location of the xml schema to the text editor:

```
<Problem
  xmlns:xsi="http://www.w3.org/2001/XMLSchema-instance"
  xsi:noNamespaceSchemaLocation="/path/to/schema.xsd" />
```

The schema contains a list of xml blocks and attributes that are supported by GEOSX, indicates whether a given object is optional or required, and defines the format of the object (string, floating point number, etc.). A copy of the schema is included in the GEOSX source code (/path/to/GEOSX/src/coreComponents/schema/schema.xsd). It can also be generated using GEOSX: `geosx -s schema.xsd`

Many text editors can use the schema to help in the construction of an xml file and to indicate whether it is valid. Using a validation tool is highly recommended for all users. The following instructions indicate how to turn on validation for a variety of tools:

### xmllint

xmllint is a command-line tool that is typically pre-installed on UNIX-like systems. To check whether an input file is valid, run the following command:

```
xmllint --schema /path/to/schema.xsd input_file.xml
```

### Sublime Text

We recommend using the [Exalt](#) or [SublimeLinter\\_xmllint](#) plug-ins to validate xml files within sublime. If you have not done so already, install the sublime [Package Control](#). To install the package, press `ctrl + shift + p`, type and select `Package Control: Install Package`, and search for `exalt` or `SublimeLinter / SublimeLinter-xmllint`. Note that, depending on the circumstances, these tools may indicate only a subset of the validation errors at a given time. Once resolved, the tools should re-check the document to look for any additional errors.

As an additional step for SublimLinter-xmlint, you will need to add a linter configuration. To do so, go to Preferences/Package Settings/SublimeLinter/Settings. In the right-hand side of the new window, add the xmlint configuration:

```
{
  "linters": {
    "xmlint": {
      {
        "args": "--schema /path/to/schema.xsd",
        "styles": [
          {
            "mark_style": "fill",
            "scope": "region.bluish",
            "types": ["error"],
            "icon": "stop",
          }
        ]
      },
    },
  }
}
```

## VS Code

We recommend using the [XML](#) for validating xml files. After installing this extension, you can associate GEOSX format xml files by adding the following entry to the user settings file (replacing *systemId* with the correct path to the schema file):

```
{
  "xml.fileAssociations": [
    {
      "pattern": "**.xml",
      "systemId": "/path/to/GEOSX/src/coreComponents/schema/schema.xsd"
    }
  ]
}
```

## Eclipse

The Eclipse Web Develop Tools includes features for validating xml files. To install them, go to Help -> Eclipse Marketplace, search for the Eclipse Web Developer Tools, install the package, and restart Eclipse. Finally, configure the xml validation preferences under Window -> Preferences -> XML -> XML Files -> Validation. Eclipse will automatically fetch the schema, and validate an active xml file. The editor will highlight any lines with errors, and underline the specific errors.

## GEOSX XML Tools

The `geosx_xml_tools` package, which is used to enable advanced features such as parameters, symbolic math, etc., contains tools for validating xml files. To do so, call the command-line script with the `-s` argument, i.e.: `preprocess_xml input_file.xml -s /path/to/schema.xsd`. After compiling the final xml file, pygeosx will fetch the designated schema, validate, and print any errors to the screen.

Note: Attributes that are using advanced xml features will likely contain characters that are not allowed by their corresponding type pattern. As such, file editors that are configured to use other validation methods will likely identify errors in the raw input file.

### 5.1.3 XML Schema

An XML schema definition (XSD) file lays out the expected structure of an input XML file. During the build process, GEOSX automatically constructs a comprehensive schema from the code's data structure, and updates the version in the source (GEOSX/src/coreComponents/schema/schema.xsd).

#### Schema Components

The first entry in the schema are a set of headers the file type and version. Following this, the set of available simple types for attributes are laid out. Each of these includes a variable type name, which mirrors those used in the main code, and a regular expression, which is designed to match valid inputs. These patterns are defined and documented in `DataTypes::typeRegex`. The final part of the schema is the file layout, beginning with the root `Problem`. Each complex type defines an element, its children, and its attributes. Each attribute defines the input name, type, default value, and/or usage. Comments preceding each attribute are used to relay additional information to the users.

#### Automatic Schema Generation

A schema may be generated by calling the main code with the `-s` argument, e.g.: `geosx -s schema.xsd` (Note: this is done automatically during the build process). To do this, GEOSX does the following:

- 1) Initialize the GEOSX data structure.
- 2) Initialize objects that are registered to catalogs via `ManagedGroup::ExpandObjectCatalogs()`.
- 3) Recursively write element and attribute definitions to the schema using information stored in GEOSX groups and wrappers.
- 4) Define any expected deviations from the schema via `ManagedGroup::SetSchemaDeviations()`.

### 5.1.4 Advanced XML Features

The `geosx_xml_tools` python package adds a set of advanced features to the GEOSX xml format: units, parameters, and symbolic expressions. See [Python Tools Setup](#) for details on setup instructions, and [GEOSX XML Tools](#) for package API details.

#### Usage

An input file that uses advanced xml features requires preprocessing before it can be used with GEOSX. The preprocessor writes a compiled xml file to the disk, which can be read directly by GEOSX and serves as a permanent record for the simulation. There are three ways to apply the preprocessor:

- 1) Automatic Preprocessing: Substituting `geosx` for `geosx_preprocessed` when calling the code will automatically apply the preprocessor to the input xml file, and then pass the remaining arguments to GEOSX. With this method, the compiled xml files will have the suffix `‘.preprocessed’`. Before running the code, the compiled xml file will also be validated against the xml schema.



```
# Serial example
geosx_preprocessed -i input.xml

# Parallel example
srun -n 2 geosx_preprocessed -i input.xml -x 2
```

- 2) Manual Preprocessing: For this approach, xml files are preprocessed manually by the user with the *preprocess\_xml* script. These files can then be submitted to GEOSX separately:

```
# The -c argument is used to manually specify the compiled name
preprocess_xml -i input.xml -c input.xml.processed
geosx -i input.xml.processed

# Otherwise, a random name will be chosen by the tool
compiled_input=$(preprocess_xml input.xml)
geosx -i $compiled_input
```

- 3) Python / pygeosx: The preprocessor can also be applied directly in python or in pygeosx simulations. An example of this is method is provided here: *GEOSX/examples/pygeosxExamples/hydraulicFractureWithMonitor/*.

Each of these options support specifying multiple input files via the command line (e.g. *geosx\_preprocessed -i input\_a.xml -i input\_b.xml*). They also support any number of command-line parameter overrides (e.g. *geosx\_preprocessed -i input\_a.xml -p parameter\_a alpha -p parameter\_b beta*).

## Included Files

Both the XML preprocessor and GEOSX executable itself provide the capability to build complex multi-file input decks by including XML files into other XML files.

The files to be included are listed via the `<Included>` block. There maybe any number of such blocks. Each block contains a list of `<File name="...">` tags, each indicating a file to include. The *name* attribute must contain either an absolute or a relative path to the included file. If the path is relative, it is treated as relative to the location of the referring file. Included files may also contain includes of their own, i.e. it is possible to have *a.xml* include *b.xml* which in turn includes *c.xml*.

---

**Note:** When creating multi-file input decks, it is considered best practice to use relative file paths. This applies both to XML includes, and to other types of file references (for example, table file names). Relative paths keep input decks both relocatable within the file system and sharable between users.

---

XML preprocessor's merging capabilities are more advanced than GEOSX built-in ones. Both are outlined below.

## XML preprocessor

The merging approach is applied recursively, allowing children to include their own files. Any potential conflicts are handled via the following scheme:

- **Merge two objects if:**
  - At the root level an object with the matching tag exists.
  - If the “name” attribute is present and an object with the matching tag and name exists.
  - Any preexisting attributes on the object are overwritten by the donor.
- Otherwise append the XML structure with the target.

## GEOSX

GEOSX's built-in processing simply inserts the included files' content (excluding the root node) into the XML element tree, at the level of `<Included>` tag. Partial merging is handled implicitly by GEOSX's data structure, which treats repeated top-level XML blocks as if they are one single block. This is usually sufficient for merging together top-level input sections from multiple files, such as multiple `<FieldSpecifications>` or `<Events>` sections, but more complex cases may require the use of preprocessor.

---

**Note:** While GEOSX's XML processing is capable of handling any number of `<Included>` block at any level, the XML schema currently produced by GEOSX only allows a single such block, and only directly within the `<Problem>` tag. Inputs that use multiple blocks or nest them deeper may run but will fail to validate against the schema. This is a known discrepancy that may be fixed in the future.

---

## Parameters

Parameters are a convenient way to build a configurable and human-readable input XML. They are defined via a block in the XML structure. To avoid conflicts with other advanced features, parameter names can include upper/lower case letters and underscores. Parameters may have any value, including:

- Numbers (with or without units)
- A path to a file
- A symbolic expression
- Other parameters
- Etc.

They can be used as part of any input xml attribute as follows:

- `$x_par$` (preferred)
- `$x_par`
- `$.x_par`
- `$.x_par$`

Attributes can be used across Included files, but cannot be used to set the names of included files themselves. The following example uses parameters to set the root path for a table function, which is then scaled by another parameter:

```
<Parameters>
  <Parameter
    name="flow_scale"
    value="0.5"/>
  <Parameter
    name="table_root"
    value="/path/to/table/root"/>
</Parameters>

<FieldSpecifications>
  <SourceFlux
    name="sourceTerm"
    objectPath="ElementRegions/Region1/block1"
    scale="$flow_scale$"
    functionName="flow_rate"
    setNames="{ source }"/>
  </SourceFlux>
</FieldSpecifications>
```

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```

</FieldSpecifications>

<Functions>
  <TableFunction
    name="flow_rate"
    inputVarNames="{time}"
    coordinateFiles="{${table_root$}/time_flow.geos}"
    voxelFile="${table_root$}/flow.geos"
    interpolation="linear"/>
</Functions>

```

Any number of parameter overrides can be issued from the command line using the *-p name value* argument in the preprocessor script. Note that if the override value contains any spaces, it may need to be surrounded by quotation marks (*-p name "paramter with spaces"*).

## Units

The units for any input values to GEOSX can be in any self-consistent system. In many cases, it is useful to override this behavior by explicitly specifying the units of the input. These are specified by appending a valid number with a unit definition in square braces. During pre-processing, these units are converted into base-SI units (we plan to support other unit systems in the future).

The unit manager supports most common units and SI prefixes, using both long- and abbreviated names (e.g.: c, centi, k, kilo, etc.). Units may include predefined composite units (dyne, N, etc.) or may be built up from sub-units using a python syntax (e.g.: [N], [kg\*m/s\*\*2]). Any (or no) amount of whitespace is allowed between the number and the unit bracket. The following shows a set of parameters with units specified:

```

<Parameters>
  <Parameter name="paramter_a" value="2[m]"/>
  <Parameter name="paramter_b" value="1.2 [cm]"/>
  <Parameter name="paramter_c" value="1.23e4 [bbl/day]"/>
  <Parameter name="paramter_d" value="1.23E-4 [km**2]"/>
</Parameters>

```

Please note that the preprocessor currently does not check whether any user-specified units are appropriate for a given input or symbolic expression.

## Symbolic Expressions

Input XML files can also include symbolic mathematical expressions. These are placed within pairs of backticks (`), and use a limited python syntax. Please note that parameters and units are evaluated before symbolic expressions. While symbolic expressions are allowed within parameters, errors may occur if they are used in a way that results in nested symbolic expressions. Also, note that residual alpha characters (e.g. *sin()*) are removed before evaluation for security. The following shows an example of symbolic expressions:

```

<Parameters>
  <Parameter name="a" value="2[m]"/>
  <Parameter name="b" value="1.2 [cm]"/>
  <Parameter name="c" value="3"/>
  <Parameter name="d" value="1.23e-4"/>
</Parameters>
<Geometry>
  <Box

```

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```
name="perf"
xMin="{`$a$ - 0.2*$b$`, -1e6, -1e6}"
xMax="{`$c$**2 / $d$`, 1e6, 1e6}" />
</Geometry>
```

## Validation

Unmatched special characters (\$, [, ', etc.) in the final xml file indicate that parameters, units, or symbolic math were not specified correctly. If the preprocessor detects these, it will throw an error and exit. Additional validation of the compiled files can be completed with *preprocess\_xml* by supplying the -s argument and the path to the GEOSX schema.

## 5.2 Meshes

The purpose of this document is to explain how users and developers interact with mesh data. This section describes how meshes are handled and stored in GEOSX.

There are two possible methods for generating a mesh: either by using GEOSX's internal mesh generator (for Cartesian meshes only), or by importing meshes from various common mesh file formats. This latter options allows one to work with more complex geometries, such as unstructured meshes comprised of a variety of element types (polyhedral elements).

### 5.2.1 Internal Mesh Generation

#### Basic Example

The Internal Mesh Generator allows one to quickly build simple cartesian grids and divide them into several regions. The following attributes are supported in the input block for InternalMesh:

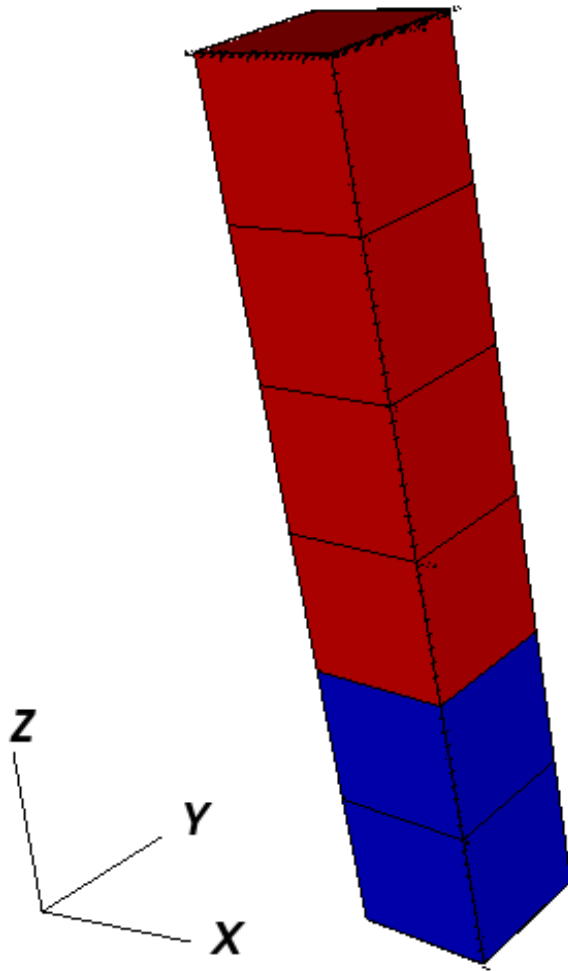
Name	Type	De- fault	Description
cellBlock-Names	string_array	re- quired	Names of each mesh block
element-Types	string_array	re- quired	Element types of each mesh block
name	string	re- quired	A name is required for any non-unique nodes
nx	inte- ger_array	re- quired	Number of elements in the x-direction within each mesh block
ny	inte- ger_array	re- quired	Number of elements in the y-direction within each mesh block
nz	inte- ger_array	re- quired	Number of elements in the z-direction within each mesh block
positionTol- erance	real64	1e-10	A position tolerance to verify if a node belong to a nodeset
trianglePat- tern	integer	0	Pattern by which to decompose the hex mesh into wedges
xBias	real64_array	{1}	Bias of element sizes in the x-direction within each mesh block ( $dx\_left=(1+b)*L/N$ , $dx\_right=(1-b)*L/N$ )
xCords	real64_array	re- quired	x-coordinates of each mesh block vertex
yBias	real64_array	{1}	Bias of element sizes in the y-direction within each mesh block ( $dy\_left=(1+b)*L/N$ , $dy\_right=(1-b)*L/N$ )
yCords	real64_array	re- quired	y-coordinates of each mesh block vertex
zBias	real64_array	{1}	Bias of element sizes in the z-direction within each mesh block ( $dz\_left=(1+b)*L/N$ , $dz\_right=(1-b)*L/N$ )
zCords	real64_array	re- quired	z-coordinates of each mesh block vertex

The following is an example XML <mesh> block, which will generate a vertical beam with two CellBlocks (one in red and one in blue in the following picture).

```
<Mesh>
  <InternalMesh name="mesh"
    elementTypes="C3D8"
    xCords="0, 1"
    yCords="0, 1"
    zCords="0, 2, 6"
    nx="1"
    ny="1"
    nz="2, 4"
    cellBlockNames="cb1 cb2"/>
</Mesh>
```

- name the name of the mesh body
- elementTypes the type of the elements that will be generated.
- xCoord List of x coordinates of the boundaries of the CellBlocks
- yCoord List of y coordinates of the boundaries of the CellBlocks
- zCoord List of z coordinates of the boundaries of the CellBlocks

- `nx` List containing the number of cells in `x` direction within the `CellBlocks`
- `ny` List containing the number of cells in `y` direction within the `CellBlocks`
- `nz` List containing the number of cells in `z` direction within the `CellBlocks`
- `cellBlockNames` List containing the names of the `CellBlocks`



## Mesh Bias

The internal mesh generator is capable of producing meshes with element sizes that vary smoothly over space. This is achieved by specifying `xBias`, `yBias`, and/or `zBias` fields. (Note: if present, the length of these must match `nx`, `ny`, and `nz`, respectively, and each individual value must be in the range  $(-1, 1)$ .)

For a given element block, the average element size will be

$$dx_{average}[i] = \frac{xCoords[i+1] - xCoords[i]}{nx[i]},$$

the element on the left-most side of the block will have size

$$dx_{left}[i] = (1 + xBias[i]) \cdot dx_{average}[i],$$

and the element on the right-most side will have size

$$dx_{right}[i] = (1 - xBias[i]) \cdot dx_{average}[i].$$

The following are the two most common scenarios that occur while designing a mesh with bias:

1. The size of the block and the element size on an adjacent region are known. Assuming that we are to the left of the target block, the appropriate bias would be:

$$xBias[i] = 1 - \frac{nx[i] \cdot dx_{left}[i + 1]}{xCoords[i + 1] - xCoords[i]}$$

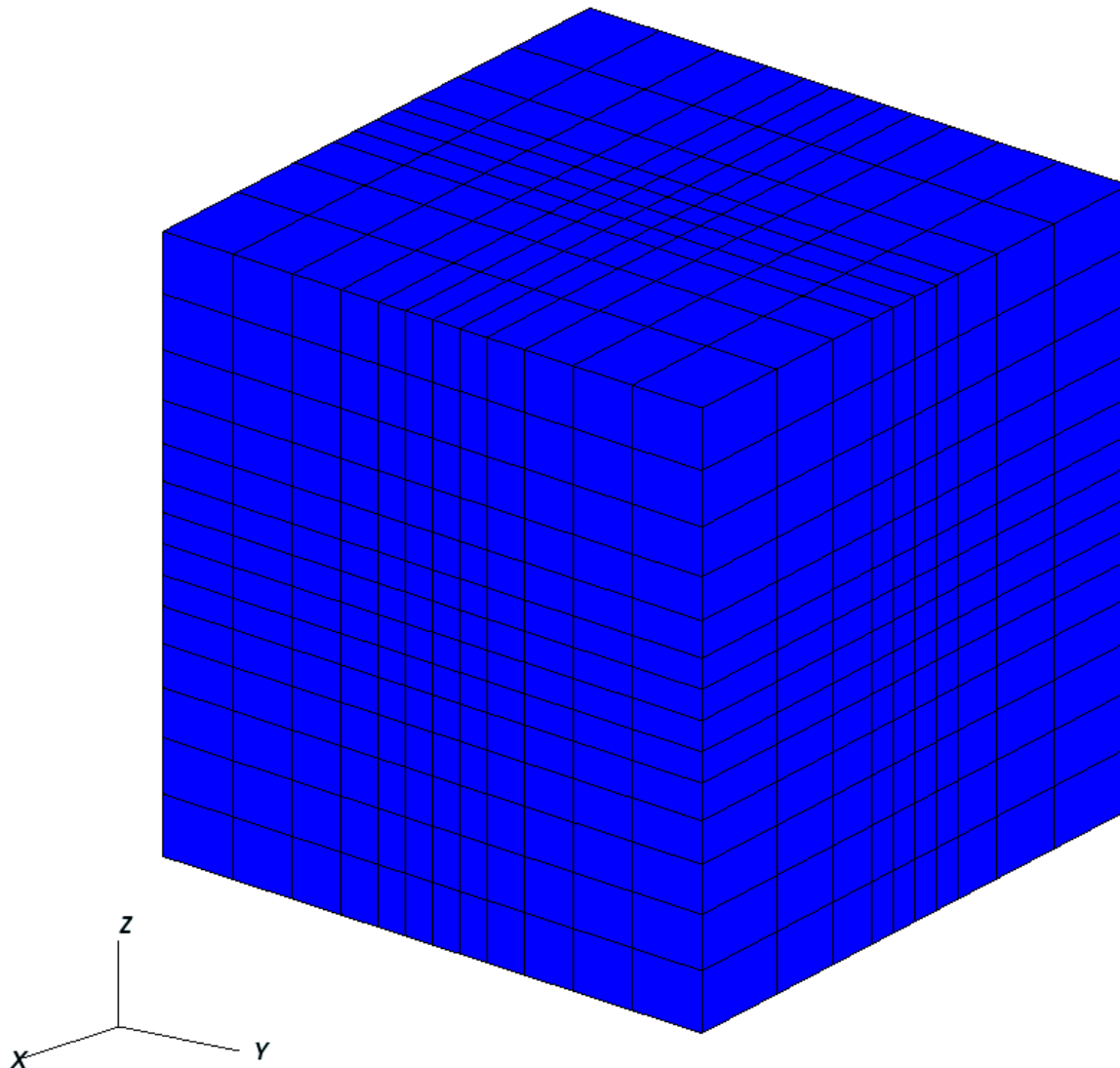
2. The bias of the block and the element size on an adjacent region are known. Again, assuming that we are to the left of the target block, the appropriate size for the block would be:

$$xCoords[i + 1] - xCoords[i] = \frac{nx[i] \cdot dx_{left}[i + 1]}{1 - xBias[i]}$$

The following is an example of a mesh block along each dimension, and an image showing the corresponding mesh. Note that there is a core region of elements with zero bias, and that the transitions between element blocks are smooth.

```
<Mesh>
  <InternalMesh
    name="mesh1"
    elementTypes="{ C3D8 }"
    xCoords="{ -10, -1, 0, 1, 10 }"
    yCoords="{ -10, -1, 0, 1, 10 }"
    zCoords="{ -10, -1, 0, 1, 10 }"
    nx="{ 4, 1, 1, 4 }"
    ny="{ 5, 1, 1, 5 }"
    nz="{ 6, 1, 1, 6 }"
    xBias="{ 0.555, 0, 0, -0.555 }"
    yBias="{ 0.444, 0, 0, -0.444 }"
    zBias="{ 0.333, 0, 0, -0.333 }"
    cellBlockNames="{ cb1 }"/>
  </InternalMesh>
</Mesh>

<Solvers>
  <SolidMechanics_LagrangianFEM
    name="lagsolve"
    strainTheory="1"
    cflFactor="0.25"
    discretization="FE1"
    targetRegions="{ Region2 }"
  />
</Solvers>
```



### Advanced Cell Block Specification

It's possible to generate more complex `CellBlock` using the `InternalMeshGenerator`. For instance, the staircase example is a model which is often used in GEOSX as an integrated test. It defines `CellBlocks` in the three directions to generate a staircase-like model with the following code.

```
<Mesh>
  <InternalMesh name="mesh1"
    elementTypes="{C3D8}"
    xCoords="{0, 5, 10}"
    yCoords="{0, 5, 10}"
    zCoords="{0, 2.5, 5, 7.5, 10}"
    nx="{5, 5}"
    ny="{5, 5}"
    nz="{3, 3, 3, 3}"
    cellBlockNames="{b00,b01,b02,b03,b04,b05,b06,b07,b08,b09,b10,b11,b12,
↪b13,b14,b15}"/>
```

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```

</Mesh>

<ElementRegions>
  <CellElementRegion name="Channel"
    cellBlocks="{b08,b00,b01,b05,b06,b14,b15,b11}"
    materialList="{fluid1, rock, relperm}"/>
  <CellElementRegion name="Barrier"
    cellBlocks="{b04,b12,b13,b09,b10,b02,b03,b07}"
    materialList="{}/>
</ElementRegions>

```

Thus, the generated mesh will be :

## 5.2.2 Using an External Mesh

### Supported Formats

GEOSX provides features to run simulations on unstructured meshes. It uses [PAMELA](#) to read the external meshes and its API to write it into the GEOSX mesh data structure.

The supported mesh format are:

- The [GMSH](#) file format (.msh v2).
- The ECLIPSE file formats (.egrid, .grdecl)

The supported mesh elements for volume elements consist of the following:

- 4 nodes tetrahedra,
- 5 nodes pyramids,
- 6 nodes wedges,
- 8 nodes hexahedra,

The mesh can be divided in several regions. These regions are intended to support different physics or to define different constitutive properties.

- For the GMSH file format, the regions are defined using the [physical entity names](#) provided by GMSH.
- For the ECLIPSE file formats, the regions have to be first defined using the ECLIPSE software.

### Importing the Mesh

#### Importing regions

Several blocks are involved to import an external mesh into GEOSX, defined in the XML input file. These are the `<Mesh>` block and the `<ElementRegions>` block.

The mesh block has the following syntax:

```

<Mesh>
  <PAMELAMesh name="MyMeshName"
    file="/path/to/the/mesh/file.msh"/>
</Mesh>

```

We advise users to use absolute path to the mesh file.

GEOSX uses `ElementRegions` to support different physics or to define different constitutive properties. An `ElementRegion` is defined as a set of `CellBlocks`. A `CellBlock` is an ensemble of elements with the same element geometry.

In the example presented above, the mesh is composed of two regions (*Top* and *Bot*). Each region contains 3 `CellBlocks`.

The `ElementRegions` are defined as below :

```
<ElementRegions>
  <ElementRegion name="Top" cellBlocks="Top_HEX Top_WEDGE Top_TETRA" materialList=
↪ "water rock"/>
  <ElementRegion name="Bot" cellBlocks="Bot_HEX Bot_WEDGE Bot_TETRA" materialList=
↪ "water rock"/>
</ElementRegions>
```

You have to use the following syntax to declare your `CellBlocks` :

```
nameOfTheRegionWithinTheMesh_typeOfTheElement
```

The keywords for the element types are :

- TETRA
- WEDGE
- PYR
- HEX

If the regions are not named in the file (it happens with all the eclipse grids and several GMSH mesh files), the name of the region is `DEFAULT`, e.g:

```
<ElementRegions>
  <ElementRegion name="Default" cellBlocks="DEFAULT_HEX" materialList="water rock"/>
</ElementRegions>
```

Using the `gmsh` file format, regions can be easily named as a preprocessed step using the `gmsh` software or directly editing the file following the syntax defined in the [documentation](#).

An example of a `gmsh` file with all the physical regions defined is used in *Tutorial 3: Regions and Property Specifications*.

### Importing surfaces

Surfaces are imported through point sets in GEOSX. This feature is supported using only the `gmsh` file format. In the same way than the regions, the surfaces of interests can be defined using the [physical entity names](#). The surfaces are automatically imported in GEOSX if they exist in the `gmsh` file. Within GEOSX, the point set will have the same name than the one given in the file. This name can be used again to impose boundary condition. For instance, if a surface is named “Bottom” and the user wants to impose a Dirichlet boundary condition of 0 on it, it can be easily done using this syntax.

```
<FieldSpecification
  name="zconstraint"
  objectPath="nodeManager"
```

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```
fieldName="Velocity"
component="2"
scale="0.0"
setNames="{ Bottom }"/>
```

The name of the surface of interest appears under the keyword `setNames`. Again, an example of a gmsh file with the surfaces fully defined is available within *Tutorial 3: Regions and Property Specifications*.

## 5.3 Physics Solvers

The `<Solvers>` section of the input file specifies one or several physics solvers to be included in the simulation.

### 5.3.1 Solution Strategy

All physics solvers share a common solution strategy for nonlinear time-dependent problems. Here, we briefly describe the nonlinear solver and the timestepping strategy employed.

#### Nonlinear Solver

At each time-step, the nonlinear system of discrete residual equations, i.e.

$$r(x) = 0$$

is solved by employing the Newton-Raphson method. Here,  $x$  is the vector of primary unknowns. Thus, each physics solver is responsible for assembling the Jacobian matrix  $J$  containing the analytical derivatives of the residual vector  $r$  with respect to the primary variables. Then, at each Newton iteration  $\nu$ , the following linear system is solved

$$J^\nu \delta x^{\nu+1} = -r^\nu,$$

where,  $\delta x^{\nu+1}$  is the Newton update. This linear system can be solved with a variety of different linear solvers described in *Linear Solvers*. The Newton update,  $\delta x^{\nu+1}$  is then applied to the primary variables:

$$x^{\nu+1} = x^\nu + \delta x^{\nu+1}.$$

This procedure is repeated until convergence is achieved or until the maximum number of iterations is reached.

#### Line Search

A line search method can be applied along with the Newton's method to facilitate Nonlinear convergence. After the Newton update, if the residual norm has increased instead of decreased, a line search algorithm is employed to correct the Newton update.

The user can choose between two different behaviors in case the line search fails to provide a reduced residual norm:

1. accept the solution and move to the next Newton iteration;
2. reject the solution and request a timestep cut;

## Timestepping Strategy

The actual timestep size employed is determined by a combination of several factors. In particular, specific output events may have timestep requirements that force a specific timestep to be used. However, physics solvers do have the possibility of requesting a specific timestep size to the event manager based on their specific requirements. In particular, in case of fast convergence indicated by a small number of Newton iterations, i.e.

$$\text{numIterations} < \text{dtIncIterLimit} \cdot \text{newtonMaxIter},$$

the physics solver will require to double the timestep size. On the other hand, if a large number of nonlinear iterations are necessary to find the solution at timestep  $n$

$$\text{numIterations} > \text{dtCutIterLimit} \cdot \text{newtonMaxIter},$$

the physics solver will request the next timestep,  $n + 1$ , to be half the size of timestep  $n$ . Here,

Additionally, in case the nonlinear solver fails to converge with the timestep provided by the event manager, the timestep size is cut, i.e.

$$\text{dt} = \text{timestepCutFactor} \cdot \text{dt},$$

and the nonlinear loop is repeated with the new timestep size.

## Parameters

All parameters defining the behavior of the nonlinear solver and determining the timestep size requested by the physics solver are defined in the `NonlinearSolverParameters` and are presented in the following table.

Name	Type	Default	Description
allowNonConverged	integer	0	Allow non-converged solution to be accepted. (i.e. exit from the Newton loop without achieving the desired tolerance)
lineSearchAction	geosx_NonlinearSolverParameters.lineSearchAction	Attempt	How the line search is to be used. Options are: <ul style="list-style-type: none"> <li>* None - Do not use line search.</li> <li>* Attempt - Use line search. Allow exit from line search without achieving smaller residual than starting residual.</li> <li>* Require - Use line search. If smaller residual than starting residual is not achieved, cut time step.</li> </ul>
lineSearchCutFactor	real64	0.5	Line search cut factor. For instance, a value of 0.5 will result in the effective application of the last solution by a factor of (0.5, 0.25, 0.125, ...)
lineSearchInterpolationType	geosx_NonlinearSolverParameters.lineSearchInterpolationType	Linear	Strategy to cut the solution update during the line search. Options are: <ul style="list-style-type: none"> <li>* Linear</li> <li>* Parabolic</li> </ul>
lineSearchMaxCuts	integer	4	Maximum number of line search cuts.
logLevel	integer	0	Log level
maxAllowedResidualNorm	real64	1e+09	Maximum value of residual norm that is allowed in a Newton loop
maxNumConfigurationAttempts	integer	10	Max number of times that the configuration can be changed
maxSubSteps	integer	10	Maximum number of time sub-steps allowed for the solver
maxTimeStepCuts	integer	2	Max number of time step cuts
newtonMaxIter	integer	5	Maximum number of iterations that are allowed in a Newton loop.
newtonMinIter	integer	1	Minimum number of iterations that are required before exiting the Newton loop.
<b>5.3. Physics Solvers</b>			
newtonTol	real64	1e-06	The required tolerance in order to exit the Newton

## 5.3.2 Solid Mechanics Solver

### List of Symbols

$i, j, k$	$\equiv$ indices over spatial dimensions
$a, b, c$	$\equiv$ indices over nodes
$l$	$\equiv$ indices over volumetric elements
$q, r, s$	$\equiv$ indices over faces
$n$	$\equiv$ indices over time
$kiter$	$\equiv$ iteration count for non-linear solution scheme
$\Omega$	$\equiv$ Volume of continuum body
$\Omega_{crack}$	$\equiv$ Volume of open crack
$\Gamma$	$\equiv$ External surface of $\Omega$
$\Gamma_t$	$\equiv$ External surface where tractions are applied
$\Gamma_u$	$\equiv$ External surface where kinematics are specified
$\Gamma_{crack}$	$\equiv$ entire surface of crack
$\Gamma_{cohesive}$	$\equiv$ surface of crack subject to cohesive tractions
$\eta_0$	$\equiv$ set of all nodes
$\eta_f$	$\equiv$ set of all nodes on flow mesh
$m$	$\equiv$ mass
$\kappa_k$	$\equiv$ all elements connected to element $k$
$\phi$	$\equiv$ porosity
$p_f$	$\equiv$ fluid pressure
$\mathbf{u}$	$\equiv$ displacement
$\mathbf{q}$	$\equiv$ volumetric flow rate
$\mathbf{T}$	$\equiv$ Cauchy stress
$\rho$	$\equiv$ density in the current configuration
$\mathbf{x}$	$\equiv$ current position
$\mathbf{w}$	$\equiv$ aperture, or gap vector

### Introduction

The *SolidMechanics\_LagrangianFEM* solver applies a Continuous Galerkin finite element method to solve the linear momentum balance equation. The primary variable is the displacement field which is discretized at the nodes.

### Theory

#### Governing Equations

The *SolidMechanics\_LagrangianFEM* solves the equations of motion as given by

$$T_{ij,j} + \rho(b_i - \ddot{x}_i) = 0,$$

which is a 3-dimensional expression for the well known expression of Newtons Second Law ( $F = ma$ ). These equations of motion are discretized using the Finite Element Method, which leads to a discrete set of residual equations:

$$(R_{solid})_{ai} = \int_{\Gamma_t} \Phi_a t_i dA - \int_{\Omega} \Phi_{a,j} T_{ij} dV + \int_{\Omega} \Phi_a \rho (b_i - \Phi_b \ddot{x}_{ib}) dV = 0$$

### Quasi-Static Time Integration

The Quasi-Static time integration option solves the equation of motion after removing the inertial term, which is expressed by

$$T_{ij,j} + \rho b_i = 0,$$

which is essentially a way to express the equation for static equilibrium ( $\Sigma F = 0$ ). Thus, selection of the Quasi-Static option will yield a solution where the sum of all forces at a given node is equal to zero. The resulting finite element discretized set of residual equations are expressed as

$$(R_{solid})_{ai} = \int_{\Gamma_t} \Phi_a t_i dA - \int_{\Omega} \Phi_{a,j} T_{ij} dV + \int_{\Omega} \Phi_a \rho b_i dV = 0,$$

Taking the derivative of these residual equations wrt. the primary variable (displacement) yields

$$\frac{\partial (R_{solid}^e)_{ai}}{\partial u_{bj}} = - \int_{\Omega^e} \Phi_{a,k} \frac{\partial T_{ik}}{\partial u_{bj}} dV,$$

And finally, the expression for the residual equation and derivative are used to express a non-linear system of equations

$$\left( \frac{\partial (R_{solid}^e)_{ai}}{\partial u_{bj}} \right) \bigg|_{kiter}^{n+1} \left( (u_{bj})_{kiter+1}^{n+1} - (u_{bj})_{kiter}^{n+1} \right) = -(R_{solid})_{ai} \big|_{kiter}^{n+1},$$

which are solved via the solver package.

### Implicit Dynamics Time Integration (Newmark Method)

For implicit dynamic time integration, we use an implementation of the classical Newmark method. This update method can be posed in terms of a simple SDOF spring/dashpot/mass model. In the following,  $M$  represents the mass,  $C$  represent the damping of the dashpot,  $K$  represents the spring stiffness, and  $F$  represents some external load.

$$Ma^{n+1} + Cv^{n+1} + Ku^{n+1} = F_{n+1},$$

and a series of update equations for the velocity and displacement at a point:

$$\begin{aligned} u^{n+1} &= u^n + v^{n+1/2} \Delta t, \\ u^{n+1} &= u^n + \left( v^n + \frac{1}{2} [(1 - 2\beta)a^n + 2\beta a^{n+1}] \Delta t \right) \Delta t, \\ v^{n+1} &= v^n + [(1 - \gamma)a^n + \gamma a^{n+1}] \Delta t. \end{aligned}$$

As intermediate quantities we can form an estimate (predictor) for the end of step displacement and midstep velocity by assuming zero end-of-step acceleration.

$$\begin{aligned} \tilde{u}^{n+1} &= u^n + \left( v^n + \frac{1}{2} (1 - 2\beta) a^n \Delta t \right) \Delta t = u^n + \hat{u} \\ \tilde{v}^{n+1} &= v^n + (1 - \gamma) a^n \Delta t = v^n + \hat{v} \end{aligned}$$

This gives the end of step displacement and velocity in terms of the predictor with a correction for the end step acceleration.

$$\begin{aligned} u^{n+1} &= \tilde{u}^{n+1} + \beta a^{n+1} \Delta t^2 \\ v^{n+1} &= \tilde{v}^{n+1} + \gamma a^{n+1} \Delta t \end{aligned}$$

The acceleration and velocity may now be expressed in terms of displacement, and ultimately in terms of the incremental displacement.

$$\begin{aligned} a^{n+1} &= \frac{1}{\beta \Delta t^2} (u^{n+1} - \tilde{u}^{n+1}) = \frac{1}{\beta \Delta t^2} (\hat{u} - \hat{\tilde{u}}) \\ v^{n+1} &= \tilde{v}^{n+1} + \frac{\gamma}{\beta \Delta t} (u^{n+1} - \tilde{u}^{n+1}) = \tilde{v}^{n+1} + \frac{\gamma}{\beta \Delta t} (\hat{u} - \hat{\tilde{u}}) \end{aligned}$$

plugging these into equation of motion for the SDOF system gives:

$$M \left( \frac{1}{\beta \Delta t^2} (\hat{u} - \hat{\tilde{u}}) \right) + C \left( \tilde{v}^{n+1} + \frac{\gamma}{\beta \Delta t} (\hat{u} - \hat{\tilde{u}}) \right) + K u^{n+1} = F_{n+1}$$

Finally, we assume Rayleigh damping for the dashpot.

$$C = a_{mass} M + a_{stiff} K$$

Of course we know that we intend to model a system of equations with many DOF. Thus the representation for the mass, spring and dashpot can be replaced by our finite element discretized equation of motion. We may express the system in context of a nonlinear residual problem

$$\begin{aligned} (R_{solid}^e)_{ai} &= \int_{\Gamma_i^e} \Phi_a t_i dA \\ &\quad - \int_{\Omega^e} \Phi_{a,j} \left( T_{ij}^{n+1} + a_{stiff} \left( \frac{\partial T_{ij}^{n+1}}{\partial \hat{u}_{bk}} \right)_{elastic} \left( \tilde{v}_{bk}^{n+1} + \frac{\gamma}{\beta \Delta t} (\hat{u}_{bk} - \hat{\tilde{u}}_{bk}) \right) \right) dV \\ &\quad + \int_{\Omega^e} \Phi_a \rho \left( b_i - \Phi_b \left( a_{mass} \left( \tilde{v}_{bi}^{n+1} + \frac{\gamma}{\beta \Delta t} (\hat{u}_{bi} - \hat{\tilde{u}}_{bi}) \right) + \frac{1}{\beta \Delta t^2} (\hat{u}_{bi} - \hat{\tilde{u}}_{bi}) \right) \right) dV, \\ \frac{\partial (R_{solid}^e)_{ai}}{\partial \hat{u}_{bj}} &= - \int_{\Omega^e} \Phi_{a,k} \left( \frac{\partial T_{ik}^{n+1}}{\partial \hat{u}_{bj}} + a_{stiff} \frac{\gamma}{\beta \Delta t} \left( \frac{\partial T_{ik}^{n+1}}{\partial \hat{u}_{bj}} \right)_{elastic} \right) dV \\ &\quad - \left( \frac{\gamma a_{mass}}{\beta \Delta t} + \frac{1}{\beta \Delta t^2} \right) \int_{\Omega^e} \rho \Phi_a \Phi_c \frac{\partial \hat{u}_{ci}}{\partial \hat{u}_{bj}} dV. \end{aligned}$$

Again, the expression for the residual equation and derivative are used to express a non-linear system of equations

$$\left( \frac{\partial (R_{solid}^e)_{ai}}{\partial \hat{u}_{bj}} \right) \Big|_{kiter}^{n+1} \left( (u_{bj})_{kiter+1}^{n+1} - (u_{bj})_{kiter}^{n+1} \right) = -(R_{solid}^e)_{ai} \Big|_{kiter}^{n+1},$$

which are solved via the solver package. Note that the derivatives involving  $u$  and  $\hat{u}$  are interchangeable, as are differences between the non-linear iterations.

### Explicit Dynamics Time Integration (Special Implementation of Newmark Method with gamma=0.5, beta=0)

For the Newmark Method, if gamma=0.5, beta=0, and the inertial term contains a diagonalized “mass matrix”, the update equations may be carried out without the solution of a system of equations. In this case, the update equations simplify to a non-iterative update algorithm.



First the mid-step velocity and end-of-step displacements are calculated through the update equations

$$v^{n+1/2} = v^n + a^n \left( \frac{\Delta t}{2} \right), \text{ and}$$

$$u^{n+1} = u^n + v^{n+1/2} \Delta t.$$

Then the residual equation/s are calculated, and acceleration at the end-of-step is calculated via

$$\left( M + \frac{\Delta t}{2} C \right) a^{n+1} = F_{n+1} - C v^{n+1/2} - K u^{n+1}.$$

Note that the mass matrix must be diagonal, and damping term may not include the stiffness based damping coefficient for this method, otherwise the above equation will require a system solve. Finally, the end-of-step velocities are calculated from the end of step acceleration:

$$v^{n+1} = v^{n+1/2} + a^{n+1} \left( \frac{\Delta t}{2} \right).$$

Note that the velocities may be stored at the midstep, resulting one less kinematic update. This approach is typically referred to as the “Leapfrog” method. However, in GEOSX we do not offer this option since it can cause some confusion that results from the storage of state at different points in time.

## Parameters

In the preceding XML block, The *SolidMechanics\_LagrangianFEM* is specified by the title of the subblock of the *Solvers* block. The following attributes are supported in the input block for *SolidMechanics\_LagrangianFEM*:

Name	Type	Default	Description
cflFactor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
contactRelationName	string	NOCONTACT	Name of contact relation to enforce constraints on fracture boundary.
discretization	string	required	Name of discretization object (defined in the <a href="#">Numerical Methods</a> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <a href="#">Finite Element Discretization</a> should be specified. If this is a Finite Volume Method, the name of a <a href="#">Finite Volume Discretization</a> discretization should be specified.
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
logLevel	integer	0	Log level
massDamping	real64	0	Value of mass based damping coefficient.
maxNumResolves	integer	10	Value to indicate how many resolves may be executed after some other event is executed. For example, if a SurfaceGenerator is specified, it will be executed after the mechanics solve. However if a new surface is generated, then the mechanics solve must be executed again due to the change in topology.
name	string	required	A name is required for any non-unique nodes
newmarkBeta	real64	0.25	Value of $\beta$ in the Newmark Method for Implicit Dynamic time integration option. This should be $\text{pow}(\text{newmarkGamma}+0.5, 2.0)/4.0$ unless you know what you are doing.
newmarkGamma	real64	0.5	Value of $\gamma$ in the Newmark Method for Implicit Dynamic time integration option.
308			<b>Chapter 5. User Guide</b>
stiffnessDamping	real64	0	Value of stiffness based damping coefficient.
strainTheory	integer	0	

The following data are allocated and used by the solver:

Name	Type	Description
maxForce	real64	The maximum force contribution in the problem domain.
maxStableDt	real64	Value of the Maximum Stable Timestep for this solver.
meshTargets	geosx_mapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >	MeshBody/Region combinations that the solver will be applied to.
Linear-SolverParameters	node	<i>Datastructure: Linear-SolverParameters</i>
Nonlinear-SolverParameters	node	<i>Datastructure: Nonlinear-SolverParameters</i>
Solver-Statistics	node	<i>Datastructure: Solver-Statistics</i>

## Example

An example of a valid XML block is given here:

```
<Solvers>
  <SolidMechanics_LagrangianFEM
    name="lagsolve"
    strainTheory="1"
    cflFactor="0.25"
    discretization="FE1"
    targetRegions="{ Region2 }"
  />
</Solvers>
```

## 5.3.3 Singlephase Flow Solver

### Introduction

Here, we describe the single-phase flow solver. The role of this solver is to implement the fully implicit finite-volume discretization (mainly, accumulation and source terms, boundary conditions) of the equations governing compressible single-phase flow in porous media. This solver can be combined with the SinglePhaseWell class which handles the discrete multi-segment well model and provides source/sink terms for the fluid flow solver.

### Theory

### Governing Equations

This is a cell-centered Finite Volume solver for compressible single-phase flow in porous media. Fluid pressure as the primary solution variable. Darcy's law is used to calculate fluid velocity from pressure gradient. The solver currently only supports Dirichlet-type boundary conditions (BC) applied on cells or faces and Neumann no-flow type BC.

The following mass balance equation is solved in the domain:

$$\frac{\partial}{\partial t}(\phi\rho) + \nabla \cdot (\rho\mathbf{u}) + q = 0,$$

where

$$\mathbf{u} = -\frac{1}{\mu}\mathbf{k}(\nabla p - \rho\mathbf{g})$$

and  $\phi$  is porosity,  $\rho$  is fluid density,  $\mu$  is fluid viscosity,  $\mathbf{k}$  is the permeability tensor,  $\mathbf{g}$  is the gravity vector, and  $q$  is the source function (currently not supported). The details on the computation of the density and the viscosity are given in [Compressible single phase fluid model](#).

When the entire pore space is filled by a single phase, we can substitute the Darcy's law into the mass balance equation to obtain the single phase flow equation

$$\frac{\partial}{\partial t}(\phi\rho) - \nabla \cdot \frac{\rho\mathbf{k}}{\mu}(\nabla p - \gamma\nabla z) + q = 0,$$

with  $\gamma\nabla z = \rho\mathbf{g}$ .

## Discretization

### Space Discretization

Let  $\Omega \subset \mathbb{R}^n$ ,  $n = 1, 2, 3$  be an open set defining the computational domain. We consider  $\Omega$  meshed by element such that  $\Omega = \cup_i V_i$  and integrate the single phase flow equation, described above, over each element  $V_i$ :

$$\int_{V_i} \frac{\partial}{\partial t}(\phi\rho)dV - \int_{V_i} \nabla \cdot \frac{\rho\mathbf{k}}{\mu}(\nabla p - \gamma\nabla z)dV + \int_{V_i} qdV = 0.$$

Applying the divergence theorem to the second term leads to

$$\int_{V_i} \frac{\partial}{\partial t}(\phi\rho)_i - \oint_{S_i} \left( \frac{\rho\mathbf{k}}{\mu}(\nabla p - \gamma\nabla z) \right) \cdot \mathbf{n}dS + \int_{V_i} qdV = 0.$$

where  $S_i$  represents the surface area of the element  $V_i$  and  $\mathbf{n}$  is a outward unit vector normal to the surface.

For the flux term, the (static) transmissibility is currently computed with a Two-Point Flux Approximation (TPFA) as described in [Finite Volume Discretization](#).

The pressure-dependent mobility  $\lambda = \frac{\rho}{\mu}$  at the interface is approximated using a first-order upwinding on the sign of the potential difference.

### Time Discretization

Let  $t_0 < t_1 < \dots < t_N = T$  be a grid discretization of the time interval  $[t_0, T]$ ,  $t_0, T \in \mathbb{R}^+$ . We use the backward Euler (fully implicit) method to integrate the single phase flow equation between two grid points  $t_n$  and  $t_{n+1}$ ,  $n < N$  to obtain the residual equation:

$$\int_{V_i} \frac{(\phi\rho)_i^{n+1} - (\phi\rho)_i^n}{\Delta t} - \oint_{S_i} \left( \frac{\rho\mathbf{k}}{\mu}(\nabla p - \gamma\nabla z) \right)^{n+1} \cdot \mathbf{n}dS + \int_{V_i} q^{n+1}dV = 0$$

where  $\Delta t = t_{n+1} - t_n$  is the time-step. The expression of this residual equation and its derivative are used to form a linear system, which is solved via the solver package.

## Parameters

The solver is enabled by adding a <SinglePhaseFVM> node in the Solvers section. Like any solver, time stepping is driven by events, see [Event Management](#).

The following attributes are supported:

Name	Type	Default	Description
cflFactor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
discretization	string	required	Name of discretization object (defined in the <a href="#">Numerical Methods</a> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <a href="#">Finite Element Discretization</a> should be specified. If this is a Finite Volume Method, the name of a <a href="#">Finite Volume Discretization</a> discretization should be specified.
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
input-FluxEstimate	real64	1	Initial estimate of the input flux used only for residual scaling. This should be essentially equivalent to the input flux * dt.
isThermal	integer	0	Flag indicating whether the problem is thermal or not.
logLevel	integer	0	Log level
name	string	required	A name is required for any non-unique nodes
targetRegions	string array	required	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
temperature	real64	0	Temperature
Linear-Solver-Parameters	node	unique	<i>Element: LinearSolverParameters</i>
Non-linear-Solver-Parameters	node	unique	<i>Element: NonlinearSolverParameters</i>

In particular:

- `discretization` must point to a Finite Volume flux approximation scheme defined in the Numerical Methods section of the input file (see [Finite Volume Discretization](#))
- `fluidName` must point to a single phase fluid model defined in the Constitutive section of the input file (see [Constitutive Models](#))
- `solidName` must point to a solid mechanics model defined in the Constitutive section of the input file (see [Constitutive Models](#))
- `targetRegions` is used to specify the regions on which the solver is applied

Primary solution field label is `pressure`. Initial conditions must be prescribed on this field in every region, and boundary conditions must be prescribed on this field on cell or face sets of interest.

### Example

```
<Solvers>
  <SinglePhaseFVM
    name="SinglePhaseFlow"
    logLevel="1"
    discretization="singlePhaseTPFA"
    targetRegions="{ mainRegion }">
    <NonlinearSolverParameters
      newtonTol="1.0e-6"
      newtonMaxIter="8"/>
    <LinearSolverParameters
      solverType="gmres"
      preconditionerType="amg"
      krylovTol="1.0e-10"/>
    </SinglePhaseFVM>
  </Solvers>
```

We refer the reader to [this page](#) for a complete tutorial illustrating the use of this solver.

## 5.3.4 Compositional Multiphase Flow Solver

### Introduction

This flow solver is in charge of implementing the finite-volume discretization (mainly, accumulation and flux terms, boundary conditions) of the equations governing compositional multiphase flow in porous media. The present solver can be combined with the [Compositional Multiphase Well Solver](#) which handles the discrete multi-segment well model and provides source/sink terms for the fluid flow solver.

Below, we first review the set of [Governing Equations](#), followed by a discussion of the choice of [Primary Variables](#) used in the global variable formulation. Then we give an overview of the [Discretization](#) and, finally, we provide a list of the solver [Parameters](#) and an input [Example](#).

### Theory

#### Governing Equations

##### Mass Conservation Equations

Mass conservation for component  $c$  is expressed as:

$$\phi \frac{\partial}{\partial t} \left( \sum_{\ell} \rho_{\ell} y_{c\ell} S_{\ell} \right) + \nabla \cdot \left( \sum_{\ell} \rho_{\ell} y_{c\ell} \mathbf{u}_{\ell} \right) - \sum_{\ell} \rho_{\ell} y_{c\ell} q_{\ell} = 0,$$

where  $\phi$  is the porosity of the medium,  $S_{\ell}$  is the saturation of phase  $\ell$ ,  $y_{c\ell}$  is the mass fraction of component  $c$  in phase  $\ell$ ,  $\rho_{\ell}$  is the phase density, and  $t$  is time. We note that the formulation currently implemented in GEOSX is isothermal.

## Darcy's Law

Using the multiphase extension of Darcy's law, the phase velocity  $\mathbf{u}_\ell$  is written as a function of the phase potential gradient  $\nabla \Phi_\ell$ :

$$\mathbf{u}_\ell := -\mathbf{k} \lambda_\ell \nabla \Phi_\ell = -\mathbf{k} \lambda_\ell (\nabla(p - P_{c,\ell}) - \rho_\ell g \nabla z).$$

In this equation,  $\mathbf{k}$  is the rock permeability,  $\lambda_\ell = k_{r\ell}/\mu_\ell$  is the phase mobility, defined as the phase relative permeability divided by the phase viscosity,  $p$  is the reference pressure,  $P_{c,\ell}$  is the the capillary pressure,  $g$  is the gravitational acceleration, and  $z$  is depth. The evaluation of the relative permeabilities, capillary pressures, and viscosities is reviewed in the section about [Constitutive Models](#).

Combining the mass conservation equations with Darcy's law yields a set of  $n_c$  equations written as:

$$\phi \frac{\partial}{\partial t} \left( \sum_\ell \rho_\ell y_{c\ell} S_\ell \right) - \nabla \cdot \mathbf{k} \left( \sum_\ell \rho_\ell y_{c\ell} \lambda_\ell \nabla \Phi_\ell \right) - \sum_\ell \rho_\ell y_{c\ell} q_\ell = 0.$$

## Constraints and Thermodynamic Equilibrium

The volume constraint equation states that the pore space is always completely filled by the phases. The constraint can be expressed as:

$$\sum_\ell S_\ell = 1.$$

The system is closed by the following thermodynamic equilibrium constraints:

$$f_{c\ell} - f_{cm} = 0.$$

where  $f_{c\ell}$  is the fugacity of component  $c$  in phase  $\ell$ . The flash calculations performed to enforce the thermodynamical equilibrium are reviewed in the section about [Constitutive Models](#).

To summarize, the compositional multiphase flow solver assembles a set of  $n_c + 1$  equations in each element, i.e.,  $n_c$  mass conservation equations and one volume constraint equation. A separate module discussed in the [Constitutive Models](#) is responsible for the enforcement of the thermodynamic equilibrium at each nonlinear iteration.

Number of equations	Equation type
$n_c$	Mass conservation equations
1	Volume constraint

## Primary Variables

The variable formulation implemented in GEOSX is a global variable formulation based on  $n_c + 1$  primary variables, namely, one pressure,  $p$ , and  $n_c$  component densities,  $\rho_c$ . By default, we use molar component densities. A flag discussed in the section [Parameters](#) can be used to select mass component densities instead of molar component densities.

Number of primary variables	Variable type
1	Pressure
$n_c$	Component densities

Assembling the residual equations and calling the *Constitutive Models* requires computing the molar component fractions and saturations. This is done with the relationship:

$$z_c := \frac{\rho_c}{\rho_T},$$

where

$$\rho_T := \sum_c \rho_c.$$

These secondary variables are used as input to the flash calculations. After the flash calculations, the saturations are computed as:

$$S_\ell := \nu_\ell \frac{\rho_T}{\rho_\ell},$$

where  $\nu_\ell$  is the global mole fraction of phase  $\ell$  and  $\rho_\ell$  is the molar density of phase  $\ell$ . These steps also involve computing the derivatives of the component fractions and saturations with respect to the pressure and component densities.

## Discretization

### Spatial Discretization

The governing equations are discretized using standard cell-centered finite-volume discretization.

In the approximation of the flux term at the interface between two control volumes, the calculation of the pressure stencil is general and will ultimately support a Multi-Point Flux Approximation (MPFA) approach. The current implementation of the transmissibility calculation is reviewed in the section about *Numerical Methods*.

The approximation of the dynamic transport coefficients multiplying the discrete potential difference (e.g., the phase mobilities) is performed with a first-order phase-per-phase single-point upwinding based on the sign of the phase potential difference at the interface.

### Temporal Discretization

The compositional multiphase solver uses a fully implicit (backward Euler) temporal discretization.

### Solution Strategy

The nonlinear solution strategy is based on Newton's method. At each Newton iteration, the solver assembles a residual vector,  $R$ , collecting the  $n_c$  discrete mass conservation equations and the volume constraint for all the control volumes.

### Parameters

The following attributes are supported:



Name	Type	Default	Description
allowLocal-CompDen-sityChop-ping	in- te- ger	1	Flag indicating whether local (cell-wise) chopping of negative compositions is allowed
cflFactor	real64	0.5	Factor to apply to the CFL condition when calculating the maximum allowable time step. Values should be in the interval (0,1]
discretiza- tion	string	re- quired	Name of discretization object (defined in the <i>Numerical Methods</i> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <i>Finite Element Discretization</i> should be specified. If this is a Finite Volume Method, the name of a <i>Finite Volume Discretization</i> discretization should be specified.
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
inputFlux- Estimate	real64	1	Initial estimate of the input flux used only for residual scaling. This should be essentially equivalent to the input flux * dt.
isThermal	in- te- ger	0	Flag indicating whether the problem is thermal or not.
logLevel	in- te- ger	0	Log level
max- CompFrac- tionChange	real64	0.5	Maximum (absolute) change in a component fraction in a Newton iteration
maxRela- tivePres- sureChange	real64	0.5	Maximum (relative) change in pressure in a Newton iteration (expected value between 0 and 1)
maxRela- tiveTem- pera- tureChange	real64	0.5	Maximum (relative) change in temperature in a Newton iteration (expected value between 0 and 1)
name	string	re- quired	A name is required for any non-unique nodes
solution- ChangeScal- ingFactor	real64	0.5	Damping factor for solution change targets
target- PhaseVol- Fraction- ChangeIn- TimeStep	real64	0.2	Target (absolute) change in phase volume fraction in a time step
targetRe- gions	string array	re- quired	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
targetRel- ativePres- sureChangeIn- TimeStep	real64	0.2	Target (relative) change in pressure in a time step (expected value between 0 and 1)
targetRel- ativeTem- pera- tureChangeIn- TimeStep	real64	0.2	Target (relative) change in temperature in a time step (expected value between 0 and 1)
temperature	real64	re- quired	Temperature
useMass	in- te- ger	0	Use mass formulation instead of molar

## Example

```
<Solvers>
  <CompositionalMultiphaseFVM
    name="compflow"
    logLevel="1"
    discretization="fluidTPFA"
    targetRelativePressureChangeInTimeStep="1"
    targetPhaseVolFractionChangeInTimeStep="1"
    targetRegions="{ Channel }"
    temperature="300">
    <NonlinearSolverParameters
      newtonTol="1.0e-10"
      newtonMaxIter="40"/>
    <LinearSolverParameters
      directParallel="0"/>
  </CompositionalMultiphaseFVM>
</Solvers>
```

We refer the reader to *Multiphase Flow* for a complete tutorial illustrating the use of this solver.

## 5.3.5 Compositional Multiphase Well Solver

### Introduction

Here, we present a description of the well solvers. These solvers are designed to be coupled with the flow solvers. Their specific task is to implement the multi-segment well discretization using the fluid model used in the corresponding flow solver – i.e., either single-phase flow or compositional multiphase flow. In particular, the perforation terms computed by the well solvers are a source/sink term for the discrete reservoir equations assembled by the flow solvers.

In the present description, we focus on the compositional multiphase well solver. The structure of the single-phase well solver is analogous and will not be described here for brevity.

### Theory

Here, we give an overview of the well formulation implemented in GEOSX. We review the set of *Discrete Equations*, and then we describe the *Primary variables* used the well solvers.

### Discrete Equations

We assume that the well is discretized into segments denoted by the index  $i$ .

### Mass Conservation

In well segment  $i$ , mass conservation for component  $c$  reads:

$$z_{c,(i-1,i)}^{upw} q_{(i-1,i)} - z_{c,(i,i+1)}^{upw} q_{(i,i+1)} + q_{c,i}^{perf} = 0,$$

where we have neglected the accumulation term. In the previous equation,  $z_{c,(i,j)}^{upw}$  is the upwinded mass fraction of component  $c$  at the interface between segments  $i$  and  $j$ ,  $q_{(i,j)}$  is the total mass flux between segments  $i$  and  $j$ , and

$q_{c,i}^{perf}$  is the source/sink term generated by the perforations – i.e., the connections between the well and the reservoir. The upwinded mass fractions are computed as:

$$z_{c,(i,j)}^{upw} = \begin{cases} z_{c,i} & \text{if } q_{(i,j)} > 0 \\ z_{c,j} & \text{otherwise.} \end{cases}$$

The perforation terms are obtained with:

$$q_{c,i}^{perf} = \begin{cases} WI z_{c,i} \rho_{m,i} \lambda_T^{res} \Delta\Phi & \text{if the well is upstream (i.e., } \Delta\Phi > 0) \\ WI x_{c,\ell}^{res} \rho_{\ell}^{res} \lambda_{\ell}^{res} \Delta\Phi & \text{otherwise,} \end{cases}$$

where  $\Delta\Phi = p_i - p^{res} + \rho_{m,i} g \Delta d_{i,perf}$  is the potential difference between the segment center and the reservoir center. In the expression of the potential difference, the mixture density is computed as  $\rho_{m,i} = \sum_{\ell} S_{\ell,i} \rho_{\ell,i}$ . The well index,  $WI$ , is currently an input of the simulator. The superscript *res* means that the variable is evaluated at the center of the reservoir element.

### Volume Constraint Equation

As in the *Compositional Multiphase Flow Solver*, the system is closed with a volume constraint equation.

### Pressure Relations

In the current implementation of the well solver, we assume a hydrostatic equilibrium:

$$p_{i+1} - p_i = \rho_{m,(i,i+1)} g \Delta d_{i,i+1},$$

where  $\rho_{m,(i,i+1)}$  is the arithmetic average of the mixture densities evaluated in segments  $i$  and  $i + 1$ . Pressure drop components due to friction and acceleration are not implemented at the moment.

### Pressure and Rate Controls

The well solver supports two types of control, namely, pressure control and rate control.

If pressure control is chosen, we add the following constraint for the pressure of the top segment of the well:

$$p_0 - p^{target} = 0$$

In this case, we check that at each iteration of the Newton solver, the rate at the top of the first segment is smaller than the maximum rate specified by the user. If this is not the case, we switch to rate control.

If rate control is used, we add the following constraint for the rate at the top of the first segment, denoted by  $q_{(-1,0)}$ :

$$q_{(-1,0)} - q^{target} = 0$$

If the pressure at the top segment becomes larger than the maximum pressure specified by the user, then we switch to pressure control.

To summarize, the compositional multiphase flow solver assembles a set of  $n_c + 2$  equations, i.e.,  $n_c$  mass conservation equations and 1 volume constraint equation in each segment, plus 1 pressure relation at the interface between a segment and the next segment in the direction of the well head. For the top segment, the pressure relation is replaced with the control equation.

Number of equations	Equation type
$n_c$	Mass conservation equations
1	Pressure relation or control equation
1	Volume constraint

### Primary variables

The well variable formulation is the same as that of the *Compositional Multiphase Flow Solver*. In a well segment, in addition to the  $n_c + 1$  primary variables of the *Compositional Multiphase Flow Solver*, namely, one pressure,  $p$ , and  $n_c$  component densities,  $\rho_c$ , we also treat the total mass flux at the interface with the next segment, denoted by  $q$ , as a primary variable.

Number of primary variables	Variable type
1	Pressure
1	Total mass flux at the interface with next segment
$n_c$	Component densities

### Parameters

The following attributes are supported:

Name	Type	Default	Description
allowLocalCompDensityChopping	integer	1	Flag indicating whether local (cell-wise) chopping of negative compositions is allowed
cflFactor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
logLevel	integer	0	Log level
maxCompFractionChange	real64	1	Maximum (absolute) change in a component fraction between two Newton iterations
maxRelativePressureChange	real64	1	Maximum (relative) change in pressure between two Newton iterations (recommended with rate control)
name	string	required	A name is required for any non-unique nodes
targetRegions	string array	required	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
useMass	integer	0	Use mass formulation instead of molar
LinearSolverParameters	node	unique	<i>Element: LinearSolverParameters</i>
NonlinearSolverParameters	node	unique	<i>Element: NonlinearSolverParameters</i>
WellControls	node		<i>Element: WellControls</i>

### Example

```

<CompositionalMultiphaseWell
  name="compositionalMultiphaseWell"
  logLevel="1"
  targetRegions="{ wellRegion1, wellRegion2, wellRegion3 }">
  <WellControls
    name="wellControls1"
    type="producer"
    control="BHP"
    referenceElevation="0.5"
    targetBHP="4e6"
    targetPhaseRate="1e-3"
    targetPhaseName="oil"/>

```

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```

<WellControls
  name="wellControls2"
  type="producer"
  control="phaseVolRate"
  referenceElevation="0.5"
  targetBHP="2e6"
  targetPhaseRate="2.5e-7"
  targetPhaseName="oil"/>
<WellControls
  name="wellControls3"
  type="injector"
  control="totalVolRate"
  referenceElevation="0.5"
  targetBHP="4e7"
  targetTotalRate="5e-7"
  injectionTemperature="297.15"
  injectionStream="{ 0.1, 0.1, 0.1, 0.7 }"/>
</CompositionalMultiphaseWell>

```

### 5.3.6 Poromechanics Solver

#### Introduction

This section describes the use of the poroelasticity models implemented in GEOSX.

#### Theory

##### Governing Equations

In our model, the geomechanics (elasticity) equation is expressed in terms of the total stress  $\sigma$ :

$$\nabla \sigma + \rho_b \mathbf{g} = 0$$

where it relates to effective stress  $\sigma'$  and pore pressure  $p$  through Biot's coefficient  $b$ :

$$\sigma = \sigma' - bp\mathbf{I}$$

The fluid mass conservation equation is expressed in terms of pore pressure and volumetric (mean) total stress:

$$\left( \frac{1}{M} + \frac{b^2}{K_{dr}} \right) \frac{\partial p}{\partial t} + \frac{b}{K_{dr}} \frac{\partial \sigma_v}{\partial t} + \nabla \cdot \mathbf{v}_f = f$$

where  $M$  is the Biot's modulus and  $K_{dr}$  is the drained bulk modulus.

Unlike the conventional reservoir model that uses Lagrange's porosity, in the coupled geomechanics and flow model, Euler's porosity  $\phi$  is adopted so the porosity variation is derived as:

$$\partial \phi = \left( \frac{b - \phi}{K_s} \right) \partial p + (b - \phi) \partial \epsilon_v$$

where  $K_s$  is the bulk modulus of the solid grain and  $\epsilon_v$  is the volumetric strain.

## Parameters

The poroelasticity model is implemented as a main solver listed in `<Solvers>` block of the input XML file that calls both `SolidMechanicsLagrangianSSLE` and `SinglePhaseFlow` solvers. In the main solver, it requires the specification of `solidSolverName`, `fluidSolverName`, and `couplingTypeOption`.

The following attributes are supported:

Name	Type	Default	Description
cflFactor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
flow-Solver-Name	string	required	Name of the flow solver used by the coupled solver
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
logLevel	integer	0	Log level
name	string	required	A name is required for any non-unique nodes
solid-Solver-Name	string	required	Name of the solid solver used by the coupled solver
targetRegions	string array	required	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
Linear-Solver-Parameters	node	unique	<i>Element: LinearSolverParameters</i>
Non-linear-Solver-Parameters	node	unique	<i>Element: NonlinearSolverParameters</i>

- `couplingTypeOption`: defines the coupling scheme.

The solid constitutive model used here is `PoroLinearElasticIsotropic`, which derives from `ElasticIsotropic` and includes an additional parameter: Biot's coefficient. The fluid constitutive model is the same as `SinglePhaseFlow` solver. For the parameter setup of each individual solver, please refer to the guideline of the specific solver.

An example of a valid XML block for the constitutive model is given here:

```
<Constitutive>
  <PorousElasticIsotropic
    name="porousRock"
    solidModelName="skeleton"
    porosityModelName="skeletonPorosity"
    permeabilityModelName="skeletonPerm"/>

  <ElasticIsotropic
```

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```

    name="skeleton"
    defaultDensity="0"
    defaultYoungModulus="1.0e4"
    defaultPoissonRatio="0.2"/>

    <CompressibleSinglePhaseFluid
      name="fluid"
      defaultDensity="1"
      defaultViscosity="1.0"
      referencePressure="0.0"
      referenceDensity="1"
      compressibility="0.0e0"
      referenceViscosity="1"
      viscosibility="0.0"/>

    <BiotPorosity
      name="skeletonPorosity"
      grainBulkModulus="1.0e27"
      defaultReferencePorosity="0.3"/>

    <ConstantPermeability
      name="skeletonPerm"
      permeabilityComponents="{ 1.0e-4, 1.0e-4, 1.0e-4 }"/>
  </Constitutive>

```

## Example

```

  <SinglePhasePoromechanics
    name="PoroelectricitySolver"
    solidSolverName="LinearElasticitySolver"
    flowSolverName="SinglePhaseFlowSolver"
    logLevel="1"
    targetRegions="{ Domain }">
    <LinearSolverParameters
      solverType="gmres"
      preconditionerType="mgr"/>
  </SinglePhasePoromechanics>

  <SolidMechanicsLagrangianSSLE
    name="LinearElasticitySolver"
    timeIntegrationOption="QuasiStatic"
    logLevel="1"
    discretization="FE1"
    targetRegions="{ Domain }"/>

  <SinglePhaseFVM
    name="SinglePhaseFlowSolver"
    logLevel="1"
    discretization="singlePhaseTPFA"
    targetRegions="{ Domain }"/>
</Solvers>

```

## 5.3.7 Proppant Transport Solver



## Introduction

The *ProppantTransport* solver applies the finite volume method to solve the equations of proppant transport in hydraulic fractures. The behavior of proppant transport is described by a continuum formulation. Here we briefly outline the usage, governing equations and numerical implementation of the proppant transport model in GEOSX.

## Theory

The following mass balance and constitutive equations are solved inside fractures,

### Proppant-fluid Slurry Flow

$$\frac{\partial}{\partial t}(\rho_m) + \nabla \cdot (\rho_m \mathbf{u}_m) = 0,$$

where the proppant-fluid mixture velocity  $\mathbf{u}_m$  is approximated by the Darcy's law as,

$$\mathbf{u}_m = -\frac{K_f}{\mu_m}(\nabla p - \rho_m \mathbf{g}),$$

and  $p$  is pressure,  $\rho_m$  and  $\mu_m$  are density and viscosity of the mixed fluid, respectively, and  $\mathbf{g}$  is the gravity vector. The fracture permeability  $K_f$  is determined based on fracture aperture  $a$  as

$$K_f = \frac{a^2}{12}$$

### Proppant Transport

$$\frac{\partial}{\partial t}(c) + \nabla \cdot (c \mathbf{u}_p) = 0,$$

in which  $c$  and  $\mathbf{u}_p$  represent the volume fraction and velocity of the proppant particles.

### Multi-component Fluid Transport

$$\frac{\partial}{\partial t}[\rho_i \omega_i (1 - c)] + \nabla \cdot [\rho_i \omega_i (1 - c) \mathbf{u}_f] = 0.$$

Here  $\mathbf{u}_f$  represents the carrying fluid velocity.  $\rho_i$  and  $\omega_i$  denote the density and concentration of  $i$ -th component in fluid, respectively. The fluid density  $\rho_f$  can now be readily written as

$$\rho_f = \sum_{i=1}^{N_c} \rho_i \omega_i,$$

where  $N_c$  is the number of components in fluid. Similarly, the fluid viscosity  $\mu_f$  can be calculated by the mass fraction weighted average of the component viscosities.

The density and velocity of the slurry fluid are further expressed as,

$$\rho_m = (1 - c)\rho_f + c\rho_p,$$

and

$$\rho_m \mathbf{u}_m = (1 - c) \rho_f \mathbf{u}_f + c \rho_p \mathbf{u}_p,$$

in which  $\rho_f$  and  $\mathbf{u}_f$  are the density and velocity of the carrying fluid, and  $\rho_p$  is the density of the proppant particles.

### Proppant Slip Velocity

The proppant particle and carrying fluid velocities are related by the slip velocity  $\mathbf{u}_{slip}$ ,

$$\mathbf{u}_{slip} = \mathbf{u}_p - \mathbf{u}_f.$$

The slip velocity between the proppant and carrying fluid includes gravitational and collisional components, which take account of particle settling and collision effects, respectively.

The gravitational component of the slip velocity  $\mathbf{u}_{slipG}$  is written as a form as

$$\mathbf{u}_{slipG} = F(c) \mathbf{u}_{settling},$$

where  $\mathbf{u}_{settling}$  is the settling velocity for a single particle,  $d_p$  is the particle diameter, and  $F(c)$  is the correction factor to the particle settling velocity in order to account for hindered settling effects as a result of particle-particle interactions,

$$F(c) = e^{-\lambda_s c},$$

with the hindered settling coefficient  $\lambda_s$  as an empirical constant set to 5.9 by default (Barree & Conway, 1995).

The settling velocity for a single particle,  $\mathbf{u}_{settling}$ , is calculated based on the Stokes drag law by default,

$$\mathbf{u}_{settling} = (\rho_p - \rho_f) \frac{d_p^2}{18\mu_f} \mathbf{g}.$$

Single-particle settling under intermediate Reynolds-number and turbulent flow conditions can also be described respectively by the Allen's equation (Barree & Conway, 1995),

$$\mathbf{u}_{settling} = 0.2 d_p^{1.18} \left[ \frac{g(\rho_p - \rho_f)}{\rho_f} \right]^{0.72} \left( \frac{\rho_f}{\mu_f} \right)^{0.45} \mathbf{e},$$

and Newton's equation (Barree & Conway, 1995),

$$\mathbf{u}_{settling} = 1.74 d_p^{0.5} \left[ \frac{g(\rho_p - \rho_f)}{\rho_f} \right]^{0.5} \mathbf{e}.$$

$\mathbf{e}$  is the unit gravity vector and  $d_p$  is the particle diameter.

The collisional component of the slip velocity is modeled by defining  $\lambda$ , the ratio of the particle velocity to the volume averaged mixture velocity as a function of the proppant concentration. From this the particle slip velocity in horizontal direction is related to the mixed fluid velocity by,

$$\mathbf{u}_{slipH} = \frac{\lambda - 1}{1 - c} \mathbf{v}_m$$

with  $\mathbf{v}_m$  denoting volume averaged mixture velocity. We use a simple expression of  $\lambda$  proposed by Barree & Conway (1995) to correct the particle slip velocity in horizontal direction,

$$\lambda = [\alpha - |c - c_{slip}|^\beta]$$

where  $\alpha$  and  $\beta$  are empirical constants,  $c_{slip}$  is the volume fraction exhibiting the greatest particle slip. By default the model parameters are set to the values given in (Barree & Conway, 1995):  $\alpha = 1.27$ ,  $c_{slip} = 0.1$  and  $\beta = 1.5$ . This model can be extended to account for the transition to the particle pack as the proppant concentration approaches the jamming transition.

## Proppant Bed Build-up and Load Transport

In addition to suspended particle flow the GEOSX has the option to model proppant settling into an immobile bed at the bottom of the fracture. As the proppant cannot settle further down the proppant bed starts to form and develop at the element that is either at the bottom of the fracture or has an underlying element already filled with particles. Such an “inter-facial” element is divided into proppant flow and immobile bed regions based on the proppant-pack height.

Although proppant becomes immobile fluid can continue to flow through the settled proppant pack. The pack permeability  $K$  is defined based on the Kozeny-Carmen relationship:

$$K = \frac{(sd_p)^2}{180} \frac{\phi^3}{(1 - \phi)^2}$$

and

$$\phi = 1 - c_s$$

where  $\phi$  is the porosity of particle pack and  $c_s$  is the saturation or maximum fraction for proppant packing,  $s$  is the sphericity and  $d_p$  is the particle diameter.

The growth of the settled pack in an “inter-facial” element is controlled by the interplay between proppant gravitational settling and shear-force induced lifting as (Hu et al., 2018),

$$\frac{dH}{dt} = \frac{cu_{\text{settling}}F(c)}{c_s} - \frac{Q_{\text{lift}}}{Ac_s},$$

where  $H$ ,  $t$ ,  $c_s$ ,  $Q_{\text{lift}}$ , and  $A$  represent the height of the proppant bed, time, saturation or maximum proppant concentration in the proppant bed, proppant-bed load (wash-out) flux, and cross-sectional area, respectively.

The rate of proppant bed load transport (or wash out) due to shear force is calculated by the correlation proposed by Wiberg and Smith (1989) and McClure (2018),

$$Q_{\text{lift}} = a \left( d_p \sqrt{\frac{gd_p(\rho_p - \rho_f)}{\rho_f}} \right) (9.64N_{sh}^{0.166})(N_{sh} - N_{sh,c})^{1.5}.$$

$a$  is fracture aperture, and  $N_{sh}$  is the Shields number measuring the relative importance of the shear force to the gravitational force on a particle of sediment (Miller et al., 1977; Biot & Medlin, 1985; McClure, 2018) as

$$N_{sh} = \frac{\tau}{d_p g(\rho_p - \rho_f)},$$

and

$$\tau = 0.125f\rho_f u_m^2$$

where  $\tau$  is the shear stress acting on the top of the proppant bed and  $f$  is the Darcy friction coefficient.  $N_{sh,c}$  is the critical Shields number for the onset of bed load transport.

## Proppant Bridging and Screenout

Proppant bridging occurs when proppant particle size is close to or larger than fracture aperture. The aperture at which bridging occurs,  $h_b$ , is defined simply by

$$h_b = \lambda_b d_p,$$

in which  $\lambda_b$  is the bridging factor.

## Slurry Fluid Viscosity

The viscosity of the bulk fluid,  $\mu_m$ , is calculated as a function of proppant concentration as (Keck et al., 1992),

$$\mu_m = \mu_f \left[ 1 + 1.25 \left( \frac{c}{1 - c/c_s} \right) \right]^2.$$

Note that continued model development and improvement are underway and additional empirical correlations or functions will be added to support the above calculations.

## Spatial Discretization

The above governing equations are discretized using a cell-centered two-point flux approximation (TPFA) finite volume method. We use an upwind scheme to approximate proppant and component transport across cell interfaces.

## Solution Strategy

The discretized non-linear slurry flow and proppant/component transport equations at each time step are separately solved by the Newton-Raphson method. The coupling between them is achieved by a time-marching sequential (operator-splitting) solution approach.

## Parameters

The solver is enabled by adding a `<ProppantTransport>` node and a `<SurfaceGenerator>` node in the Solvers section. Like any solver, time stepping is driven by events, see [Event Management](#).

The following attributes are supported:

Name	Type	De- fault	Description
bridg- ingFac- tor	real64	0	Bridging factor used for bridging/screen-out calculation
cflFac- tor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
critical- Shield- sNum- ber	real64	0	Critical Shields number
dis- cretiza- tion	string	re- quired	Name of discretization object (defined in the <a href="#">Numerical Methods</a> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <a href="#">Finite Element Discretiza- tion</a> should be specified. If this is a Finite Volume Method, the name of a <a href="#">Finite Volume Discretization</a> discretization should be specified.
fric- tion- Coeffi- cient	real64	0.03	Friction coefficient
ini- tialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
input- FluxEs- timate	real64	1	Initial estimate of the input flux used only for residual scaling. This should be essentially equivalent to the input flux * dt.
isTher- mal	in- te- ger	0	Flag indicating whether the problem is thermal or not.
logLevel	in- te- ger	0	Log level
max- Prop- ant- Con- centra- tion	real64	0.6	Maximum proppant concentration
name	string	re- quired	A name is required for any non-unique nodes
prop- ant- Density	real64	2500	Proppant density
prop- antDi- ameter	real64	0.0004	Proppant diameter
targe- tRe- gions	string_array	re- quired	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will beapplied to rests in the EventManager.
up- dateProp- ant- Pack- ing	in- te- ger	0	Flag that enables/disables proppant-packing update
Linear- Solver- Param- eters	node	unique	<i>Element: LinearSolverParameters</i>
Non- linear-	node	unique	<i>Element: NonlinearSolverParameters</i>

In particular:

- `discretization` must point to a Finite Volume flux approximation scheme defined in the Numerical Methods section of the input file (see *Finite Volume Discretization*)
- `proppantName` must point to a particle fluid model defined in the Constitutive section of the input file (see *Constitutive Models*)
- `fluidName` must point to a slurry fluid model defined in the Constitutive section of the input file (see *Constitutive Models*)
- `solidName` must point to a solid mechanics model defined in the Constitutive section of the input file (see *Constitutive Models*)
- `targetRegions` attribute is currently not supported, the solver is always applied to all regions.

Primary solution field labels are `proppantConcentration` and `pressure`. Initial conditions must be prescribed on these field in every region, and boundary conditions must be prescribed on these fields on cell or face sets of interest. For static (non-propagating) fracture problems, the fields `ruptureState` and `elementAperture` should be provided in the initial conditions.

In addition, the solver declares a scalar field named `referencePorosity` and a vector field named `permeability`, that contains principal values of the symmetric rank-2 permeability tensor (tensor axis are assumed aligned with the global coordinate system). These fields must be populated via *Element: FieldSpecification* section and `permeability` should be supplied as the value of `coefficientName` attribute of the flux approximation scheme used.

## Example

First, we specify the proppant transport solver itself and apply it to the fracture region:

```
<ProppantTransport
  name="ProppantTransport"
  logLevel="1"
  discretization="singlePhaseTPFA"
  targetRegions="{ Fracture }">
  <NonlinearSolverParameters
    newtonTol="1.0e-8"
    newtonMaxIter="8"
    lineSearchAction="None"/>
  <LinearSolverParameters
    directParallel="0"/>
</ProppantTransport>
```

Then, we specify a compatible flow solver (currently a specialized `SinglePhaseProppantFVM` solver must be used):

```
<SinglePhaseProppantFVM
  name="SinglePhaseFVM"
  logLevel="1"
  discretization="singlePhaseTPFA"
  targetRegions="{ Fracture }">
  <NonlinearSolverParameters
    newtonTol="1.0e-8"
    newtonMaxIter="8"
    lineSearchAction="None"/>
  <LinearSolverParameters
    solverType="gmres"
```

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```
krylovTol="1.0e-12"/>
</SinglePhaseProppantFVM>
```

Finally, we couple them through a coupled solver that references the two above:

```
<FlowProppantTransport
  name="FlowProppantTransport"
  proppantSolverName="ProppantTransport"
  flowSolverName="SinglePhaseFVM"
  targetRegions="{ Fracture }"
  logLevel="1"/>
```

## References

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- M. C. Miller, I. N. McCave, & P. D. Komar. “Threshold of sediment motion under unidirectional currents”, Sedimentology 24 (4), 507527, 1977.
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## 5.4 Constitutive Models

Constitutive models describe relations between various physical quantities. In a physics simulation they are used to model the response or state of material (solid, fluid, or a mixture) as a function of input variables. There are many types of constitutive models available in GEOSX. These models are grouped together based on their input/output interface.

### 5.4.1 Solid Models

A solid model governs the relationship between deformation and stress in a solid (or porous) material. GEOSX provides interfaces for both small and large deformation models, as well as specific implementations of a number of well known models.

### Deformation Theories

## Table of Contents

- *Deformation Theories*
  - *Introduction*
  - *Small Strain Models*
  - *Finite Deformation Models with Hypo-Materials*
  - *Finite Deformation Models with Hyper-Materials*

## Introduction

The solid mechanics solvers in GEOSX work in a time-discrete setting, in which the system state at time  $t^n$  is fully known, and the goal of the solution procedure is to advance forward one timestep to  $t^{n+1} = t^n + \Delta t$ . As part of this process, calls to a solid model must be made to compute the updated stress  $\sigma^{n+1}$  resulting from incremental deformation over the timestep. History-dependent models may also need to compute updates to one or more internal state variables  $Q^{n+1}$ .

The exact nature of the incremental update will depend, however, on the kinematic assumptions made. Appropriate measures of deformation and stress depend on assumptions of *infinitesimal* or *finite* strain, as well as other factors like rate-dependence and material anisotropy.

This section briefly reviews three main classes of solid models in GEOSX, grouped by their kinematic assumptions. The presentation is deliberately brief, as much more extensive presentations can be found in almost any textbook on linear and nonlinear solid mechanics.

## Small Strain Models

Let  $u$  denote the displacement field, and  $\nabla u$  its gradient. In small strain theory, one assumes the displacement gradients  $\nabla u \ll 1$ . In this case, it is sufficient to use the linearized strain tensor

$$\epsilon = \frac{1}{2} (\nabla u + u \nabla)$$

as the deformation measure. Higher-order terms present in finite strain theories are neglected. For inelastic problems, this strain is additively decomposed into elastic and inelastic components as

$$\epsilon = \epsilon^e + \epsilon^i.$$

Inelastic strains can arise from a number of sources: plasticity, damage, etc. Most constitutive models (including nonlinear elastic and inelastic models) can then be generically expressed in rate form as

$$\dot{\sigma} = c : \dot{\epsilon}^e$$

where  $\dot{\sigma}$  is the Cauchy stress rate and  $c$  is the tangent stiffness tensor. Observe that the stress rate is driven by the elastic component  $\dot{\epsilon}^e$  of the strain rate.

In the time-discrete setting (as implemented in the code) the incremental constitutive update for stress is computed from a solid model update routine as

$$\sigma^{n+1} = \sigma(\Delta\epsilon, \Delta t, Q^n),$$

where  $\Delta\epsilon = \epsilon^{n+1} - \epsilon^n$  is the incremental strain,  $\Delta t$  is the timestep size (important for rate-dependent models), and  $Q^n$  is a collection of material state variables (which may include the previous stress and strain).



For path and rate independent models, such as linear elasticity, a simpler constitutive update may be formulated in terms of the total strain:

$$\sigma^{n+1} = \sigma(\epsilon^{n+1}).$$

GEOSX will use this latter form in specific, highly-optimized solvers when we know in advance that a linear elastic model is being applied. The more general interface is the the default, however, as it can accommodate a much wider range of constitutive behavior within a common interface.

When implicit timestepping is used, the solid models must also provide the stiffness tensor,

$$c^{n+1} = \frac{\partial \sigma^{n+1}}{\partial \epsilon^{n+1}},$$

in order to accurately linearize the governing equations. In many works, this fourth-order tensor is referred to as the algorithmic or consistent tangent, in the sense that it must be “consistent” with the discrete timestepping scheme being used (Simo and Hughes 1987). For inelastic models, it depends not only on the intrinsic material stiffness, but also the incremental nature of the loading process. The correct calculation of this stiffness can have a dramatic impact on the convergence rate of Newton-type solvers used in the implicit solid mechanics solvers.

## Finite Deformation Models with Hypo-Materials

In the finite deformation regime, there are two broad classes of constitutive models frequently used:

- Hypo-elastic models (and inelastic extensions)
- Hyper-elastic models (and inelastic extensions)

Hypo-materials typically rely on a rate-form of the constitutive equations expressed in the spatial configuration. Let  $v(x, t)$  denote the spatial velocity field. It can be decomposed into symmetric and anti-symmetric components as

$$d = \frac{1}{2} (\nabla v + v \nabla) \quad \text{and} \quad w = \frac{1}{2} (\nabla v - v \nabla),$$

where  $d$  is the rate of deformation tensor and  $w$  is the spin tensor. A hypo-material model can be written in rate form as

$$\dot{\tau} = c : d^e$$

where  $\dot{\tau}$  is an **objective rate** of the Kirchhoff stress tensor,  $c$  is the tangent stiffness tensor, and  $d^e$  is the elastic component of the deformation rate. We see that the structure is similar to the rate form in the small strain regime, except the rate of Cauchy stress is replaced with an objective rate of Kirchhoff stress, and the linearized strain rate is replaced with the rate of deformation tensor.

The key difference separating most hypo-models is the choice of the objective stress rate. In GEOSX, we adopt the incrementally objective integration algorithm proposed by Hughes & Winget (1980). This method relies on the concept of an incrementally rotating frame of reference in order to preserve objectivity of the stress rate. In particular, the stress update sequence is

$$\begin{aligned} \Delta R &= (I - \frac{1}{2} \Delta t w)^{-1} (I + \frac{1}{2} \Delta t w) && \text{(compute incremental rotation),} \\ \bar{\tau}^n &= \Delta R \tau^n \Delta R^T && \text{(rotate previous stress),} \\ \tau^{n+1} &= \bar{\tau}^n + \Delta \tau && \text{(call constitutive model to update stress).} \end{aligned}$$

First, the previous timestep stress is rotated to reflect any rigid rotations occurring over the timestep. If the model has tensor-valued state variables besides stress, these must also be rotated. Then, a standard constitutive update routine can be called, typically driven by the incremental strain  $\Delta \epsilon = \Delta t d$ . In fact, an identical update routine as used for small strain models can be re-used at this point.

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**Note:** Hypo-models suffer from several well known deficiencies. Most notably, the energy dissipation in a closed loading cycle of a hypo-elastic material is not guaranteed to be zero, as one might desire from thermodynamic considerations.

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## Finite Deformation Models with Hyper-Materials

Hyper-elastic models (and inelastic extensions) attempt to correct the thermodynamic deficiencies of their hypo-elastic cousins. The constitutive update can be generically expressed at

$$S^{n+1} = S(\Delta \mathbf{F}, Q^n, \Delta t),$$

where  $S$  is the second Piola-Kirchhoff stress and  $\Delta \mathbf{F}$  is the incremental deformation gradient. Depending on the model, the deformation gradient can be converted to different deformation measures as needed. Similarly, different stress tensors can be recovered through appropriate push-forward and pull-back operations.

In a hyperelastic material, the elastic response is expressed in terms of a stored strain-energy function that serves as the potential for stress, e.g.

$$\mathbf{S} = \frac{\partial \psi(C)}{\partial C},$$

where  $\psi$  is the stored energy potential, and  $C$  is the right Cauchy-Green deformation tensor. This potential guarantees that the energy dissipated or gained in a closed elastic cycle is zero.

## Voigt Notation

In GEOSX we express rank-two symmetric tensors using [Voigt notation](#). Stress tensors are represented as an “un-rolled” six-component vector storing only the unique component values. For strain tensors, note that *engineering strains* are used such that the shear components of strain are multiplied by a factor of two. With this representation, the strain energy density is, conveniently, the inner product of the stress and strain vectors.

Voigt representation of common rank-2 symmetric tensors are:

$$\sigma = \begin{bmatrix} \sigma_{11} \\ \sigma_{22} \\ \sigma_{33} \\ \sigma_{23} \\ \sigma_{13} \\ \sigma_{12} \end{bmatrix}, \mathbf{S} = \begin{bmatrix} S_{11} \\ S_{22} \\ S_{33} \\ S_{23} \\ S_{13} \\ S_{12} \end{bmatrix}, \epsilon = \begin{bmatrix} \epsilon_{11} \\ \epsilon_{22} \\ \epsilon_{33} \\ 2\epsilon_{23} \\ 2\epsilon_{13} \\ 2\epsilon_{12} \end{bmatrix}, \mathbf{D} = \begin{bmatrix} D_{11} \\ D_{22} \\ D_{33} \\ 2D_{23} \\ 2D_{13} \\ 2D_{12} \end{bmatrix}, \mathbf{E} = \begin{bmatrix} E_{11} \\ E_{22} \\ E_{33} \\ 2E_{23} \\ 2E_{13} \\ 2E_{12} \end{bmatrix}, \mathbf{B} = \begin{bmatrix} B_{11} \\ B_{22} \\ B_{33} \\ 2B_{23} \\ 2B_{13} \\ 2B_{12} \end{bmatrix}, \mathbf{C} = \begin{bmatrix} C_{11} \\ C_{22} \\ C_{33} \\ 2C_{23} \\ 2C_{13} \\ 2C_{12} \end{bmatrix},$$

where  $\sigma$  is the Cauchy stress,  $\mathbf{S}$  is the second Piola-Kirchhoff stress,  $\epsilon$  is the small strain tensor,  $\mathbf{D}$  is the rate of deformation tensor,  $\mathbf{E}$  is the Lagrangian finite strain tensor,  $\mathbf{B}$  is the left Cauchy-Green tensor,  $\mathbf{C}$  is the right Cauchy-Green deformation tensor.

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**Note:** The factor of two in the shear components of strain (and strain-like) quantities is a frequent source of confusion, even for expert modelers. It can be particularly challenging to use in nuanced situations like stiffness tensor calculations or invariant decompositions. If you plan to implement new models within GEOSX, please pay extra attention to this detail. We also provide many common operations in centralized helper functions to avoid re-inventing the wheel.

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## Plasticity Notation

### Table of Contents

- *Plasticity Notation*
  - *Overview*
  - *Two-Invariant Models*
  - *Three-Invariant Models*

### Overview

According to the [theory of plasticity](#) in the small strain regime, the total strain  $\epsilon$  can be additively split into elastic ( $\epsilon^e$ ) and plastic ( $\epsilon^p$ ) strains:

$$\epsilon = \epsilon^e + \epsilon^p.$$

The plastic strain tensor is obtained from the flow rule:

$$\dot{\epsilon}^p = \dot{\lambda} \frac{\partial g}{\partial \sigma},$$

in which  $\dot{\lambda} \geq 0$  is the magnitude of plastic strain rate and  $g$  is the plastic potential. The elastic strain is related to Cauchy stress tensor in rate form as:

$$\dot{\sigma} = c^e : \dot{\epsilon}^e,$$

where  $c^e$  is the fourth order elastic stiffness tensor. The Cauchy stress tensor is related to the total strain as

$$\dot{\sigma} = c^{ep} : \dot{\epsilon},$$

in which  $c^{ep}$  is the fourth order elasto-plastic stiffness tensor.

### Two-Invariant Models

Two-invariant plasticity models use the first invariant of the Cauchy stress tensor and the second invariant of the deviatoric stress tensor to describe the yield surface.

Here we use the following stress invariants to define the yield surface: the von Mises stress  $q = \sqrt{3J_2} = \sqrt{3/2}\|s\|$  and mean normal stress  $p = I_1/3$ . Here,  $I_1$  and  $J_2$  are the first invariant of the stress tensor and second invariant of the deviatoric stress, defined as

$$I_1 = \text{tr}(\sigma)/3, \quad J_2 = \frac{1}{2}\|s\|^2, \quad s = \sigma - p\mathbf{1},$$

in which  $\mathbf{1}$  is the identity tensor.

Similarly, we can define invariants of strain tensor, namely, volumetric strain  $\epsilon_v$  and deviatoric strain  $\epsilon_s$ .

$$\epsilon_v = \text{tr}(\epsilon), \quad \epsilon_s = \sqrt{\frac{2}{3}}\|e\|, \quad \text{where} \quad e = \epsilon - \frac{1}{3}\epsilon_v\mathbf{1}.$$

Stress and strain tensors can then be recomposed from the invariants as:

$$\sigma = p\mathbf{1} + \sqrt{\frac{2}{3}}q\hat{n}$$

$$\boldsymbol{\epsilon} = \frac{1}{3}\epsilon_v \mathbf{1} + \sqrt{\frac{3}{2}}\epsilon_s \hat{\mathbf{n}}$$

in which  $\hat{\mathbf{n}} = \mathbf{e}/\|\mathbf{e}\|$ .

The following two-invariant models are currently implemented in GEOSX:

- *DruckerPrager*
- *J2Plasticity*
- *ModifiedCamClay*
- *DelftEgg*

## Three-Invariant Models

Several three-invariant models are under active development, but are not yet available in develop. If you are interested in helping to add additional material models, please submit a feature request.

## Triaxial Driver

### Table of Contents

- *Triaxial Driver*
  - *Introduction*
  - *XML Structure*
  - *Test Modes*
  - *Output Format*
  - *Model Convergence*
  - *Unit Testing*

## Introduction

When calibrating solid material parameters to experimental data, it can be a hassle to launch a full finite element simulation to mimic experimental loading conditions. Instead, GEOSX provides a `TriaxialDriver` allowing the user to run loading tests on a single material point. This makes it easy to understand the material response and fit it to lab data. The driver itself is launched like any other GEOSX simulation, but with a particular XML structure:

```
./bin/geosx -i myTest.xml
```

## XML Structure

A typical XML file to run the triaxial driver will have the following key elements. We present the whole file first, before digging into the individual blocks.

```

<?xml version="1.0" ?>

<Problem>
  <Tasks>
    <TriaxialDriver
      name="triaxialDriver"
      material="drucker"
      mode="mixedControl"
      axialControl="strainFunction"
      radialControl="stressFunction"
      initialStress="-1.0"
      steps="40"
      output="none"
      baseline="testTriaxial_druckerPragerExtended.txt"
      logLevel="0"/>
    </Tasks>

    <Events
      maxTime="1">
        <SoloEvent
          name="triaxialDriver"
          target="/Tasks/triaxialDriver"/>
        </Events>

    <Constitutive>
      <ExtendedDruckerPrager
        name="drucker"
        defaultDensity="2700"
        defaultBulkModulus="500"
        defaultShearModulus="300"
        defaultCohesion="0.0"
        defaultInitialFrictionAngle="15.27"
        defaultResidualFrictionAngle="23.05"
        defaultDilationRatio="1.0"
        defaultHardening="0.001"/>
      </Constitutive>

    <Functions>
      <TableFunction
        name="strainFunction"
        inputVarNames="{ time }"
        coordinates="{ 0.0, 3.0, 4.0, 7.0, 8.0 }"
        values="{ 0, -0.003, -0.002, -0.005, -0.004 }"/>

      <TableFunction
        name="stressFunction"
        inputVarNames="{ time }"
        coordinates="{ 0.0, 8.0 }"
        values="{ -1.0, -1.0 }"/>
      </Functions>

    <!-- Mesh is not used, but GEOSX throws an error without one. Will resolve soon-->
    <Mesh>
      <InternalMesh
        name="mesh1"
        elementTypes="{ C3D8 }"
        xCoords="{ 0, 1 }"

```

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```

yCoords="{ 0, 1 }"
zCoords="{ 0, 1 }"
nx="{ 1 }"
ny="{ 1 }"
nz="{ 1 }"
cellBlockNames="{ cellBlock01 }"/>
</Mesh>

<ElementRegions>
  <CellElementRegion
    name="dummy"
    cellBlocks="{ cellBlock01 }"
    materialList="{ dummy }"/>
  </ElementRegions>
</Problem>

```

The first thing to note is that the XML structure is identical to a standard GEOSX input deck. In fact, once the constitutive block is calibrated, one could start adding solver and discretization blocks to the same file to create a proper field simulation. This makes it easy to go back and forth between calibration and simulation.

The `TriaxialDriver` is added as a `Task`, a particular type of executable event often used for simple actions. It is added as a `SoloEvent` to the event queue. This leads to a trivial event queue, since all we do is launch the driver and then quit.

Internally, the triaxial driver uses a simple form of time-stepping to advance through the loading steps, allowing for both rate-dependent and rate-independent models to be tested. This timestepping is handled independently from the more complicated time-stepping pattern used by physics `Solvers` and coordinated by the `EventManager`. In particular, in the XML file above, the `maxTime` parameter in the `Events` block is an event manager control, controlling when/if certain events occur. Once launched, the triaxial driver internally determines its own max time and timestep size using a combination of the strain function's time coordinates and the requested number of loadsteps. It is therefore helpful to think of the driver as an instantaneous *event* (from the event manager's point of view), but one which has a separate, internal clock.

The key parameters for the `TriaxialDriver` are:

Name	Type	Default	Description
axialControl	string	required	Function controlling axial stress or strain (depending on test mode)
baseline	path	none	Baseline file
initialStress	real64	required	Initial stress (scalar used to set an isotropic stress state)
logLevel	integer	0	Log level
material	string	required	Solid material to test
mode	string	required	Test mode [stressControl, strainControl, mixedControl]
name	string	required	A name is required for any non-unique nodes
output	string	none	Output file
radialControl	string	required	Function controlling radial stress or strain (depending on test mode)
steps	integer	required	Number of load steps to take

**Note:** GEOSX uses the *engineering* sign convention where compressive stresses and strains are *negative*. This is one of the most frequent issues users make when calibrating material parameters, as stress- and strain-like quantities often need to be negative to make physical sense. You may note in the XML above, for example, that `stressFunction` and `strainFunction` have negative values for a compressive test.

## Test Modes

The most complicated part of the driver is understanding how the stress and strain functions are applied in different testing modes. The driver mimics laboratory core tests, with loading controlled in the axial and radial directions. These conditions may be either strain-controlled or stress-controlled, with the user providing time-dependent functions to describe the loading. The following table describes the available test modes in detail:

mode	axial loading	radial loading	initial stress
strainControl	axial strain controlled with axialControl	radial strain controlled with radialControl	isotropic stress using initialStress
stressControl	axial stress controlled with axialControl	radial stress controlled with radialControl	isotropic stress using initialStress
mixedControl	axial strain controlled with axialControl	radial stress controlled with radialControl	isotropic stress using initialStress

Note that a classical triaxial test can be described using either the `stressControl` or `mixedControl` mode. We recommend using the `mixedControl` mode when possible, because this almost always leads to well-posed loading conditions. In a pure stress controlled test, it is possible for the user to request that the material sustain a load beyond its intrinsic strength envelope, in which case there is no feasible solution and the driver will fail to converge. Imagine, for example, a perfectly plastic material with a yield strength of 10 MPa, but the user attempts to load it to 11 MPa.

A volumetric test can be created by setting the axial and radial control functions to the same time history function. Similarly, an oedometer test can be created by setting the radial strain to zero.

The user should be careful to ensure that the initial stress set via the `initialStress` value is consistent any applied stresses set through axial or radial loading functions. Otherwise, the material may experience sudden and unexpected deformation at the first timestep because it is not in static equilibrium.

## Output Format

The `output` key is used to identify a file to which the results of the simulation are written. If this key is omitted, or the user specifies `output="none"`, file output will be suppressed. The file is a simple ASCII format with a brief header followed by test data:

```
# column 1 = time
# column 2 = axial_strain
# column 3 = radial_strain_1
# column 4 = radial_strain_2
# column 5 = axial_stress
# column 6 = radial_stress_1
# column 7 = radial_stress_2
# column 8 = newton_iter
# column 9 = residual_norm
0.0000e+00 0.0000e+00 0.0000e+00 0.0000e+00 -1.0000e+00 -1.0000e+00 -1.0000e+00 0.
↪0000e+00 0.0000e+00
1.6000e-01 -1.6000e-04 4.0000e-05 4.0000e-05 -1.1200e+00 -1.0000e+00 -1.0000e+00 2.
↪0000e+00 0.0000e+00
3.2000e-01 -3.2000e-04 8.0000e-05 8.0000e-05 -1.2400e+00 -1.0000e+00 -1.0000e+00 2.
↪0000e+00 0.0000e+00
...
```

This file can be readily plotted using any number of plotting tools. Each row corresponds to one timestep of the driver, starting from initial conditions in the first row.

We note that the file contains two columns for radial strain and two columns for radial stress. For an isotropic material, the stresses and strains along the two radial axes will usually be identical. We choose to output this way, however, to accommodate both anisotropic materials and true-triaxial loading conditions. In these cases, the stresses and strains in the radial directions could potentially differ.

These columns can be added and subtracted to produce other quantities of interest, like mean stress or deviatoric stress. For example, we can plot the output produce stress / strain curves (in this case for a plastic rather than simple elastic material):

Fig. 5.1: Stress/strain behavior for a plastic material.

In this plot, we have reversed the sign convention to be consistent with typical experimental plots. Note also that the `strainFunction` includes two unloading cycles, allowing us to observe both plastic loading and elastic unloading.

## Model Convergence

The last two columns of the output file contain information about the convergence behavior of the material driver. In `triaxial` mode, the mixed nature of the stress/strain control requires using a Newton solver to converge the solution. This last column reports the number of Newton iterations and final residual norm. Large values here would be indicative of the material model struggling (or failing) to converge. Convergence failures can result from several reasons, including:

1. Inappropriate material parameter settings
2. Overly large timesteps
3. Infeasible loading conditions (i.e. trying to load a material to a physically-unreachable stress point)
4. Poor model implementation

We generally spend a lot of time vetting the material model implementations (#4). When you first encounter a problem, it is therefore good to explore the other three scenarios first. If you find something unusual in the model implementation or are just really stuck, please submit an issue on our issue tracker so we can help resolve any bugs.

## Unit Testing

The development team also uses the Triaxial Driver to perform unit testing on the various material models within GEOSX. The optional argument `baseline` can be used to point to a previous output file that has been validated (e.g. against analytical or experimental benchmarks). If such a file is specified, the driver will perform a loading run and then compare the new results against the baseline. In this way, any regressions in the material models can be quickly identified.

Developers of new models are encouraged to add their own baselines to `src/coreComponents/constitutive/unitTests`. Adding additional tests is straightforward:

1. Create a new xml file for your test in `src/coreComponents/constitutive/unitTests`. There are several examples in this directory already to use as a template. We suggest using the naming convention `testTriaxial_myTest.xml`, so that all triaxial tests will be grouped together alphabetically. Set the output file to `testTriaxial_myTest.txt`, and run your test. Validate the results however is appropriate.
2. This output file will now become your new baseline. Replace the `output` key with `baseline` so that the driver can read in your file as a baseline for comparison. Make sure there is no remaining `output` key, or set `output=none`, to suppress further file output. While you can certainly write a new output for debugging purposes, during our automated unit tests we prefer to suppress file output. Re-run the triaxial driver to confirm that the comparison test passes.



3. Modify `src/coreComponents/constitutive/unitTests/CMakeLists.txt` to enable your new test in the unit test suite. In particular, you will need to add your new XML file to the existing list in the `gtest_triaxial_xmls` variable:

```
set( gtest_triaxial_xmls
    testTriaxial_elasticIsotropic.xml
    testTriaxial_druckerPragerExtended.xml
    testTriaxial_myTest.xml
)
```

4. Run `make` in your build directory to make sure the CMake syntax is correct
5. Run `ctest -V -R Triax` to run the triaxial unit tests. Confirm your test is included and passes properly.

If you run into troubles, do not hesitate to contact the development team for help.

## Model: Elastic Isotropic

### Overview

This model may be used for solid materials with a linear elastic isotropic behavior. The relationship between stress and strain is given by [Hooke's Law](#), expressed as:

$$\sigma_{ij} = \lambda \epsilon_{kk} + 2\mu \epsilon_{ij},$$

where  $\sigma_{ij}$  is the  $ij$  component of the Cauchy stress tensor,  $\epsilon_{ij}$  is the  $ij$  component of the strain tensor,  $\lambda$  is the first Lamé elastic constant, and  $\mu$  is the elastic shear modulus.

Hooke's Law may also be expressed using [Voigt notation](#) for stress and strain vectors as:

$$\sigma = C \cdot \epsilon,$$

or,

$$\begin{bmatrix} \sigma_{11} \\ \sigma_{22} \\ \sigma_{33} \\ \sigma_{23} \\ \sigma_{13} \\ \sigma_{12} \end{bmatrix} = \begin{bmatrix} 2\mu + \lambda & \lambda & \lambda & 0 & 0 & 0 \\ \lambda & 2\mu + \lambda & \lambda & 0 & 0 & 0 \\ \lambda & \lambda & 2\mu + \lambda & 0 & 0 & 0 \\ 0 & 0 & 0 & \mu & 0 & 0 \\ 0 & 0 & 0 & 0 & \mu & 0 \\ 0 & 0 & 0 & 0 & 0 & \mu \end{bmatrix} \begin{bmatrix} \epsilon_{11} \\ \epsilon_{22} \\ \epsilon_{33} \\ 2\epsilon_{23} \\ 2\epsilon_{13} \\ 2\epsilon_{12} \end{bmatrix}.$$

### Variations

For finite deformation solvers, the elastic isotropic model can be called within a hypo-elastic update routine. See [Finite Deformation Models with Hypo-Materials](#)

### Parameters

The following attributes are supported. Note that any two elastic constants can be provided, and the other two will be internally calculated. The “default” keyword in front of certain properties indicates that this is the default value adopted for a region unless the user separately specifies a heterogeneous field via the `FieldSpecification` mechanism.

Name	Type	Default	Description
defaultBulkModulus	real64	-1	Default Bulk Modulus Parameter
defaultDensity	real64	required	Default Material Density
defaultPoissonRatio	real64	-1	Default Poisson's Ratio
defaultShearModulus	real64	-1	Default Shear Modulus Parameter
defaultYoungModulus	real64	-1	Default Young's Modulus
name	string	required	A name is required for any non-unique nodes

## Example

A typical `Constitutive` block will look like:

```
<Constitutive>
  <ElasticIsotropic
    name="shale"
    defaultDensity="2700"
    defaultBulkModulus="60.0e6"
    defaultShearModulus="30.0e6" />
</Constitutive>
```

## Model: Elastic Isotropic Pressure Dependent

### Overview

This model may be used for solid materials with a pressure-dependent elastic isotropic behavior. The relationship between stress and strain is given by a [hyperelastic law](#). The elastic constitutive equations for the volumetric and deviatoric stresses and strain are expressed as:

$$p = p_0 \exp \left( \frac{\epsilon_{v0} - \epsilon_v^e}{c_r} \right), \quad q = 3\mu\epsilon_s^e$$

where  $p$  and  $q$  are the volumetric and deviatoric components of the Cauchy stress tensor.  $\epsilon_v^e$  and  $\epsilon_s^e$  are the volumetric and deviatoric components of the strain tensor.  $\epsilon_{v0}$  and  $p_0$  are the initial volumetric strain and initial pressure.  $c_r$  denotes the elastic compressibility index, and  $\mu$  is the elastic shear modulus. In this model, the shear modulus is constant and the bulk modulus,  $K$ , varies linearly with pressure as follows:

$$K = -\frac{p}{c_r}$$

### Parameters

The following attributes are supported. Note that two elastic constants  $c_r$  and  $\mu$ , as well as the initial volumetric strain and initial pressure need to be provided. The “default” keyword in front of certain properties indicates that this is the default value adopted for a region unless the user separately specifies a heterogeneous field via the `FieldSpecification` mechanism.

## Example

A typical `Constitutive` block will look like:

```

<Constitutive>
  <ElasticIsotropicPressureDependent
    name="elasticPressure"
    defaultDensity="2700"
    defaultRefPressure="-1.0"
    defaultRefStrainVol="1"
    defaultRecompressionIndex="0.003"
    defaultShearModulus="200"/>
</Constitutive>

```

## Model: Elastic Transverse-Isotropic

### Overview

This model may be used for solid materials with a linear elastic, transverse-isotropic behavior. This is most readily expressed in Voight notation as

$$\begin{bmatrix} \sigma_{11} \\ \sigma_{22} \\ \sigma_{33} \\ \sigma_{23} \\ \sigma_{13} \\ \sigma_{12} \end{bmatrix} = \begin{bmatrix} C_{11} & C_{12} & C_{13} & 0 & 0 & 0 \\ C_{12} & C_{11} & C_{13} & 0 & 0 & 0 \\ C_{13} & C_{13} & C_{33} & 0 & 0 & 0 \\ 0 & 0 & 0 & C_{44} & 0 & 0 \\ 0 & 0 & 0 & 0 & C_{44} & 0 \\ 0 & 0 & 0 & 0 & 0 & (C_{11} - C_{12})/2 \end{bmatrix} \begin{bmatrix} \epsilon_{11} \\ \epsilon_{22} \\ \epsilon_{33} \\ 2\epsilon_{23} \\ 2\epsilon_{13} \\ 2\epsilon_{12} \end{bmatrix} .$$

This system contains five independent constants. These constants are calculated from the input parameters indicated below.

### Parameters

The following attributes are supported. The “default” keyword in front of certain properties indicates that this is the default value adopted for a region unless the user separately specifies a heterogeneous field via the FieldSpecification mechanism.

Name	Type	Default	Description
defaultC11	real64	-1	Default Stiffness Parameter C11
defaultC13	real64	-1	Default Stiffness Parameter C13
defaultC33	real64	-1	Default Stiffness Parameter C33
defaultC44	real64	-1	Default Stiffness Parameter C44
defaultC66	real64	-1	Default Stiffness Parameter C66
defaultDensity	real64	required	Default Material Density
defaultPoissonRatioAxialTransverse	real64	-1	Default Axial-Transverse Poisson’s Ratio
defaultPoissonRatioTransverse	real64	-1	Default Transverse Poisson’s Ratio
defaultShearModulusAxialTransverse	real64	-1	Default Axial-Transverse Shear Modulus
defaultYoungModulusAxial	real64	-1	Default Axial Young’s Modulus
defaultYoungModulusTransverse	real64	-1	Default Transverse Young’s Modulus
name	string	required	A name is required for any non-unique nodes

### Example

A typical Constitutive block will look like:

```
<Constitutive>
  <ElasticTransverseIsotropic
    name="shale"
    defaultDensity="2700"
    defaultPoissonRatioAxialTransverse="0.20"
    defaultPoissonRatioTransverse="0.30"
    defaultYoungModulusAxial="50.0e6"
    defaultYoungModulusTransverse="60.0e6"
    defaultShearModulusAxialTransverse="30.0e6" />
</Constitutive>
```

## Model: Elastic Orthotropic

### Overview

This model may be used for solid materials with a linear elastic, orthotropic behavior. This is most readily expressed in Voight notation as

$$\begin{bmatrix} \sigma_{11} \\ \sigma_{22} \\ \sigma_{33} \\ \sigma_{23} \\ \sigma_{13} \\ \sigma_{12} \end{bmatrix} = \begin{bmatrix} C_{11} & C_{12} & C_{13} & 0 & 0 & 0 \\ C_{12} & C_{22} & C_{23} & 0 & 0 & 0 \\ C_{13} & C_{23} & C_{33} & 0 & 0 & 0 \\ 0 & 0 & 0 & C_{44} & 0 & 0 \\ 0 & 0 & 0 & 0 & C_{55} & 0 \\ 0 & 0 & 0 & 0 & 0 & C_{66} \end{bmatrix} \begin{bmatrix} \epsilon_{11} \\ \epsilon_{22} \\ \epsilon_{33} \\ 2\epsilon_{23} \\ 2\epsilon_{13} \\ 2\epsilon_{12} \end{bmatrix}.$$

This system contains nine independent constants. These constants are calculated from the input parameters indicated below.

### Parameters

The following attributes are supported. The “default” keyword in front of certain properties indicates that this is the default value adopted for a region unless the user separately specifies a heterogeneous field via the `FieldSpecification` mechanism.

Name	Type	Default	Description
defaultC11	real64	-1	Default C11 Component of Voigt Stiffness Tensor
defaultC12	real64	-1	Default C12 Component of Voigt Stiffness Tensor
defaultC13	real64	-1	Default C13 Component of Voigt Stiffness Tensor
defaultC22	real64	-1	Default C22 Component of Voigt Stiffness Tensor
defaultC23	real64	-1	Default C23 Component of Voigt Stiffness Tensor
defaultC33	real64	-1	Default C33 Component of Voigt Stiffness Tensor
defaultC44	real64	-1	Default C44 Component of Voigt Stiffness Tensor
defaultC55	real64	-1	Default C55 Component of Voigt Stiffness Tensor
defaultC66	real64	-1	Default C66 Component of Voigt Stiffness Tensor
defaultDensity	real64	required	Default Material Density
defaultE1	real64	-1	Default Young's Modulus E1
defaultE2	real64	-1	Default Young's Modulus E2
defaultE3	real64	-1	Default Young's Modulus E3
defaultG12	real64	-1	Default Shear Modulus G12
defaultG13	real64	-1	Default Shear Modulus G13
defaultG23	real64	-1	Default Shear Modulus G23
defaultNu12	real64	-1	Default Poission's Ratio Nu12
defaultNu13	real64	-1	Default Poission's Ratio Nu13
defaultNu23	real64	-1	Default Poission's Ratio Nu23
name	string	required	A name is required for any non-unique nodes

## Example

A typical Constitutive block will look like:

```
<Constitutive>
  <ElasticOrthotropic
    name="shale"
    defaultDensity="2700"
    defaultNu12="0.20"
    defaultNu13="0.25"
    defaultNu23="0.30"
    defaultE1="40.0e6"
    defaultE2="50.0e6"
    defaultE3="60.0e6"
    defaultG12="20.0e6"
    defaultG13="30.0e6"
    defaultG23="40.0e6" />
  </Constitutive>
```

## Model: Drucker-Prager

### Overview

This model may be used to represent a solid material with plastic response to loading according to the [Drucker-Prager](#) yield criterion below:

$$f(p, q) = q + bp - a = 0.$$

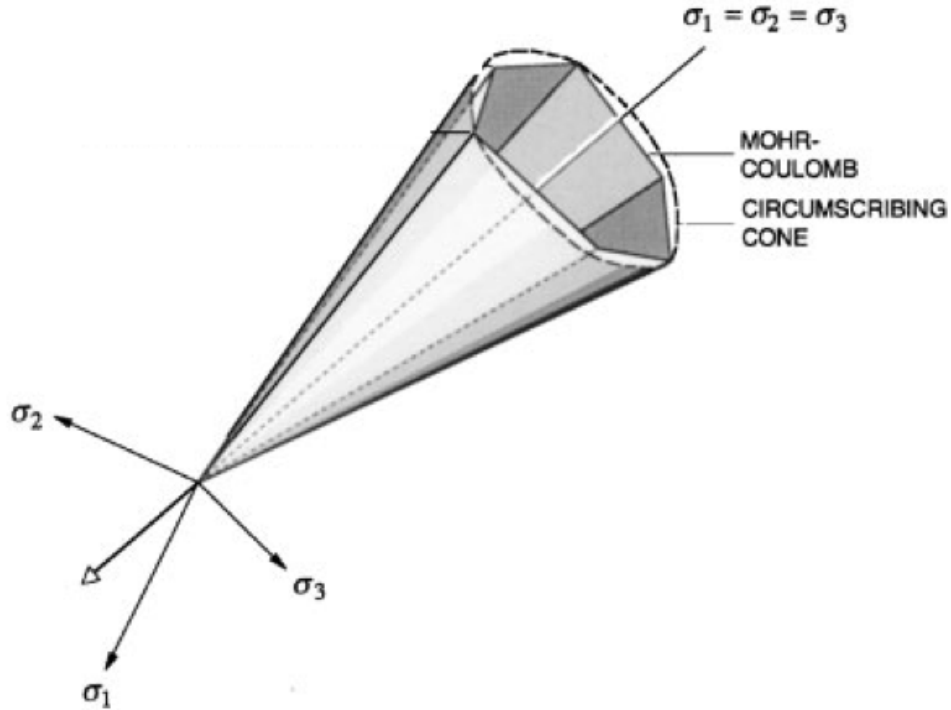


Fig. 5.2: Mohr-Coulomb and Drucker-Prager yield surfaces in principal stress axes (Borja, 2002).

The material behavior is linear elastic (see [Model: Elastic Isotropic](#)) for  $f < 0$ , and plastic for  $f = 0$ . The two material parameters  $a$  and  $b$  are derived by approximating the Mohr-Coulomb surface with a cone. Figure 3 shows the Mohr-Coulomb yield surface and circumscribing Drucker-Prager surface in principal stress space. The Drucker-Prager yield surface has a circular cross-section in the deviatoric plane that passes through the tension or compression corners of the Mohr-Coulomb yield surface, as shown in the Figure 4. The material parameters  $a$  and  $b$  are derived as:

$$a = \frac{6c \cos \phi}{3 \pm \sin \phi}, \quad b = \frac{6 \sin \phi}{3 \pm \sin \phi}$$

where plus signs are for circles passing through the tension corners, and minus signs are for circles passing through compression corners. Also,  $\phi$  and  $c$  denote friction angle and cohesion, respectively, as defined by the Mohr-Coulomb failure envelope shown in Figure 5. In GEOSX, we use a compression corner fit (minus signs) to convert the user-specified friction angle and cohesion to  $a$  and  $b$ .

We consider a non-associative plastic potential to determine the direction of plastic flow.

$$g(p, q) = q + b' p,$$

where  $b' \leq b$  is the dilatancy parameter. Setting  $b' = b$  leads to associative flow rule, while for  $b' < b$  non-associative flow is obtained. The parameter  $b'$  is related to [dilation angle](#) as:

$$b' = \frac{6 \sin \psi}{3 \pm \sin \psi},$$

where  $\psi \leq \phi$  is the dilation angle. If  $\psi > 0$ , then the plastic flow is dilative. Again, we use a compression corner fit (minus sign).

A hardening rule is defined which determines how the yield surface will change as a result of plastic deformations. Here we use linear hardening for the cohesion parameter,  $a$ ,

$$\dot{a} = h \dot{\lambda},$$

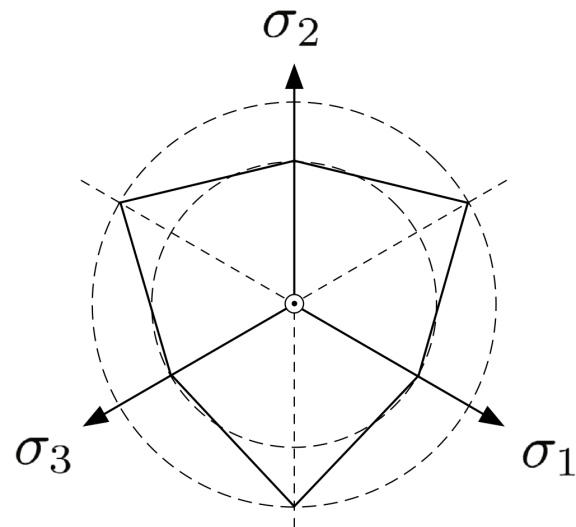


Fig. 5.3: Mohr-Coulomb and Drucker-Prager yield surfaces on the deviatoric plane (Borja, 2013).

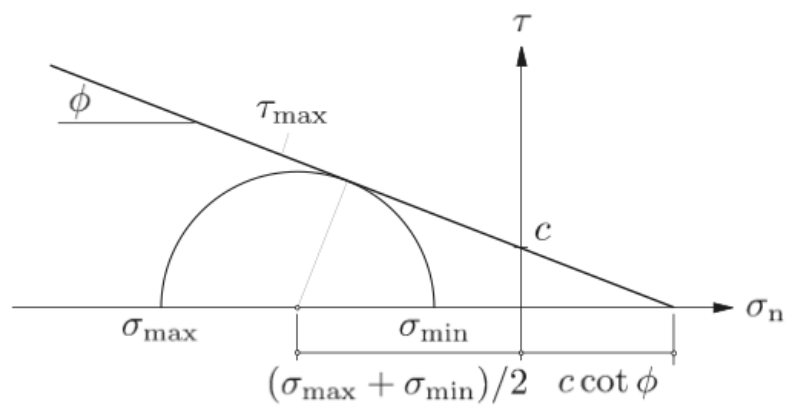


Fig. 5.4: The Mohr-Coulomb failure envelope (Borja, 2013).

where  $h$  is the hardening parameter. A positive hardening parameter will allow the cohesion to grow, shifting the cohesion intercept vertically on the  $q$ -axis. A negative hardening parameter will cause the cohesion to shrink, though negative cohesion values are not allowed. Once all cohesion has been lost, the cohesion will remain at zero, so the cone vertex is fixed at the origin. In either case, the friction and dilation angles remain constant. See the *Drucker-PragerExtended* model for an alternative version of hardening behavior.

## Parameters

The following attributes are supported:

Name	Type	Default	Description
defaultBulkModulus	real64	-1	Default Bulk Modulus Parameter
defaultCohesion	real64	0	Initial cohesion
defaultDensity	real64	required	Default Material Density
defaultDilationAngle	real64	30	Dilation angle (degrees)
defaultFrictionAngle	real64	30	Friction angle (degrees)
defaultHardeningRate	real64	0	Cohesion hardening/softening rate
defaultPoissonRatio	real64	-1	Default Poisson's Ratio
defaultShearModulus	real64	-1	Default Shear Modulus Parameter
defaultYoungModulus	real64	-1	Default Young's Modulus
name	string	required	A name is required for any non-unique nodes

## Example

```
<Constitutive>
  <DruckerPrager name="drucker"
    defaultDensity="2700"
    defaultBulkModulus="1000.0"
    defaultShearModulus="1000.0"
    defaultFrictionAngle="30.0"
    defaultDilationAngle="20.0"
    defaultHardeningRate="0.0"
    defaultCohesion="10.0" />
</Constitutive>
```

## Variant: J2 plasticity

J2 yield criterion can be obtained as a special case of the Drucker-Prager model by setting the friction and dilation angles to zero, i.e.  $\phi = \psi = 0$ .

## Model: Extended Drucker-Prager

### Overview

This model implements a more sophisticated version of the Drucker-Prager model (see *Model: Drucker-Prager*) allowing for both cohesion and friction hardening / softening. We implement the specific hardening model reported in Liu et al. (2020). The yield surface is given by

$$f(p, q) = q + b \left( p - \frac{a_i}{b_i} \right) = 0,$$



where  $b$  is the current yield surface slope,  $b_i$  is the initial slope, and  $a_i$  is the initial cohesion intercept in  $p$ - $q$  space. The vertex of the Drucker-Prager cone is fixed at  $p = a_i/b_i$ . Let  $\lambda$  denote the accumulated plastic strain measure. The current yield surface slope is given by the hyperbolic relationship

$$b = b_i + \frac{\lambda}{m + \lambda} (b_r - b_i)$$

with  $m$  a parameter controlling the hardening rate. Here,  $b_r$  is the residual yield surface slope. If  $b_r < b_i$ , hardening behavior will be observed, while for  $b_r > b_i$  softening behavior will occur.

In the resulting model, the yield surface begins at an initial position defined by the initial cohesion and friction angle. As plastic deformation occurs, the friction angle hardens (or softens) so that it asymptotically approaches a residual friction angle. The vertex of the cone remains fixed in  $p$ - $q$  space, but the cohesion intercept evolves in tandem with the friction angle. See *Liu et al. (2020)* <<https://doi.org/10.1007/s00603-019-01992-5>> for complete details.

In order to allow for non-associative behavior, we define a “dilation ratio” parameter  $\theta \in [0, 1]$  such that  $b' = \theta b$ , where  $b'$  is the slope of the plastic potential surface, while  $b$  is the slope of the yield surface. Choosing  $\theta = 1$  leads to associative behavior, while  $\theta = 0$  implies zero dilatancy.

## Parameters

The supported attributes will be documented soon.

## Example

```
<Constitutive>
  <ExtendedDruckerPrager
    name="edp"
    defaultDensity="2700"
    defaultBulkModulus="500"
    defaultShearModulus="300"
    defaultCohesion="0.0"
    defaultInitialFrictionAngle="15.0"
    defaultResidualFrictionAngle="23.0"
    defaultDilationRatio="1.0"
    defaultHardening="0.001"
  />
</Constitutive>
```

## Model: Modified Cam-Clay

This model may be used to represent a solid material with plastic response to loading according to the [Modified Cam-Clay \(MCC\)](#) critical state model. The MCC yield function is defined as:

$$f = q^2 + M^2 [p(p - p_c)] = 0,$$

where  $p_c$  is the preconsolidation pressure, and  $M$  is the slope of the critical state line (CSL).  $M$  can be related to the critical state friction angle  $\phi_{cs}$  as

$$M = \frac{6 \sin \phi_{cs}}{3 - \sin \phi_{cs}}.$$

Here  $f$  represents the yield surface, as shown in Figure 6.

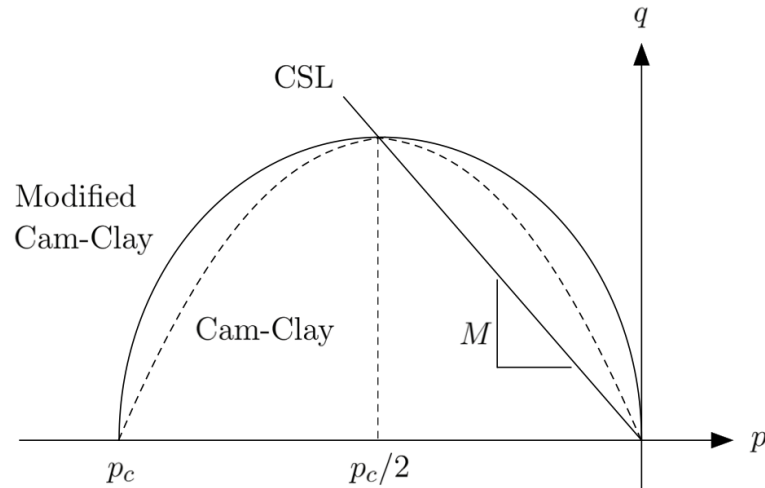


Fig. 5.5: Cam-Clay and Modified Cam-Clay yield surfaces in p-q space (Borja, 2013).

Here we use a hyper-elastic constitutive law using the following elastic rate constitutive equation

$$\dot{p} = -\frac{p}{c_r} \dot{\epsilon}_v^e,$$

where  $c_r > 0$  is the elastic compressibility index. The tangential elastic bulk modulus is  $K = -\frac{p}{c_r}$  and varies linearly with pressure. We assume a constant shear modulus, and can write stress invariants  $p$  and  $q$  as

$$p = p_0 \exp\left(\frac{\epsilon_{v0} - \epsilon_v^e}{c_r}\right), \quad q = 3\mu\epsilon_s^e,$$

where  $p_0$  is the reference pressure and  $\epsilon_{v0}$  is the reference volumetric strain. The hardening law is derived from the linear relationship between logarithm of specific volume and logarithm of preconsolidation pressure, as show in Figure 7.

The hardening law describes evolution of the preconsolidation pressure  $p_c$  as

$$\dot{p}_c = -\frac{tr(\dot{\epsilon}^p)}{c_c - c_r} p_c,$$

where  $c_c$  is the virgin compressibility index and we have  $0 < c_r < c_c$ .

## Parameters

The supported attributes will be documented soon.

## Example

```
<Constitutive>
  <ModifiedCamClay name="mcc"
    defaultDensity="2700"
    defaultRefPressure="-1.0"
    defaultRefStrainVol="0"
    defaultShearModulus="200.0"
```

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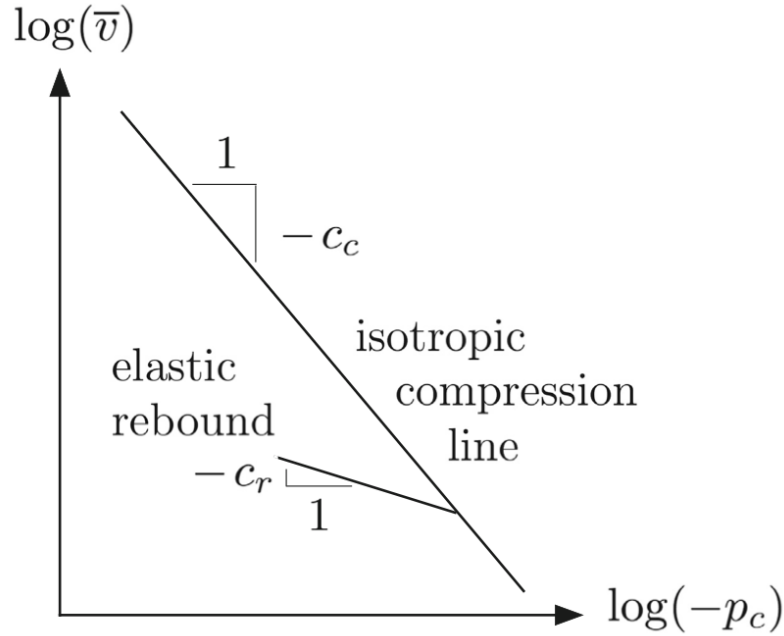


Fig. 5.6: Bilogarithmic hardening law derived from isotropic compression tests (Borja, 2013).

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```
defaultPreConsolidationPressure="-1.5"
defaultCslSlope="1.2"
defaultRecompressionIndex="0.002"
defaultVirginCompressionIndex="0.003" />
</Constitutive>
```

### Model: Delft Egg

The **Delft-Egg** plasticity model uses a generalization of the Modified Cam-Clay yield surface, defined as

$$f = q^2 - M^2 \left[ \alpha^2 p \left( \frac{2\alpha}{\alpha+1} p_c - p \right) - \frac{\alpha^2(\alpha-1)}{\alpha+1} p_c^2 \right] = 0 \quad \left( \text{for } p_c > \frac{\alpha}{\alpha+1} \right)$$

$$f = q^2 - M^2 p \left( \frac{2\alpha}{\alpha+1} p_c - p \right) = 0 \quad \left( \text{for } p_c \leq \frac{\alpha}{\alpha+1} \right)$$

where  $\alpha \geq 1$  is the shape parameter. For  $\alpha = 1$ , this model leads to a Modified Cam-Clay (MCC) type model with an ellipsoidal yield surface. For  $\alpha > 1$ , an egg-shaped yield surface is obtained. The additional parameter makes it easier to fit the cap behavior of a broader range of soils and rocks.

Because Delft-Egg is frequently used for hard rocks, GEOSX uses a linear model for the elastic response, rather than the hyper-elastic model used for MCC. This is a slight deviation from the original formulation proposed in the reference above. For reservoir applications, the ability to use a simpler linear model was a frequent user request.

### Parameters

The supported attributes will be documented soon.

## Example

```
<Constitutive>
  <DelftEgg
    name="DE"
    defaultDensity="2700"
    defaultBulkModulus="10.0e9"
    defaultShearModulus="6.0e9"
    defaultPreConsolidationPressure="-20.0e6"
    defaultShapeParameter="6.5"
    defaultCslSlope="1.2"
    defaultVirginCompressionIndex="0.005"
    defaultRecompressionIndex="0.001"/>
</Constitutive>
```

## Damage Models

The damage models are in active development, and documentation will be added when they are ready for production release.

## 5.4.2 Fluid Models

These models provide density, viscosity, and composition relationships for single fluids and fluid mixtures.

### Compressible single phase fluid model

#### Overview

This model represents a compressible single-phase fluid with constant compressibility and pressure-dependent viscosity. These assumptions are valid for slightly compressible fluids, such as water, and some types of oil with negligible amounts of dissolved gas.

Specifically, fluid density is computed as

$$\rho(p) = \rho_0 e^{c_\rho(p-p_0)}$$

where  $c_\rho$  is compressibility,  $p_0$  is reference pressure,  $\rho_0$  is density at reference pressure. Similarly,

$$\mu(p) = \mu_0 e^{c_\mu(p-p_0)}$$

where  $c_\mu$  is viscosibility (viscosity compressibility),  $\mu_0$  is reference viscosity.

Either exponent may be approximated by linear (default) or quadratic terms of Taylor series expansion. Currently there is no temperature dependence in the model, although it may be added in future.

#### Parameters

The model is represented by `<CompressibleSinglePhaseFluid>` node in the input.

The following attributes are supported:

Name	Type	Default	Description
compressibility	real64	0	Fluid compressibility
defaultDensity	real64	required	Default value for density.
defaultViscosity	real64	required	Default value for viscosity.
densityModelType	geosx_constitutive_ExponentialApproximationType		Type of density model. Valid options: * exponential * linear * quadratic
name	string	required	A name is required for any non-unique nodes
referenceDensity	real64	1000	Reference fluid density
referencePressure	real64	0	Reference pressure
referenceViscosity	real64	0.001	Reference fluid viscosity
viscosibility	real64	0	Fluid viscosity exponential coefficient
viscosityModelType	geosx_constitutive_ExponentialApproximationType		Type of viscosity model. Valid options: * exponential * linear * quadratic

## Example

```

<Constitutive>
  <CompressibleSinglePhaseFluid name="water"
    referencePressure="2.125e6"
    referenceDensity="1000"
    compressibility="1e-19"
    referenceViscosity="0.001"
    viscosibility="0.0"/>
</Constitutive>

```

## Black-oil fluid model

### Overview

In the black-oil model three pseudo-components, oil (o), gas (g) and water (w) are considered. These are assumed to be partitioned across three fluid phases, named liquid (l), vapor (v) and aqueous (a).

Phase behavior is characterized by the following quantities which are used to relate properties of the fluids in the reservoir to their properties at surface conditions.

- $B_o$ : oil formation volume factor

- $B_g$ : gas formation volume factor
- $R_s$ : gas/oil ratio
- $R_v$ : oil/gas ratio

By tables, that tabulate saturated and undersaturated oil and gas properties as functions of pressure and solution ratios.

## Dead oil

In **dead-oil** each component occupies only one phase. Thus, the following partition matrix determines the components distribution within the three phases:

$$\begin{bmatrix} y_{gv} & y_{gl} & y_{ga} \\ y_{ov} & y_{ol} & y_{oa} \\ y_{wv} & y_{wl} & y_{wa} \end{bmatrix} = \begin{bmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{bmatrix}$$

and the phase densities are

$$\rho_l = \frac{\rho_o^{STC}}{B_o}$$

$$\rho_v = \frac{\rho_g^{STC}}{B_g}.$$

## Live oil

The live oil fluid model make no assumptions about the partitioning of the hydrocarbon components and the following composition matrix can be used

$$\begin{bmatrix} y_{gv} & y_{gl} & y_{ga} \\ y_{ov} & y_{ol} & y_{oa} \\ y_{wv} & y_{wl} & y_{wa} \end{bmatrix} = \begin{bmatrix} \frac{\rho_g^{STC}}{\rho_g^{STC} + \rho_o^{STC} r_s} & \frac{\rho_g^{STC} R_s}{\rho_o^{STC} + \rho_g^{STC} R_s} & 0 \\ \frac{\rho_o^{STC} r_s}{\rho_g^{STC} + \rho_o^{STC} r_s} & \frac{\rho_o^{STC}}{\rho_o^{STC} + \rho_g^{STC} R_s} & 0 \\ 0 & 0 & 1 \end{bmatrix}$$

whereas the densities of the two hydrocarbon phases are

$$\rho_l = \frac{\rho_o^{STC} + \rho_g^{STC} R_s}{B_o}$$

$$\rho_v = \frac{\rho_g^{STC} + \rho_o^{STC} R_v}{B_g}$$

See [Petrowiki](#) for more information.

## Parameters

Both types are represented by <BlackOilFluid> node in the input. Under the hood this is a wrapper around PVTPackage library, which is included as a submodule. In order to use the model, GEOSX must be built with -DENABLE\_PVTPACKAGE=ON (default).

The following attributes are supported:

Name	Type	Default	Description
componentMolarWeight	real64_array	required	Component molar weights
componentNames	string_array	{}	List of component names
hydrocarbonFormationVolumeFactorTableNames	string_array	{}	<p>List of formation volume factor TableFunction names from the Functions block.</p> <p>The user must provide one TableFunction per hydrocarbon phase, in the order provided in “phaseNames”.</p> <p>For instance, if “oil” is before “gas” in “phaseNames”, the table order should be: oilTableName, gasTableName</p>
hydrocarbonViscosityTableNames	string_array	{}	<p>List of viscosity TableFunction names from the Functions block.</p> <p>The user must provide one TableFunction per hydrocarbon phase, in the order provided in “phaseNames”.</p> <p>For instance, if “oil” is before “gas” in “phaseNames”, the table order should be: oilTableName, gasTableName</p>
name	string	required	A name is required for any non-unique nodes
phaseNames	string_array	required	List of fluid phases
surfaceDensities	real64_array	required	List of surface mass densities for each phase
tableFiles	path_array	{}	List of filenames with input PVT tables (one per phase)
waterCompressibility	real64	0	Water compressibility
waterFormationVolumeFactor	real64	0	Water formation volume factor
waterReferencePressure	real64	0	Water reference pressure
waterViscosity	real64	0	Water viscosity

Supported phase names are:

Value	Comment
oil	Oil phase
gas	Gas phase
water	Water phase

## Example

```
<Constitutive>
  <BlackOilFluid name="fluid1"
    fluidType="LiveOil"
    phaseNames="{ oil, gas, water }"
    surfaceDensities="{ 800.0, 0.9907, 1022.0 }"
    componentMolarWeight="{ 114e-3, 16e-3, 18e-3 }"
    tableFiles="{ pvt.o.txt, pvt.g.txt, pvt.w.txt }"/>
</Constitutive>
```

## Compositional multiphase fluid model

### Overview

This model represents a full composition description of a multiphase multicomponent fluid. Phase behavior is modeled by an Equation of State (EOS) and partitioning of components into phases is computed based on instantaneous chemical equilibrium via a two- or three-phase flash. Each component (species) is characterized by molar weight and critical properties that serve as input parameters for the EOS. See [Petrowiki](#) for more information.

### Parameters

The model represented by `<CompositionalMultiphaseFluid>` node in the input. Under the hood this is a wrapper around `PVTPackage` library, which is included as a submodule. In order to use the model, GEOSX must be built with `-DENABLE_PVTPACKAGE=ON` (default).

The following attributes are supported:

Name	Type	Default	Description
componentAcentricFactor	real64_array	required	Component acentric factors
componentBinaryCoeff	real64_array2d	{{0}}	Table of binary interaction coefficients
componentCriticalPressure	real64_array	required	Component critical pressures
componentCriticalTemperature	real64_array	required	Component critical temperatures
componentMolarWeight	real64_array	required	Component molar weights
componentNames	string_array	required	List of component names
componentVolumeShift	real64_array	{0}	Component volume shifts
equationsOfState	string_array	required	List of equation of state types for each phase
name	string	required	A name is required for any non-unique nodes
phaseNames	string_array	required	List of fluid phases

Supported phase names are:



Value	Comment
oil	Oil phase
gas	Gas phase
water	Water phase

Supported Equation of State types:

Value	Comment
PR	Peng-Robinson EOS
SRK	Soave-Redlich-Kwong EOS

## Example

```
<Constitutive>
  <CompositionalMultiphaseFluid name="fluid1"
    phaseNames="{ oil, gas }"
    equationsOfState="{ PR, PR }"
    componentNames="{ N2, C10, C20, H2O }"
    componentCriticalPressure="{ 34e5, 25.3e5, 14.6e5, ↵
↵220.5e5 }"
    componentCriticalTemperature="{ 126.2, 622.0, 782.0, ↵
↵647.0 }"
    componentAcentricFactor="{ 0.04, 0.443, 0.816, 0.344 }
↵"
    componentMolarWeight="{ 28e-3, 134e-3, 275e-3, 18e-3 }
↵"
    componentVolumeShift="{ 0, 0, 0, 0 }"
    componentBinaryCoeff="{ { 0, 0, 0, 0 },
                             { 0, 0, 0, 0 },
                             { 0, 0, 0, 0 },
                             { 0, 0, 0, 0 } }"/>
</Constitutive>
```

## CO2-brine model

### Summary

The CO2-brine model implemented in GEOSX includes two components (CO2 and H2O) that are transported by one or two fluid phases (the brine phase and the CO2 phase). We refer to the brine phase with the subscript  $\ell$  and to the CO2 phase with the subscript  $g$  (although the CO2 phase can be in supercritical, liquid, or gas state). The water component is only present in the brine phase, while the CO2 component can be present in the CO2 phase as well as in the brine phase. Thus, considering the molar phase component fractions,  $y_{c,p}$  (i.e., the fraction of the molar mass of phase  $p$  represented by component  $c$ ) the following partition matrix determines the component distribution within the two phases:

$$\begin{bmatrix} y_{H2O,\ell} & y_{CO2,\ell} \\ 0 & 1 \end{bmatrix}$$

The update of the fluid properties is done in two steps:

- 1) The phase fractions ( $\nu_p$ ) and phase component fractions ( $y_{c,p}$ ) are computed as a function of pressure ( $p$ ), temperature ( $T$ ), component fractions ( $z_c$ ), and a constant salinity.

- 2) The phase densities ( $\rho_p$ ) and phase viscosities ( $\mu_p$ ) are computed as a function of pressure, temperature, the updated phase component fractions, and a constant salinity.

Once the phase fractions, phase component fractions, phase densities, phase viscosities—and their derivatives with respect to pressure, temperature, and component fractions—have been computed, the *Compositional Multiphase Flow Solver* proceeds to the assembly of the accumulation and flux terms. Note that the current implementation of the flow solver is isothermal and that the derivatives with respect to temperature are therefore discarded.

The models that are used in steps 1) and 2) are reviewed in more details below.

### Step 1: Computation of the phase fractions and phase component fractions (flash)

At initialization, GEOSX performs a preprocessing step to construct a two-dimensional table storing the values of CO2 solubility in brine as a function of pressure, temperature, and a constant salinity. The user can parameterize the construction of the table by specifying the salinity and by defining the pressure ( $p$ ) and temperature ( $T$ ) axis of the table in the form:

FlashModel	CO2Solubility	$p_{min}$	$p_{max}$	$\Delta p$	$T_{min}$	$T_{max}$	$\Delta T$	Salinity
------------	---------------	-----------	-----------	------------	-----------	-----------	------------	----------

Note that the pressures are in Pascal, temperatures are in Kelvin, and the salinity is a molality (moles of NaCl per kg of brine). The temperature must be between 283.15 and 623.15 Kelvin. The table is populated using the model of Duan and Sun (2003). Specifically, we solve the following nonlinear CO2 equation of state (equation (A1) in Duan and Sun, 2003) for each pair ( $p, T$ ) to obtain the reduced volume,  $V_r$ .

$$\frac{p_r V_r}{T_r} = 1 + \frac{a_1 + a_2/T_r^2 + a_3/T_r^3}{V_r} + \frac{a_4 + a_5/T_r^2 + a_6/T_r^3}{V_r^2} + \frac{a_7 + a_8/T_r^2 + a_9/T_r^3}{V_r^4} + \frac{a_{10} + a_{11}/T_r^2 + a_{12}/T_r^3}{V_r^5} + \frac{a_{13}}{T_r^3 V_r^2} \left( a_{14} + \frac{a_{15}}{V_r^2} \right) \exp\left(-\frac{a_{15}}{V_r^2}\right)$$

where  $p_r = p/p_{crit}$  and  $T_r = T/T_{crit}$  are respectively the reduced pressure and the reduced temperature. We refer the reader to Table (A1) in Duan and Sun (2003) for the definition of the coefficients  $a_i$  involved in the previous equation. Using the reduced volume,  $V_r$ , we compute the fugacity coefficient of CO2,  $\ln_\phi(p, T)$ , using equation (A6) of Duan and Sun (2003). To conclude this preprocessing step, we use the fugacity coefficient of CO2 to compute and store the solubility of CO2 in brine,  $s_{CO2}$ , using equation (6) of Duan and Sun (2003):

$$\ln \frac{y_{CO2} P}{s_{CO2}} = \frac{\Phi_{CO2}}{RT} - \ln_\phi(p, T) + \sum_c 2\lambda_c m + \sum_a 2\lambda_a m + \sum_{a,c} \zeta_{a,c} m^2$$

where  $\Phi_{CO2}$  is the chemical potential of the CO2 component,  $R$  is the gas constant, and  $m$  is the salinity. The mole fraction of CO2 in the vapor phase,  $y_{CO2}$ , is computed with equation (4) of Duan and Sun (2003). Note that the first, third, fourth, and fifth terms in the equation written above are approximated using equation (7) of Duan and Sun (2003) as recommended by the authors.

During the simulation, Step 1 starts with a look-up in the precomputed table to get the CO2 solubility,  $s_{CO2}$ , as a function of pressure and temperature. Then, we compute the phase fractions as:

$$\nu_\ell = \frac{1 + s_{CO2}}{1 + z_{CO2}/(1 - z_{CO2})}$$

$$\nu_g = 1 - \nu_\ell$$

We conclude Step 1 by computing the phase component fractions as:

$$y_{CO2,\ell} = \frac{s_{CO2}}{1 + s_{CO2}}$$

$$y_{H2O,\ell} = 1 - y_{CO2,\ell}$$

$$y_{CO2,g} = 1$$

$$y_{H2O,g} = 0$$

## Step 2: Computation of the phase densities and phase viscosities

### CO2 phase density and viscosity

In GEOSX, the computation of the CO2 phase density and viscosity is entirely based on look-up in precomputed tables. The user defines the pressure (in Pascal) and temperature (in Kelvin) axis of the density table in the form:

DensityFun	SpanWagnerCO2Density	$p_{min}$	$p_{max}$	$\Delta p$	$T_{min}$	$T_{max}$	$\Delta T$
------------	----------------------	-----------	-----------	------------	-----------	-----------	------------

This correlation is valid for pressures less than  $8 \times 10^8$  Pascal and temperatures less than 1073.15 Kelvin. Using these parameters, GEOSX internally constructs a two-dimensional table storing the values of density as a function of pressure and temperature. This table is populated as explained in the work of Span and Wagner (1996) by solving the following nonlinear Helmholtz energy equation for each pair  $(p, T)$  to obtain the value of density,  $\rho_g$ :

$$\frac{p}{RT\rho_g} = 1 + \delta\phi_\delta^r(\delta, \tau)$$

where  $R$  is the gas constant,  $\delta := \rho_g/\rho_{crit}$  is the reduced CO2 phase density, and  $\tau := T_{crit}/T$  is the inverse of the reduced temperature. The definition of the residual part of the energy equation, denoted by  $\phi_\delta^r$ , can be found in equation (6.5), page 1544 of Span and Wagner (1996). The coefficients involved in the computation of  $\phi_\delta^r$  are listed in Table (31), page 1544 of Span and Wagner (1996). These calculations are done in a preprocessing step.

The pressure and temperature axis of the viscosity table can be parameterized in a similar fashion using the format:

ViscosityFun	FenghourCO2Viscosity	$p_{min}$	$p_{max}$	$\Delta p$	$T_{min}$	$T_{max}$	$\Delta T$
--------------	----------------------	-----------	-----------	------------	-----------	-----------	------------

This correlation is valid for pressures less than  $3 \times 10^8$  Pascal and temperatures less than 1493.15 Kelvin. This table is populated as explained in the work of Fenghour and Wakeham (1998) by computing the CO2 phase viscosity,  $\mu_g$ , as follows:

$$\mu_g = \mu_0(T) + \mu_{excess}(\rho_g, T) + \mu_{crit}(\rho_g, T)$$

The “zero-density limit” viscosity,  $\mu_0(T)$ , is computed as a function of temperature using equations (3), (4), and (5), as well as Table (1) of Fenghour and Wakeham (1998). The excess viscosity,  $\mu_{excess}(\rho_g, T)$ , is computed as a function of temperature and CO2 phase density (computed as explained above) using equation (8) and Table (3) of Fenghour and Wakeham (1998). We currently neglect the critical viscosity,  $\mu_{crit}$ . These calculations are done in a preprocessing step.

During the simulation, the update of CO2 phase density and viscosity is simply done with a look-up in the precomputed tables.

### Brine density and viscosity using Phillips correlation

The computation of the brine density involves a tabulated correlation presented in Phillips et al. (1981). The user specifies the (constant) salinity and defines the pressure and temperature axis of the brine density table in the form:

DensityFun	PhillipsBrineDensity	$p_{min}$	$p_{max}$	$\Delta p$	$T_{min}$	$T_{max}$	$\Delta T$	Salinity
------------	----------------------	-----------	-----------	------------	-----------	-----------	------------	----------

The pressure must be in Pascal and must be less than  $5 \times 10^7$  Pascal. The temperature must be in Kelvin and must be between 283.15 and 623.15 Kelvin. The salinity is a molality (moles of NaCl per kg of brine). Using these parameters,

GEOSX performs a preprocessing step to construct a two-dimensional table storing the brine density,  $\rho_{\ell,table}$  for the specified salinity as a function of pressure and temperature using the expression:

$$\rho_{\ell,table} = A + Bx + Cx^2 + Dx^3$$

$$x = c_1 \exp(a_1 m) + c_2 \exp(a_2 T) + c_3 \exp(a_3 P)$$

We refer the reader to Phillips et al. (1981), equations (4) and (5), pages 14 and 15 for the definition of the coefficients involved in the previous equation. This concludes the preprocessing step.

Then, during the simulation, the brine density update proceeds in two steps. First, a table look-up is performed to retrieve the value of density,  $\rho_{\ell,table}$ . Then, in a second step, the density is modified using the method of Garcia (2001) to account for the presence of CO2 dissolved in brine as follows:

$$\rho_{\ell} = \rho_{\ell,table} + M_{CO_2} c_{CO_2} - c_{CO_2} \rho_{\ell,table} V_{\phi}$$

where  $M_{CO_2}$  is the molecular weight of CO2,  $c_{CO_2}$  is the concentration of CO2 in brine, and  $V_{\phi}$  is the apparent molar volume of dissolved CO2. The CO2 concentration in brine is obtained as:

$$c_{CO_2} = \frac{y_{CO_2,\ell} \rho_{\ell,table}}{M_{H_2O}(1 - y_{CO_2,\ell})}$$

where  $M_{H_2O}$  is the molecular weight of water. The apparent molar volume of dissolved CO2 is computed as a function of temperature using the expression:

$$V_{\phi} = 37.51 - 9.585 \times 10^{-2} T + 8.740 \times 10^{-4} T^2 - 5.044 \times 10^{-7} T^3$$

The brine viscosity is controlled by a salinity parameter provided by the user in the form:

ViscosityFun	PhillipsBrineViscosity	Salinity
--------------	------------------------	----------

During the simulation, the brine viscosity is updated as a function of temperature using the analytical relationship of Phillips et al. (1981):

$$\mu_{\ell} = aT + b$$

where the coefficients  $a$  and  $b$  are defined as:

$$a = \mu_w(T) \times 0.000629(1.0 - \exp(-0.7m))$$

$$b = \mu_w(T)(1.0 + 0.0816m + 0.0122m^2 + 0.000128m^3)$$

where  $\mu_w$  is the pure water viscosity computed as a function of temperature, and  $m$  is the user-defined salinity (in moles of NaCl per kg of brine).

## Brine density and viscosity using Ezrokhi correlation

Brine density  $\rho_l$  is computed from pure water density  $\rho_w$  at specified pressure and temperature corrected by Ezrokhi correlation presented in Zaytsev and Aseyev (1993):

$$\log_{10}(\rho_l) = \log_{10}(\rho_w(P, T)) + A(T)x_{CO_2,\ell}$$

$$A(T) = a_0 + a_1 T + a_2 T^2,$$

where  $a_0, a_1, a_2$  are correlation coefficients defined by user:

DensityFun	EzrokhiBrineDensity	$a_0$	$a_1$	$a_2$
------------	---------------------	-------	-------	-------

While  $x_{CO_2,\ell}$  is mass fraction of CO2 component in brine, computed from molar fractions as

$$x_{CO_2,\ell} = \frac{M_{CO_2} y_{CO_2,\ell}}{M_{CO_2} y_{CO_2,\ell} + M_{H_2O} y_{H_2O,\ell}},$$

Pure water density is computed according to:

$$\rho_w = \rho_{w,sat}(T) e^{c_w (P - P_{w,sat}(T))},$$

where  $c_w$  is water compressibility defined as a constant  $4.5 \times 10^{-10} Pa^{-1}$ , while  $\rho_{w,sat}(T)$  and  $P_{w,sat}(T)$  are density and pressure of saturated water at a given temperature. Both are obtained through internally constructed tables tabulated as functions of temperature and filled with the steam table data from Engineering ToolBox (2003, 2004).

Brine viscosity  $\mu_\ell$  is computed from pure water viscosity  $\mu_w$  similarly:

$$\log_{10}(\mu_\ell) = \log_{10}(\mu_w(P, T)) + B(T) x_{CO_2,\ell}$$

$$B(T) = b_0 + b_1 T + b_2 T^2,$$

where  $b_0, b_1, b_2$  are correlation coefficients defined by user:

ViscosityFun	EzrokhiBrineViscosity	$b_0$	$b_1$	$b_2$
--------------	-----------------------	-------	-------	-------

Mass fraction of CO2 component in brine  $x_{CO_2,\ell}$  is exactly as in density calculation. The dependency of pure water viscosity from pressure is ignored, and it is approximated as saturated pure water viscosity:

$$\mu_w(P, T) = \mu_{w,sat}(T),$$

which is tabulated using internal table as a function of temperature based on steam table data Engineering ToolBox (2004).

## Parameters

The models are represented by <CO2BrinePhillipsFluid>, <CO2BrineEzrokhiFluid> nodes in the input.

The following attributes are supported:

Name	Type	De- fault	Description
componentMolar- Weight	real64_array	{0}	Component molar weights
componentNames	string_array	{ }	List of component names
flashModelParaFile	path	re- quired	Name of the file defining the parameters of the flash model
name	string	re- quired	A name is required for any non-unique nodes
phaseNames	string_array	{ }	List of fluid phases
phasePVTParaFiles	path_array	re- quired	Names of the files defining the parameters of the viscosity and density models

Supported phase names are:

Value	Comment
gas	CO2 phase
water	Water phase

Supported component names are:

Value	Component
co2,CO2	CO2 component
water,liquid	Water component

### Example

```
<Constitutive>
  <CO2BrinePhillipsFluid
    name="fluid"
    phaseNames="{ gas, water }"
    componentNames="{ co2, water }"
    componentMolarWeight="{ 44e-3, 18e-3 }"
    phasePVTParaFiles="{ pvtgas.txt, pvtliquid.txt }"
    flashModelParaFile="co2flash.txt"/>
  </Constitutive>
```

```
<Constitutive>
  <CO2BrineEzrokhiFluid
    name="fluid"
    phaseNames="{ gas, water }"
    componentNames="{ co2, water }"
    componentMolarWeight="{ 44e-3, 18e-3 }"
    phasePVTParaFiles="{ pvtgas.txt, pvtliquid.txt }"
    flashModelParaFile="co2flash.txt"/>
  </Constitutive>
```

In the XML code listed above, “co2flash.txt” parameterizes the CO2 solubility table constructed in Step 1. The file “pvtgas.txt” parameterizes the CO2 phase density and viscosity tables constructed in Step 2, the file “pvtliquid.txt” parameterizes the brine density and viscosity tables according to Phillips or Ezrokhi correlation, depending on chosen fluid model.

### References

- Z. Duan and R. Sun, An improved model calculating CO2 solubility in pure water and aqueous NaCl solutions from 273 to 533 K and from 0 to 2000 bar., Chemical Geology, vol. 193.3-4, pp. 257-271, 2003.
- R. Span and W. Wagner, A new equation of state for carbon dioxide covering the fluid region from the triple-point temperature to 1100 K at pressure up to 800 MPa, J. Phys. Chem. Ref. Data, vol. 25, pp. 1509-1596, 1996.
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- S. L. Phillips et al., A technical databook for geothermal energy utilization, Lawrence Berkeley Laboratory report, 1981.
- J. E. Garcia, Density of aqueous solutions of CO2. No. LBNL-49023. Lawrence Berkeley National Laboratory, Berkeley, CA, 2001.
- Zaytsev, I.D. and Aseyev, G.G. Properties of Aqueous Solutions of Electrolytes, Boca Raton, Florida, USA CRC Press, 1993.
- Engineering ToolBox, Water - Density, Specific Weight and Thermal Expansion Coefficients, 2003

- Engineering ToolBox, [Water - Dynamic \(Absolute\) and Kinematic Viscosity](#), 2004

## PVT Driver

### Table of Contents

- *PVT Driver*
  - *Introduction*
  - *XML Structure*
  - *Parameters*
  - *Output Format*
  - *Unit Testing*

## Introduction

When calibrating fluid material parameters to experimental or other reference data, it can be a hassle to launch a full flow simulation just to confirm density, viscosity, and other fluid properties are behaving as expected. Instead, GEOSX provides a `PVTDriver` allowing the user to test fluid property models for a well defined set of pressure, temperature, and composition conditions. The driver itself is launched like any other GEOSX simulation, but with a particular XML structure:

```
./bin/geosx -i myFluidTest.xml
```

This driver will work for any multi-phase fluid model (e.g. black-oil, co2-brine, compositional multiphase) enabled within GEOSX.

## XML Structure

A typical XML file to run the driver will have several key elements. Here, we will walk through an example file included in the source tree at

```
src/coreComponents/unitTests/constitutiveTests/testPVT_docExample.xml
```

The first thing to note is that the XML file structure is identical to a standard GEOSX input deck. In fact, once the constitutive block is calibrated, one could start adding solver and discretization blocks to the same file to create a proper field simulation. This makes it easy to go back and forth between calibration and simulation.

The first step is to define a parameterized fluid model to test. Here, we create a particular type of CO<sub>2</sub>-Brine mixture:

```
<Constitutive>
  <CO2BrinePhillipsFluid
    name="co2Mixture"
    phaseNames="{ gas, water }"
    componentNames="{ co2, water }"
    componentMolarWeight="{ 44e-3, 18e-3 }"
    phasePVTParaFiles="{ testPVT_data/carbonDioxidePVT.txt, testPVT_data/brinePVT.
↪txt }"
    flashModelParaFile="testPVT_data/carbonDioxideFlash.txt"/>
  </Constitutive>
```

We also define two time-history functions for the pressure (Pascal units) and temperature (Kelvin units) conditions we want to explore.

```
<Functions>
  <TableFunction
    name="pressureFunction"
    inputVarNames="{ time }"
    coordinates="{ 0.0, 1.0 }"
    values="{ 1e6, 50e6 }"/>

  <TableFunction
    name="temperatureFunction"
    inputVarNames="{ time }"
    coordinates="{ 0.0, 1.0 }"
    values="{ 350, 350 }"/>
</Functions>
```

Note that the time-axis here is just a pseudo-time, allowing us to [parameterize](#) arbitrarily complicated paths through a (pressure,temperature) diagram. The actual time values have no impact on the resulting fluid properties. Here, we fix the temperature at 350K and simply ramp up pressure from 1 MPa to 50 MPa:

A PVTDriver is then added as a Task, a particular type of executable event often used for simple actions.

```
<Tasks>
  <PVTDriver
    name="testCO2"
    fluid="co2Mixture"
    feedComposition="{ 1.0, 0.0 }"
    pressureControl="pressureFunction"
    temperatureControl="temperatureFunction"
    steps="49"
    output="pvtOutput.txt"
    logLevel="1"/>
</Tasks>
```

The driver itself takes as input the fluid model, the pressure and temperature control functions, and a “feed composition.” The latter is the mole fraction of each component in the mixture to be tested. The `steps` parameter controls how many steps are taken along the parametric (P,T) path. Results will be written in a simple ASCII table format (described below) to the file `output`. The `logLevel` parameter controls the verbosity of log output during execution.

The driver task is added as a `SoloEvent` to the event queue. This leads to a trivial event queue, since all we do is launch the driver and then quit.

```
<Events
  maxTime="1">
  <SoloEvent
    name="eventA"
    target="/Tasks/testCO2"/>
</Events>
```

Internally, the driver uses a simple form of time-stepping to advance through the (P,T) steps. This timestepping is handled independently of the more complicated time-stepping pattern used by physics `Solvers` and coordinated by the `EventManager`. In particular, in the XML file above, the `maxTime` parameter in the `Events` block is an event manager control, controlling when/if certain events occur. Once launched, the `PVTDriver` internally determines its own max time and timestep size using a combination of the input functions’ time coordinates and the requested number of loadsteps. It is therefore helpful to think of the driver as an instantaneous *event* (from the event manager’s point of view), but one which has a separate, internal clock.



## Parameters

The key XML parameters for the PVTDriver are summarized in the following table:

Name	Type	Default	Description
baseline	path	none	Baseline file
feedComposition	real64_array	required	Feed composition array [mol fraction]
fluid	string	required	Fluid to test
logLevel	integer	0	Log level
name	string	required	A name is required for any non-unique nodes
output	string	none	Output file
pressureControl	string	required	Function controlling pressure time history
steps	integer	required	Number of load steps to take
temperatureControl	string	required	Function controlling temperature time history

## Output Format

The `output` key is used to identify a file to which the results of the simulation are written. If this key is omitted, or the user specifies `output="none"`, file output will be suppressed. The file is a simple ASCII format with a brief header followed by test data:

```
# column 1 = time
# column 2 = pressure
# column 3 = temperature
# column 4 = density
# columns 5-6 = phase fractions
# columns 7-8 = phase densities
# columns 9-10 = phase viscosities
0.0000e+00 1.0000e+06 3.5000e+02 1.5581e+01 1.0000e+00 4.1138e-11 1.5581e+01 1.
↪0033e+03 1.7476e-05 9.9525e-04
2.0408e-02 2.0000e+06 3.5000e+02 3.2165e+01 1.0000e+00 4.1359e-11 3.2165e+01 1.
↪0050e+03 1.7601e-05 9.9525e-04
4.0816e-02 3.0000e+06 3.5000e+02 4.9901e+01 1.0000e+00 4.1563e-11 4.9901e+01 1.
↪0066e+03 1.7778e-05 9.9525e-04
...
```

Note that the number of columns will depend on how many phases are present. In this case, we have a two-phase, two-component mixture. The total density is reported in column 4, while phase fractions, phase densities, and phase viscosities are reported in subsequent columns. The phase order will match the one defined in the input XML (here, the `co2`-rich phase followed by the water-rich phase). This file can be readily plotted using any number of plotting tools. Each row corresponds to one timestep of the driver, starting from initial conditions in the first row.

## Unit Testing

The development team also uses the PVTDriver to perform unit testing on the various fluid models within GEOSX. The optional argument `baseline` can be used to point to a previous output file that has been validated (e.g. against experimental benchmarks or reference codes). If such a file is specified, the driver will perform a testing run and then compare the new results against the baseline. In this way, any regressions in the fluid models can be quickly identified.

Developers of new models are encouraged to add their own baselines to `src/coreComponents/constitutive/unitTests`. Adding additional tests is straightforward:

1. Create a new xml file for your test in `src/coreComponents/constitutive/unitTests` or (easier) add extra blocks to the existing XML at `src/coreComponents/constitutive/unitTests/testPVT.xml`. For new XMLs, we suggest using the naming convention `testPVT_myTest.xml`, so that all tests will be grouped together alphabetically. Set the output file to `testPVT_myTest.txt`, and run your test. Validate the results however is appropriate. If you have reference data available for this validation, we suggest archiving it in the `testPVT_data/` subdirectory, with a description of the source and formatting in the file header. Several reference datasets are included already as examples. This directory is also a convenient place to store auxiliary input files like PVT tables.

2. This output file will now become your new baseline. Replace the output key with `baseline` so that the driver can read in your file as a baseline for comparison. Make sure there is no remaining output key, or set `output=none`, to suppress further file output. While you can certainly write a new output for debugging purposes, during our automated unit tests we prefer to suppress file output. Re-run the driver to confirm that the comparison test passes.

3. Modify `src/coreComponents/constitutive/unitTests/CMakeLists.txt` to enable your new test in the unit test suite. In particular, you will need to add your new XML file to the existing list in the `gtest_pvt_xmls` variable. Note that if you simply added your test to the existing `testPVT.xml` file, no changes are needed.

```
set( gtest_pvt_xmls
    testPVT.xml
    testPVT_myTest.xml
)
```

4. Run `make` in your build directory to make sure the CMake syntax is correct

5. Run `ctest -V -R PVT` to run the PVT unit tests. Confirm your test is included and passes properly.

If you run into troubles, do not hesitate to contact the development team for help.

### 5.4.3 Relative Permeability Models

There are two ways to specify relative permeabilities in GEOSX. The user can either select an analytical relative permeability model (e.g., Brooks-Corey or Van Genuchten) or provide relative permeability tables. This is explained in the following sections.

#### Brooks-Corey relative permeability model

##### Overview

The following paragraphs explain how the Brooks-Corey model is used to compute the phase relative permeabilities as a function of volume fraction (i.e., saturation) with the expression:

$$k_{r\ell} = k_{r\ell,max} S_{\ell,scaled}^{\lambda_{\ell}}$$

where the scaled volume fraction of phase  $\ell$  is computed as:

$$S_{\ell,scaled} = \frac{S_{\ell} - S_{\ell,min}}{1 - \sum_{m=1}^{n_p} S_{m,min}}.$$

The minimum phase volume fractions  $S_{\ell,min}$  are model parameters specified by the user.

## Parameters

The relative permeability constitutive model is listed in the `<Constitutive>` block of the input XML file. The relative permeability model must be assigned a unique name via `name` attribute. This name is used to assign the model to regions of the physical domain via a `materialList` attribute of the `<ElementRegions>` node.

The following attributes are supported:

Name	Type	De- fault	Description
name	string	re- quired	A name is required for any non-unique nodes
phaseMinVolumeFrac- tion	real64_array	{0}	Minimum volume fraction value for each phase
phaseNames	string_array	re- quired	List of fluid phases
phaseRelPermExpo- nent	real64_array	{1}	Minimum relative permeability power law exponent for each phase
phaseRelPermMax- Value	real64_array	{0}	Maximum relative permeability value for each phase

Below are some comments on the model parameters.

- `phaseNames` - The number of phases can be either two or three. Note that for three-phase flow, this model does not apply a special treatment to the intermediate phase relative permeability (no Stone or Baker interpolation). Supported phase names are:

Value	Phase
oil	Oil phase
gas	Gas phase
water	Water phase

- `phaseMinVolFraction` - The list of minimum volume fractions  $S_{\ell,min}$  for each phase is specified in the same order as in `phaseNames`. Below this volume fraction, the phase is assumed to be immobile.
- `phaseRelPermExponent` - The list of exponents  $\lambda_{\ell}$  for each phase is specified in the same order as in `phaseNames`.
- `phaseMaxValue` - The list of maximum values  $k_{r\ell,max}$  for each phase is specified in the same order as in `phaseNames`.

## Examples

For a two-phase water-gas system (for instance in the CO2-brine fluid model), a typical relative permeability input looks like:

```
<Constitutive>
...
<BrooksCoreyRelativePermeability
  name="relPerm"
  phaseNames="{ water, gas }"
  phaseMinVolumeFraction="{ 0.02, 0.015 }"
  phaseRelPermExponent="{ 2, 2.5 }"
  phaseRelPermMaxValue="{ 0.8, 1.0 }"/>
```

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```
...
</Constitutive>
```

For a three-phase oil-water-gas system (for instance in the Black-Oil fluid model), a typical relative permeability input looks like:

```
<Constitutive>
...
  <BrooksCoreyRelativePermeability
    name="relPerm"
    phaseNames="{ water, oil, gas }"
    phaseMinVolumeFraction="{ 0.02, 0.1, 0.015 }"
    phaseRelPermExponent="{ 2, 2, 2.5 }"
    phaseRelPermMaxValue="{ 0.8, 1.0, 1.0 }"/>
...
</Constitutive>
```

## Three-phase relative permeability model

### Overview

For the simulation of three-phase flow in porous media, it is common to use a specific treatment (i.e., different from the typical two-phase procedure) to evaluate the oil relative permeability. Specifically, the three-phase oil relative permeability is obtained by interpolation of oil-water and oil-gas experimental data measured independently in two-phase displacements.

Let  $k_{rw,wo}$  and  $k_{ro,wo}$  be the water-oil two-phase relative permeabilities for the water phase and the oil phase, respectively. Let  $k_{rg,go}$  and  $k_{ro,go}$  be the oil-gas two-phase relative permeabilities for the gas phase and the oil phase, respectively. In the current implementation, the two-phase relative permeability data is computed analytically using the *Brooks-Corey relative permeability model*.

The water and gas three-phase relative permeabilities are simply given by two-phase data and only depend on  $S_w$  and  $S_g$ , respectively. That is,

$$k_{rw,wog}(S_w) = k_{rw,wo}(S_w),$$

$$k_{rg,wog}(S_g) = k_{rg,go}(S_g).$$

The oil three-phase relative permeability is obtained using a variant of the saturation-weighted interpolation procedure initially proposed by [Baker](#). Specifically, we compute:

$$k_{ro,wog}(S_w, S_g) = \frac{(S_w - S_{w,min})k_{ro,wo}(S_w) + S_g k_{rg,go}(S_g)}{(S_w - S_{w,min}) + S_g}.$$

This procedure provides a simple but effective formula avoiding the problems associated with the other interpolation methods (negative values).

### Parameters

The relative permeability constitutive model is listed in the `<Constitutive>` block of the input XML file. The relative permeability model must be assigned a unique name via `name` attribute. This name is used to assign the model to regions of the physical domain via a `materialList` attribute of the `<ElementRegion>` node.

The following attributes are supported:

Name	Type	Default	Description
gasOilRelPermExponent	real64_array	{1}	Rel perm power law exponent for the pair (gas phase, oil phase) at residual water saturation The expected format is “{ gasExp, oilExp }”, in that order
gasOilRelPermMaxValue	real64_array	{0}	Maximum rel perm value for the pair (gas phase, oil phase) at residual water saturation The expected format is “{ gasMax, oilMax }”, in that order
name	string	required	A name is required for any non-unique nodes
phaseMinVolumeFraction	real64_array	{0}	Minimum volume fraction value for each phase
phaseNames	string_array	required	List of fluid phases
waterOilRelPermExponent	real64_array	{1}	Rel perm power law exponent for the pair (water phase, oil phase) at residual gas saturation The expected format is “{ waterExp, oilExp }”, in that order
waterOilRelPermMaxValue	real64_array	{0}	Maximum rel perm value for the pair (water phase, oil phase) at residual gas saturation The expected format is “{ waterMax, oilMax }”, in that order

Below are some comments on the model parameters.

- `phaseNames` - The number of phases should be 3. Supported phase names are:

Value	Phase
oil	Oil phase
gas	Gas phase
water	Water phase

- `phaseMinVolFraction` - The list of minimum volume fractions  $S_{\ell,min}$  for each phase is specified in the same order as in `phaseNames`. Below this volume fraction, the phase is assumed to be immobile.
- `waterOilRelPermExponent` - The list of exponents  $\lambda_{\ell,wo}$  for the two-phase water-oil relative permeability data, with the water exponent first and the oil exponent next. These exponents are then used to compute  $k_{r\ell,wo}$  in the *Brooks-Corey relative permeability model*.
- `waterOilRelPermMaxValue` - The list of maximum values  $k_{r\ell,wo,max}$  for the two-phase water-oil relative permeability data, with the water max value first and the oil max value next. These exponents are then used to compute  $k_{r\ell,wo}$  in the *Brooks-Corey relative permeability model*.
- `gasOilRelPermExponent` - The list of exponents  $\lambda_{\ell,go}$  for the two-phase gas-oil relative permeability data, with the gas exponent first and the oil exponent next. These exponents are then used to compute  $k_{r\ell,go}$  in the *Brooks-Corey relative permeability model*.
- `gasOilRelPermMaxValue` - The list of maximum values  $k_{r\ell,go,max}$  for the two-phase gas-oil relative permeability data, with the gas max value first and the oil max value next. These exponents are then used to compute  $k_{r\ell,go}$  in the *Brooks-Corey relative permeability model*.

### Example

```
<Constitutive>
...
<BrooksCoreyBakerRelativePermeability name="relperm"
    phaseNames="{oil, gas, water}"
    phaseMinVolumeFraction="{0.05, 0.05, 0.05}"
    waterOilRelPermExponent="{2.5, 1.5}"
    waterOilRelPermMaxValue="{0.8, 0.9}"
    gasOilRelPermExponent="{3, 3}"
    gasOilRelPermMaxValue="{0.4, 0.9}"/>
...
</Constitutive>
```

## Table relative permeability

### Overview

The user can specify the relative permeabilities using tables describing a piecewise-linear relative permeability function of volume fraction (i.e., saturation) for each phase. Depending on the number of fluid phases, this model is used as follows:

- For two-phase flow, the user must specify two relative permeability tables, that is, one for the wetting-phase relative permeability, and one for the non-wetting phase relative permeability. During the simulation, the relative permeabilities are then obtained by interpolating in the tables as a function of phase volume fraction.
- For three-phase flow, following standard reservoir simulation practice, the user must specify four relative permeability tables. Specifically, two relative permeability tables are required for the pair wetting-phase–intermediate phase (typically, water-oil), and two relative permeability tables are required for the pair non-wetting-phase–intermediate phase (typically, gas-oil). During the simulation, the relative permeabilities of the wetting

and non-wetting phases are computed by interpolating in the tables as a function of their own phase volume fraction. The intermediate phase relative permeability is obtained by interpolating the two-phase relative permeabilities using the Baker interpolation procedure.

## Parameters

The relative permeability constitutive model is listed in the `<Constitutive>` block of the input XML file. The relative permeability model must be assigned a unique name via `name` attribute. This name is used to assign the model to regions of the physical domain via a `materialList` attribute of the `<ElementRegions>` node.

The following attributes are supported:

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes
nonWettingIntermediateRelPermTableNames	string_array	{}	<p>List of relative permeability tables for the pair (non-wetting phase, intermediate phase)</p> <p>The expected format is “{ nonWetting-PhaseRelPermTable-Name, intermediatePhaseRelPermTable-Name }”, in that order</p> <p>Note that this input is only used for three-phase flow.</p> <p>If you want to do a two-phase simulation, please use instead wettingNonWettingRelPermTableNames to specify the table names</p>
phaseNames	string_array	required	List of fluid phases
wettingIntermediateRelPermTableNames	string_array	{}	<p>List of relative permeability tables for the pair (wetting phase, intermediate phase)</p> <p>The expected format is “{ wetting-PhaseRelPermTable-Name, intermediatePhaseRelPermTable-Name }”, in that order</p> <p>Note that this input is only used for three-phase flow.</p> <p>If you want to do a two-phase simulation, please use instead wettingNonWettingRelPermTableNames to specify the table names</p>
wettingNonWettingRelPermTableNames	string_array	{}	List of relative permeability tables for the pair (wetting phase, non-wetting phase)
370			<p><b>Chapter 5. User Guide</b></p> <p>The expected format is “{ wetting-</p>



Below are some comments on the model parameters.

- `phaseNames` - The number of phases can be either two or three. For three-phase flow, this model applies a Baker interpolation to the intermediate phase relative permeability. Supported phase names are:

Value	Phase
oil	Oil phase
gas	Gas phase
water	Water phase

- `wettingNonWettingRelPermTableNames` - The list of relative permeability table names for two-phase systems, starting with the name of the wetting-phase relative permeability table, followed by the name of the non-wetting phase relative permeability table. Note that this keyword is only valid for two-phase systems, and is not allowed for three-phase systems (for which the user must specify instead `wettingIntermediateRelPermTableNames` and `nonWettingIntermediateRelPermTableNames`).
- `wettingIntermediateRelPermTableNames` - The list of relative permeability table names for the pair wetting-phase–intermediate-phase, starting with the name of the wetting-phase relative permeability table, and continuing with the name of the intermediate phase relative permeability table. Note that this keyword is only valid for three-phase systems, and is not allowed for two-phase systems (for which the user must specify instead `wettingNonWettingRelPermTableNames`).
- `nonWettingIntermediateRelPermTableNames` - The list of relative permeability table names for the pair non-wetting-phase–intermediate-phase, starting with the name of the non-wetting-phase relative permeability table, and continuing with the name of the intermediate phase relative permeability table. Note that this keyword is only valid for three-phase systems, and is not allowed for two-phase systems (for which the user must specify instead `wettingNonWettingRelPermTableNames`).

---

**Note:** We remind the user that the relative permeability must be a strictly increasing function of phase volume fraction. GEOSX throws an error when this condition is not satisfied.

---

## Examples

For a two-phase water-gas system (for instance in the CO<sub>2</sub>-brine fluid model), a typical relative permeability input looks like:

```
<Constitutive>
...
<TableRelativePermeability
  name="relPerm"
  phaseNames="{ water, gas }"
  wettingNonWettingRelPermTableNames="{ waterRelativePermeabilityTable,
↪gasRelativePermeabilityTable }"/>
...
</Constitutive>
```

---

**Note:** The name of the wetting-phase relative permeability table must be specified before the name of the non-wetting phase relative permeability table.

---

For a three-phase oil-water-gas system (for instance in the Black-Oil fluid model), a typical relative permeability input looks like:

```
<Constitutive>
...
  <TableRelativePermeability
    name="relPerm"
    phaseNames="{ water, oil, gas }"
    wettingIntermediateRelPermTableNames="{ waterRelativePermeabilityTable,
    ↪oilRelativePermeabilityTableForWO }"
    nonWettingIntermediateRelPermTableNames="{ gasRelativePermeabilityTable,
    ↪oilRelativePermeabilityTableForGO }"/>
...
</Constitutive>
```

**Note:** For the wetting-phase–intermediate-phase pair, the name of the wetting-phase relative permeability table must be specified first. For the non-wetting-phase–intermediate-phase pair, the name of the non-wetting-phase relative permeability table must be specified first. If the results look incoherent, this is something to double-check.

The tables mentioned above by name must be defined in the <Functions> block of the XML file using the <TableFunction> keyword.

## 5.4.4 Capillary Pressure Models

There are two ways to specify capillary pressures in GEOSX. The user can either select an analytical capillary pressure model (e.g., Brooks-Corey or Van Genuchten) or provide capillary pressure tables. This is explained in the following sections.

### Brooks-Corey capillary pressure model

#### Overview

In GEOSX, the oil-phase pressure is assumed to be the primary pressure. The following paragraphs explain how the Brooks-Corey capillary pressure model is used to compute the water-phase and gas-phase pressures as:

$$p_w = p_o - P_{c,w}(S_w),$$

and

$$p_g = p_o + P_{c,g}(S_g).$$

In the Brooks-Corey model, the water-phase capillary pressure is computed as a function of the water-phase volume fraction with the following expression:

$$P_{c,w}(S_w) = p_{e,w} S_{w,scaled}^{-1/\lambda_w},$$

where the scaled water-phase volume fraction is computed as:

$$S_{w,scaled} = \frac{S_w - S_{w,min}}{1 - S_{w,min} - S_{o,min} - S_{g,min}}.$$

The gas capillary pressure is computed analogously.

## Parameters

The capillary pressure constitutive model is listed in the `<Constitutive>` block of the input XML file. The capillary pressure model must be assigned a unique name via `name` attribute. This name is used to assign the model to regions of the physical domain via a `materialList` attribute of the `<ElementRegions>` node.

The following attributes are supported:

Name	Type	De-fault	Description
capPressureEpsilon	real64	1e-06	Wetting-phase saturation at which the max cap. pressure is attained; used to avoid infinite cap. pressure values for saturations close to zero
name	string	required	A name is required for any non-unique nodes
phaseCapPressureExponentInv	real64_array	{2}	Inverse of capillary power law exponent for each phase
phaseEntryPressure	real64_array	{1}	Entry pressure value for each phase
phaseMinVolumeFraction	real64_array	{0}	Minimum volume fraction value for each phase
phaseNames	string_array	required	List of fluid phases

Below are some comments on the model parameters:

- `phaseNames` - The number of phases can be either 2 or 3. The names entered for this attribute should match the phase names specified in the relative permeability block, either in *Brooks-Corey relative permeability model* or in *Three-phase relative permeability model*. The capillary pressure model assumes that oil is always present. Supported phase names are:

Value	Phase
oil	Oil phase
gas	Gas phase
water	Water phase

- `phaseMinVolFraction` - The list of minimum volume fractions  $S_{\ell,min}$  for each phase is specified in the same order as in `phaseNames`. Below this volume fraction, the phase is assumed to be immobile. The values entered for this attribute have to match those of the same attribute in the relative permeability block.
- `phaseCapPressureExponentInv` - The list of exponents  $\lambda_{\ell}$  for each phase is specified in the same order as in `phaseNames`. The parameter corresponding to the oil phase is currently not used.
- `phaseEntryPressure` - The list of entry pressures  $p_{e,\ell}$  for each phase is specified in the same order as in `phaseNames`. The parameter corresponding to the oil phase is currently not used.
- `capPressureEpsilon` - This parameter is used for both the water-phase and gas-phase capillary pressure. To avoid extremely large, or infinite, capillary pressure values, we set  $P_{c,w}(S_w) := P_{c,w}(\epsilon)$  whenever  $S_w < \epsilon$ . The gas-phase capillary pressure is treated analogously.

## Example

```
<Constitutive>
...
```

(continues on next page)

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```

<BrooksCoreyCapillaryPressure name="capPressure"
    phaseNames="{oil, gas}"
    phaseMinVolumeFraction="{0.01, 0.015}"
    phaseCapPressureExponentInv="{0, 6}"
    phaseEntryPressure="{0, 1e8}"
    capPressureEpsilon="1e-8"/>
...
</Constitutive>

```

## Van Genuchten capillary pressure model

### Overview

In GEOSX, the oil-phase pressure is assumed to be the primary pressure. The following paragraphs explain how the Van Genuchten capillary pressure model is used to compute the water-phase and gas-phase pressures as:

$$p_w = p_o - P_{c,w}(S_w),$$

and

$$p_g = p_o + P_{c,g}(S_g),$$

The Van Genuchten model computes the water-phase capillary pressure as a function of the water-phase volume fraction as:

$$P_c(S_w) = \alpha_w (S_{w,scaled}^{-1/m_w} - 1)^{(1-m_w)/2},$$

where the scaled water-phase volume fraction is computed as:

$$S_{w,scaled} = \frac{S_w - S_{w,min}}{1 - S_{w,min} - S_{o,min} - S_{g,min}}.$$

The gas-phase capillary pressure is computed analogously.

### Parameters

The capillary pressure constitutive model is listed in the `<Constitutive>` block of the input XML file. The capillary pressure model must be assigned a unique name via `name` attribute. This name is used to assign the model to regions of the physical domain via a `materialList` attribute of the `<ElementRegions>` node.

The following attributes are supported:

Name	Type	Default	Description
capPressureEpsilon	real64	1e-06	Saturation at which the extremum capillary pressure is attained; used to avoid infinite capillary pressure values for saturations close to 0 and 1
name	string	required	A name is required for any non-unique nodes
phaseCapPressureExponentInv	real64_array	{0.5}	Inverse of capillary power law exponent for each phase
phaseCapPressureMultiplier	real64_array	{1}	Entry pressure value for each phase
phaseMinVolumeFraction	real64_array	{0}	Minimum volume fraction value for each phase
phaseNames	string_array	required	List of fluid phases

Below are some comments on the model parameters:

- `phaseNames` - The number of phases can be either 2 or 3. The phase names entered for this attribute should match the phase names specified in the relative permeability block, either in *Brooks-Corey relative permeability model* or in *Three-phase relative permeability model*. The capillary model assumes that oil is always present. Supported phase names are:

Value	Phase
oil	Oil phase
gas	Gas phase
water	Water phase

- `phaseMinVolFraction` - The list of minimum volume fractions  $S_{\ell,min}$  for each phase is specified in the same order as in `phaseNames`. Below this volume fraction, the phase is assumed to be immobile. The values entered for this attribute have to match those of the same attribute in the relative permeability block.
- `phaseCapPressureExponentInv` - The list of exponents  $m_{\ell}$  for each phase is specified in the same order as in `phaseNames`. The parameter corresponding to the oil phase is not used.
- `phaseCapPressureMultiplier` - The list of multipliers  $\alpha_{\ell}$  for each phase is specified in the same order as in `phaseNames`. The parameter corresponding to the oil phase is not used.
- `capPressureEpsilon` - The parameter  $\epsilon$ . This parameter is used for both the water-phase and gas-phase capillary pressure. To avoid extremely large, or infinite, capillary pressure values, we set  $P_{c,w}(S_w) := P_{c,w}(\epsilon)$  whenever  $S_w < \epsilon$ . The gas-phase capillary pressure is treated analogously.

### Example

```
<Constitutive>
...
  <VanGenuchtenCapillaryPressure name="capPressure"
    phaseNames="{ water, oil }"
    phaseMinVolumeFraction="{ 0.1, 0.015 }"
    phaseCapPressureExponentInv="{ 0.55, 0 }"
    phaseCapPressureMultiplier="{ 1e6, 0 }"
    capPressureEpsilon="1e-7"/>
...
</Constitutive>
```

## Table capillary pressure

### Overview

The user can specify the capillary pressures using tables describing a piecewise-linear capillary pressure function of volume fraction (i.e., saturation) for each phase, except the reference phase for which capillary pressure is assumed to be zero. Depending on the number of fluid phases, this model is used as follows:

- For two-phase flow, the user must specify one capillary pressure table. During the simulation, the capillary pressure of the non-reference phase is computed by interpolating in the table as a function of the non-reference phase saturation.
- For three-phase flow, the user must specify two capillary pressure tables. One capillary pressure table is required for the pair wetting-phase–intermediate-phase (typically, water–oil), and one capillary pressure table is required for the pair non-wetting-phase–intermediate-phase (typically, gas–oil). During the simulation, the former is used to compute the wetting-phase capillary pressure as a function of the wetting-phase volume fraction and the latter is used to compute the non-wetting-phase capillary pressure as a function of the non-wetting-phase volume fraction. The intermediate phase is assumed to be the reference phase, and its capillary pressure is set to zero.

Below is a table summarizing the choice of reference pressure for the various phase combinations:

Phases present in the model	Reference phase
oil, water, gas	Oil phase
oil, water	Oil phase
oil, gas	Oil phase
water, gas	Gas phase

In all cases, the user-provided capillary pressure is used in GEOSX to compute the phase pressure using the formula:

$$P_c = p_{nw} - p_w.$$

where  $p_{nw}$  and  $p_w$  are respectively the non-wetting-phase and wetting-phase pressures.

### Parameters

The capillary pressure constitutive model is listed in the `<Constitutive>` block of the input XML file. The capillary pressure model must be assigned a unique name via `name` attribute. This name is used to assign the model to regions of the physical domain via a `materialList` attribute of the `<ElementRegions>` node.

The following attributes are supported:

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes
nonWettingIntermediateCapPressureTableName	string		<p>Capillary pressure table [Pa] for the pair (non-wetting phase, intermediate phase)</p> <p>Note that this input is only used for three-phase flow.</p> <p>If you want to do a two-phase simulation, please use instead wettingNonWettingCapPressureTableName to specify the table names</p>
phaseNames	string_array	required	List of fluid phases
wettingIntermediateCapPressureTableName	string		<p>Capillary pressure table [Pa] for the pair (wetting phase, intermediate phase)</p> <p>Note that this input is only used for three-phase flow.</p> <p>If you want to do a two-phase simulation, please use instead wettingNonWettingCapPressureTableName to specify the table names</p>
wettingNonWettingCapPressureTableName	string		<p>Capillary pressure table [Pa] for the pair (wetting phase, non-wetting phase)</p> <p>Note that this input is only used for two-phase flow.</p> <p>If you want to do a three-phase simulation, please use instead wettingIntermediateCapPressureTableName and nonWettingIntermediateCapPressureTableName to specify the table names</p>

Below are some comments on the model parameters.

- `phaseNames` - The number of phases can be either two or three. Supported phase names are:

Value	Phase
oil	Oil phase
gas	Gas phase
water	Water phase

- `wettingNonWettingCapPressureTableName` - The name of the capillary pressure table for two-phase systems. Note that this keyword is only valid for two-phase systems, and is not allowed for three-phase systems (for which the user must specify instead `wettingIntermediateCapPressureTableName` and `nonWettingIntermediateCapPressureTableName`). This capillary pressure must be a strictly decreasing function of the water-phase volume fraction (for oil-water systems and gas-water systems), or a strictly increasing function of the gas-phase volume fraction (for oil-gas systems).
- `wettingIntermediateCapPressureTableName` - The name of the capillary pressure table for the pair wetting-phase–intermediate-phase. This capillary pressure is applied to the wetting phase, as a function of the wetting-phase volume fraction. Note that this keyword is only valid for three-phase systems, and is not allowed for two-phase systems (for which the user must specify instead `wettingNonWettingCapPressureTableName`). This capillary pressure must be a strictly decreasing function of the wetting-phase volume fraction.
- `nonWettingIntermediateCapPressureTableName` - The name of the capillary pressure table for the pair non-wetting-phase–intermediate-phase. Note that this keyword is only valid for three-phase systems, and is not allowed for two-phase systems (for which the user must specify instead `wettingNonWettingCapPressureTableName`). This capillary pressure must be a strictly increasing function of the non-wetting-phase volume fraction.

## Examples

For a two-phase water-gas system (for instance in the CO<sub>2</sub>-brine fluid model), a typical capillary pressure input looks like:

```
<Constitutive>
...
<TableCapillaryPressure
  name="capPressure"
  phaseNames="{ water, gas }"
  wettingNonWettingCapPressureTableNames="waterCapillaryPressureTable"/>
...
</Constitutive>
```

For a three-phase oil-water-gas system (for instance in the Black-Oil fluid model), a typical capillary pressure input looks like:

```
<Constitutive>
...
<TableCapillaryPressure
  name="capPressure"
  phaseNames="{ water, oil, gas }"
  wettingIntermediateCapPressureTableName="waterCapillaryPressureTable"
  nonWettingIntermediateCapPressureTableName="gasCapillaryPressureTable"/>
...
</Constitutive>
```



The tables mentioned above by name must be defined in the `<Functions>` block of the XML file using the `<TableFunction>` keyword.

## 5.4.5 Porosity models

### Pressure dependent porosity

#### Overview

This model assumes a simple exponential law for the porosity as function of pressure, i.e.

$$\phi = \phi_{ref} \exp(c \cdot (p - p_{ref}))$$

where  $\phi_{ref}$  is the reference porosity at reference pressure,  $p_{ref}$ ,  $p$  is the pressure and,  $c$  is the compressibility.

#### Parameters

The following attributes are supported:

Name	Type	Default	Description
compressibility	real64	required	Solid compressibility
defaultReferencePorosity	real64	required	Default value of the reference porosity
name	string	required	A name is required for any non-unique nodes
referencePressure	real64	required	Reference pressure for solid compressibility

#### Example

```
<Constitutive>
...
  <PressurePorosity name="rockPorosity"
    referencePressure="1.0e27"
    defaultReferencePorosity="0.3"
    compressibility="1.0e-9"/>
...
</Constitutive>
```

### Biot Porosity Model

#### Overview

According to the poroelasticity theory, the porosity (pore volume),  $\phi$ , can be computed as

$$\phi = \phi_{ref} + \alpha(\epsilon_v - \epsilon_{v,ref}) + (p - p_{ref})/N.$$

Here,  $\phi_{ref}$  is the porosity at a reference state with pressure  $p_{ref}$  and volumetric strain  $\epsilon_{v,ref}$ . Additionally,  $\alpha$  is the Biot coefficient,  $\epsilon_v$  is the volumetric strain,  $p$  is the fluid pressure and  $N = \frac{K_s}{\alpha - \phi_{ref}}$ , where  $K_s$  is the grain bulk modulus.

## Parameters

The Biot Porosity Model can be called in the `<Constitutive>` block of the input XML file. This porosity model must be assigned a unique name via the `name` attribute. This name is used to assign the model to regions of the physical domain via a `materialList` attribute in the `<ElementRegions>` block.

The following attributes are supported:

Name	Type	Default	Description
defaultReferencePorosity	real64	required	Default value of the reference porosity
grainBulkModulus	real64	required	Grain bulk modulus
name	string	required	A name is required for any non-unique nodes

## Example

```
<Constitutive>
...
  <BiotPorosity name="rockPorosity"
    grainBulkModulus="1.0e27"
    defaultReferencePorosity="0.3"/>
...
</Constitutive>
```

## 5.4.6 Permeability models

### Constant Permeability Model

#### Overview

This model is used to define a diagonal permeability tensor that does not depend on any primary variable.

## Parameters

The following attributes are supported:

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes
permeabilityComponents	R1Tensor	required	xx, yy and zz components of a diagonal permeability tensor.

## Example

```
<Constitutive>
...
  <ConstantPermeability name="matrixPerm"
    permeabilityComponents="{1.0e-12, 1.0e-12, 1.0e-12}"/>
...
</Constitutive>
```

## Kozeny-Carman Permeability Model

### Overview

In the Kozeny-Carman model (see [ref](#)), the permeability of a porous medium is governed by several key parameters, including porosity, grain size, and grain shape:

$$k = \frac{(s_{\epsilon} D_p)^2 \phi^3}{150(1 - \phi)^2}$$

where  $s_{\epsilon}$  is the sphericity of the particles,  $D_p$  is the particle diameter,  $\phi$  is the porosity of the porous medium.

### Parameters

The Kozeny-Carman Permeability Model can be called in the `<Constitutive>` block of the input XML file. This permeability model must be assigned a unique name via the `name` attribute. This name is used to assign the model to regions of the physical domain via a `materialList` attribute in the `<ElementRegions>` block.

The following attributes are supported:

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes
particleDiameter	real64	required	Diameter of the spherical particles.
sphericity	real64	required	Sphericity of the particles.

### Example

```
<Constitutive>
...
  <CarmanKozenyPermeability name="matrixPerm"
    particleDiameter="0.0002"
    sphericity="1.0"/>
...
</Constitutive>
```

## Parallel Plates Permeability Model

### Overview

The parallel plates permeability model defines the relationship between the hydraulic fracture aperture and its corresponding permeability following the classic lubrication model (Witherspoon et al. ). In this model, the two fracture walls are assumed to be smooth and parallel to each other and separated by a uniform aperture.

$$k = \frac{a^3}{12}$$

where  $a$  denotes the hydraulic fracture aperture.

Remark:  $k$ , dimensionally, is not a permeability (as it is expressed in  $m^3$ ).

## Parameters

The Parallel Plates Permeability Model can be called in the `<Constitutive>` block of the input XML file. This permeability model must be assigned a unique name via the `name` attribute. This name is used to assign the model to regions of the physical domain via a `materialList` attribute in the `<ElementRegions>` block.

The following attributes are supported:

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes

## Slip Dependent Permeability Model

### Overview

The slip dependent permeability model defines the relationship between the relative shear displacement and fracture permeability. In this model, fractures/faults are represented as slip interfaces.

$$k = k_i \left[ (M_{mult} - 1) \tanh \left( 3 \frac{U_s}{U_{threshold}} \right) + 1 \right]$$

where  $k_i$  is the initial fracture permeability;  $M_{mult}$  is the maximum permeability multiplier;  $U_s$  is the relative shear displacement;  $U_{threshold}$  is the slip threshold.

## Parameters

The Slip Dependent Permeability model can be called in the `<Constitutive>` block of the input XML file. This permeability model must be assigned a unique name via the `name` attribute. This name is used to assign the model to regions of the physical domain via a `permeabilityModelName` attribute in the `<CompressibleSolidSlipDependentPermeability>` block.

The following attributes are supported:

Name	Type	Default	Description
initialPermeability	R1Tensor	required	initial permeability of the fracture.
maxPermMultiplier	real64	required	Maximum permeability multiplier.
name	string	required	A name is required for any non-unique nodes
shearDispThreshold	real64	required	Threshold of shear displacement.

## Example

```
<Constitutive>
...
  <SlipDependentPermeability
    name="fracturePerm"
    shearDispThreshold="0.005"
    maxPermMultiplier="1000.0"
    initialPermeability="{1e-15, 1e-15, 1e-15}"/>
...
</Constitutive>
```

### 5.4.7 Porous Solids

#### Overview

Simulation of fluid flow in porous media and of poromechanics, requires to define, along with fluid properties, the hydrodynamical properties of the solid matrix. Thus, for porous media flow and poromechanical simulation in GEOSX, two types of composite constitutive models can be defined to specify the characteristics of a porous material: (1) a *CompressibleSolid* model, used for flow-only simulations and which assumes that all poromechanical effects can be represented by the pressure dependency of the porosity; (2) a *PorousSolid* model which, instead, allows to couple any solid model with a *BiotPorosity* model and to include permeability's dependence on the mechanical response.

Both these composite models require the names of the solid, porosity and permeability models that, combined, define the porous material. The following sections outline how these models can be defined in the Constitutive block of the xml input files and which type of submodels they allow for.

#### CompressibleSolid

This composite constitutive model requires to define a *NullModel* as solid model (since no mechanical properties are used), a *PressurePorosity* model and any type of *Permeability* model.

To define this composite model the keyword *CompressibleSolid* has to be appended to the name of the permeability model of choice, as shown in the following example for the *ConstantPermeability* model.

```
<Constitutive>
  <CompressibleSolidConstantPermeability name="porousRock"
    solidModelName="nullSolid"
    porosityModelName="rockPorosity"
    permeabilityModelName="rockPermeability"/>

  <NullModel name="nullSolid"/>

  <PressurePorosity name="rockPorosity"
    referencePressure="1.0e27"
    defaultReferencePorosity="0.3"
    compressibility="1.0e-9"/>

  <ConstantPermeability name="rockPermeability"
    permeabilityComponents="{ 1.0e-4, 1.0e-4, 1.0e-4 }"/>

</Constitutive>
```

#### PorousSolid

To run poromechanical problems, the total stress is decomposed into an “effective stress” (driven by mechanical deformations) and a pore fluid pressure component, following the [Biot theory of poroelasticity](#). For single-phase flow, or multiphase problems with no capillarity, this decomposition reads

$$\sigma_{ij} = \sigma'_{ij} - bp\delta_{ij}$$

where  $\sigma_{ij}$  is the  $ij$  component of the total stress tensor,  $\sigma'_{ij}$  is the  $ij$  component of the effective (Cauchy) stress tensor,  $b$  is Biot's coefficient,  $p$  is fluid pressure, and  $\delta$  is the Kronecker delta.

The *PorousSolid* models simply append the keyword *Porous* in front of the solid model they contain, e.g., *PorousElasticIsotropic*, *PorousDruckerPrager*, and so on. Additionally, they require to define a *BiotPorosity* model and a *ConstantPermeability* model. For example, a Poroelastic material with a certain permeability can be defined as

```

<Constitutive>
  <PorousElasticIsotropic name="porousRock"
    porosityModelName="rockPorosity"
    solidModelName="rockSkeleton"
    permeabilityModelName="rockPermeability"/>

  <ElasticIsotropic name="rockSkeleton"
    defaultDensity="0"
    defaultYoungModulus="1.0e4"
    defaultPoissonRatio="0.2"/>

  <BiotPorosity name="rockPorosity"
    grainBulkModulus="1.0e27"
    defaultReferencePorosity="0.3"/>

  <ConstantPermeability name="rockPermeability"
    permeabilityComponents="{ 1.0e-4, 1.0e-4, 1.0e-4 }"/>
</Constitutive>

```

Note that any of the previously described solid models is used by the *PorousSolid* model to compute the effective stress, leading to either poro-elastic, poro-plastic, or poro-damage behavior depending on the specific model chosen.

In an input XML file, constitutive models are listed in the `<Constitutive>` block. Each parameterized model has its own XML tag, and each must be assigned a unique name via the `name` attribute. Names are used to assign models to regions of the physical domain via the `materialList` attribute of the `<ElementRegion>` node (see [Element: ElementRegions](#)). In some cases, physics solvers must also be assigned specific constitutive models to use (see [Physics Solvers](#)).

A typical `<Constitutive>` and `<ElementRegions>` block will look like:

```

<Problem>

  <Constitutive>

    <!-- Define a compressible, single-phase fluid called "water"-->
    <CompressibleSinglePhaseFluid
      name="water"
      referencePressure="2.125e6"
      referenceDensity="1000"
      compressibility="1e-19"
      referenceViscosity="0.001"
      viscosibility="0.0"/>

  </Constitutive>

  <ElementRegions>

    <!--Add water to the material list for region 1-->
    <ElementRegion
      name="region1"
      cellBlocks="cellBlock1"
      materialList="water"/>

  </ElementRegions>

  ... remainder of problem definition here ...

</Problem>

```

## 5.5 Initial and Boundary Conditions

### 5.5.1 Hydrostatic Equilibrium Initial Condition

#### Overview

The user can request an initialization procedure enforcing a hydrostatic equilibrium for flow simulations and for coupled flow and mechanics simulations. The hydrostatic initialization is done by placing one or more **HydrostaticEquilibrium** tag(s) in the **FieldSpecifications** block of the XML input file. This initialization procedure is described below in the context of single-phase and compositional multiphase flow. At the end of this document, we compare the hydrostatic equilibrium method to another initialization method, based on the input of x-y-z tables.

#### Single-phase flow parameters

For single-phase flow, the **HydrostaticEquilibrium** initialization procedure requires the following user input parameters:

- **datumElevation**: the elevation (in meters) at which the datum pressure is enforced. The user must ensure that the datum elevation is within the elevation range defined by the input mesh. GEOSX issues a warning if this is not the case.
- **datumPressure**: the pressure value (in Pascal) enforced by GEOSX at the datum elevation.
- **objectPath**: the path defining the groups on which the hydrostatic equilibrium is computed. We recommend using **ElementRegions** to apply the hydrostatic equilibrium to all the cells in the mesh. Alternatively, the format **ElementRegions/NameOfRegion/NameOfCellBlock** can be used to select only a cell block on which the hydrostatic equilibrium is computed.

---

**Note:** In GEOSX, the z-axis is positive going upward, this is why the attributes listed in this page are expressed as a function of elevation, not depth.

---

Using these parameters and the pressure-density constitutive relationship, GEOSX uses a fixed-point iteration scheme to populate a table of hydrostatic pressures as a function of elevation. The fixed-point iteration scheme uses two optional attributes: **equilibriumTolerance**, the absolute tolerance to declare that the algorithm has converged, and **maxNumberOfEquilibrationTolerance**, the maximum number of iterations for a given elevation in the fixed point iteration scheme.

In addition, the elevation spacing of the hydrostatic pressure table is set with the optional **elevationIncrementInHydrostaticPressureTable** parameter (in meters), whose default value is 0.6096 meters. Then, once the table is fully constructed, the hydrostatic pressure in each cell is obtained by interpolating in the hydrostatic pressure table using the elevation at the center of the cell.

---

**Note:** The initialization algorithm assumes that the **gravityVector** (defined in the **Solvers** XML tag) is aligned with the z-axis. If this is not the case, GEOSX terminates the simulation when the **HydrostaticEquilibrium** tag is detected in the XML file.

---

#### Compositional multiphase flow parameters

For compositional multiphase flow, the **HydrostaticEquilibrium** initialization procedure follows the same logic but requires more input parameters. In addition to the required **datumElevation**, **datumPressure**, and **objectPath** parameters listed above, the user must specify:

- `componentNames`: the names of the components present in the fluid model. This field is used to make sure that the components provided to **HydrostaticEquilibrium** are consistent with the components listed in the fluid model of the **Constitutive** block.
- `componentFractionVsElevationTableNames`: the names of  $n_c$  tables (where  $n_c$  is the number of components) specifying the component fractions as a function of elevation. There must be one table name per component, and the table names must be listed in the same order as the components in `componentNames`.
- `temperatureVsElevationTableName`: the names of the table specifying the temperature (in Kelvin) as a function of elevation.
- `initialPhaseName`: the name of the phase initially saturating the domain. The other phases are assumed to be at residual saturation at the beginning of the simulation.

These parameters are used with the fluid density model (depending for compositional flow on pressure, component fractions, and in some cases, temperature) to populate the hydrostatic pressure table, and later initialize the pressure in each cell.

---

**Note:** The current initialization algorithm has an important limitation and does not support initial phase contacts (e.g., water-oil, gas-oil, or water-gas contacts). The implementation assumes only one mobile phase in the initial system, identified by the `initialPhaseName` attribute. The other phases are assumed at residual saturation. As a result, the system may not be at equilibrium if there is initially more than one mobile phase in the system (for instance if the domain is saturated with gas at the top, and water at the bottom, for instance).

---



---

**Note:** As in the single-phase flow case, GEOSX terminates the simulation if **HydrostaticEquilibrium** tag is present in an XML file defining a `gravityVector` not aligned with the z-axis.

---

The full list of parameters is provided below:



Name	Type	Default	Description
bcApplicationTableName	string		Name of table that specifies the on/off application of the boundary condition.
beginTime	real64	-1e+99	Time at which the boundary condition will start being applied.
componentFractionVsElevationTableNames	string_array		Names of the tables specifying the (component fraction vs elevation) relationship for each component
componentNames	string_array		Names of the fluid components
datumElevation	real64	required	Datum elevation [m]
datumPressure	real64	required	Datum pressure [Pa]
direction	RTensor	{0,0,0}	Direction to apply boundary condition to.
elevationIncrementInHydrostaticPressureTable	real64	0.6096	Elevation increment [m] in the hydrostatic pressure table constructed internally
endTime	real64	1e+99	Time at which the boundary condition will stop being applied.
equilibrationTolerance	real64	0.001	Tolerance in the fixed-point iteration scheme used for hydrostatic initialization
functionName	string		Name of function that specifies variation of the boundary condition.
initialPhaseName	string		Name of the phase initially saturating the reservoir
logLevel	integer	0	Log level
maxNumberOfEquilibrationIterations	integer	5	Maximum number of equilibration iterations
name	string	required	A name is required for any non-unique nodes
objectPath	string		Path to the target field
scale	real64	0	Scale factor for value of the boundary condition.
temperatureVsElevationTableName	string		Name of the table specifying the (temperature [K] vs elevation) relationship

## Examples

For single-phase flow, a typical hydrostatic equilibrium input looks like:

```
<FieldSpecifications>

  <HydrostaticEquilibrium
    name="equil"
    objectPath="ElementRegions"
    datumElevation="5"
    datumPressure="1e6"/>

</FieldSpecifications>
```

For compositional multiphase flow, using for instance the CO2-brine flow model, a typical hydrostatic equilibrium input looks like:

```
<FieldSpecifications>

  <HydrostaticEquilibrium
```

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(continued from previous page)

```

name="equil"
objectPath="ElementRegions"
datumElevation="28.5"
datumPressure="1.1e7"
initialPhaseName="water"
componentNames="{ co2, water }"
componentFractionVsElevationTableNames="{ initCO2CompFracTable,
                                           initWaterCompFracTable }"
temperatureVsElevationTableName="initTempTable"/>

```

&lt;/FieldSpecifications&gt;

In this case, a possible way to provide the three required tables is:

&lt;Functions&gt;

&lt;TableFunction

```

name="initCO2CompFracTable"
coordinates="{ 0.0, 10.0, 20.0, 30.0 }"
values="{ 0.04, 0.045, 0.05, 0.055 }"/>

```

&lt;TableFunction

```

name="initWaterCompFracTable"
coordinates="{ 0.0, 10.0, 20.0, 30.0 }"
values="{ 0.96, 0.955, 0.95, 0.945 }"/>

```

&lt;TableFunction

```

name="initTempTable"
coordinates="{ 0.0, 15.0, 30.0 }"
values="{ 358.15, 339.3, 333.03 }"/>

```

&lt;/Functions&gt;

Note that the spacing of the two component fraction tables must be the same, but the spacing of the temperature table can be different.

## Expected behavior and comparison with another initialization method

As illustrated in *Tutorial 3: Regions and Property Specifications*, users can also use multiple **FieldSpecification** tags to impose initial fields, such as the pressure, component fractions, and temperature fields. To help users select the initialization method that best meets their needs, we summarize and compare below the two possible ways to initialize complex, non-uniform initial fields for compositional multiphase simulations in GEOSX.

## Initialization using HydrostaticEquilibrium

This is the initialization procedure that we have described in the first sections of this page. In **HydrostaticEquilibrium**, the initial component fractions and temperatures are provided as a function of elevation only, and the hydrostatic pressure is computed internally before the simulation starts. The typical input was illustrated for a CO<sub>2</sub>-brine fluid model in the previous paragraph.

Expected behavior:

- If **FieldSpecification** tags specifying initial pressure, component fractions, and/or temperature are included in an XML input file that also contains the **HydrostaticEquilibrium** tag, the **FieldSpecification** tags are ignored

by GEOSX. In other words, only the pressure, component fractions, and temperature fields defined with the **HydrostaticEquilibrium** tag as a function of elevation are taken into account.

- In the absence of source/sink terms and wells, the initial flow residual should be smaller than  $10^{-6}$ . Similarly, in coupled simulations, the residual of the mechanical problem should be close to zero.

### Initialization using FieldSpecification tags

This is the initialization method illustrated in *Tutorial 3: Regions and Property Specifications*. The user can impose initial pressure, component fractions, and temperature fields using **FieldSpecification** tags, such as, for a two-component CO2-brine case:

```
<FieldSpecifications>

  <FieldSpecification
    name="initialPressure"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions"
    fieldName="pressure"
    scale="1"
    functionName="initialPressureTableXYZ"/>

  <FieldSpecification
    name="initialCO2CompFraction"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions"
    fieldName="globalCompFraction"
    component="0"
    scale="1"
    functionName="initialCO2CompFracTableXYZ"/>

  <FieldSpecification
    name="initialWaterCompFrac"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions"
    fieldName="globalCompFraction"
    component="1"
    scale="1"
    functionName="initialWaterCompFracTableXYZ"/>

  <FieldSpecification
    name="initialTemperature"
    initialCondition="1"
    setNames="{ all }"
    objectPath="ElementRegions"
    fieldName="temperature"
    scale="1"
    functionName="initialTemperatureTableXYZ"/>

</FieldSpecifications>
```

In this input method, `initialPressureTableXYZ`, `initialCO2CompFracTableXYZ`, `initialWaterCompFracTableXYZ`, and `initialTemperatureTableXYZ` are tables describing these initial fields as a function of the x, y, and z spatial coordinates. Then, the cell-wise values are determined by

interpolating in these tables using the coordinates of the center of the cells.

Expected behavior:

- In this approach, it is the responsibility of the user to make sure that these initial fields satisfy a hydrostatic equilibrium. If not, the model will equilibrate itself during the first time steps of the simulation, possibly causing large initial changes in pressure, component fractions, and temperature.
- If the initial state imposed by the **FieldSpecification** tags is not at equilibrium, the displacements produced by coupled flow and mechanics simulations should be interpreted with caution, as these displacements are computed with respect to a non-equilibrium initial state.
- This method is suited to impose initial fields in complex cases currently not supported by the **HydrostaticEquilibrium** tag (e.g., in the presence of phase contacts, capillary pressure, etc). Specifically, the user can equilibrate the model using other means (such as using another simulator, or running a few steps of GEOSX), retrieve the equilibrated values, convert them into x-y-z tables, and impose them in the new GEOSX simulations using **FieldSpecification** tags.

## 5.5.2 Aquifer Boundary Condition

### Overview

Aquifer boundary conditions allow simulating flow between the computational domain (the reservoir) and one or multiple aquifers. In GEOSX, we use a Carter-Tracy aquifer model parameterized in **Aquifer** tags of the **FieldSpecifications** XML input file blocks.

### Aquifer model

An aquifer  $A$  is a source of volumetric flow rate  $q_f^A$ , where  $f$  is the index of a face connecting the aquifer and the reservoir. We use a Carter-Tracy model in GEOSX to compute this volumetric flow rate.

Once  $q_f^A$  is computed, the aquifer mass contribution  $F_f^A$  is assembled and added to the mass conservation equations of the reservoir cell  $K$  connected to face  $f$ . The computation of  $F_f^A$  depends on the sign of the volumetric flow rate  $q_f^A$ .

The upwinding procedure is done as follows: if the sign of  $q_f^A$  indicates that flow goes from the aquifer to the reservoir, the aquifer contribution to the conservation equation of component  $c$  is:

$$F_{f,c}^A = \rho_w^A y_{w,c}^A q_f^A$$

where  $\rho_w^A$  is the aquifer mass/molar water phase density and  $y_{w,c}^A$  is the aquifer mass/molar fraction of component  $c$  in the water phase. We assume that the aquifer is fully saturated with the water phase.

If the sign of  $q_f^A$  indicates that flow goes from the reservoir into the aquifer, the aquifer contribution to the mass/molar conservation equation of component  $c$  is computed as:

$$F_{f,c}^A = \sum_{\ell=1}^{n_p} (\rho_{\ell} S_{\ell} y_{\ell,c})_K q_f^A$$

where  $n_p$  is the number of fluid phases,  $(\rho_{\ell})_K$  is the reservoir cell mass/molar density of phase  $\ell$ ,  $(S_{\ell})_K$  is the reservoir cell saturation of phase  $\ell$ , and  $(y_{\ell,c})_K$  is the reservoir cell mass/molar fraction of component  $c$  in phase  $\ell$ .

In the next section, we review the computation of the aquifer volumetric flow rate  $q_f^A$ .

## Carter-Tracy analytical aquifer

The Carter-Tracy aquifer model is a simplified approximation to a fully transient model (see R. D. Carter and G. W. Tracy, [An improved method for calculating water influx](#), Transactions of the AIME, 1960).

Although the theory was developed for a radially symmetric reservoir surrounded by an annular aquifer, this method applies to any geometry where the dimensionless pressure can be expressed as a function of a dimensionless time.

The two main parameters that govern the behavior of the aquifer are the time constant and the influx constant. These two parameters are precomputed at the beginning of the simulation and are later used to compute the aquifer volumetric flow rate.

### Time constant

The time constant,  $T_c$ , has the dimension of time (in seconds).

It is computed as:

$$T_c = \frac{\mu_w^A \phi^A c_t^A (r_0^A)^2}{k^A}$$

where  $\mu_w^A$  is the aquifer water phase viscosity,  $\phi^A$  is the aquifer porosity,  $c_t^A$  is the aquifer total compressibility (fluid and rock),  $r_0^A$  is the inner radius of the aquifer, and  $k^A$  is the aquifer permeability.

The time constant is used to convert time ( $t$ , in seconds) into dimensionless time,  $t_D$  using the following expression:

$$t_D = \frac{t}{T_c}$$

### Influx constant

The influx constant,  $\beta$ , has the dimension of  $m^3.Pa^{-1}$ .

It is computed as:

$$\beta = 6.283 h^A \theta^A \phi^A c_t^A (r_0^A)^2$$

where  $h^A$  is the aquifer thickness,  $\theta^A$  is the aquifer angle,  $\phi^A$  is the aquifer porosity,  $c_t^A$  is the aquifer total compressibility (fluid and rock), and  $r_0^A$  is the inner radius of the aquifer.

### Aquifer volumetric flow rate

Let us consider a reservoir cell  $K$  connected to aquifer  $A$  through face  $f$ , and the corresponding aquifer volumetric flow rate  $q_f^A$  over time interval  $[t^n, t^{n+1}]$ .

The computation of  $q_f^A$  proceeds as follows:

$$q_f^A = \alpha_f^A (a - b(p_K(t^{n+1}) - p_K(t^n)))$$

where  $\alpha_f^A$  is the area fraction of face  $f$ , and  $p_K(t^{n+1})$  and  $p_K(t^n)$  are the pressures in cell  $K$  at time  $t^{n+1}$  and time  $t^n$ , respectively.

The area fraction of face  $f$  with area  $|f|$  is computed as:

$$\alpha_f^A = \frac{|f|}{\sum_{f_i \in A} |f_i|}$$

The coefficient  $a$  is computed as:

$$a = \frac{1}{T_c} \frac{\beta \Delta \Phi_K^A(t_D^n) - W^A(t_D^n) P_D'(t_D^{n+1})}{P_D(t_D^{n+1}) - t_D^{n+1} P_D'(t_D^{n+1})}$$

and the coefficient  $b$  is given by the formula:

$$b = \frac{1}{T_c} \frac{\beta}{P_D(t_D^{n+1}) - t_D^{n+1} P_D'(t_D^{n+1})}$$

where  $\Delta \Phi_K^A(t_D^n) := p^A - p_K(t^n) - \rho_w^A g(z_K - z^A)$  is the potential difference between the reservoir cell and the aquifer at time  $t^n$ ,  $P_D(t_D)$  is the dimensionless pressure evaluated at dimensionless time  $t_D$ ,  $P_D'(t_D)$  is the derivative of the dimensionless pressure with respect to dimensionless time, evaluated at dimensionless time  $t_D$ .

The functional relationship of dimensionless pressure,  $P_D$ , as a function of dimensionless time is provided by the user. A default table is also available, as shown below. The cumulative aquifer flow rate,  $W^A(t_D^n)$ , is an explicit quantity evaluated at  $t_D^n$  and updated at the end of each converged time step using the formula:

$$W^A(t_D^{n+1}) = W^A(t_D^n) + (t^{n+1} - t^n) \sum_{f \in A} q_f^A$$

with  $W^A(0) := 0$ .

## Parameters

The main Carter-Tracy parameters and the expected units are listed below:

- *aquiferPorosity*: the aquifer porosity  $\phi^A$ .
- *aquiferPermeability*: the aquifer permeability  $k^A$  (in m2).
- *aquiferInitialPressure*: the aquifer initial pressure  $p^A$  (in Pa), used to compute  $\Delta \Phi_K^A$ .
- *aquiferWaterViscosity*: the aquifer water viscosity  $\mu_w^A$  (in Pa.s).
- *aquiferWaterDensity*: the aquifer water mass/molar density  $\rho_w^A$  (in kg/m3 or mole/m3).
- *aquiferWaterPhaseComponentNames*: the name of the components in the water phase. These names must match the component names listed in the fluid model of the **Constitutive** block. This parameter is ignored in single-phase flow simulations.
- *aquiferWaterPhaseComponentFraction*: the aquifer component fractions in the water phase,  $y_{w,c}^A$ . The components must be listed in the order of the components in *aquiferWaterPhaseComponentNames*. This parameter is ignored in single-phase flow simulations.
- *aquiferTotalCompressibility*: the aquifer total compressibility (for the fluid and the solid)  $c_t^A$  (in 1/Pa).
- *aquiferElevation*: the elevation of the aquifer (in m).
- *aquiferThickness*: the thickness of the aquifer (in m).
- *aquiferInnerRadius*: the aquifer inner radius (in m).
- *aquiferAngle*: the angle subtended by the aquifer boundary from the center of reservoir (in degrees, must be between 0 and 360).
- *allowAllPhasesIntoAquifer*: flag controlling the behavior of the aquifer when there is flow from the reservoir to the aquifer. If the flag is equal to 1, all phases can flow into the aquifer. If the flag is equal to 0, only the water phase can flow into the aquifer. The default value of this optional parameter is 0.
- *pressureInfluenceFunctionName*: the name of the table providing the dimensionless pressure as a function of dimensionless time. This table must be defined as a **TableFunction** in the **Functions** block of the XML file. If this optional parameter is omitted, a default pressure influence table is used.

- *setNames*: the names of the face sets on which the aquifer boundary condition is applied.

---

**Note:** Following the GEOSX convention, the z-coordinate is increasing upward. This convention must be taken into account when providing the *aquiferElevation*. In other words, the z-value is not a depth.

---

The full list of parameters is provided below:

Name	Type	Default	Description
allowAllPhasesIntoAquifer	integer	0	<p>Flag to allow all phases to flow into the aquifer. This flag only matters for the configuration in which flow is from reservoir to aquifer.</p> <ul style="list-style-type: none"> <li>- If the flag is equal to 1, then all phases, including non-aqueous phases, are allowed to flow into the aquifer.</li> <li>- If the flag is equal to 0, then only the water phase is allowed to flow into the aquifer.</li> </ul> <p>If you are in a configuration in which flow is from reservoir to aquifer and you expect non-aqueous phases to saturate the reservoir cells next to the aquifer, set this flag to 1.</p> <p>This keyword is ignored for single-phase flow simulations</p>
aquiferAngle	real64	required	Angle subtended by the aquifer boundary from the center of the reservoir [degrees]
aquiferElevation	real64	required	Aquifer elevation (positive going upward) [m]
aquiferInitialPressure	real64	required	Aquifer initial pressure [Pa]
aquiferInnerRadius	real64	required	Aquifer inner radius [m]
aquiferPermeability	real64	required	Aquifer permeability [m <sup>2</sup> ]
aquiferPorosity	real64	required	Aquifer porosity
aquiferThickness	real64	required	Aquifer thickness [m]
aquiferTotalCompressibility	real64	required	Aquifer total compressibility (rock and fluid) [Pa <sup>-1</sup> ]
aquiferWaterDensity	real64	required	Aquifer water density [kg.m <sup>-3</sup> ]
aquiferWaterPhaseComponentFraction	real64 array	{0}	Aquifer water phase component fraction. This keyword is ignored for single-phase flow simulations.
aquiferWaterPhaseComponentNames	string array	{}	Aquifer water phase component names. This keyword is ignored for single-phase flow simulations.



## Examples

Setting up the **Aquifer** boundary condition requires two additional pieces of information in the XML input file: a set of faces to specify where the aquifer boundary conditions will apply, and an aquifer tag that specifies the physical characteristics of the aquifer and determines how the boundary condition is applied.

- 1) To specify a set of faces: on simple grids, in the **Geometry** block of the XML file, we can define a **Box** that selects and assigns a name to a set of faces. To be included in a set, the faces must be fully enclosed in the **Box** (all vertices of a face must be inside the box for the face to be included to the set). The name of this box is a user-defined string, and it will be used in the aquifer tag to locate the face set. Here is an example of XML code to create such a face set from a box:

```
<Geometry>
...
<Box
  name="aquifer"
  xMin="{ 999.99, 199.99, 3.99 }"
  xMax="{ 1010.01, 201.01, 6.01 }"/>
...
</Geometry>
```

**Note:** This step captures *faces*, not *cells*. For now, the user must ensure that the box actually contains faces (GEOSX will proceed even if the face set is empty).

For more complex meshes, such as those imported using the **PAMELAMesh**, using a **Box** to perform a face selection is challenging. We recommend using a tagged `PhysicalEntity` in the `.msh` file instead. This physical entity will be used to locate the face set.

- 2) To specify the aquifer characteristics: in the **FieldSpecifications** block of the XML file, we include an **Aquifer** tag. For single-phase flow, the aquifer definition looks like:

```
<FieldSpecifications>
...
<Aquifer
  name="aquiferBC"
  aquiferPorosity="2e-1"
  aquiferPermeability="3e-13"
  aquiferInitialPressure="9e6"
  aquiferWaterViscosity="0.00089"
  aquiferWaterDensity="962.81"
  aquiferTotalCompressibility="1e-10"
  aquiferElevation="4"
  aquiferThickness="18"
  aquiferInnerRadius="2000"
  aquiferAngle="20"
  setNames="{ aquifer }"/>
...
</FieldSpecifications>
```

For compositional multiphase flow, the user must include additional parameters to specify the water composition. We have additional influx controls over the aquifer with `allowAllPhasesIntoAquifer`. This is illustrated below for the CO<sub>2</sub>-brine fluid model:

```
<FieldSpecifications>
...
<Aquifer
```

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```

    name="aquiferBC"
    aquiferPorosity="2e-1"
    aquiferPermeability="3e-13"
    aquiferInitialPressure="9e6"
    aquiferWaterViscosity="0.00089"
    aquiferWaterDensity="962.81"
    aquiferWaterPhaseComponentFraction="{ 0.0, 1.0 }"
    aquiferWaterPhaseComponentNames="{ co2, water }"
    aquiferTotalCompressibility="1e-10"
    aquiferElevation="4"
    aquiferThickness="18"
    aquiferInnerRadius="2000"
    aquiferAngle="20"
    allowAllPhasesIntoAquifer="1"
    setNames="{ aquifer }"/>
    ...
</FieldSpecifications>

```

Finally, for both single-phase and multiphase flow, if a `pressureInfluenceFunctionName` attribute is specified in the **Aquifer** tag, a **TableFunction** must be included in the **Functions** block of the XML file as follows:

```

<Functions>
    ...
    <TableFunction
        name="pressureInfluenceFunction"
        coordinates="{ 0.01, 0.05, 0.1, 0.15, 0.2, 0.25, 0.3, 0.4, 0.5, 0.6, 0.7, 0.8, 0.
↪9, 1.0, 1.5,
                        2.0, 2.5, 3.0, 4.0, 5.0, 6.0, 7.0, 8.0, 9.0, 10.0, 15.0, 20.0, 25.
↪0, 30.0, 40.0,
                        50.0, 60.0, 70.0, 80.0, 90.0, 100.0, 200.0, 800.0, 1600.0, 3200.0, ↪
↪6400.0, 12800.0 }"
        values="{ 0.112, 0.229, 0.315, 0.376, 0.424, 0.469, 0.503, 0.564, 0.616, 0.659, 0.
↪702, 0.735, 0.772, 0.802,
                  0.927, 1.02, 1.101, 1.169, 1.275, 1.362, 1.436, 1.5, 1.556, 1.604, 1.
↪651, 1.829, 1.96, 2.067, 2.147,
                  2.282, 2.388, 2.476, 2.55, 2.615, 2.672, 2.723, 3.0537, 3.7468, 4.0934, ↪
↪4.44, 4.7866, 5.1331 }"/>
    ...
</Functions>

```

**Note:** The values provided in the table above are the default values used internally in GEOSX when the user does not specify the pressure influence function in the XML file.

## 5.6 Event Management

The goal of the GEOSX event manager is to be flexible with regards to event type, application order, and method of triggering. The event manager is configured via the **Event** block in an input .xml file, i.e.:

```

<Events maxTime="1.0e-2">
    <PeriodicEvent name="event_a"
        target="/path/to/event"
        forceDt="1" />

```

(continues on next page)

(continued from previous page)

```

<HaltEvent name="event_b"
            target="/path/to/halt_target"
            maxRunTime="1e6" />
</Events>

```

### 5.6.1 Event Execution Rules

The EventManager will repeatedly iterate through a list of candidate events specified via the Events block **in the order they are defined in the xml**. When certain user-defined criteria are met, they will trigger and perform a task. The simulation `cycle` denotes the number of times the primary event loop has completed, `time` denotes the simulation time at the beginning of the loop, and `dt` denotes the global timestep during the loop.

During each cycle, the EventManager will do the following:

1. Loop through each event and obtain its timestep request by considering:
  - a. The maximum `dt` specified via the target's `GetTimestepRequest` method
  - b. The time remaining until user-defined points (e.g. application start/stop times)
  - c. Any timestep overrides (e.g. user-defined maximum `dt`)
  - d. The timestep request for any of its children
2. Set the cycle `dt` to the smallest value requested by any event
3. Loop through each event and:
  - a. Calculate the event `forecast`, which is defined as the expected number of cycles until the event is expected to execute.
  - b. `if (forecast == 1)` the event will signal its target to prepare to execute. This is useful for preparing time-consuming I/O operations.
  - c. `if (forecast <= 0)` the event will call the `Execute` method on its target object
4. Check to see if the EventManager exit criteria have been met

After exiting the main event loop, the EventManager will call the `Cleanup` method for each of its children (to produce final plots, etc.). Note: if the code is resuming from a restart file, the EventManager will pick up exactly where it left off in the execution loop.

### 5.6.2 Event Manager Configuration

#### Event

The children of the Event block define the events that may execute during a simulation. These may be of type `HaltEvent`, `PeriodicEvent`, or `SoloEvent`. The exit criteria for the global event loop are defined by the attributes `maxTime` and `maxCycle` (which by default are set to their max values). If the optional `logLevel` flag is set, the EventManager will report additional information with regards to timestep requests and event forecasts for its children.

Name	Type	Default	Description
logLevel	integer	0	Log level
maxCycle	integer	2147483647	Maximum simulation cycle for the global event loop.
maxTime	real64	1.79769e+308	Maximum simulation time for the global event loop.
minTime	real64	0	Start simulation time for the global event loop.
HaltEvent	node		<i>Element: HaltEvent</i>
PeriodicEvent	node		<i>Element: PeriodicEvent</i>
SoloEvent	node		<i>Element: SoloEvent</i>

## PeriodicEvent

This is the most common type of event used in GEOSX. As its name suggests, it will execute periodically during a simulation. It can be triggered based upon a user-defined `cycleFrequency` or `timeFrequency`.

If `cycleFrequency` is specified, the event will attempt to execute every X cycles. Note: the default behavior for a `PeriodicEvent` is to execute every cycle. The event forecast for this case is given by: `forecast = cycleFrequency - (cycle - lastCycle)`.

If `timeFrequency` is specified, the event will attempt to execute every X seconds (this will override any cycle-dependent behavior). By default, the event will attempt to modify its timestep requests to respect the `timeFrequency` (this can be turned off by specifying `targetExactTimestep="0"`). The event forecast for this case is given by: `if (dt > 0), forecast = (timeFrequency - (time - lastTime)) / dt, otherwise forecast=max`

By default, a `PeriodicEvent` will execute throughout the entire simulation. This can be restricted by specifying the `beginTime` and/or `endTime` attributes. Note: if either of these values are set, then the event will modify its timestep requests so that a cycle will occur at these times (this can be turned off by specifying `targetExactStartStop="0"`).

The timestep request event is typically determined via its target. However, this value can be overridden by setting the `forceDt` or `maxEventDt` attributes.

Name	Type	Default	Description
beginTime	real64	0	Start time of this event.
cycleFrequency	integer	1	Event application frequency (cycle, default)
endTime	real64	1e+100	End time of this event.
finalDtStretch	real64	0.001	Allow the final dt request for this event to grow by this percentage to match the endTime exactly.
forceDt	real64	-1	While active, this event will request this timestep value (ignoring any children/targets requests).
function	string		Name of an optional function to evaluate when the time/cycle criteria are met. If the result is greater than the specified eventThreshold, the function will continue to execute.
logLevel	integer	0	Log level
maxEventDt	real64	-1	While active, this event will request a timestep $\leq$ this value (depending upon any child/target requests).
name	string	required	A name is required for any non-unique nodes
object	string		If the optional function requires an object as an input, specify its path here.
set	string		If the optional function is applied to an object, specify the setname to evaluate (default = everything).
stat	integer	0	If the optional function is applied to an object, specify the statistic to compare to the eventThreshold. The current options include: min, avg, and max.
target	string		Name of the object to be executed when the event criteria are met.
targetExactStartStop	integer	1	If this option is set, the event will reduce its timestep requests to match any specified beginTime/endTimes exactly.
targetExactTimestep	integer	1	If this option is set, the event will reduce its timestep requests to match the specified timeFrequency perfectly: $dt\_request = \min(dt\_request, t\_last + time\_frequency - time)$ .
threshold	real64	0	If the optional function is used, the event will execute if the value returned by the function exceeds this threshold.
timeFrequency	real64	-1	Event application frequency (time). Note: if this value is specified, it will override any cycle-based behavior.
HaltEvent	node		<i>Element: HaltEvent</i>
PeriodicEvent	node		<i>Element: PeriodicEvent</i>
SoloEvent	node		<i>Element: SoloEvent</i>

## SoloEvent

This type of event will execute once once the event loop reaches a certain cycle (targetCycle) or time (targetTime). Similar to the PeriodicEvent type, this event will modify its timestep requests so that a cycle occurs at the exact time requested (this can be turned off by specifying targetExactTimestep="0"). The forecast calculations follow an similar approach to the PeriodicEvent type.

Name	Type	De- fault	Description
beginTime	real64	0	Start time of this event.
endTime	real64	1e+100	End time of this event.
finalDt-Stretch	real64	0.001	Allow the final dt request for this event to grow by this percentage to match the end-Time exactly.
forceDt	real64	-1	While active, this event will request this timestep value (ignoring any children/targets requests).
logLevel	in- te- ger	0	Log level
max-EventDt	real64	-1	While active, this event will request a timestep $\leq$ this value (depending upon any child/target requests).
name	string	re- quired	A name is required for any non-unique nodes
target	string		Name of the object to be executed when the event criteria are met.
targetCy- cle	in- te- ger	-1	Targeted cycle to execute the event.
targetEx- actStart- Stop	in- te- ger	1	If this option is set, the event will reduce its timestep requests to match any specified beginTime/endTimes exactly.
targe- tExact- Timestep	in- te- ger	1	If this option is set, the event will reduce its timestep requests to match the specified execution time exactly: $dt\_request = \min(dt\_request, t\_target - time)$ .
targetTime	real64	-1	Targeted time to execute the event.
HaltEvent	node		<i>Element: HaltEvent</i>
Peri- odicEvent	node		<i>Element: PeriodicEvent</i>
SoloEvent	node		<i>Element: SoloEvent</i>

## HaltEvent

This event type is designed to track the wall clock. When the time exceeds the value specified via maxRunTime, the event will trigger and set a flag that instructs the main EventManager loop to cleanly exit at the end of the current cycle. The event for cast for this event type is given by:  $forecast = (maxRuntime - (currentTime - startTime)) / realDt$

Name	Type	De- fault	Description
beginTime	real64	0	Start time of this event.
endTime	real64	1e+100	End time of this event.
finalDtStretch	real64	0.001	Allow the final dt request for this event to grow by this percentage to match the endTime exactly.
forceDt	real64	-1	While active, this event will request this timestep value (ignoring any children/targets requests).
logLevel	integer	0	Log level
maxEventDt	real64	-1	While active, this event will request a timestep $\leq$ this value (depending upon any child/target requests).
maxRuntime	real64	re- quired	The maximum allowable runtime for the job.
name	string	re- quired	A name is required for any non-unique nodes
target	string		Name of the object to be executed when the event criteria are met.
targetExact- StartStop	integer	1	If this option is set, the event will reduce its timestep requests to match any specified beginTime/endTimes exactly.
HaltEvent	node		<i>Element: HaltEvent</i>
PeriodicEvent	node		<i>Element: PeriodicEvent</i>
SoloEvent	node		<i>Element: SoloEvent</i>

### 5.6.3 Other Event Features

#### Event Progress Indicator

Because the event manager allows the user to specify the order of events, it could introduce ambiguity into the timestamps of output files. To resolve this, we pass two arguments to the target's Execute method:

1. eventCounter (integer) - the application index for the event (or sub-event)
2. eventProgress (real64) - the percent completion of the event loop, paying attention to events whose targets are associated with physics (from the start of the event, indicated via target->GetTimestepBehavior())

For example, consider the following Events block:

```
<Events maxTime="1.0e-2">
  <PeriodicEvent name="outputs"
    timeFrequency="1e-6"
    targetExactTimestep="0"
    target="/Outputs/siloOutput">
  <PeriodicEvent name="solverApplications_a"
    forceDt="1.0e-5"
    target="/Solvers/lagsolve" />
  <PeriodicEvent name="solverApplications_b"
    target="/Solvers/otherSolver" />
  <PeriodicEvent name="restarts"
    timeFrequency="5.0e-4"
    targetExactTimestep="0"
    target="/Outputs/restartOutput"/>
</Events>
```

In this case, the events solverApplications\_a and solverApplications\_b point target physics events. The eventCounter,

eventProgress pairs will be: outputs (0, 0.0), solverApplications\_a (1, 0.0), solverApplications\_b (2, 0.5), and restarts (3, 1.0). These values are supplied to the target events via their Execute methods for use. For example, for the name of a silo output file will have the format: “%s\_%06d%02d” % (name, cycle, eventCounter), and the time listed in the file will be  $\text{time} = \text{time} + \text{dt} * \text{eventProgress}$

## Nested Events

The event manager allows its child events to be nested. If this feature is used, then the manager follows the basic execution rules, with the following exception: When its criteria are met, an event will first execute its (optional) target. It will then estimate the forecast for its own sub-events, and execute them following the same rules as in the main loop. For example:

```
<Events maxTime="1.0e-2">
  <PeriodicEvent name="event_a"
    target="/path/to/target_a" />

  <PeriodicEvent name="event_b"
    timeFrequency="100">

    <PeriodicEvent name="subevent_b_1"
      target="/path/to/target_b_1"/>

    <PeriodicEvent name="subevent_b_2"
      target="/path/to/target_b_2"/>

  </PeriodicEvent/>
</Events>
```

In this example, event\_a will trigger during every cycle and call the Execute method on the object located at /path/to/target\_a. Because it is time-driven, event\_b will execute every 100 s. When this occurs, it will execute it will execute its own target (if it were defined), and then execute subevent\_b\_1 and subevent\_b\_2 in order. Note: these are both cycle-driven events which, by default would occur every cycle. However, they will not execute until each of their parents, grandparents, etc. execution criteria are met as well.

## 5.7 Tasks Manager

The GEOSX tasks manager allows a user to specify tasks to be executed. These tasks are compatible targets for the *Event Management*.

The tasks manager is configured via the Tasks block in an input .xml file, i.e.:

```
<Tasks>
  <PackCollection name="historyCollection" objectPath="nodeManager" fieldName=
    "Velocity" />
</Tasks>
```

### 5.7.1 Tasks Manager Configuration

#### Task

The children of the Tasks block define different Tasks to be triggered by events specified in the *Event Management* during the execution of the simulation. At present the only supported task is the PackCollection used to collect time history data for output by a TimeHistory output.



Name	Type	Default	Description
CompositionalMultiphaseStatistics	node		<i>Element: CompositionalMultiphaseStatistics</i>
PVTDriver	node		<i>Element: PVTDriver</i>
PackCollection	node		<i>Element: PackCollection</i>
SinglePhaseStatistics	node		<i>Element: SinglePhaseStatistics</i>
SolidMechanicsStateReset	node		<i>Element: SolidMechanicsStateReset</i>
SolidMechanicsStatistics	node		<i>Element: SolidMechanicsStatistics</i>
TriaxialDriver	node		<i>Element: TriaxialDriver</i>

## PackCollection

The `PackCollection` Task is used to collect time history information from fields. Either the entire field or specified named sets of indices in the field can be collected.

Name	Type	De- fault	Description
fieldName	string	re- quired	The name of the (packable) field associated with the specified object to retrieve data from
name	string	re- quired	A name is required for any non-unique nodes
objectPath	string	re- quired	The name of the object from which to retrieve field values.
onlyOn-SetChange	integer	0	Whether or not to only collect when the collected sets of indices change in any way.
setNames	string_array	{ }	The set(s) for which to retrieve data.

Note: The time history information collected via this task is buffered internally until it is output by a linked `TimeHistory` Output.

## Triggering the Tasks

Tasks can be triggered using the *Event Management*. Recurring tasks should use a `<PeriodicEvent>` and one-time tasks should use a `<SoloEvent>`:

```
<PeriodicEvent name="historyCollectEvent"
  timeFrequency="1.0"
  targetExactTimeset="1"
  target="/Tasks/historyCollection" />
```

The keyword `target` has to match the name of a Task specified as a child of the `<Tasks>` block.

## 5.8 Functions

Functions are the primary avenue for specifying values that change in space, time, or any other dimension. These are specified in the `Functions` block, and may be referenced by name throughout the rest of the .xml file. For example:

```
<Functions>
  <TableFunction name="q"
    inputVarNames="time"
```

(continues on next page)

```
        coordinates="0 60 1000"
        values="0 1 1" />
</Functions>
<FieldSpecifications>
  <SourceFlux name="sourceTerm"
    objectPath="ElementRegions/Region1/block1"
    scale="0.001"
    functionName="q"
    setNames="{source}" />
</FieldSpecifications>
```

## 5.8.1 Function Inputs and Application

The inputs to each function type are specified via the `inputVarName` attribute. These can either be the name of an array (e.g. “Pressure”) or the special keyword “time” (time at the beginning of the current cycle). If any of the input variables are vectors (e.g. “referencePosition”), the components will be given as function arguments in order.

In the .xml file, functions are referenced by name. Depending upon the application, the functions may be applied in one of three ways:

1. Single application: The function is applied to get a single scalar value. For example, this could define the flow rate applied via a BC at a given time.
2. Group application: The function is applied to a (user-specified) `ManagedGroup` of size `N`. When called, it will iterate over the `inputVarNames` list and build function inputs from the group’s wrappers. The resulting value will be a wrapper of size `N`. For example, this could be used to apply a user-defined constitutive relationship or specify a complicated boundary condition.
3. Statistics mode: The function is applied in the same manner as the group application, except that the function will return an array that contains the minimum, average, and maximum values of the results.

## 5.8.2 Function Types

There are three types of functions available for use: `TableFunction`, `SymbolicFunction`, and `CompositeFunction`. **Note: the symbolic and composite function types are currently only available for x86-64 systems.**

### TableFunction

A table function uses a set of pre-computed values defined at points on a structured grid to represent an arbitrary-dimensional function. Typically, the axes of the table will represent time and/or spatial dimensions; however, these can be applied to represent phase diagrams, etc.

Name	Type	Default	Description
coordinateFiles	path_array	{}	List of coordinate file names for ND Table
coordinates	real64_array	{0}	Coordinates inputs for 1D tables
inputVarNames	string_array	{}	Name of fields are input to function.
interpolation	geosx_TableFunction_InterpolationType		Interpolation method. Valid options: * linear * nearest * upper * lower
name	string	required	A name is required for any non-unique nodes
values	real64_array	{0}	Values for 1D tables
voxelFile	path		Voxel file name for ND Table

## 1D Table

For 1D tables, the function may be defined using the coordinates and values attributes. These represent the location of the grid nodes (ordered from smallest to largest) and the function values at those points, respectively. For example, the following function defines a simple ramp function with a rise-time of 60 seconds:

```
<TableFunction name="q"
  inputVarNames="time"
  coordinates="0 60 1000"
  values="0 1 1" />
```

## ND Table

For ND tables, the grid coordinates and values may be defined using a set of .csv files. The `coordinateFiles` attribute specifies the file names that define the coordinates for each axis. The values in each coordinate file must be comma-delimited and ordered from smallest to largest. The dimensionality of the table is defined by the number of coordinate files (`coordinateFiles="x.csv"` would indicate a 1D table, `coordinateFiles="x.csv y.csv z.csv t.csv"` would indicate a 4D table, etc.). The `voxelFile` attribute specifies name of the file that defines the value of the function at each point along the grid. These values must be comma-delimited (line-breaks are allowed) and be specified in Fortran order, i.e., column-major order (where the index of the first dimension changes the fastest, and the index of the last dimension changes slowest).

The following would define a simple 2D function  $c = a + 2*b$ :

```
<TableFunction name="c"
  inputVarNames="a b"
  coordinateFiles="a.csv b.csv"
  voxelFile="c.csv" />
```

- a.csv: "0, 1"

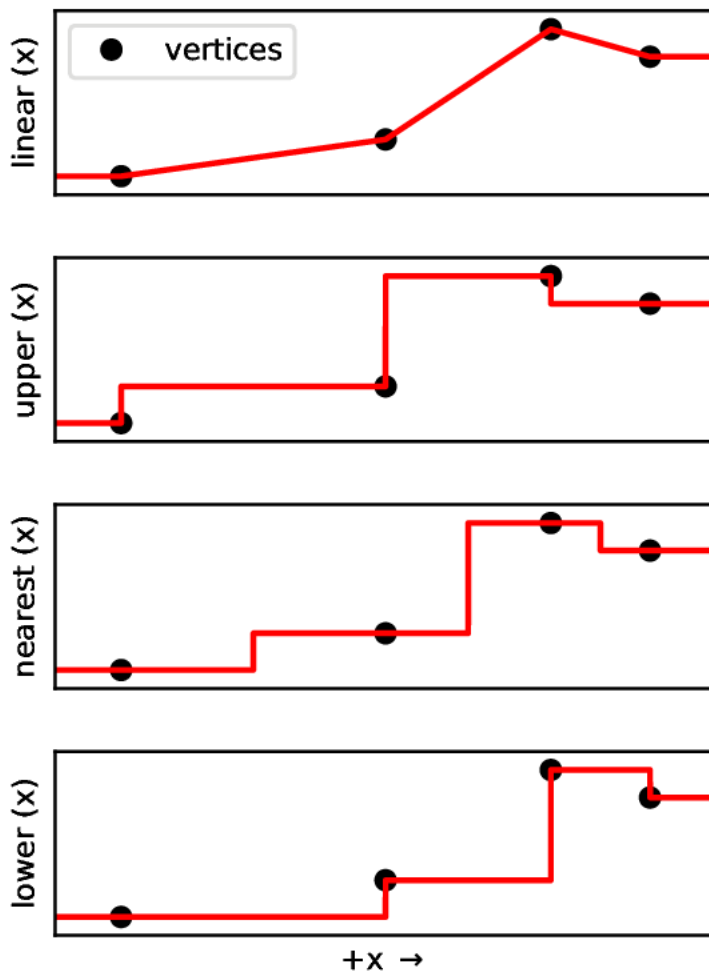
- b.csv: “0, 0.5, 1”
- c.csv: “0, 1, 1, 2, 2, 3”

## Interpolation Methods

There are four interpolation methods available for table functions. Within the table axes, these will return a value:

- linear: using piecewise-linear interpolation
- upper: equal to the value of the next table vertex
- nearest: equal to the value of the nearest table vertex
- lower: equal to the value of the previous table vertex

Outside of the table axes, these functions will return the edge-values. The following figure illustrates how each of these methods work along a single dimension, given identical table values:



## Table Generation Example

The following is an example of how to generate the above tables in Python:

```
import numpy as np

# Define table axes
a = np.array([0.0, 1.0])
b = np.array([0.0, 0.5, 1.0])

# Generate table values (note: the indexing argument is important)
A, B = np.meshgrid(a, b, indexing='ij')
C = A + 2.0*B

# Write axes, value files
np.savetxt('a.csv', a, fmt='%1.2f', delimiter=',')
np.savetxt('b.csv', b, fmt='%1.2f', delimiter=',')
values = np.reshape(C, (-1), order='F')
np.savetxt('c.csv', values, fmt='%1.2f', delimiter=',')
```

## SymbolicFunction

This function leverages the symbolic expression library mathpresso to define and evaluate functions. These functions are processed using an x86-64 JIT compiler, so are nearly as efficient as natively compiled C++ expressions.

Name	Type	De- fault	Description
expression	string	re- quired	Symbolic math expression
inputVar- Names	string_array	{ }	Name of fields are input to function.
name	string	re- quired	A name is required for any non-unique nodes
variable- Names	string_array	re- quired	List of variables in expression. The order must match the evaluate argument

The `variableNames` attribute defines a set of single-character names for the inputs to the symbolic function. There should be a definition for each scalar input and for each component of a vector input. For example if `inputVarName="time, ReferencePosition"`, then `variableNames="t, x, y, z"`. The `expression` attribute defines the symbolic expression to be executed. Aside from the following exceptions, the syntax mirrors python:

- The function string cannot contain any spaces
- The power operator is specified using the C-style expression (e.g. `pow(x,3)` instead of `x**3`)

The following would define a simple 2D function `c = a + 2*b`:

```
<SymbolicFunction name="c"
    inputVarNames="a b"
    variableNames="x y"
    expression="x+(2*y)"/>
```

## CompositeFunction

This function is derived from the symbolic function. However, instead of using the time or object as inputs, it is used to combine the outputs of other functions using a symbolic expression.

Name	Type	De- fault	Description
expression	string		Composite math expression
function-Names	string_array	{ }	List of source functions. The order must match the variableNames argument.
inputVar-Names	string_array	{ }	Name of fields are input to function.
name	string	re- quired	A name is required for any non-unique nodes
variable-Names	string_array	{ }	List of variables in expression

The `functionNames` attribute defines the set of input functions to use (these may be of any type, and may each have any number of inputs). The `variableNames` attribute defines a set of single-character names for each function. The `expression` attribute defines the symbolic expression, and follows the same rules as above. The `inputVarNames` attribute is ignored for this function type.

The following would define a simple 1D table function  $f(t) = 1 + t$ , a 3D symbolic function  $g(x, y, z) = x^2 + y^2 + z^2$ , and a 4D composite function  $h = \sin(f(t)) + g(x, y, z)$ :

```
<Functions>
  <TableFunction name="f"
    inputVarNames="time"
    coordinates="0 1000"
    values="1 1001" />

  <SymbolicFunction name="g"
    inputVarNames="ReferencePosition"
    variableNames="x y z"
    expression="pow(x,2)+pow(y,2)+pow(z,2)"/>

  <CompositeFunction name="h"
    inputVarNames="ignored"
    functionNames="f g"
    variableNames="x y"
    expression="sin(x)+y"/>
</Events>
```

## 5.9 Linear Solvers

### 5.9.1 Introduction

Any physics solver relying on standard finite element and finite volume techniques requires the solution of algebraic linear systems, which are obtained upon linearization and discretization of the governing equations, of the form:

$$Ax = b$$

with a  $A$  a square sparse matrix,  $x$  the solution vector, and  $b$  the right-hand side. For example, in a classical linear elastostatics problem  $A$  is the stiffness matrix, and  $x$  and  $b$  are the displacement and nodal force vectors, respectively.

This solution stage represents the most computationally expensive portion of a typical simulation. Solution algorithms generally belong to two families of methods: direct methods and iterative methods. In GEOSX both options are made available wrapping around well-established open-source linear algebra libraries, namely [HYPRE](#), [PETSc](#), [SuperLU](#), and [Trilinos](#).

### 5.9.2 Direct methods

The major advantages are their reliability, robustness, and ease of use. However, they have large memory requirements and exhibit poor scalability. Direct methods should be used in a prototyping stage, for example when developing a new formulation or algorithm, when the dimension of the problem, namely the size of matrix  $A$ , is small. Irrespective of the selected direct solver implementation, three stages can be identified:

- (1) **Setup Stage:** the matrix is first analyzed and then factorized
- (2) **Solve Stage:** the solution to the linear systems involving the factorized matrix is computed
- (3) **Finalize Stage:** the systems involving the factorized matrix have been solved and the direct solver lifetime ends

The default option in GEOSX relies on [SuperLU](#), a general purpose library for the direct solution of large, sparse, nonsymmetric systems of linear equations, that is called taking advantage of the interface provided in [HYPRE](#).

### 5.9.3 Iterative methods

As the problem size (number of computational cells) increases, global iterative solution strategies are the method of choice—typically nonsymmetric Krylov solvers. Because of the possible poor conditioning of  $A$ , preconditioning is essential to solve such systems efficiently. *“Preconditioning is simply a means of transforming the original linear system into one which has the same solution, but which is likely to be easier to solve with an iterative solver”* [Saad (2003)].

The design of a robust and efficient preconditioner is based on a trade-off between two competing objectives:

- **Robustness:** reducing the number of iterations needed by the preconditioned solver to achieve convergence;
- **Efficiency:** limiting the time required to construct and apply the preconditioner.

Assuming a preconditioning matrix  $M$  is available, three standard approaches are used to apply the preconditioner:

- (1) **Left preconditioning:** the preconditioned system is  $M^{-1}Ax = M^{-1}b$
- (2) **Right preconditioning:** the preconditioned system is  $AM^{-1}y = b$ , with  $x = M^{-1}y$
- (3) **Split preconditioning:** the preconditioned system is  $M_L^{-1}AM_R^{-1}y = M_L^{-1}b$ , with  $x = M_R^{-1}y$

### 5.9.4 Summary

The following table summarizes the available input parameters for the linear solver.

Name	Type	Default	Description
amgAggressiveCoarseningLevel	integer	0	AMG number levels for aggressive coarsening Available options are: TODO
amgCoarseSolver	geosx_LinearSolverParameters	direct AMG_CoarseType	AMG coarsest level solver/smoother type. Available options are: default\ jacobi\ llt\ jacobi\ fgs\
amgCoarseningType	string	HMIS	AMG coarsening algorithm Available options are: TODO
amgInterpolationType	integer	6	AMG interpolation algorithm Available options are: TODO
amgNullSpaceType	geosx_LinearSolverParameters	constantModes	AMG near null space approximation. Available options are: constantModes\ rigidBodyModes\
amgNumFunctions	integer	1	AMG number of functions Available options are: TODO
amgNumSweeps	integer	2	AMG smoother sweeps
amgSmootherType	geosx_LinearSolverParameters	fgs AMG_SmootherType	AMG smoother type. Available options are: default\ jacobi\ llt\ jacobi\ fgs\
amgThreshold	real64	0	AMG strength-of-connection threshold
directCheckResidual	integer	0	Whether to check the linear system solution residual
directColPerm	geosx_LinearSolverParameters	none Direct_ColPerm	How to permute the columns. Available options are: none\ MMD_AtA\ colMMD_AtA\ col
directEquil	integer	1	Whether to scale the rows and columns of the matrix
directIterRef	integer	1	Whether to perform iterative refinement
directParallel	integer	1	Whether to use a parallel solver (instead of a serial one)
410			<b>Chapter 5. User Guide</b>
directReplTinyPivot	integer	1	Whether to replace tiny pivots by $\sqrt{\epsilon} \cdot \text{norm}(A)$



### 5.9.5 Preconditioner descriptions

This section provides a brief description of the available preconditioners.

- **None:** no preconditioning is used, i.e.,  $M^{-1} = I$ .
- **Jacobi:** diagonal scaling preconditioning, with  $M^{-1} = D^{-1}$ , with D the matrix diagonal. Further details can be found in:
  - [HYPRE documentation](#),
  - [PETSc documentation](#),
  - [Trilinos documentation](#).
- **ILUK:** incomplete LU factorization with fill level k of the original matrix:  $M^{-1} = U^{-1}L^{-1}$ . Further details can be found in:
  - [HYPRE documentation](#),
  - [PETSc documentation](#),
  - [Trilinos documentation](#).
- **ILUT:** a dual threshold incomplete LU factorization:  $M^{-1} = U^{-1}L^{-1}$ . Further details can be found in:
  - [HYPRE documentation](#),
  - not yet available through PETSc interface,
  - [Trilinos documentation](#).
- **ICC:** incomplete Cholesky factorization of a symmetric positive definite matrix:  $M^{-1} = L^{-T}L^{-1}$ . Further details can be found in:
  - not yet available through *hypre* interface,
  - [PETSc documentation](#),
  - [Trilinos documentation](#).
- **AMG:** algebraic multigrid (can be classical or aggregation-based according to the specific package). Further details can be found in:
  - [HYPRE documentation](#),
  - [PETSc documentation](#),
  - [Trilinos documentation](#).
- **MGR:** multigrid reduction. Available through *hypre* interface only. Specific documentation coming soon. Further details can be found in [MGR documentation](#).
- **Block:** custom preconditioner designed for a 2 x 2 block matrix.

### 5.9.6 HYPRE MGR Preconditioner

MGR stands for multigrid reduction, a multigrid method that uses the interpolation, restriction operators, and the Galerkin triple product, to reduce a linear system to a smaller one, similar to a Schur complement approach. As such, it is designed to target block linear systems resulting from discretizations of multiphysics problems. GEOSX uses MGR through an implementation in [HYPRE](#). More information regarding MGR can be found [here](#). Currently, MGR strategies are implemented for hydraulic fracturing, poroelastic, compositional flow with and without wells. More multiphysics solvers with MGR will be enabled in the future.

To use MGR for a specific block system, several components need to be specified.

- (1) **The number of reduction levels and the coarse points (corresponding to fields) for each level.** For example, for single-phase hydraulic fracturing, there are two fields, i.e. displacement and fluid pressure, a two-level MGR strategy can be used with the fluid pressure being the coarse degrees of freedom.
- (2) **Interpolation/restriction operators and the coarse-grid computation strategy.** A simple but effective strategy is to use Jacobi diagonalization for interpolation and injection for restriction. For most cases, a Galerkin coarse grid strategy can be used, but for special cases such as poroelastic, a non-Galerkin approach is preferable.
- (3) **Global smoother.** Depending on the problem, a global relaxation step could be beneficial. Some options include ILU(k), (block) Jacobi, (block) Gauss-Seidel.
- (4) **Solvers for F-relaxation and coarse-grid correction.** These solvers should be chosen carefully for MGR to be effective. The choice of these solvers should correspond to the properties of the blocks specified by the C- and F-points. For example, if the  $A_{FF}$  block is hyperbolic, a Jacobi smoother is sufficient while for an elliptic operator an AMG V-cycle might be required. For the single-phase hydraulic fracturing case, an AMG V-cycle is needed for both F-relaxation and coarse-grid correction.

Note that these are only general guidelines for designing a working MGR recipe. For complicated multiphysics problems, experimentation with different numbers of levels, choices of C- and F-points, and smoothers/solvers, etc., is typically needed to find the best strategy. Currently, these options are only available to developers. We are working on exposing these functionalities to the users in future releases.

## 5.9.7 Block preconditioner

This framework allows the user to design a block preconditioner for a  $2 \times 2$  block matrix. The key component is the Schur complement  $S = A_{11} - A_{10}\tilde{A}_{00}^{-1}A_{01}$  computation, that requires an approximation of the leading block. Currently, available options for  $\tilde{A}_{00}^{-1}$  are:

- diagonal with diagonal values (essentially, a Jacobi preconditioner);
- diagonal with row sums as values (e.g., used for CPR-like preconditioners).

Once the Schur complement is computed, to properly define the block preconditioner we need:

- the preconditioner for  $A_{00}$  (any of the above listed single-matrix preconditioner);
- the preconditioner for  $S$  (any of the above listed single-matrix preconditioner);
- the application strategy. This can be:
  - diagonal: none of the coupling terms is used;
  - upper triangular: only the upper triangular coupling term is used;
  - lower-upper triangular: both coupling terms are used.

Moreover, a block scaling is available. Feasible options are:

- none: keep the original scaling;
- Frobenius norm: equilibrate Frobenius norm of the diagonal blocks;
- user provided.

## 5.10 Numerical Methods

This section describes the specification of numerical methods used by solvers.

Name	Type	Default	Description
FiniteElements	node	unique	<i>Element: FiniteElements</i>
FiniteVolume	node	unique	<i>Element: FiniteVolume</i>

### 5.10.1 Finite Element Discretization

We are currently refactoring the finite element infrastructure, and will update the documentation soon to reflect the new structure.

### 5.10.2 Finite Volume Discretization

Two different finite-volume discretizations are available to simulate single-phase flow in GEOSX, namely, a standard cell-centered TPFA approach, and a hybrid finite-volume scheme relying on both cell-centered and face-centered degrees of freedom. The key difference between these two approaches is the computation of the flux, as detailed below.

#### Standard cell-centered TPFA FVM

This is the standard scheme implemented in the *SinglePhaseFVM* flow solver. It only uses cell-centered degrees of freedom and implements a Two-Point Flux Approximation (TPFA) for the computation of the flux. The numerical flux is obtained using the following expression for the mass flux between cells  $K$  and  $L$ :

$$F_{KL} = \Upsilon_{KL} \frac{\rho^{upw}}{\mu^{upw}} (p_K - p_L - \rho^{avg} g(d_K - d_L)),$$

where  $p_K$  is the pressure of cell  $K$ ,  $d_K$  is the depth of cell  $K$ , and  $\Upsilon_{KL}$  is the standard TPFA transmissibility coefficient at the interface. The fluid density,  $\rho^{upw}$ , and the fluid viscosity,  $\mu^{upw}$ , are upwinded using the sign of the potential difference at the interface.

This is currently the only available discretization in the *Compositional Multiphase Flow Solver*.

#### Hybrid FVM

This discretization scheme overcomes the limitations of the standard TPFA on non K-orthogonal meshes. The hybrid finite-volume scheme—equivalent to the well-known hybrid Mimetic Finite Difference (MFD) scheme—remains consistent with the pressure equation even when the mesh does not satisfy the K-orthogonality condition. This numerical scheme is currently implemented in the *SinglePhaseHybridFVM* solver.

The hybrid FVM scheme uses both cell-centered and face-centered pressure degrees of freedom. The one-sided face flux,  $F_{K,f}$ , at face  $f$  of cell  $K$  is computed as:

$$F_{K,f} = \frac{\rho^{upw}}{\mu^{upw}} \tilde{F}_{K,f},$$

where  $\tilde{F}_{K,f}$  reads:

$$\tilde{F}_{K,f} = \sum_{f'} \Upsilon_{ff'} (p_K - \pi_f - \rho_K g(d_K - d_f)).$$

In the previous equation,  $p_K$  is the cell-centered pressure,  $\pi_f$  is the face-centered pressure,  $d_K$  is the depth of cell  $K$ , and  $d_f$  is the depth of face  $f$ . The fluid density,  $\rho^{upw}$ , and the fluid viscosity,  $\mu^{upw}$ , are upwinded using the sign of  $\tilde{F}_{K,f}$ . The local transmissibility  $\Upsilon$  of size  $n_{local\ faces} \times n_{local\ faces}$  satisfies:

$$NK = \Upsilon C$$

Above,  $N$  is a matrix of size  $n_{local\ faces} \times 3$  storing the normal vectors to each face in this cell,  $C$  is a matrix of size  $n_{local\ faces} \times 3$  storing the vectors from the cell center to the face centers, and  $K$  is the permeability tensor. The local transmissibility matrix,  $\Upsilon$ , is currently computed using the quasi-TPFA approach described in Chapter 6 of this [book](#). The scheme reduces to the TPFA discretization on K-orthogonal meshes but remains consistent when the mesh does not satisfy this property. The mass flux  $F_{K,f}$  written above is then added to the mass conservation equation of cell  $K$ .

In addition to the mass conservation equations, the hybrid FVM involves algebraic constraints at each mesh face to enforce mass conservation. For a given interior face  $f$  between two neighboring cells  $K$  and  $L$ , the algebraic constraint reads:

$$\tilde{F}_{K,f} + \tilde{F}_{L,f} = 0.$$

We obtain a numerical scheme with  $n_{cells}$  cell-centered degrees of freedom and  $n_{faces}$  face-centered pressure degrees of freedom. The system involves  $n_{cells}$  mass conservation equations and  $n_{faces}$  face-based constraints. The linear systems can be efficiently solved using the MultiGrid Reduction (MGR) preconditioner implemented in the Hypre linear algebra package.

The implementation of the hybrid FVM scheme for *Compositional Multiphase Flow Solver* is in progress.

## 5.11 Parallel Partitioning

Parallel GEOSX simulations involves multiple `partitions` and there are `ghost` objects in each partition. Users need to understand these concepts to effectively design models and visualize results.

### 5.11.1 Partition and ghosting : simple examples

A model, or more strictly, a computational domain, is stored in a distributed fashion among many processors. In the following simple example, the computational domain has 10 cells and the simulation involves two processors. The first processor, “partition 0” (“0” is called the “rank” of the processor) owns the first five cells (0 to 4) while “partition 1” owns 5 to 9. When the whole domain is divided into partitions, each partition will number the cells and other objects such as nodes, faces, and edges in the partition. Therefore, in both partitions, the cells IDs start from zero. Element 0 in partition 1 is cell 5 (or the sixth cell) of the original domain. In parallel computing, each partition does not only need to know information about its own cells, but it also needs to know information about some cells owned by the neighbor partitions if these cells are directly connected to objects in this partition. For example, cell 0 in partition 1 (i.e. cell 5 in the original whole domain) is connected to cell 4 in partition 0. Therefore, partition 0 will keep a copy of this cell (including the data associated with this cell) which is synchronized with the corresponding information in the partition that actually owns this cell.

In summary, a partition owns a number of cells and other objects (e.g. faces) and also keeps copies of objects from neighbor partitions. Partitioning is automatically handled by GEOSX once the user specifies how the domain should be divided.

The following figure show the partitioning of a simple mesh. Real nodes appear as solid red circles in the owning partition and ghost nodes are shown as hollow circles.

This concept of ghosting and communications between owned cells and ghost cells can also be applied to the other types of elements in GEOSX (Faces, Edges, Nodes). The next figure summarizes the way nodes, edges, faces and cells are ghosted.

## 5.11.2 Specifying partitioning pattern

### Cartesian partitioning

In the command line to run GEOSX, the user can specify the partitioning pattern by adding the following switches:

- `-x`, `--x-partitions` - Number of partitions in the x-direction
- `-y`, `--y-partitions` - Number of partitions in the y-direction
- `-z`, `--z-partitions` - Number of partitions in the z-direction

### Graph-based partitioning

The Graph-based partitioning is used only when importing external meshes using the `PAMELAMesh` (see [Tutorial 3: Regions and Property Specifications](#) section for more details using external meshes). While importing the mesh, PAMELA computes the graph of connectivity between all the volume elements of the mesh. The partitioning is then done using the `METIS` library. The graph is not weighted so the expected result is as mesh divided in  $n$  parts, with  $n$  being the number of MPI ranks used for simulation containing a similar amount of cells.

## 5.11.3 Ghost ranks

Each object (node, edge, face, or cell) has a `ghost_rank` attribute, stored in the `ghostRank` field. If a object does not appear in any other partition as a ghost, its ghost rank is a large negative number, `-2.14e9` in a typical system. If a object is real (owned by the current partition) but exists in other partitions as ghosts, its ghost rank is `-1`. The ghost rank of a ghost object is the rank of the partition that owns the corresponding real object.

## 5.11.4 Considerations for visualization

In VisIt, a partition is called a `domain`. The ID of a domain is the rank of the corresponding partition in GEOSX plus one. VisIt would display all elements/objects regardless if they are real or ghosts. As information about a ghost is synchronized with the real object, VisIt just overlaying the same images on top of each other. The user would not perceive the overlapping between partitions unless the models are shown as semi-transparent entities. Note that if ghosts are not hidden, results from a `query` operation, such as summation of variable values, would be wrong due to double-counting. Therefore, it is a good practice or habit to hide ghost objects using `ghostRank` as a filter.

If the visualization method involves interpolation, such as interpolating a zonal field into a nodal field or generating contours, the interpretation near partition boundaries is not accurate.

## 5.12 Outputs

This section describes how outputs are handled by GEOSX

The outputs are defined in a `<Outputs>` XML block.

There are three available formats to output the results of a simulation: `SILO`, `VTK`, and Time History output into simple dataset `HDF5` files which are consumable by post-processing scripts..

## 5.12.1 Defining an output

### SILO Output

The SILO output is defined through the `<Silo>` XML node (subnode of `<Outputs>` XML block) as shown here:

```
<Outputs>
  <Silo name="siloOutput" />
</Outputs>
```

The parameter options are listed in the following table:

Name	Type	De- fault	Description
childDirectory	string		Child directory path
fieldNames	string array		Names of the fields to output. If this attribute is specified, GEOSX outputs all (and only) the fields specified by the user, regardless of their <i>plotLevel</i>
name	string	required	A name is required for any non-unique nodes
onlyPlotSpecifiedFieldNames	integer	0	If this flag is equal to 1, then we only plot the fields listed in <i>fieldNames</i> . Otherwise, we plot all the fields with the required <i>plotLevel</i> , plus the fields listed in <i>fieldNames</i>
parallelThreads	integer	1	Number of plot files.
plotFileRoot	string	plot	(no description available)
plotLevel	integer	1	(no description available)
writeCellElementMesh	integer	1	(no description available)
writeEdgeMesh	integer	0	(no description available)
writeFEM-Faces	integer	0	(no description available)
writeFaceElementMesh	integer	1	(no description available)

### VTK Output

The VTK output is defined through the `<VTK>` XML node (subnode of `<Outputs>` XML block) as shown here:

```
<Outputs>
  <VTK name="vtkOutput" />
</Outputs>
```

The parameter options are listed in the following table:

Name	Type	Default	Description
childDirectory	string		Child directory path
fieldNames	string_array	{ }	Names of the fields to output. If this attribute is specified, GEOSX outputs all the fields specified by the user, regardless of their <i>plotLevel</i>
format	geosx_vtk_VTKOutputFormat	binary	Output data format. Valid options: binary, ascii
name	string	required	A name is required for any non-unique nodes
onlyPlot-Specified-FieldNames	integer	0	If this flag is equal to 1, then we only plot the fields listed in <i>fieldNames</i> . Otherwise, we plot all the fields with the required <i>plotLevel</i> , plus the fields listed in <i>fieldNames</i>
outputRegionType	geosx_vtk_VTKRegionType	cell	Output region types. Valid options: cell, well, surface, all
parallelThreads	integer	1	Number of plot files.
plotFileRoot	string	VTK	Name of the root file for this output.
plotLevel	integer	1	Level detail plot. Only fields with lower of equal plot level will be output.
writeFEM-Faces	integer	0	(no description available)

## TimeHistory Output

The TimeHistory output is defined through the `<TimeHistory>` XML node (subnode of `<Outputs>` XML block) as shown here:

```
<Outputs>
  <TimeHistory name="timeHistoryOutput" sources="{/Tasks/collectionTask}" filename=
    ↪ "timeHistory" />
</Outputs>
```

The parameter options are listed in the following table:

Name	Type	Default	Description
childDirectory	string		Child directory path
filename	string	TimeHistory	The filename to which to write time history output.
format	string	hdf	The output file format for time history output.
name	string	required	A name is required for any non-unique nodes
parallelThreads	integer	1	Number of plot files.
sources	string_array	required	A list of collectors from which to collect and output time history information.

In order to properly collect and output time history information the following steps must be accomplished:

1. Specify one or more collection tasks using the *Tasks Manager*.
2. Specify a *TimeHistory Output* using the collection task(s) as source(s).
3. Specify an event in the *Event Management* to trigger the collection task(s).

4. Specify an event in the *Event Management* to trigger the output.

Note: Currently if the collection and output events are triggered at the same simulation time, the one specified first will also trigger first. Thus in order to output time history for the current time in this case, always specify the time history collection events prior to the time history output events.

### 5.12.2 Triggering the outputs

The outputs can be triggered using the *Event Management*. It is recommended to use a `<PeriodicEvent>` to output results with a defined frequency:

```
<PeriodicEvent name="outputs"
               timeFrequency="5000.0"
               targetExactTimestep="1"
               target="/Outputs/siloOutput" />
```

The keyword `target` has to match with the name of the `<Silo>`, `<VTK>`, or `<TimeHistory>` node.

### 5.12.3 Visualisation of the outputs

We suggest the use of [VisIT](#), [Paraview](#), and [Matplotlib](#) to visualize the outputs.

#### Visualizing Silo outputs with VisIT

If the `<Silo>` XML node was defined, GEOSX writes the results in a folder called `siloFiles`.

In VisIT :

1. File > Open file...
2. On the right panel, browse to the `siloFiles` folder.
3. On the left panel, select the file(s) you want to visualize. Usually, one file is written according the frequency defined in the `timeFrequency` keyword of the Event that has triggered the output.
4. To load fields, use the “Add” button and browse to the fields you want to plot.
5. To plot fields, use the “Draw” button.

Please consult the [VisIT](#) documentation for further explanations on its usage.

#### Visualizing VTK outputs with VisIT

If the `<VTK>` XML node was defined, GEOSX writes the results in a folder named after the `plotFileRoot` attribute (default = `vtkOutputs`). For problems with multiple active regions (e.g. *Hydraulic Fracturing*), additional work may be required to ensure that `vtk` files can be read by VisIt. Options include:

1. Using a VisIt macro / python script to convert default multi-region `vtk` files (see `GEOSX/src/coreComponents/python/visitMacros/visitVTKConversion.py`).
2. Using the `outputRegionType` attribute to output separate sets of files per region. For example:

```
<Problem>
<Events>
  <!-- Use nested events to trigger both vtk outputs -->
  <PeriodicEvent
```

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```

name="outputs"
timeFrequency="2.0"
targetExactTimestep="0">
  <PeriodicEvent
    name="outputs_cell"
    target="/Outputs/vtkOutput_cell"/>
  <PeriodicEvent
    name="outputs_surface"
    target="/Outputs/vtkOutput_surface"/>
  </PeriodicEvent>
</Events>

<Outputs>
  <VTK
    name="vtkOutput_cell"
    outputRegionType="cell"
    plotFileRoot="vtk_cell"/>

  <VTK
    name="vtkOutput_surface"
    outputRegionType="surface"
    plotFileRoot="vtk_surface"/>
</Outputs>
</Problem>

```

In VisIt, the prepared files can be visualized as follows:

1. Click the Open icon (or select File/Open from the top menu).
2. Enter the vtk output folder in the left-hand panel (these will be separate if you use option 2 above).
3. Select the desired \*.vtm group in the right-hand panel (by double-clicking or selecting OK).
4. (Repeat for any additional datasets/regions.)
5. In the main window, select the desired dataset from the Active Sources dropdown menu.
6. Click on the Add icon, and select a parameter to visualize from the mesh, pseudocolor, or vector plot options
7. At some point you may be prompted to create a Database Correlation. If you select Yes, then VisIt will create a new time slider, which can be used to change the time state while keeping the various aspects of the simulation synchronized.
8. Finally, click the Draw button and use the time slider to visualize the output.

Please consult the [VisIT](#) documentation for further explanations on its usage.

## Visualizing VTK outputs with Paraview

If the <VTK> XML node was defined, GEOSX writes a folder and a .pvd file named after the string defined in name keyword.

The .pvd file contains references to the .pvtu files. One .pvtu file is output according the frequency defined in the timeFrequency keyword of the Event that has triggered the output.

One .pvtu contains references to .vtu files. There is as much .vtu file as there were MPI processes used for the computation.

All these files can be opened with paraview. To have the whole results for every output time steps, you can open the .pvd file.

## Visualizing TimeHistory outputs with Matplotlib

If the `<TimeHistory>` XML node was defined, GEOSX writes a file named after the string defined in the `filename` keyword and formatted as specified by the string defined in the `format` keyword (only `HDF5` is currently supported).

The TimeHistory file contains the collected time history information from each specified time history collector. This information includes datasets for the time itself, any metadata sets describing index association with specified collection sets, and the time history information itself.

It is recommended to use `Matplotlib` and format-specific accessors (like `H5PY` for `HDF5`) to access and easily plot the time history data.

## 5.13 pygeosx — GEOSX in Python

GEOSX can be manipulated and executed through a Python script.

High-level control of GEOSX is managed through the top-level `pygeosx` functions, like `initialize` and `run`. GEOSX's data can be manipulated by getting `pylvarray` views of `LvArray` objects living in GEOSX's data repository. These `pylvarray` views are fetched by calling `Wrapper.value()` after getting a `Wrapper` out of the data repository.

**Warning:** The `pygeosx` module provides plenty of opportunities to crash Python. See the Segmentation Faults section below.

Only Python 3 is supported.

### 5.13.1 Module Functions

`pygeosx.initialize(rank, args)`

Initialize GEOSX for the first time, with a rank and command-line arguments.

This function should only be called **once**. To reinitialize, use the `reinit` function.

Generally the rank is obtained from the `mpi4py` module and the arguments are obtained from `sys.argv`.

Returns a `Group` representing the `ProblemManager` instance.

`pygeosx.reinit(args)`

Reinitialize GEOSX with a new set of command-line arguments.

Returns a `Group` representing the `ProblemManager` instance.

`pygeosx.apply_initial_conditions()`

Apply the initial conditions.

`pygeosx.finalize()`

Finalize GEOSX. After this no calls into `pygeosx` or to MPI are allowed.

`pygeosx.run()`

Enter the GEOSX event loop.

Runs until hitting a breakpoint defined in the input deck, or until the simulation is complete.

Returns one of the state constants defined below.

### 5.13.2 GEOSX State

`pygeosx.UNINITIALIZED`

`pygeosx.INITIALIZED`

`pygeosx.READY_TO_RUN`

This state indicates that GEOSX still has time steps left to run.

`pygeosx.COMPLETED`

This state indicates that GEOSX has completed the current simulation.

### 5.13.3 Module Classes

**class** `pygeosx.Group`

Python interface to `geosx::dataRepository::Group`.

Used to get access to other groups, and ultimately to get wrappers and convert them into Python views of C++ objects.

**groups** ()

Return a list of the subgroups.

**wrappers** ()

Return a list of the wrappers.

**get\_group** (*path*)

**get\_group** (*path*, *default*)

Return the `Group` at the relative path *path*; *default* is optional. If no group exists and *default* is not given, raise a `ValueError`; otherwise return *default*.

**get\_wrapper** (*path*)

**get\_wrapper** (*path*, *default*)

Return the `Wrapper` at the relative path *path*; *default* is optional. If no `Wrapper` exists and *default* is not given, raise a `ValueError`; otherwise return *default*.

**register** (*callback*)

Register a callback on the physics solver.

The callback should take two arguments: the `CRSMatrix` and the array.

Raise `TypeError` if the group is not the `Physics` solver.

**class** `pygeosx.Wrapper`

Python interface to `geosx::dataRepository::WrapperBase`.

Wraps a generic C++ object. Use `repr` to get a description of the type.

**value** ()

Return a view of the wrapped value, or `None` if it cannot be exported to Python.

A breakdown of the possible return types:

- Instance of a `pylvarray` class If the wrapped type is one of the `LvArray` types that have a Python wrapper type.
- 1D `numpy.ndarray` If the wrapped type is a numeric constant. The returned array is a shallow copy and has a single entry.
- `str` If the wrapped type is a `std::string` this returns a copy of the string.

- list of str If the wrapped type is a `LvArray::Array< std::string, 1, ... >` or a `std::vector< std::string >`. This is a copy.
- None If the wrapped type is not covered by any of the above.

### 5.13.4 Segmentation Faults

Improper use of this module and associated programs can easily cause Python to crash. There are two main causes of crashes. Both can be avoided by following some general guidelines.

#### Stale Numpy Views

The `pylvarray` classes (which may be returned from `Wrapper.value()`) provide various ways to get Numpy views of their data. However, those views are only valid as long as the `LvArray` object's buffer is not reallocated. The buffer may be reallocated by invoking methods (the ones that require the `pylarray.RESIZEABLE` permission) or by calls into `pygeosx`. It is strongly recommended that you do not keep Numpy views of `LvArray` objects around after calls to `pygeosx`.

```
my_array = pygeosx.get_wrapper("path").value()
view = my_array.to_numpy()
my_array.resize(1000)
print(view) # segfault
```

#### Destroyed LvArray C++ objects

As mentioned earlier, the classes defined in this module cannot be created in Python; `pygeosx` must create an `LvArray` object in C++, then create a `pylvarray` view of it. However, the Python view will only be valid as long as the underlying `LvArray` C++ object is kept around. If that is destroyed, the Python object will be left holding an invalid pointer and subsequent attempts to use the Python object will cause undefined behavior. Unfortunately, `pygeosx` may destroy `LvArray` objects without warning. It is therefore strongly recommended that you do not keep `pylvarray` objects around after calls to `pygeosx`. The following code snippet, for instance, could segfault:

```
my_array = pygeosx.get_wrapper("path").value()
pygeosx.run()
view = my_array.to_numpy() # segfault
```

## 5.14 Indices and tables

- [genindex](#)
- [modindex](#)
- [search](#)

Welcome to the GEOSX developer guide.

## 6.1 Contributing

### 6.1.1 Code style

#### Introduction

GEOSX is written in standard c++14. In general, target platforms are:

- Linux
- Mac OS X

Currently, our CI/CD system tests on these platforms:

- Ubuntu 18.04, with gcc 8.0 and clang 8.0.0 + cuda10.1.243
- Centos 7.6.1810, with gcc 8.3.1 + cuda10.1.243
- Centos 7.7, with clang 9.0.0
- Mac OS X, with xcode 11.2

#### Naming Conventions

##### File Names

- File names should be [PascalCase](#).
- C++ header files are always named with a file extension of \*.hpp.
- C++ header implementation files, which contain templated or inline function definitions, are always named \*Helpers.hpp.

- C++ source files are always named with a file extension of \*.cpp.
- C++ class declarations and definitions are contained files with identical names, except for the extensions.
- C++ free function headers and source files are declared/defined in files with identical names, except for the extension.

For example, a class named “Foo” may be declared in a file named “Foo.hpp”, with inline/templated functions defined in “FooHelpers.hpp”, with the source implementation contained in Foo.cpp.

**Warning:** There should not be identical filenames that only differ by case. Some filesystems are not case-sensitive, and worse, some filesystems such as MacOSX are case-preserving but not case sensitive.

### Function Names

Function and member function names should be `camelCase`.

### Variable Names

Variables should be `camelCase`.

### Member Names

Member data should be `camelCase` prefix with “m\_” (i.e. `double m_dataVariable;`)

### Class/Struct Names

Please use `PascalCase` for typenames (i.e. classes)

```
class MyClass;
```

```
class MyClass
{
    double m_doubleDataMember;
    int m_integerDataMember;
}
```

### Alias/Typedef Names

Alias and typedefs should be the case of the underlying type that they alias. If no clear format is apparent, as is the case with `double`, then use `camelCase`

### Namespace Names

Namespaces names are all lower `camel case`.

## Example

One example of would be a for a class named “Foo”, the declaration would be in a header file named “Foo.hpp”

```
/*
 * Foo.hpp
 */

namespace bar
{

class Foo
{
public:
    Foo();
private:
    double m_myDouble;
}
}
```

and a source file named “Foo.cpp”

```
/*
 * Foo.cpp
 */

namespace bar
{
    Foo::Foo() :
        m_myDouble(0.0)
    {
        // some constructor stuff
    }
}
```

## Const Keyword

1. All functions and accessors should be declared as “const” functions unless modification to the class is required.
2. In the case of accessors, both a “const” and “non-const” version should be provided.
3. The const keyword should be placed in the location read by the compiler, which is right to left.

The following examples are provided:

```
int a=0; // regular int
int const b = 0; // const int
int * const c = &a; // const pointer to non const int
int const * const d = &b; // const pointer to const int
int & e = a; // reference to int
int const & f = b; // reference to const int
```

## Code Format

GEOSX applies a variant of the [BSD/Allman Style](#). Key points to the GEOSX style are:

1. Opening braces (i.e. “{”) go on the next line of any control statement, and are not indented from the control statement.

2. NO TABS. Only spaces. In case it isn't clear ... NO TABS!
3. 2-space indentation

```
for( int i=0 ; i<10 ; ++i )
{
    std::cout << "blah" << std::endl;
}
```

4. Try to stay under 100 character line lengths. To achieve this apply these rules in order
5. Align function declaration/definitions/calls on argument list
6. Break up return type and function definition on new line
7. Break up scope resolution operators

```
void
SolidMechanics_LagrangianFEM::
TimeStepExplicit( real64 const& time_n,
                  real64 const& dt,
                  const int cycleNumber,
                  DomainPartition * const domain )
{
    code here
}
```

As part of the continuous integration testing, this GEOSX code style is enforced via the uncrustify tool. While quite extensive, uncrustify does not enforce every example of the preferred code style. In cases where uncrustify is unable to enforce code style, it will ignore formatting rules. In these cases it is acceptable to proceed with pull requests, as there is no logical recourse.

## Header Guards

Header guard names should consist of the name *GEOSX*, followed by the component name (e.g. *dataRepository*), and finally the name of the header file. All characters in the macro should be capitalized.

### 6.1.2 Git Workflow

The GEOSX project is hosted on github [here](#). For instructions on how to clone and build GEOSX, please refer to the *Quick Start Guide*. Consider consulting <https://try.github.io/> for practical references on how to use git.

## Git Credentials

Those who want to contribute to GEOSX should setup SSH keys for authentication, and connect to github through SSH as discussed in [this article](#). Before going further, you should [test your ssh connection](#). If it fails (perhaps because of your institution's proxy), you may consider the [personal access token option](#) as an alternative.

## Downloading the Code

Once you have created an `ssh-key` and you have added it to your *Github* account you can download the code through SSH. The following steps clone the repository into `your_geosx_dir`:



```
git clone git@github.com:GEOSX/GEOSX.git your_geosx_dir
cd your_geosx_dir
git lfs install
git submodule init
git submodule update
```

If all goes well, you should have a complete copy of the GEOSX source at this point. The most common errors people encounter here have to do with Github not recognizing their authentication settings.

## Branching Model

The branching model used in GEOSX is a modified [Gitflow](#) approach, with some modifications to the merging strategy, and the treatment of release branches, and hotfix branches.

In GEOSX, there are two main branches, `release` and `develop`. The `develop` branch serves as the main branch for the development of new features. The `release` branch serves as the “stable release” branch. The remaining branch types are described in the following subsections.

---

**Note:** The early commits in GEOSX (up to version 0.2) used a pure [Gitflow](#) approach for merging feature branches into `develop`. This was done without cleaning the commit history in each feature branch prior to the merge into `develop`, resulting in an overly verbose history. Furthermore, as would be expected, having many active feature branches resulted in a fairly wide (spaghetti) history. At some point in the development process, we chose to switch primarily to a squash-merge approach which results in a linear `develop` history. While this fixes the spaghetti history, we do potentially lose important commit history during the development process. Options for merging are discussed in the following sections.

---

## Feature Branches

New developments (new features or modifications to features) are branched off of `develop` into a feature branch. The naming of feature branches should follow `feature/[developer]/[branch-description]` if you expect that only a single developer will contribute to the branch, or `feature/[branch-description]` if you expect it will be a collaborative effort. For example, if a developer named `neo` were to add or modify a code feature expecting that they would be the only contributor, they would create a branch using the following commands to create the local branch and push it to the remote repository:

```
git checkout -b feature/neo/freeYourMind
git push -u origin feature/neo/freeYourMind
```

However if the branch is a collaborative branch amongst many developers, the appropriate commands would be:

```
git checkout -b feature/freeYourMind
git push -u origin feature/freeYourMind
```

When feature branches are ready to be merged into `develop`, a Pull Request should be created to perform the review and merging process.

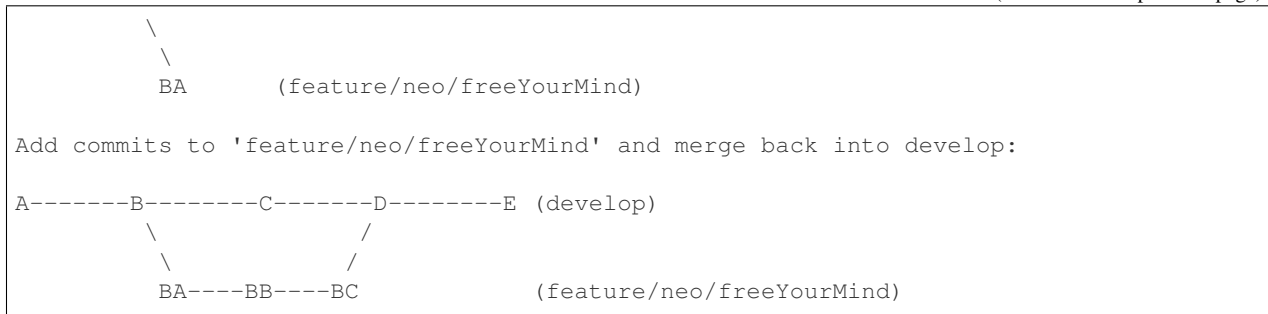
An example lifecycle diagram for a feature branch:

```
create new feature branch:
git checkout -b feature/neo/freeYourMind

A-----B-----C (develop)
```

(continues on next page)

(continued from previous page)



See below for details about [Submitting a Pull Request](#).

## Bugfix Branches

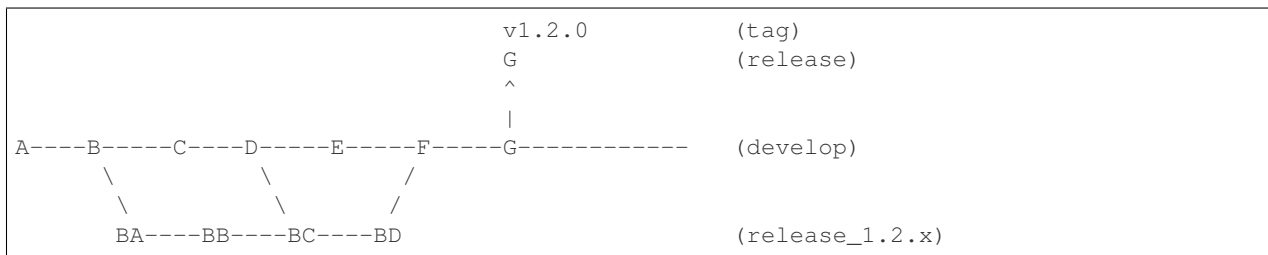
Bugfix branches are used to fix bugs that are present in the develop branch. A similar naming convention to that of the feature branches is used, replacing “feature” with “bugfix” (i.e. bugfix/neo/squashAgentSmith). Typically, bugfix branches are completed by a single contributor, but just as with the feature branches, a collaborative effort may be required resulting in dropping the developer name from the branch name.

When bugfix branches are ready to be merged into develop, a Pull Request should be created to perform the review and merging process. See below for details about [Submitting a Pull Request](#).

## Release Candidate Branches

When develop has progressed to a point where we would like to create a new release, we will create a release candidate branch with the name consisting of release\_major.minor.x number, where the x represents the sequence of patch tags that will be applied to the branch. For instance if we were releasing version 1.2.0, we would name the branch release\_1.2.x. Once the release candidate is ready, it is merged back into develop. Then the develop branch is merged into the release branch and tagged. From that point the release branch exists to provide a basis for maintaining a stable release version of the code. Note that the absence of hotfix branches, the history for release and develop would be identical.

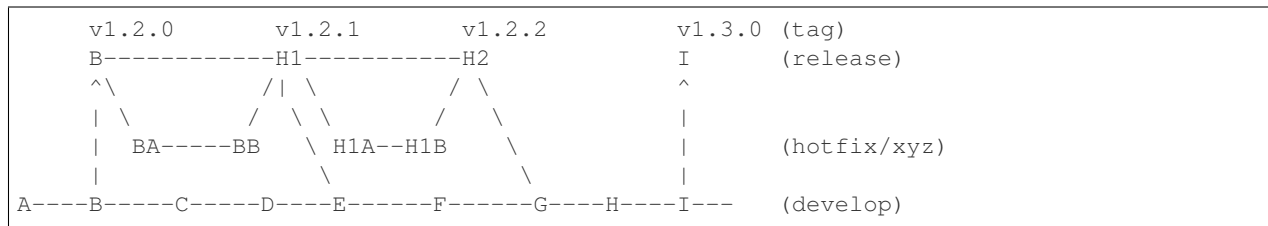
An example lifecycle diagram for a release candidate branch:



## Hotfix Branches

A hotfix branch fixes a bug in the release branch. It uses the same naming convention as a bugfix branch. The main difference with a bugfix branch is that the primary target branch is the release branch instead of develop. As a soft policy, merging a hotfix into a release branch should result in a patch increment for the release sequence of tags. So if a hotfix was merged into release with a most recent tag of 1.2.1, the merged commit would be tagged with 1.2.2. Finally, at some point prior to the next major/minor release, the release branch should be merged back into develop to incorporate any hotfix changes into develop.

An example lifecycle diagram for hotfix branches:



## Documentation Branches

A `docs` branch is focused on writing and improving the documentation for GEOSX. The use of the `docs` branch name root applies to both sphinx documentation and doxygen documentation. The `docs` branch follows the same naming conventions as described in the [Feature Branches](#) section. The html produced by a documentation branch should be proofread using sphinx/doxygen prior to merging into `develop`.

## Keeping Your Branch Current

Over the course of a long development effort in a single feature branch, a developer may need to either merge `develop` into their feature branch, or rebase their feature branch on `develop`. We do not have a mandate on how you keep your branch current, but we do have guidelines on the branch history when merging your branch into `develop`. Typically, merging `develop` into your branch is the easiest approach, but will lead to a complex relationship with `develop` with multiple interactions... which can lead to a confusing history. Conversely, rebasing your branch onto `develop` is more difficult, but will lead to a linear history within the branch. For a complex history, we will perform a squash merge into `develop`, thereby the work from the branch will appear as a single commit in `develop`. For clean branch histories where the individual commits are meaningful and should be preserved, we have the option to perform a merge commit in with the PR is merged into `develop`, with the addition of a merge commit, thus maintaining the commit history.

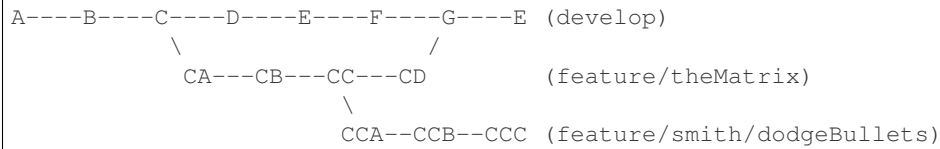
## Branching off of a Branch

During the development processes, sometimes it is appropriate to create a branch off of a branch. For instance, if there is a large collaborative development effort on the branch `feature/theMatrix`, and a developer would like to add a self-contained and easily reviewable contribution to that effort, he/she should create a branch as follows:

```
git checkout feature/theMatrix
git checkout -b feature/smith/dodgeBullets
git push -u origin feature/smith/dodgeBullets
```

If `feature/smith/dodgeBullets` is intended to be merged into `feature/theMatrix`, and the commit history of `feature/theMatrix` is not changed via `git rebase`, then the process of merging the changes back into `feature/theMatrix` is fairly standard.

However, if `feature/theMatrix` is merged into `develop` via a squash merge, and then smith would like to merge `feature/smith/dodgeBullets` into `develop`, there is a substantial problem due to the diverged history of the branches. Specifically, `feature/smith/dodgeBullets` branched off a commit in `feature/theMatrix` that does not exist in `develop` (because it was squash-merged). For simplicity, let us assume that the commit hash that `feature/smith/dodgeBullets` originated from is CC, and that there were commits CA, CB, CC, CD in `feature/theMatrix`. When `feature/theMatrix` was squash-merged, all of the changes appear in `develop` as commit G. To further complicate the situation, perhaps a complex PR was merged after G, resulting in E on `develop`. The situation is illustrated by:



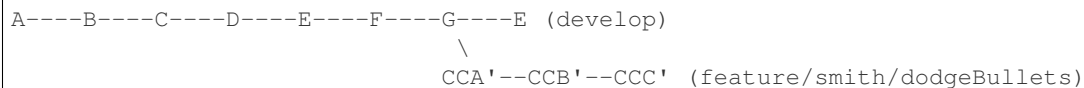
In order to successfully merge `feature/smith/dodgeBullets` into `develop`, all commits present in `feature/smith/dodgeBullets` after `CC` must be included, while discarding `CA`, `CB`, which exist in `feature/smith/dodgeBullets` as part of its history, but not in `develop`.

One “solution” is to perform a `git rebase --onto` of `feature/smith/dodgeBullets` onto `develop`. Specifically, we would like to rebase `CCA`, `CCB`, `CCC` onto `G`, and proceed with our development of `feature/smith/dodgeBullets`. This would look like:

```

git checkout develop
git pull
git checkout feature/smith/dodgeBullets
git rebase -onto G CC
  
```

As should be apparent, we have specified the starting point as `G`, and the point at which we replay the commits in `feature/smith/dodgeBullets` as all commits AFTER `CC`. The result is:



Now you may proceed with standard methods for keeping `feature/smith/dodgeBullets` current with `develop`.

## Submitting a Pull Request

Once you have created your branch and pushed changes to Github, you can create a [Pull Request](#) on Github. The PR creates a central place to review and discuss the ongoing work on the branch. Creating a pull request early in the development process is preferred as it allows for developers to collaborate on the branch more readily.

---

**Note:** When initially creating a pull request (PR) on GitHub, always create it as a *draft* PR while work is ongoing and the PR is not ready for testing, review, and merge consideration.

---

When you create the initial draft PR, please ensure that you apply appropriate labels. Applying labels allows other developers to more quickly filter the live PRs and access those that are relevant to them. Always add the *new* label upon PR creation, as well as to the appropriate *type*, *priority*, and *effort* labels. In addition, please also add any appropriate *flags*.

---

**Note:** If your branch and PR will resolve any open issues, be sure to *link* them to the PR to ensure they are appropriately resolved once the PR is merged. In order to *link* the issue to the PR for automatic resolution, you must use one of the keywords followed by the issue number (e.g. resolves #1020) in either the main description of the PR, or a commit message. Entries in PR comments that are not the main description or a commit message will be ignored, and the issue will not be automatically closed. A complete list of keywords are:

- close
- closes
- closed

- fix
- fixes
- fixed
- resolve
- resolves
- resolved

For more details, see the [Github Documentation](#).

Once you are satisfied with your work on the branch, you may promote the PR out of draft status, which will allow our integrated testing suite to execute on the PR branch to ensure all tests are passing prior to merging.

Once the tests are passing – or in some cases immediately – add the *flag: ready for review* label to the PR, and be sure to tag any relevant developers to review the PR. The PR *must* be approved by reviewers in order to be merged.

Note that whenever a pull request is merged into `develop`, commits are either *squashed*, or preserved depending on the cleanliness of the history.

## Keeping Submodules Current

Whenever you switch between branches locally, pull changes from `origin` and/or merge from the relevant branches, it is important to update the submodules to move the head to the proper commit.

```
git submodule update --recursive
```

You may also wish to modify your *git pull* behavior to update your submodules recursively for you in one command, though you forfeit some control granularity to do so. The method for accomplishing this varies between git versions, but as of git 2.15 you should be able to globally configure git to accomplish this via:

```
git config --global submodule.recurse true
```

In some cases, code changes will require to rebaseline the `Integrated Tests`. If that is the case, you will need to modify the `integrated tests submodule`. Instructions on how to modify a submodule are presented in the following section.

## Working on the Submodules

Sometimes it may be necessary to modify one of the submodules. In order to do so, you need to create a pull request on the submodule repository. The following steps can be followed in order to do so.

Move to the folder of the `submodule` that you intend to modify.

```
cd submodule-folder
```

Currently the `submodule` is in detached head mode, so you first need to move to the main branch (either `develop` or `master`) on the submodule repository, pull the latest changes, and then create a new branch.

```
git checkout <main-branch>
git pull
git checkout -b <branch-name>
```

You can perform some work on this branch, *add* and *commit* the changes and then push the newly created branch to the submodule repository on which you can eventually create a pull request using the same process discussed above in *Submitting a Pull Request*.

```
git push --set-upstream origin <branch-name>
```

### Resolving Submodule Changes in Primary Branch PRs

When you conduct work on a submodule during work on a primary GEOSX branch with an open PR, the merging procedure requires that the submodule referenced by the GEOSX PR branch be consistent with the submodule in the main branch of the project. This is checked and enforced via TravisCI.

Thus, in order to merge a PR that includes modifications to submodules, the various PRs for each repository should be staged and finalized, to the point they are all ready to be merged, with higher-level PRs in the merge hierarchy having the correct submodule references for the current main branch for their repository.

Starting from the bottom of the submodule hierarchy, the PRs are resolved, after which the higher-level PRs with reference to a resolved PR must update their submodule references to point to the new main branch of the submodule with the just-resolved PR merged. After any required automated tests pass, the higher-level PRs can then be merged.

The name of the main branch of each submodule is presented in the table below.

Submodule	Main branch
blt	develop
LvArray	develop
integratedTests	develop
hdf5_interface	master
PAMELA	master
PVTPackage	master

### 6.1.3 Sphinx Documentation

#### Generating the documentation

- To generate the documentation files, you will need to install Sphinx using

```
sudo apt install python-sphinx
```

Then you can generate the documentation files with the following command

```
cd GEOSX/build-your-platform-release
make geosx_docs
```

- That will create a new folder

```
GEOSX/build-your-platform-release/html/docs/sphinx
```

which contains all the html files generated.

#### Documenting the code

The documentation is generated from restructured text files (*.rst*). Most files can be found in `src/docs/sphinx`. Files which are specific to parts of the code, like those describing a specific class, can instead be found in `docs`

subdirectory in the folder containing the source code.

Information about how to write `rst` files can be found [here](#).

## 6.1.4 Doxygen Documentation

Developer documentation of code is provided in the form of Doxygen-style comment blocks. [Doxygen](#) is a tool for generating html/xml/latex documentation for C++ code from specially marked code comments. Having concise but high quality documentation of public APIs helps both users and developers of these APIs. We use Doxygen and Ctest to enforce documentation coverage. If Doxygen produces any warnings, your pull request will fail CI checks! See [Git Workflow](#) for more on pull requests and CI.

### Accessing

There are two ways to access Doxygen documentation.

### Build locally

Prior to configuring a GEOSX build, have Doxygen installed:

```
sudo apt install doxygen
```

---

**Note:** Eventually, doxygen (version 1.8.13) is provided within the *thirdPartyLibs* repository.

---

Configure GEOSX and go the build directory:

```
cd GEOSX/build-your-platform-release
```

Build doxygen docs only:

```
make geosx_doxygen
```

Or build all docs:

```
make geosx_docs
```

Open in browser:

```
google-chrome html/doxygen_output/html/index.html
```

### On readthedocs

Go to [GEOSX documentation](#), select the version of interest, and follow the Doxygen link at the left-hand-side.

### Guidelines

#### What to document

The following entities declared in project header files within *geosx* namespace require documentation:

- all classes and structs, including public nested ones
- global functions, variables and type aliases
- public and protected member functions, variables and type aliases in classes
- preprocessor macros

Exceptions are made for:

- overrides of virtual functions in derived types
- implementation details nested in namespace *internal*
- template specializations in some cases

## How to document

The following rules and conventions are used. Some are stricter than others.

1. We use @-syntax for all Doxygen commands (e.g. *@brief* instead of *\brief*).
2. Entities such as type aliases and member variables that typically only require a brief description, can have a single-line documentation starting with *///*.
  - *@brief* is not required for single-line comments.
3. Entities such as classes and functions that typically require either detailed explanation or parameter documentation, are documented with multiline comment blocks.
  - *@brief* is required for comment blocks.
4. Brief and detailed descriptions should be complete sentences (i.e. start with a capital letter and end with a dot).
5. Prefer concise wording in *@brief*, e.g. “Does X.” instead of “This is a function that does X.”
6. All functions parameters and return values must be explicitly documented via *@param* and *@return*.
  - An exception to this rule seem to be copy/move constructor/assignment, where parameter documentation can be omitted.
7. Add *[in]* and *[out]* tags to function parameters, as appropriate.
8. Function and template parameter descriptions are not full sentences (i.e. not capitalized nor end with a dot).
9. For hierarchies with virtual inheritance, document base virtual interfaces rather than overriding implementations.
10. Documented functions cannot use *GEOSX\_UNUSED\_ARG()* in their declarations.
11. For empty virtual base implementations that use *GEOSX\_UNUSED\_ARG(x)* to remove compiler warnings, use one of two options:
  - move empty definition away (e.g. out of class body) and keep *GEOSX\_UNUSED\_ARG(x)* in definition only;
  - put *GEOSX\_UNUSED\_VAR(x)* into the inline empty body.
12. For large classes, logically group functions using member groups via *///@{* and *///@}* and give them group names and descriptions (if needed) via a *@name* comment block. Typical groups may include:
  - constructors/destructor/assignment operators;
  - getter/setter type functions;
  - overridable virtual functions;
  - any other logically coherent groups (functions related to the same aspect of class behavior).



13. In-header implementation details (e.g. template helpers) often shouldn't appear in user documentation. Wrap these into *internal* namespace.
14. Use `/// @cond DO_NOT_DOCUMENT` and `/// @endcond` tags to denote a section of public API that should not be documented for some reason. This should be used rarely and selectively. An example is in-class helper structs that must be public but that user should not refer to explicitly.

## Example

```

/// This is a documented macro
#define USEFUL_MACRO

/**
 * @brief Short description.
 * @tparam T type of input value
 * @param[in] x input value explanation
 * @return return value explanation
 *
 * Detailed description goes here.
 *
 * @note A note warning users of something unexpected.
 */
template<typename T>
int Foo( T const & x );

/**
 * @brief Class for showing Doxygen.
 * @tparam T type of value the class operates on
 *
 * This class does nothing useful except show how to use Doxygen.
 */
template<typename T>
class Bar
{
public:

    /// A documented member type alias.
    using size_type = typename std::vector<T>::size_type;

    /**
     * @name Constructors/destructors.
     */
    //@{

    /**
     * @brief A documented constructor.
     * @param value to initialize the object
     */
    explicit Bar( T t );

    /**
     * @brief A deleted, but still documented copy constructor.
     * @param an optionally documented parameter
     */
    Bar( Bar const & source ) = delete;

    /**

```

(continues on next page)

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```

    * @brief A defaulted, but still documented move constructor.
    * @param an optionally documented parameter
    */
    Bar( Bar const & source ) = default;

    /**
     * @brief A documented desctructor.
     * virtual ~Bar() = default;
     */

    ///@}

    /**
     * @name Getters for stored value.
     */
    ///@{

    /**
     * @brief A documented public member function.
     * @return a reference to contained value
     */
    T & getValue();

    /**
     * @copydoc getValue()
     */
    T const & getValue() const;

    ///@}

protected:

    /**
     * @brief A documented protected pure virtual function.
     * @param[in] x the input value
     * @param[out] y the output value
     *
     * Some detailed explanation for users and implementers.
     */
    virtual void doSomethingOverridable( int const x, T & y ) = 0;

    /// @cond DO_NOT_DOCUMENT
    // Some stuff we don't want showing up in Doxygen
    struct BarHelper
    {
    };
    /// @endcond

private:

    /// An optionally documented (not enforced) private member.
    T m_value;

};

```

## Current Doxygen

[Link to Doxygen](#)

### 6.1.5 Unit Testing

Unit testing is integral to the GEOSX development process. While not all components naturally lend themselves to unit testing (for example a physics solver) every effort should be made to write comprehensive quality unit tests.

Each sub-directory in `coreComponents` should have a `unitTests` directory containing the test sources. Each test consists of a `cpp` file whose name begins with `test` followed by a name to describe the test. Please read over the `LvArray` unit test documentation as it gives an intro to the Google Test framework and a set of best practices.

### GEOSX Specific Recommendations

An informative example is `testSinglePhaseBaseKernels` which tests the single phase flow mobility and accumulation kernels on a variety of inputs.

```
TEST( SinglePhaseBaseKernels, mobility )
{
    int constexpr NTEST = 3;

    real64 const dens[NTEST]      = { 800.0, 1000.0, 1500.0 };
    real64 const dDens_dPres[NTEST] = { 1e-5, 1e-10, 0.0 };
    real64 const visc[NTEST]       = { 5.0, 2.0, 1.0 };
    real64 const dVisc_dPres[NTEST] = { 1e-7, 0.0, 0.0 };

    for( int i = 0; i < NTEST; ++i )
    {
        SCOPED_TRACE( "Input # " + std::to_string( i ) );

        real64 mob;
        real64 dMob_dPres;

        MobilityKernel::compute( dens[i], dDens_dPres[i], visc[i], dVisc_dPres[i], mob,
        ↪ dMob_dPres );

        // compute etalon
        real64 const mob_et = dens[i] / visc[i];
        real64 const dMob_dPres_et = mob_et * (dDens_dPres[i] / dens[i] - dVisc_dPres[i] /
        ↪ visc[i]);

        EXPECT_DOUBLE_EQ( mob, mob_et );
        EXPECT_DOUBLE_EQ( dMob_dPres, dMob_dPres_et );
    }
}
```

[Source: `coreComponents/physicsSolvers/fluidFlow/unitTests/testSinglePhaseBaseKernels.cpp`]

What makes this such a good test is that it depends on very little other than kernels themselves. There is no need to involve the data repository or parse an XML file. Sometimes however this is not possible, or at least not without a significant duplication of code. In this case it is better to embed the XML file into the test source as a string instead of creating a separate XML file and passing it to the test as a command line argument or hard coding the path. One example of this is `testLaplaceFEM` which tests the laplacian solver. The embedded XML is shown below.

```

char const * xmlInput =
"<Problem>\n"
"  <Solvers gravityVector=\"{ 0.0, 0.0, -9.81 }\">\n"
"    <CompositionalMultiphaseFVM name=\"compflow\">\n"
"      logLevel=\"0\">\n"
"        discretization=\"fluidTPFA\">\n"
"          targetRegions=\"{region}\">\n"
"            temperature=\"297.15\">\n"
"              useMass=\"1\">\n"
"                \n"
"          <NonlinearSolverParameters newtonTol=\"1.0e-6\">\n"
"            newtonMaxIter=\"2\"/>\n"
"          <LinearSolverParameters solverType=\"gmres\">\n"
"            krylovTol=\"1.0e-10\"/>\n"
"        </CompositionalMultiphaseFVM>\n"
"      </Solvers>\n"
"    <Mesh>\n"
"      <InternalMesh name=\"mesh\">\n"
"        elementTypes=\"{C3D8}\">\n"
"          xCoords=\"{0, 3}\">\n"
"            yCoords=\"{0, 1}\">\n"
"              zCoords=\"{0, 1}\">\n"
"                nx=\"{3}\">\n"
"                  ny=\"{1}\">\n"
"                    nz=\"{1}\">\n"
"                      cellBlockNames=\"{cb1}\"/>\n"
"            </Mesh>\n"
"          <NumericalMethods>\n"
"            <FiniteVolume>\n"
"              <TwoPointFluxApproximation name=\"fluidTPFA\"/>\n"
"            </FiniteVolume>\n"
"          </NumericalMethods>\n"
"        <ElementRegions>\n"
"          <CellElementRegion name=\"region\" cellBlocks=\"{cb1}\" materialList=\"{fluid,
↪ rock, relperm, cappressure}\" />\n"
"        </ElementRegions>\n"
"        <Constitutive>\n"
"          <CompositionalMultiphaseFluid name=\"fluid\">\n"
"            phaseNames=\"{oil, gas}\">\n"
"              equationsOfState=\"{PR, PR}\">\n"
"                componentNames=\"{N2, C10, C20, H2O}\">\n"
"                  componentCriticalPressure=\"{34e5, 25.3e5, 14.
↪ 6e5, 220.5e5}\">\n"
"                    componentCriticalTemperature=\"{126.2, 622.0,
↪ 782.0, 647.0}\">\n"
"                      componentAcentricFactor=\"{0.04, 0.443, 0.816, 0.
↪ 344}\">\n"
"                        componentMolarWeight=\"{28e-3, 134e-3, 275e-3,
↪ 18e-3}\">\n"
"                          componentVolumeShift=\"{0, 0, 0, 0}\">\n"
"                            componentBinaryCoeff=\"{ {0, 0, 0, 0},\n"
"                              {0, 0, 0, 0},\n"
"                                {0, 0, 0, 0},\n"
"                                  {0, 0, 0, 0} }\"/>\n"
"                          <CompressibleSolidConstantPermeability name=\"rock\">\n"
"                            solidModelName=\"nullSolid\">\n"
"                              porosityModelName=\"rockPorosity\">\n"

```

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```

"         permeabilityModelName=\"rockPerm\"/>\n"
" <NullModel name=\"nullSolid\"/> \n"
" <PressurePorosity name=\"rockPorosity\" \n"
"         defaultReferencePorosity=\"0.05\" \n"
"         referencePressure = \"0.0\" \n"
"         compressibility=\"1.0e-9\"/>\n"
" <BrooksCoreyRelativePermeability name=\"relperm\" \n"
"         phaseNames=\"{oil, gas}\" \n"
"         phaseMinVolumeFraction=\"{0.1, 0.15}\" \n"
"         phaseRelPermExponent=\"{2.0, 2.0}\" \n"
"         phaseRelPermMaxValue=\"{0.8, 0.9}\"/>\n"
" <BrooksCoreyCapillaryPressure name=\"cappressure\" \n"
"         phaseNames=\"{oil, gas}\" \n"
"         phaseMinVolumeFraction=\"{0.2, 0.05}\" \n"
"         phaseCapPressureExponentInv=\"{4.25, 3.5}\" \n"
"         phaseEntryPressure=\"{0., 1e8}\" \n"
"         capPressureEpsilon=\"0.0\"/> \n"
" <ConstantPermeability name=\"rockPerm\" \n"
"         permeabilityComponents=\"{2.0e-16, 2.0e-16, 2.0e-16}\"/> \n
"
" </Constitutive>\n"
" <FieldSpecifications>\n"
"   <FieldSpecification name=\"initialPressure\" \n"
"       initialCondition=\"1\" \n"
"       setNames=\"{all}\" \n"
"       objectPath=\"ElementRegions/region/cb1\" \n"
"       fieldName=\"pressure\" \n"
"       functionName=\"initialPressureFunc\" \n"
"       scale=\"5e6\"/>\n"
"   <FieldSpecification name=\"initialComposition_N2\" \n"
"       initialCondition=\"1\" \n"
"       setNames=\"{all}\" \n"
"       objectPath=\"ElementRegions/region/cb1\" \n"
"       fieldName=\"globalCompFraction\" \n"
"       component=\"0\" \n"
"       scale=\"0.099\"/>\n"
"   <FieldSpecification name=\"initialComposition_C10\" \n"
"       initialCondition=\"1\" \n"
"       setNames=\"{all}\" \n"
"       objectPath=\"ElementRegions/region/cb1\" \n"
"       fieldName=\"globalCompFraction\" \n"
"       component=\"1\" \n"
"       scale=\"0.3\"/>\n"
"   <FieldSpecification name=\"initialComposition_C20\" \n"
"       initialCondition=\"1\" \n"
"       setNames=\"{all}\" \n"
"       objectPath=\"ElementRegions/region/cb1\" \n"
"       fieldName=\"globalCompFraction\" \n"
"       component=\"2\" \n"
"       scale=\"0.6\"/>\n"
"   <FieldSpecification name=\"initialComposition_H20\" \n"
"       initialCondition=\"1\" \n"
"       setNames=\"{all}\" \n"
"       objectPath=\"ElementRegions/region/cb1\" \n"
"       fieldName=\"globalCompFraction\" \n"
"       component=\"3\" \n"
"       scale=\"0.001\"/>\n"

```

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```

" </FieldSpecifications>\n"
" <Functions>\n"
"   <TableFunction name=\"initialPressureFunc\" \n"
"               inputVarNames=\"{elementCenter}\" \n"
"               coordinates=\"{0.0, 3.0}\" \n"
"               values=\"{1.0, 0.5}\" />\n"
" </Functions>"
"</Problem>";

```

[Source: *coreComponents/physicsSolvers/fluidFlow/unitTests/testCompMultiphaseFlow.cpp*]

## MPI

Often times it makes sense to write a unit test that is meant to be run with multiple MPI ranks. This can be accomplished by simply adding the `NUM_MPI_TASKS` parameter to `blt_add_test` in the CMake file. For example

```

blt_add_test( NAME testWithMPI
              COMMAND testWithMPI
              NUM_MPI_TASKS ${NUMBER_OF_MPI_TASKS} )

```

With this addition make test or calling `ctest` directly will run `testWithMPI` via something analogous to `mpirun -n NUMBER_OF_MPI_TASKS testWithMPI`.

## 6.1.6 Contributing Input Files

As part of the development cycle, functional input files should be introduced into the repository in order to provide 1) testing, 2) examples of proper use of the code. The input files should be modularized such that the majority of the input resides in a base file, and the variations in input are contained in specific files for integrated/smoke testing, and benchmarking. For example if we had a single input file named `myProblem.xml`, we would break it up into `myProblem_base.xml`, `myProblem_smoke.xml`, and `myProblem_benchmark.xml`. Each of the smoke/benchmark files should include the base file using an include block as follows:

```

<Included>
  <File name="./myProblem_base.xml"/>
</Included>

```

The files should be placed in the appropriate application specific subdirectory under the `GEOSX/inputFiles` directory. For example, the `beamBending` problem input files reside in the `inputFiles/solidMechanics` directory. The files then be linked to from the appropriate location in the `integratedTests` repository as described in the following section.

## 6.1.7 Integrated Tests

### About

*integratedTests* is a submodule of *GEOSX* residing at the top level of the directory structure. It will run *GEOSX* with various *.xml* files and compare the output against a baseline.

Notes: you may need to install `h5py` and `mpi4py`.

Do not forget to retrieve the baselines with the git LFS (large file storage) plugin. If the content of your baselines is something like:

```
[somebody@somewhere:/tmp]$ cat /path/to/GEOSX/integratedTests/update/run/sedov/baselines/sedov_8/0to100_restart_000000
version https://git-lfs.github.com/spec/v1 oid sha256:09bbe1e92968852cb915b971af35b0bba519fae7cf5934f3abc7b709ea76308c
size 1761208
```

then it means that the LFS plugin was not activated. `git lfs install` and `git lfs pull` should help you fix this.

## Structure

The *integratedTests* directory is composed of two main directories, *integratedTests/geosxats* and *integratedTests/update/run*. The *integratedTests/geosxats* directory contains all the machinery involved in running the tests including the main executable *integratedTests/geosxats/geosxats*. The *integratedTests/update/run* directory contains all the actual tests themselves, including the *.xml* files and the baselines.

```
- integratedTests/
  - geosxats/
    - geosxats
  - update/
    - run/
      - sedov/
        - beamBending/
          - baselines/
            - beamBending/
              - <baseline-files>
            - beamBending.ats
            - beamBending.xml
```

## Arguments

The program takes a number of arguments, the most important ones are

- The path to the *GEOSX* binary directory (`<build-dir>/bin`)
- `--workingDir WORKINGDIR` which sets the working (test) directory.
- `-N NUMNODES` specifies the number of nodes to use the default is the minimum number that the tests require.
- `-a {veryclean, rebaseline, ...}` specify a specific action, *veryclean* deletes all output files, and *rebaseline* lets you rebaseline all or some of the tests.
- `-h` prints out the *geosxats* help.

## How to Run the Tests

To run all the tests from the top level *GEOSX* directory you would do

```
integratedTests/geosxats/geosxats <build-path>/bin --workingDir integratedTests/
↪update/run
```

To run only the *sedov* tests you would do

```
integratedTests/geosxats/geosxats <build-path>/bin --workingDir integratedTests/
↪update/run/sedov
```

However if you want to run all the tests there's an easier way. In the build directory there's a symbolic link to *integratedTests/update/run* called *integratedTests* and a bash script *geosxats.sh* that wraps *integratedTests/geosxats/geosxats*

and passes it the path to the binary directory and the *integratedTests/update/run* directory along with any other command line arguments. To run this script do the following

```
cd <build-path>
./geosxats.sh
```

If the script or symbolic link is not present you will need to reconfigure by running the `config-build.py` script.

When the program has finished running you will see something like this

```
FAIL RUN : 0

UNEXPECTEDPASS : 0

FAIL RUN (OPTIONAL STEP) : 0

FAIL CHECK : 1
( beamBending )

FAIL CHECK (MINOR) : 0

TIMEOUT : 0

NOT RUN : 0

INPROGRESS : 0

FILTERED : 0

RUNNING : 0

PASSED : 3
( sedov 2D_100x100_incompr_linear 2D_100x100_incompr_linear_faceBC )

EXPECTEDFAIL : 0

SKIPPED : 0

BATCHED : 0

NOT BUILT : 0

TOTAL TIME          : 0:01:13
TOTAL PROCESSOR-TIME : 0:02:13
AVAIL PROCESSOR-TIME : 1:28:26
RESOURCE UTILIZATION : 2.51%

LONGEST RUNNING TESTS:
0:01:13 sedov
0:00:23 2D_100x100_incompr_linear_faceBC
0:00:22 2D_100x100_incompr_linear
0:00:14 beamBending

Status      : TestCase      : Directory                  : Elapsed : Resources :
↪TestStep
----- : ----- : ----- : ----- : ----- :
↪-----
FAIL CHECK : beamBending : beamBending/beamBending : 0:00:14 : 0:00:14 :
↪restartcheck
```

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```

----- : ----- : ----- : ----- : ----- : -
↪-----

Generating HTML documentation files (running 'atddoc')...
Failed to create HTML documentation in /g/g14/corbett5/geosx/mirror/integratedTests/
↪update/doc

Undocumented test problems:
beamBending 2D_100x100_incompr_linear 2D_100x100_incompr_linear_faceBC sedov

```

Ignore the error regarding the failure to create a HTML documentation file and the warning about the undocumented test problems. The only important thing is if any of the tests aren't in the *PASSED* category. For a nice summary of the results open the *test\_results.html* file in the *geosxats* working directory.

When running the tests multiple times in a row, only tests that failed to pass will run. If you would like to run all the tests again call

```
./geosxats.sh -a veryclean
```

which will delete all the generated files. Furthermore these generated files are not ignored by *git*, so until you run *veryclean* the *integratedTests* repo will register changes.

**Note:** On some development machines *geosxats* won't run parallel tests by default (e.g. on an linux laptop or workstation), and as a result many tests will be skipped. We highly recommend running tests on an MPI-aware platform.

## Output Created By a Test

Since the *beamBending* test failed let's look at it's output. The output for the *beamBending* test is stored in *integratedTests/update/run/beamBending/beamBending* directory. In addition to any files *GEOSX* itself creates you will find

- *beamBending.data* which holds all of the standard output of the various steps.
- *beamBending.err* which holds all of the standard error output of the various steps.
- *beamBending.geosx.out* which holds all of the standard output for only the *geosx* step.
- *beamBending\_restart\_000000003.restartcheck* which holds all of the standard output for only the *restartcheck* step.
- *beamBending\_restart\_000000003\_diff.hdf5* which mimics the hierarchy of the restart file and has links to the differing data datasets.

## The RestartCheck File

Currently the only manner of check that we support is a restart check, this check compares a restart file output at the end of a run against a baseline. The program that does the diff is *integratedTests/geosxats/helpers/restartcheck.py*. The program compares the two restart files and writes out a *.restart\_check* file with the results, as well as exiting with an error code if the files compare differently.

This program takes a number of arguments and they are as follows

- Regex specifying the restart file. If the regex matches multiple files the one with the greater string is selected. For example *restart\_100.hdf5* wins out over *restart\_088.hdf5*.
- Regex specifying the baseline file.
- *-r* The relative tolerance for floating point comparison, the default is 0.0.

- -a The absolute tolerance for floating point comparison, the default is 0.0.
- -e A list of regex expressions that match paths in the restart file tree to exclude from comparison. The default is [\*/commandLine].
- -w Force warnings to be treated as errors, default is false.
- -s Suppress output to stdout, default is False.

The `.restart_check` file itself starts off with a summary of the arguments. The program then compares the `.root` files and if they are similar proceeds to compare all the `.hdf5` data files.

If the program encounters any differences it will spit out an error message. An error message for scalar values looks as follows

```
Error: /datagroup_0000000/sidre/external/ProblemManager/domain/ConstitutiveManager/
↳shale/YoungsModulus
  Scalar values of types float64 and float64 differ: 22500000000.0, 10000022399.9.
```

Where the first value is the value in the test's restart file and the second is the value in the baseline.

An example of an error message for arrays is

```
Error: /datagroup_0000000/sidre/external/ProblemManager/domain/MeshBodies/mesh1/
↳Level0/nodeManager/TotalDisplacement
  Arrays of types float64 and float64 have 1836 values of which 1200 have differing_
↳values.
  Statistics of the differences greater than 0:
    max_index = (1834,), max = 2.47390764755, mean = 0.514503482629, std = 0.
↳70212888881
```

This means that the max absolute difference is 2.47 which occurs at value 1834. Of the values that are not equal the mean absolute difference is 0.514 and the standard deviation of the absolute difference is 0.702.

When the tolerances are non zero the comparison is a bit more complicated. From the *FileComparison.compareFloatArrays* method documentation

```
Entries x1 and x2 are considered equal iff
  |x1 - x2| <= ATOL or |x1 - x2| <= RTOL * |x2|.
To measure the degree of difference a scaling factor q is introduced. The goal is now_
↳to minimize q such that
  |x1 - x2| <= ATOL * q or |x1 - x2| <= RTOL * |x2| * q.
If RTOL * |x2| > ATOL
  q = |x1 - x2| / (RTOL * |x2|)
else
  q = |x1 - x2| / ATOL.
If the maximum value of q over all the entries is greater than 1.0 then the arrays_
↳are considered different and an error message is produced.
```

An sample error message is

```
Error: /datagroup_0000000/sidre/external/ProblemManager/domain/MeshBodies/mesh1/
↳Level0/nodeManager/TotalDisplacement
  Arrays of types float64 and float64 have 1836 values of which 1200 fail both the_
↳relative and absolute tests.
  Max absolute difference is at index (1834,): value = 2.07474948094, base_value =_
↳4.54865712848
  Max relative difference is at index (67,): value = 0.00215842135281, base_value =_
↳0.00591771127792
  Statistics of the q values greater than 1.0 defined by the absolute tolerance: N =_
↳1200
```

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```
max = 16492717650.3, mean = 3430023217.52, std = 4680859258.74
Statistics of the q values greater than 1.0 defined by the relative tolerance: N = 0
```

The restart check step can be run in parallel using mpi via

```
mpirun -n NUM_PROCESSES python -m mpi4py restartcheck.py ...
```

In this case rank zero reads in the restart root file and then each rank parses a subset of the data files creating a `.$RANK.restartcheck` file. Rank zero then merges the output from each of these files into the main `.restartcheck` file and prints it to standard output.

## The `.diff.hdf5` File

Each error generated in the `restartcheck` step creates a group with three children in the `_diff.df5` file. For example the error given above will generate a hdf5 group

```
/FILENAME/datagroup_0000000/sidre/external/ProblemManager/domain/MeshBodies/mesh1/
↳Level0/nodeManager/TotalDisplacement
```

with datasets `baseline`, `run` and `message` where `FILENAME` is the name of the restart data file being compared. The `message` dataset contains a copy of the error message while `baseline` is a symbolic link to the baseline dataset and `run` is a symbolic link to the dataset generated by the run. This allows for easy access to the raw data underlying the diff without data duplication. For example if you want to extract the datasets into python you could do this:

```
import h5py
file_path = "beamBending_restart_000000003_diff.hdf5"
path_to_data = "/beamBending_restart_000000011_0000000.hdf5/datagroup_0000000/sidre/
↳external/ProblemManager/domain/MeshBodies/mesh1/Level0/nodeManager/TotalDisplacement
↳"
f = h5py.File("file_path", "r")
error_message = f["path_to_data/message"]
run_data = f["path_to_data/run"][:]
baseline_data = f["path_to_data/baseline"][:]

# Now run_data and baseline_data are numpy arrays that you may use as you see fit.
rtol = 1e-10
atol = 1e-15
absolute_diff = np.abs(run_data - baseline_data) < atol
hybrid_diff = np.close(run_data, baseline_data, rtol, atol)
```

When run in parallel each rank creates a `.$RANK.diff.hdf5` file which contains the diff of each data file processed by that rank.

## The `.ats` File

The `.ats` file is a python script that describes the `TestCases` to run and steps for each `TestCase`. Each `.ats` file needs to have at least one `TestCase` and each `TestCase` needs to have at least one step.

A simple example is the `beamBending.ats` file

```
TestCase(
    name = "beamBending",
    desc = "Tests beam bending.",
```

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```
label = "auto",
owner = "Ben Corbett",
independent = True,
steps = (geosx(deck="beamBending.xml"),)
```

This creates a *TestCase* called *beamBending* with a single step that runs *GEOSX* with the *beamBending.xml* input file, a *restartcheck* step automatically follows each *geosx* step. So this file describes a test that runs the *beamBending* problem and compares the restart file against the baseline.

A slightly more complicated example is the *singlePhaseFlow.ats* file.

```
decks = ("2D_100x100_incompr_linear",
         "2D_100x100_incompr_linear_faceBC")
descriptions = ("Testing the single phase incompressible flow solver.",
               "Testing the single phase incompressible flow solver with face_
↳boundary conditions.")

for i in range(len(decks)):
    deck = decks[i]
    description = descriptions[i]
    TestCase(
        name = deck,
        desc = description,
        label = "auto",
        owner = "Ben Corbett",
        independent = True,
        steps = (geosx(deck=deck + ".xml"),)
    )
```

This creates two *TestCases* each of which runs a different problem. The *independent* parameter means that the two *TestCases* can be executed independently of each other. When a *TestCase* executes it uses its name to create a directory where all the output files are stored so if you have multiple *TestCases* in an *.ats* file it's imperative that they have unique names.

Finally there's the *sedov.ats* file which tests that starting from a restart file has no impact on the final solution.

```
import os

TestCase(
    name = "sedov",
    desc = "Test the basic sedov problem and restart capabilities.",
    label = "auto",
    owner = "Ben Corbett",
    independent = True,
    steps = (geosx(deck="sedov.xml",
                  name="0to100"),
            geosx(deck="sedov.xml",
                  name="50to100",
                  restart_file=os.path.join(testcase_name, "0to100_restart_000000050.
↳root"),
                  baseline_pattern="0to100_restart_[0-9]+\.\root",
                  allow_rebaseline=False)
            )
)
```

This creates a single *TestCase* That executes *GEOSX* twice. The first step does 100 time steps followed by a *restartcheck* step. The second *geosx* step executes the original 100 time step *xml* file but restarts using the restart

file output half way through the first run. Each *geosx* step gets its name from the *xml* file, but this can be overridden by the *name* parameter. Furthermore the default behavior is to look for a baseline in the *baselines/<TestCaseName>* directory named *TestStepName\_restart\_[0-9]+.root*, however the second step overrides this to instead compare against the “0to100” baseline. Because of this it does not allow rebaselining.

You can pass parameters to the *restartcheck* step in a dictionary passed as an argument to the *geosx* step. For example to set the tolerance you would do

```
restartcheck_params={}
restartcheck_params["atol"] = 1.5E-10
restartcheck_params["rtol"] = 1E-12

TestCase(
    name = "sedov",
    desc = "Test the basic sedov problem and restart capabilities.",
    label = "auto",
    owner = "Ben Corbett",
    independent = True,
    steps = (geosx(deck="sedov.xml",
                  name="0to100",
                  restartcheck_params=restartcheck_params))
)
```

For more info see *integratedTests/geosxats/GeosxAtsTestSteps.py* and *integratedTests/geosxats/GeosxAtsTestCase.py*

## Adding a Test

To add a new test create a new folder in the *integratedTests/update/run\** directory. At a minimum this new folder needs to include an *.ats* file. Using the *beamBending* example, after creating *beamBending.ats* the directory should look like

```
- integratedTests/update/run/beamBending/
- beamBending.ats
- beamBending.xml
```

At this point you should run the test. Assuming the *geosx* step is successful the *restartcheck* step will fail because there are no baselines. At this point the directory should look like

```
- integratedTests/update/run/beamBending/
- beamBending/
  - <geosx files>...
  - <ats files>...
- beamBending.ats
- beamBending.xml
- <ats files>...
```

Now run

```
./geosxats.sh -a rebaseline
```

and rebaseline your test. Finally run the test a second time and confirm that it passes. Note that unless you disable the *restartcheck* step you will need to output a restart file. Although not strictly necessary it is best to put the *xml* file in the main *GEOSX* repo and create an relative symbolic link to it in the test directory.

### Rebaselining Tests

Occasionally it is necessary to rebaseline one or more tests due to feature changes in the code. We suggest the following workflow:

In the GEOSX repository, create a branch with your modifications:

```
cd <GEOSX-path>
git checkout -b user/feature/newFeature
```

Add your changes, confirm it passes all the continuous integration tests, and get approval for a pull request.

Now, confirm that your integratedTests submodule is up to date:

```
git submodule
```

This will list the commit hash for all submodules. Check that the integrated tests submodule is on develop and that the commit hash is the same one as the latest GEOSX develop branch points to. If you have somehow fallen behind, go into integratedTests, checkout develop, and pull.

Now go to the integratedTests submodule and check out a branch for your new baselines. It is a good idea to name branch something similar to your feature branch so it is obvious the two branches are related.

```
cd <integratedTests-path>
git checkout -b user/rebase/newFeature
```

Go back to your GEOSX build directory and run the integrated tests

```
cd <build-path>
./geosxats.sh
```

Confirm that any tests that fail need to be **legitimately** rebaselined. Arbitrarily changing baselines defeats the purpose of the integrated tests. In your PR discussion, please identify which tests will change and any unusual behavior.

We can now actually rebaseline the tests

```
./geosxats -a rebaseline
```

You'll be prompted to confirm whether rebaselining is required for every integrated test, one at a time, via a [y/n] prompt. Make sure to only answer y to the tests that you actually want to rebaseline, otherwise correct baselines for already passing tests will still be updated and bloat your pull request and repository size.

Confirm that the rebaselines are working as expected, by cleaning the test dir and re-running the checks:

```
./geosxats -a veryclean
./geosxats
```

At this point you should pass all the integratedTests. Clean the branch and commit your changes to the baseline branch.

```
./geosxats -a veryclean
cd <integratedTests-path>
git status
git add *
git commit -m "Updating baselines"
git push
```

If you haven't already set up your local branch to point to a remote branch, you will be prompted to do so when attempting to push. You will then want to create a pull request in the integratedTests repository. Once you have merge approval for your PR, you can merge your rebase branch into integratedTests/develop.

At this point, you need to get your GEOSX `user/feature/newFeature` branch pointing to the head commit on `integratedTests/develop`. We will check out the latest version of the test repo and add it to our feature branch:

```
cd <integratedTests-path>
git checkout develop
git pull

cd <GEOSX-path>
git add integratedTests
git commit -m "Updating integratedTests hash"
git push
```

You may also want to run `git submodule` to confirm the submodule hash is what we expect, the last commit in `integratedTests/develop`. Once your feature branch passes all Continuous Integration tests, it can be successfully merged into `GEOSX/develop`.

## Tips

**Parallel Tests:** On some development machines `geosxats` won't run parallel tests by default (e.g. on an linux laptop or workstation), and as a result many baselines will be skipped. We highly recommend running tests and rebaselining on an MPI-aware platform.

**Filtering Checks:** A common reason for rebaselining is that you have changed the name of an XML node in the input files. While the baselines may be numerically identical, the restarts will fail because they contain different node names. In this situation, it can be useful to add a filter to the restart check script. If you open `integratedTests/geosxats/helpers/restartcheck.py`, at line 12 you will find a line:

```
EXCLUDE_DEFAULT = [".*/commandLine", ".*schema$", ".*globalToLocalMap"]
```

This variable contains paths to be excluded from the restart checks. For example, we recently renamed the XML block `<SystemSolverParameters/>` to `<LinearSolverParameters/>`. In doing so, we had to rebaseline every test even though we expected no numerical differences. Temporarily adding the following filter helped us rapidly check this was indeed the case:

```
EXCLUDE_DEFAULT = [".*/SystemSolverParameters", ".*LinearSolverParameters", ".*/"
↪commandLine", ".*schema$", ".*globalToLocalMap"]
```

You may find this approach useful for quickly filtering tests to distinguish between expected and unexpected failures.

## 6.1.8 Benchmarks

In addition to the integrated tests which track code correctness we have a suite of benchmarks that track performance.

### Running the benchmarks

Because performance is system specific we currently only support running the benchmarks on the LLNL machines Quartz and Lassen. If you are on either of these machines the script `benchmarks/runBenchmarks.py` can be used to run the benchmarks.

```
> python ../benchmarks/runBenchmarks.py --help
usage: runBenchmarks.py [-h] [-t TIMELIMIT] [-o TIMINGCOLLECTIONDIR]
                        [-e ERRORCOLLECTIONDIR]
```

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```

                                geosxPath outputDirectory

positional arguments:
  geosxPath          The path to the GEOSX executable to benchmark.
  outputDirectory    The parent directory to run the benchmarks in.

optional arguments:
  -h, --help          show this help message and exit
  -t TIMELIMIT, --timeLimit TIMELIMIT
                        Time limit for the entire script in minutes, the
                        default is 60.
  -o TIMINGCOLLECTIONDIR, --timingCollectionDir TIMINGCOLLECTIONDIR
                        Directory to copy the timing files to.
  -e ERRORCOLLECTIONDIR, --errorCollectionDir ERRORCOLLECTIONDIR
                        Directory to copy the output from any failed runs to.

```

At a minimum you need to pass the script the path to the GEOSX executable and a directory to run the benchmarks in. This directory will be created if it doesn't exist. The script will collect a list of benchmarks to be run and submit a job to the system's scheduler for each benchmark. This means that you don't need to be in an allocation to run the benchmarks. Note that this is different from the integrated tests where you need to already be in an allocation and an internal scheduler is used to run the individual tests. Since a benchmark is a measure of performance to get consistent results it is important that each time a benchmark is run it has access to the same resources. Using the system scheduler guarantees this.

In addition to whatever outputs the input would normally produce (plot files, restart files, ...) each benchmark will produce an output file `output.txt` containing the standard output and standard error of the run and a `.cali` file containing the Caliper timing data in a format that [Spot](#) can read.

---

**Note:** A future version of the script will be able to run only a subset of the benchmarks.

---

## Specifying a benchmark

A group of benchmarks is specified with a standard GEOSX input XML file with an extra `Benchmarks` block added at the top level. This block is ignored by GEOSX itself and only used by the `runBenchmarks.py` script.

```

<Benchmarks>
  <quartz>
    <Run
      name="OMP "
      nodes="1 "
      tasksPerNode="1"
      timeLimit="10"
      autoPartition="On"/>
    <Run
      name="MPI_OMP "
      nodes="1 "
      tasksPerNode="2"
      autoPartition="On"
      timeLimit="10"
      strongScaling="{ 1, 2, 4, 8 }"/>
    <Run
      name="MPI "
      nodes="1 "

```

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```

    tasksPerNode="36"
    autoPartition="On"
    timeLimit="10"
    strongScaling="{ 1, 2, 4, 8 }"/>
</quartz>

<lassen>
  <Run
    name="OMP_CUDA"
    nodes="1"
    tasksPerNode="1"
    autoPartition="On"
    timeLimit="10"/>
  <Run
    name="MPI_OMP_CUDA"
    nodes="1"
    tasksPerNode="4"
    autoPartition="On"
    timeLimit="10"
    strongScaling="{ 1, 2, 4, 8 }"/>
</lassen>
</Benchmarks>

```

[Source: *benchmarks/SSLE-small.xml*]

The Benchmarks block consists of a block for each machine the benchmarks are to run on. Currently the only options are quartz and lassen.

## The Run block

Each machine block contains a number of Run blocks each of which specify a family of benchmarks to run. Each Run block must have the following required attributes

- **name:** The name of the family of benchmarks, must be unique among all the other Run blocks on that system.
- **nodes:** An integer which specifies the base number of nodes to run the benchmark with.
- **tasksPerNode:** An integer that specifies the number of tasks to launch per node.

Each Run block may contain the following optional attributes

- **threadsPerTask:** An integer specifying the number of threads to allocate each task.
- **timeLimit:** An integer specifying the time limit in minutes to pass to the system scheduler when submitting the benchmark.
- **args:** containing any extra command line arguments to pass to GEOSX.
- **autoPartition:** Either On or Off, not specifying autoPartition is equivalent to autoPartition="Off". When auto partitioning is enabled the script will compute the number of x, y and z partitions such that the the resulting partition is close to a perfect cube as possible, ie with 27 tasks  $x = 3, y = 3, z = 3$  and with 36 tasks  $x = 4, y = 3, z = 3$ . This is optimal when the domain itself is a cube, but will be suboptimal otherwise.
- **strongScaling:** A list of unique integers specifying the factors to scale the number of nodes by. If N number are provided then N benchmarks are run and benchmark i uses nodes \* strongScaling[ i ] nodes. Not specifying strongScaling is equivalent to strongScaling="{ 1 }".

Looking at the example `Benchmarks` block above on Lassen one benchmark from the `OMP_CUDA` family will be run with one node and one task. Four benchmarks from the `MPI_OMP_CUDA` family will be run with one, two, four and eight nodes and four tasks per node.

Note that specifying a time limit for each benchmark family can greatly decrease the time spent waiting in the scheduler's queue. A good rule of thumb is that the time limit should be twice as long as it takes to run the longest benchmark in the family.

### Adding a benchmark problem

To add a new group of benchmarks you need to create an XML input file describing the problem to be run. Then you need to add the `Benchmarks` block described above which specifies the specific benchmarks. Finally add a symbolic link to the input file in `benchmarks` and run the benchmarks to make sure everything works as expected.

### Viewing the results

Each night the `NightlyTests` repository runs the benchmarks on both Quartz and Lassen, the `timingFiles` directory contains all of the resulting caliper output files. If you're on LC then these files are duplicated at `/usr/gapps/GEOSX/timingFiles/` and if you have LC access you can view them in `Spot`. You can also open these files in Python and analyse them (See *Opening Spot caliper files in Python*).

If you want to run the benchmarks on your local branch and compare the results with develop you can use the `benchmarks/compareBenchmarks.py` python script. This requires that you run the benchmarks on your branch and on develop. It will print out a table with the initialization time speed up and run time speed up, so a run speed up of 2x means your branch runs twice as fast as develop where as a initialization speed up of 0.5x means the set up takes twice as long.

---

**Note:** A future version of the script will be able to pull timing results straight from the `.cali` files so that if you have access to the `NightlyTests` timing files you won't need to run the benchmarks on develop. Furthermore it will be able to provide more detailed information than just initialization and run times.

---

## 6.1.9 Basic profiling with CALIPER

GEOSX is equipped with `Caliper` timers. We integrate Caliper into GEOSX by marking source-code sections of interest such as computational kernels or initialization steps. Caliper is included in the GEOSX TPL library and is built by adding the following cmake configuration to a host-config file.

```
option( ENABLE_CALIPER "Enables CALIPER" On )
```

### GEOSX/Caliper Annotation Macros

The following macros may be used to annotate GEOSX:

- `GEOSX_MARK_SCOPE (name)` - Marks a scope with the given name.
- `GEOSX_MARK_FUNCTION` - Marks a function with the name of the function. The name includes the namespace the function is in but not any of the template arguments or parameters. Therefore overloaded function all show up as one entry. If you would like to mark up a specific overload use `GEOSX_MARK_SCOPE` with a unique name.
- `GEOSX_MARK_BEGIN (name)` - Marks the beginning of a user defined code region.

- `GEOSX_MARK_END (name)` - Marks the end of user defined code region.

The 2 first macros also generate annotations for NVTX if `ENABLE_CUDA_NVTOOLSEXT` is activated through CMake.

## Configuring Caliper

Caliper configuration is done by specifying a string to initialize Caliper with via the `-t` option. A few options are listed below but we refer the reader to [Caliper Config](#) for the full Caliper tutorial.

- `-t runtime-report,max_column_width=200` Will make Caliper print aggregated timing information to standard out, with a column width large enough that it doesn't truncate most function names.
- `-t runtime-report,max_column_width=200,profile.cuda` Does the same as the above, but also instruments CUDA API calls. This is only an option when building with CUDA.
- `-t runtime-report,aggregate_across_ranks=false` Will make Caliper write per rank timing information to standard out. This isn't useful when using more than one rank but it does provide more information for single rank runs.
- `-t spot()` Will make Caliper output a `.cali` timing file that can be viewed in the Spot web server.

## Using Adiak

Adiak is a library that allows the addition of meta-data to the Caliper Spot output, it is enabled with Caliper. This meta-data allows you to easily slice and dice the timings available in the Spot web server. To export meta-data use the `adiak::value` function.

See [Adiak API](#) for the full Adiak documentation.

## Using Spot

To use Spot you will need an LC account and a directory full of `.cali` files you would like to analyse. Point your browser to [Spot](#) and open up the directory containing the timing files.

## Opening Spot caliper files in Python

An example Python program for analyzing Spot Caliper files in Python is provided below. Note that it requires `pandas` and `hatchet` both of which can be installed with a package manager. In addition it requires that `cali-query` is in the `PATH` variable, this is built with Caliper so we can just point it into the TPLs.

```
import sys
import subprocess
import json
import os

import pandas as pd
from IPython.display import display, HTML

# Import hatchet, on LC this can be done by adding hatchet to PYTHONPATH
sys.path.append('/usr/gapps/spot/live/hatchet')
import hatchet as ht

# Add cali-query to PATH
cali_query_path = "/usr/gapps/GEOSX/thirdPartyLibs/2020-06-12/install-quartz-gcc@8.1.
↳0-release/caliper/bin"
```

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```

os.environ["PATH"] += os.pathsep + cali_query_path

CALI_FILES = [
{ "cali_file": "/usr/gapps/GEOSX/timingFiles/200612-04342891243.cali", "metric_name":
↪ "avg#inclusive#sum#time.duration"},
{ "cali_file": "/usr/gapps/GEOSX/timingFiles/200611-044740108300.cali", "metric_name
↪ ": "avg#inclusive#sum#time.duration"},
]

grouping_attribute = "prop:nested"
default_metric = "avg#inclusive#sum#time.duration"
query = "select %s,sum(%s) group by %s format json-split" % (grouping_attribute,
↪ default_metric, grouping_attribute)

gf1 = ht.GraphFrame.from_caliper(CALI_FILES[0]['cali_file'], query)
gf2 = ht.GraphFrame.from_caliper(CALI_FILES[1]['cali_file'], query)

# Print the tree representation using the default metric
# Also print the resulting dataframe with metadata
print(gf1.tree(color=True, metric="sum#" + default_metric))
display(HTML(gf1.dataframe.to_html()))

# Print the tree representation using the default metric
# Also print the resulting dataframe with metadata
print(gf2.tree(color=True, metric="sum#" + default_metric))
display(HTML(gf2.dataframe.to_html()))

# Compute the speedup between the first two cali files (exclusive and inclusive,
↪ metrics only)
gf3 = (gf1 - gf2) / gf2
print(gf3.tree(color=True, metric="sum#" + default_metric))

# Compute the difference between the first two cali files (exclusive and inclusive,
↪ metrics only)
# Print the resulting tree
gf4 = gf1 - gf2
print(gf4.tree(color=True, metric="sum#" + default_metric))

# Compute the sum of the first two cali files (exclusive and inclusive metrics only)
# Print the resulting tree
gf5 = gf1 + gf2
print(gf5.tree(color=True, metric="sum#" + default_metric))

```

### 6.1.10 [Unsupported] Developing inside Docker with precompiled TPL binaries

For development purposes, you may want to use the publicly available docker images or the OSX tarball instead of compiling them yourself. While this is possible and this page will help you in through this journey, please note that *this is not officially supported by the GEOSX team that reserves the right to modify its workflow or delete elements on which you may have build your own workflow.*

There are multiple options to use the exposed docker images.

- A lot of IDE now provide remote development modes (e.g. [CLion](#), [VS Code](#), [Eclipse Che](#) and surely others). Depending on your choice, please read their documentation carefully so you can add their own requirements on top the TPL images that are already available.

- Another option is to develop directly inside the container (*i.e.* not remotely). Install your favorite development inside the image (be mindful of X display issues), connect to the running container and start hacking!

You must first [install docker](#) on your machine. Note that there now exists a [rootless install](#) that may help you in case you are not granted extended permissions on your environment. Also be aware that [nvidia](#) provides its own [nvidia-docker](#) that grants access to GPUs.

Once you've installed docker, you must select from our [docker registry](#) the target environment you want to develop into.

- You can select the distribution you are comfortable with, or you may want to mimic (to some extent) a production environment.
- Our containers are built with a relative CPU agnosticism (still x86\_64), so you should be fine.
- Our GPU containers are built for a dedicated `compute capability` that may not match yours. Please dive into our configuration files and refer to the [official nvidia page](#) to see what matches your needs.
- There may be risks of kernel inconsistency between the container and the host, but if you have relatively modern systems (and/or if you do not interact directly with the kernel like `perf`) it should be fine.
- You may have noticed that our docker containers are tagged like 155-669. Please refer to [Continuous Integration process](#) for further information.

Now that you've selected your target environment, you must be aware that just running a TPL docker image is not enough to let you develop. You'll have to add extra tools.

The following *example* is for our `ubuntu` flavors. You'll notice the arguments `IMG`, `VERSION`, `ORG`. While surely overkill for most cases, if you develop in GEOSX on a regular basis you'll appreciate being able to switch containers easily. For example, simply create the image `remote-dev-ubuntu18.04-gcc8:156-642` by running

```
export VERSION=156-642
export IMG=ubuntu18.04-gcc8
export REMOTE_DEV_IMG=remote-dev-${IMG}
docker build --build-arg ORG=geosx --build-arg IMG=${IMG} --build-arg VERSION=$
↪{VERSION} -t ${REMOTE_DEV_IMG}:${VERSION} -f /path/to/Dockerfile .
```

And the Dockerfile is the following (comments are embedded)

```
1 # Define you base image for build arguments
2 ARG IMG
3 ARG VERSION
4 ARG ORG
5 FROM ${ORG}/${IMG}:${VERSION}
6
7 # Uninstall some packages, install others.
8 # I use those for clion, but VS code would have different requirements.
9 # Use yum's equivalent commands for centos/red-hat images.
10 # Feel free to adapt.
11 RUN apt-get update
12 RUN apt-get remove --purge -y texlive graphviz
13 RUN apt-get install --no-install-recommends -y openssh-server gdb rsync gdbserver ↪
↪ninja-build
14
15 # You will need cmake to build GEOSX.
16 ARG CMAKE_VERSION=3.16.8
17 RUN apt-get install -y --no-install-recommends curl ca-certificates && \
18     curl -fsSL https://cmake.org/files/v${CMAKE_VERSION%.[0-9]*}/cmake-${CMAKE_
↪VERSION}-Linux-x86_64.tar.gz | tar --directory=/usr/local --strip-components=1 -xzf ↪
↪- && \
```

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```

19     apt-get purge --auto-remove -y curl ca-certificates
20 RUN apt-get autoremove -y
21
22 # You'll most likely need ssh/sshd too (e.g. CLion and VSCode allow remote dev_
  ↳ through ssh).
23 # This is the part where I configure sshd.
24
25 # I'm developing in a version of docker that requires root.
26 # So by default I use root. Feel free to adapt.
27 RUN echo "PermitRootLogin prohibit-password" >> /etc/ssh/sshd_config
28 RUN echo "PermitUserEnvironment yes" >> /etc/ssh/sshd_config
29 RUN mkdir -p -m 700 /root/.ssh
30 # Put your own public key here!
31 RUN echo "ssh-rsa [a public ssh key]" > /root/.ssh/authorized_keys
32
33 # Some important variables are provided through the environment.
34 # You need to explicitly tell sshd to forward them.
35 # Using these variables and not paths will let you adapt to different installation_
  ↳ locations in different containers.
36 # Feel free to adapt to your own convenience.
37 RUN touch /root/.ssh/environment &&\
38     echo "CC=${CC}" >> /root/.ssh/environment &&\
39     echo "CXX=${CXX}" >> /root/.ssh/environment &&\
40     echo "MPICC=${MPICC}" >> /root/.ssh/environment &&\
41     echo "MPICXX=${MPICXX}" >> /root/.ssh/environment &&\
42     echo "MPIEXEC=${MPIEXEC}" >> /root/.ssh/environment &&\
43     echo "OMPI_CC=${CC}" >> /root/.ssh/environment &&\
44     echo "OMPI_CXX=${CXX}" >> /root/.ssh/environment &&\
45     echo "GEOSX_TPL_DIR=${GEOSX_TPL_DIR}" >> /root/.ssh/environment
46
47 # This is the default ssh port that we do not need to modify.
48 EXPOSE 22
49 # sshd's option -D prevents it from detaching and becoming a daemon.
50 # Otherwise, sshd would not block the process and `docker run` would quit.
51 RUN mkdir -p /run/sshd
52 ENTRYPOINT ["/usr/sbin/sshd", "-D"]

```

Now that you've created the image, you must instantiate it as a container. I like to do

```

docker run --cap-add=SYS_PTRACE -d --name ${REMOTE_DEV_IMG}-${VERSION} -p 64000:22 $
  ↳ ${REMOTE_DEV_IMG}:${VERSION}

```

that creates the container `remote-dev-ubuntu18.04-gcc8-156-642`, running instance of `remote-dev-ubuntu18.04-gcc8:156-642`.

- Note that you'll have to access your remote development instance though port 64000 (forwarded to standard port 22 by docker).
- Please be aware of how to retrieve your code back: you may want to bind mount volumes and store you code there (`-v/--volume=` options of [docker run](#)).
- Change docker to `nvidia-docker` and add the `--gpus=...` option for GPUs.

You can stop and restart your container with

```

docker stop ${REMOTE_DEV_IMG}-${VERSION}
docker start ${REMOTE_DEV_IMG}-${VERSION}

```

Now hack.

### 6.1.11 [Unsupported] Installing GEOSX on Windows machines using Docker

In this section, we will install GEOSX on a Windows machine using a Docker container with a precompiled version of GEOSX's third party libraries (TPL). These steps are an adaptation of *ref:UsingDocker* for the Windows environment. In the following sections, we will be using *Ubuntu* based image as an example.

#### 1. Install *Docker Desktop*

On your Windows machine, follow these [steps](#). Download the most recent installer for *Docker Desktop*. Before installation please check the current status of Windows Subsystem for Linux (WSL) on your machine as Docker will use WSL2 as a [backend](#). To do that, open a PowerShell (Admin)

```
PS > wsl --install
PS > wsl --status
PS > wsl --set-default-version 2
PS > wsl --status
```

The first command should install WSL2, download an *Ubuntu* distribution for it and ask for a restart. The following commands are used to check the status, and if the WSL is still the default one, change it to WSL2. More details on the installation procedure can be found [here](#).

Once the WSL2 is set as default, proceed with the *Docker Desktop* installation.

#### 2. Start *Docker Desktop*

When launching Docker Desktop for the first time, you should be prompted with a message informing you that it uses WSL2. Using PowerShell, you can check that Docker and WSL2 are actually running in the background:

```
PS > Get-Process docker
Handles  NPM(K)    PM(K)      WS(K)      CPU(s)      Id  SI ProcessName
-----  -
123      11      26084      25608        0.42    13960  1 docker

PS > Get-Process wsl
Handles  NPM(K)    PM(K)      WS(K)      CPU(s)      Id  SI ProcessName
-----  -
146       8       1412       6848        0.05    14816  1 wsl
146       7       1356       6696        0.02    15048  1 wsl
145       7       1368       6704        0.02    15100  1 wsl
145       7       1352       6716        0.14    15244  1 wsl
146       7       1396       6876        0.02    16156  1 wsl
```

You should be able to see one docker process and several *wsl* processes.

#### 3. Preparing *DockerFile*

Let us now prepare the installation, picking a destination folder and editing our Dockerfile:

```
PS > cd D:/
PS > mkdir install-geosx-docker
PS > cd install-geosx-docker/
PS > notepad.exe Dockerfile
```

Let us edit the Dockerfile, which is the declarative file for our container:

```

1 # Define you base image for build arguments
2 ARG IMG
3 ARG VERSION
4 ARG ORG
5 FROM ${ORG}/${IMG}:${VERSION}
6
7 # Uninstall some packages, install others.
8 # I use those for clion, but VS code would have different requirements.
9 # Use yum's equivalent commands for centos/red-hat images.
10 # Feel free to adapt.
11 RUN apt-get update
12 RUN apt-get remove --purge -y texlive graphviz
13 RUN apt-get install --no-install-recommends -y openssh-server gdb rsync gdbserver_
    ↪ ninja-build
14
15 # You will need cmake to build GEOSX.
16 ARG CMAKE_VERSION=3.16.8
17 RUN apt-get install -y --no-install-recommends curl ca-certificates && \
18     curl -fsSL https://cmake.org/files/v${CMAKE_VERSION%.[0-9]*}/cmake-${CMAKE_
    ↪ VERSION}-Linux-x86_64.tar.gz | tar --directory=/usr/local --strip-components=1 -xzf_
    ↪ - && \
19     apt-get purge --auto-remove -y curl ca-certificates
20 RUN apt-get autoremove -y
21
22 # You'll most likely need ssh/sshd too (e.g. CLion and VSCode allow remote dev_
    ↪ through ssh).
23 # This is the part where I configure sshd.
24
25 # I'm developing in a version of docker that requires root.
26 # So by default I use root. Feel free to adapt.
27 RUN echo "PermitRootLogin prohibit-password" >> /etc/ssh/sshd_config
28 RUN echo "PermitUserEnvironment yes" >> /etc/ssh/sshd_config
29 RUN mkdir -p -m 700 /root/.ssh
30 # Put your own public key here!
31 RUN echo "ssh-rsa [a public ssh key]" > /root/.ssh/authorized_keys
32
33 # Some important variables are provided through the environment.
34 # You need to explicitly tell sshd to forward them.
35 # Using these variables and not paths will let you adapt to different installation_
    ↪ locations in different containers.
36 # Feel free to adapt to your own convenience.
37 RUN touch /root/.ssh/environment &&\
38     echo "CC=${CC}" >> /root/.ssh/environment &&\
39     echo "CXX=${CXX}" >> /root/.ssh/environment &&\
40     echo "MPICC=${MPICC}" >> /root/.ssh/environment &&\
41     echo "MPICXX=${MPICXX}" >> /root/.ssh/environment &&\
42     echo "MPIEXEC=${MPIEXEC}" >> /root/.ssh/environment &&\
43     echo "OMPI_CC=${CC}" >> /root/.ssh/environment &&\
44     echo "OMPI_CXX=${CXX}" >> /root/.ssh/environment &&\
45     echo "GEOSX_TPL_DIR=${GEOSX_TPL_DIR}" >> /root/.ssh/environment
46
47 # This is the default ssh port that we do not need to modify.
48 EXPOSE 22
49 # sshd's option -D prevents it from detaching and becoming a daemon.
50 # Otherwise, sshd would not block the process and `docker run` would quit.
51 RUN mkdir -p /run/sshd

```

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```
ENTRYPOINT ["/usr/sbin/sshd", "-D"]
```

This file enriches a base image already containing the GEOSX's TPL as well as extra utils, such as `cmake` and preparing for ssh connexion. In the end, we will be able to run it in a detached mode, and connect to it to run and develop in GEOSX.

There are two things you may have noticed reading through the `Dockerfile` :

- It has environment variables to be passed to it to select the proper image to pull, namely `${ORG}`, `${IMG}` and `${VERSION}`, we'll then have to declare them

```
PS> $env:VERSION='164-677'
PS> $env:IMG='ubuntu18.04-gcc8'
PS> $env:REMOTE_DEV_IMG="remote-dev-${env:IMG}"
```

Please note the preposition of `env :` in the windows formalisme. The `${ORG}` variable will be hard-coded as `geosx`. The last variable will be use as the image name. `164-677` refers to a specific version of the TPLs which may not be up to date. Please refer to [Continuous Integration process](#) for further info.

- You'll need to generate a ssh-key to be able to access the container without the need for defining a password. This can be done from the *PowerShell*,

```
PS > ssh-keygen.exe
PS > cat [path-to-gen-key]/[your-key].pub
```

The first command will prompt you with a message asking you to complete the desired path for the key as well as a passphrase, with confirmation. More details on [ssh-key generation](#).

## 4. Build the image and run the container

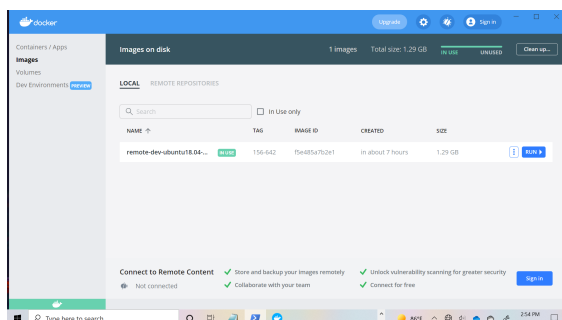
The preliminary tasks are now done. Let us build the image that will be containerized.

```
PS> cd [path-to-dockerfile-folder]/
PS > docker build --build-arg ORG=geosx --build-arg IMG=${env:IMG} --build-arg
↪VERSION=${env:VERSION} -t ${env:REMOTE_DEV_IMG}:${env:VERSION} -f Dockerfile
```

As described above, we are passing our environment variables in the building stage, which offer the flexibility of changing the version or image by a simple redefinition. A log updating or pulling the different layers should be displayed afterwards and on the last line the *image id*. We can check that the image is created using *PowerShell* CLI:

```
PS > docker images
```

or using the *Docker Desktop*

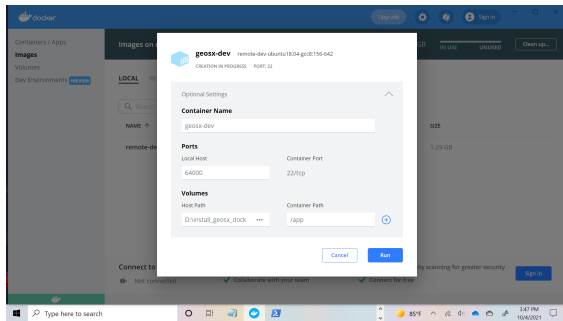


Now that we have the image build, let us run a container from,

```
PS > docker run --cap-add=SYS_PTRACE -d --name ${env:REMOTE_DEV_IMG}-${env:VERSION} -
  ↳p 64000:22 --mount 'type=bind,source=D:/install_geosx_docker/,target=/app' $
  ↳{env:REMOTE_DEV_IMG}:${env:VERSION}
```

Note that in addition to the detached flag (`-d`) and the name tag (`--name`), we provide Docker with the port the container should be associated to communicate with ssh port 22, as well as a binding between a host mount point (`D:/install_geosx_docker/`) and a container mount point (`/app`) to have a peristent storage for our development/geosx builds. More details on the `--mount` options

A similar step can be achieved using the *Docker Desktop* GUI in the image tabs, clicking on the *run* button and filling the same information in the interface,



Coming back to our PowerShell terminal, we can check that our container is running and trying to ssh to it.

```
PS > docker ps -a
```

CONTAINER ID	IMAGE	COMMAND	CREATED
1efffac66c4c	remote-dev-ubuntu18.04-gcc8:156-642	"/usr/sbin/sshd -D"	Less than a second ago
Up 18 seconds	0.0.0.0:64000->22/tcp, :::64000->22/tcp		remote-dev-ubuntu18.04-gcc8-156-642

```
PS > ssh root@localhost -p 64000
Enter passphrase for key 'C:\*****\.ssh\id_rsa':
Welcome to Ubuntu 18.04.5 LTS (GNU/Linux 5.10.16.3-microsoft-standard-WSL2 x86_64)

 * Documentation:  https://help.ubuntu.com
 * Management:    https://landscape.canonical.com
 * Support:       https://ubuntu.com/advantage
This system has been minimized by removing packages and contents that are
not required on a system that users do not log into.

To restore this content, you can run the 'unminimize' command.

The programs included with the Ubuntu system are free software;
the exact distribution terms for each program are described in the
individual files in /usr/share/doc/*/copyright.

Ubuntu comes with ABSOLUTELY NO WARRANTY, to the extent permitted by
applicable law.

root@b105f9ead860:~# cd /app && ls
```

We are now logged into our container and can start *Quick Start Guide*.

**Note:** Note that :

1. You might be prompted that you miss certificates to clone, this can be resolved by installing *ca-certificates* and updating them
2. It might occur that *git-lfs* is missing then install it

```
PS > apt install ca-certificates && update-ca-certificates
PS > apt install git-lfs
```

From there you should be able to develop in your container or access it from an IDE, e.g. [VSCode](#) or [MSVC19](#).

## 5. Running a case

Once the code is configured and compiled, let us check the status of the build,

```
root@b105f9ead860:~# cd [path-to-build]/ && ./bin/geosx --help
```

Trying to launch a case using *mpirun*, you might get the following warning

```
root@b105f9ead860:/tmp# mpirun -np 4 /app/code/GEOSX/build-environment-debug/bin/
↳ geosx -i [geosx-case].xml -x 4 -y 1 -z 1
-----
mpirun has detected an attempt to run as root.
Running at root is *strongly* discouraged as any mistake (e.g., in
defining TMPDIR) or bug can result in catastrophic damage to the OS
file system, leaving your system in an unusable state.

You can override this protection by adding the --allow-run-as-root
option to your cmd line. However, we reiterate our strong advice
against doing so - please do so at your own risk.
-----
```

A possible workaround is to create a new user account and a run folder from this account

```
root@b105f9ead860:~# adduser runner
root@b105f9ead860:~# su runner
runner@b105f9ead860:/root$ mkdir -p /tmp/geosx && cd /tmp/geosx/
runner@b105f9ead860:/tmp/geosx$ cp [path-to-case]/[geosx-case].xml .
runner@b105f9ead860:/tmp/geosx$ ${MPIEXEC} -np 4 /app/code/GEOSX/build-environment-
↳ debug/bin/geosx -i [geosx-case].xml -x 4 -y 1 -z 1
```

## 6.2 Code Components

The main code components are described here.

### 6.2.1 Data Repository

The GEOSX “Data Repository” is intended to provide the building blocks for the code structure within GEOSX. The “Data Repository” provides a general capability to store arbitrary data and objects in a hierarchical structure, similar to a standard file system.

The components/classes of the data structure that a developer will require some knowledge of are:

## Mapped Vector

A superposition of a contiguous and an associative container.

### Description

The container stores pointers to objects (which are themselves heap-allocated). Each element may be optionally *owned* by the container, in which case it will be deleted upon removal or container destruction. The pointers are stored in a contiguous memory allocation, and thus are accessible through an integral index lookup. In addition, there is a map that provides a key lookup capability to the container if that is the preferred interface.

The container template has four type parameters:

- `T` is the object type pointed to by container entries
- `T_PTR` is a pointer-to-`T` type which must be either `T *` (default) or `std::unique_ptr<T>`
- `KEY_TYPE` is the type of key used in associative lookup
- `INDEX_TYPE` is the type used in index lookup

### Element access

`MappedVector` provides three main types of data access using `[]` operator:

- **Index lookup** is the fastest way of element random access if the ordinal index is known.
- **Key lookup** is similar to key lookup of any associative container and incurs similar cost.
- **KeyIndex lookup** uses a special type, `KeyIndex`, that contains both a key and an index. Initially the index is unknown and the key is used for the lookup. The `KeyIndex` is modified during lookup, storing the index located. If the user persists the `KeyIndex` object, they may reuse it in subsequent accesses and get the benefit of direct index access.

In addition to these, an STL-conformant iterator interface is available via `begin()` and `end()` methods. The type iterated over is a key-pointer pair (provided as *value\_type* alias).

## API documentation

### MappedVector

#### Group

`dataRepository::Group` serves as a base class for most objects in GEOSX. In GEOSX, the `Group` may be thought of as an analogy to the file folder in a hierarchical filesystem-like structure. As such, a `Group` is used as a container class that holds a collection of other `Groups`, or sub-`Groups`, a pointer to the parent of the `Group`, and a collection of `Wrappers`. The `Group` also defines a general capability to create and traverse/access the objects in the hierarchy. The `Wrappers` contained in a `Group` may be of arbitrary type, but in the case of an `LvArray` object, a `Group` size and capacity may be translated down to array, thereby keeping a collection of wrapped `Array` objects that will resize in unison with the `Group`. Each group has a string “name” that defines its key in the parent `Group`. This key string must be unique in the scope of the parent `Group`.

## Implementation Details

Some noteworthy implementation details inside the declaration of `dataRepository::Group` are:

```
/// The default key type for entries in the hierarchy.
using keyType = string;

/// The default index type for entries the hierarchy.
using indexType = localIndex;
```

- In the GEOSX repository, the `keyType` is specified to be a `string` for all collection objects, while the `indexType` is specified to be a `localIndex`. The types are set in the `common/DataTypes.hpp` file, but are typically a `string` and a `std::ptrdiff_t` respectively.

```
/// The template specialization of MappedVector to use for the collection of sub-
↳Group objects.
using subGroupMap = MappedVector< Group, Group *, keyType, indexType >;

/// The template specialization of MappedVector to use for the collection wrappers
↳objects.
using wrapperMap = MappedVector< WrapperBase, WrapperBase *, keyType, indexType >;
```

- The `subGroupMap` and `wrapperMap` aliases represent the type of container that the collection of `sub-Group`s and `Wrapper`s are stored in for each `Group`. These container types are template specializations of the `MappedVector` class, which store a pointer to a type, and provides functionality for a key or index based lookup. More details may be found in the documentation for `MappedVector`.

```
/// The parent Group that contains "this" Group in its "sub-Group" collection.
Group * m_parent = nullptr;

/// Specification that this group will have the same m_size as m_parent.
integer m_sizedFromParent;

/// The container for the collection of all wrappers continued in "this" Group.
wrapperMap m_wrappers;

/// The container for the collection of all sub-groups contained in "this" Group.
subGroupMap m_subGroups;

/// The size/length of this Group...and all Wrapper<> that are are specified to
↳have the same size as their
/// owning group.
indexType m_size;

/// The capacity for wrappers in this group...and all Wrapper<> that are specified
↳to have the same size as their
/// owning group.
indexType m_capacity;

/// The name/key of this Group in its parent collection of sub-Groups.
string m_name;

/// Verbosity flag for group logs
integer m_logLevel;
```

- The `m_parent` member is a pointer to the `Group` that contains the current `Group` as part of its collection of `sub-Group`s.

**Warning:** The existence of the non-const `m_parent` gives the current `Group` access to alter the parent `Group`. Special care should be taken to avoid using this access whenever possible. Remember... with great power comes great responsibility.

- The `m_wrappers` member is the collection of `Wrappers` contained in the current `Group`.
- The `m_subGroups` member is the collection of `Group`s contained in the current `Group`.
- The `m_size` and `m_capacity` members are used to set the size and capacity of any objects contained in the `m_wrappers` collection that have been specified to be set by their owning `Group`. This is typically only useful for `Array` types and is implemented within the `WrapperBase` object.
- The `m_name` member is the key of this `Group` in the collection of `m_parent->m_subGroups`. This key is unique in the scope of `m_parent`, so some is required when constructing the hierarchy.

## Interface Functions

The public interface for `dataRepository::Group` provides functionality for constructing a hierarchy, and traversing that hierarchy, as well as accessing the contents of objects stored in the `Wrapper` containers stored within a `Group`.

## Adding New Groups

To add new sub-`Group`s there are several `registerGroup` functions that add a new `Group` under the calling `Group` scope. A listing of these functions is provided:

```
/**
 * @name Sub-group registration interface
 */
///

```
@{

/**
 * @brief Register a new Group as a sub-group of current Group.
 *
 * @tparam T The type of the Group to add/register. This should be a type that
↳ derives from Group.
 * @param[in] name The name of the group to use as a string key.
 * @param[in] newObject A unique_ptr to the object that is being registered.
 * @return A pointer to the newly registered Group.
 *
 * Registers a Group or class derived from Group as a subgroup of this Group and
↳ takes ownership.
 */
template< typename T = Group >
T & registerGroup( string const & name, std::unique_ptr< T > newObject )
{
    newObject->m_parent = this;
    return dynamicCast< T & >( *m_subGroups.insert( name, newObject.release(), true ) );
↳ };
}

/**
 * @brief @copybrief registerGroup(string const &,std::unique_ptr<T>)
 */
```


```

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```

    * @tparam T The type of the Group to add/register. This should be a type that
↳derives from Group.
    * @param[in] name          The name of the group to use as a string key.
    * @param[in] newObject     A unique_ptr to the object that is being registered.
    * @return                  A pointer to the newly registered Group.
    *
    * Registers a Group or class derived from Group as a subgroup of this Group but
↳does not take ownership.
    */
    template< typename T = Group >
    T & registerGroup( string const & name, T * newObject )
    { return dynamicCast< T &>( *m_subGroups.insert( name, newObject, false ) ); }

    /**
    * @brief @copybrief registerGroup(string const &,std::unique_ptr<T>)
    *
    * @tparam T The type of the Group to add/register. This should be a type that
↳derives from Group.
    * @param[in] name The name of the group to use as a string key.
    * @return          A pointer to the newly registered Group.
    *
    * Creates and registers a Group or class derived from Group as a subgroup of this
↳Group.
    */
    template< typename T = Group >
    T & registerGroup( string const & name )
    { return registerGroup< T >( name, std::make_unique< T >( name, this ) ); }

    /**
    * @brief @copybrief registerGroup(string const &,std::unique_ptr<T>)
    *
    * @tparam T The type of the Group to add/register. This should be a type that
↳derives from Group.
    * @param keyIndex A KeyIndexT object that will be used to specify the name of
    * the new group. The index of the KeyIndex will also be set.
    * @return A pointer to the newly registered Group, or @c nullptr if no group was
↳registered.
    *
    * Creates and registers a Group or class derived from Group as a subgroup of this
↳Group.
    */
    template< typename T = Group >
    T & registerGroup( subGroupMap::KeyIndex const & keyIndex )
    {
        T & rval = registerGroup< T >( keyIndex.key(), std::make_unique< T >( keyIndex.
↳key(), this ) );
        keyIndex.setIndex( m_subGroups.getIndex( keyIndex.key() ) );
        return rval;
    }

    /**
    * @brief @copybrief registerGroup(string const &,std::unique_ptr<T>)
    *
    * @tparam T The type of the Group to add/register. This should be a type that
↳derives from Group.
    * @tparam TBASE The type whose type catalog will be used to look up the new sub-
↳group type

```

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```

    * @param[in] name          The name of the group to use as a string key.
    * @param[in] catalogName The catalog name of the new type.
    * @return                  A pointer to the newly registered Group.
    *
    * Creates and registers a Group or class derived from Group as a subgroup of this_
    ↪Group.
    */
    template< typename T = Group, typename TBASE = Group >
    T & registerGroup( string const & name, string const & catalogName )
    {
        std::unique_ptr< TBASE > newGroup = TBASE::CatalogInterface::Factory( catalogName,
    ↪ name, this );
        return registerGroup< T >( name, std::move( newGroup ) );
    }

    /**
    * @brief Removes a child group from this group.
    * @param name the name of the child group to remove from this group.
    */
    void deregisterGroup( string const & name );

    /**
    * @brief Creates a new sub-Group using the ObjectCatalog functionality.
    * @param[in] childKey The name of the new object type's key in the
    *                    ObjectCatalog.
    * @param[in] childName The name of the new object in the collection of
    *                    sub-Groups.
    * @return A pointer to the new Group created by this function.
    */
    virtual Group * createChild( string const & childKey, string const & childName );

    //@}

```

These functions all take in a name for the new Group, which will be used as the key when trying to access the Group in the future. Some variants create a new Group, while some variants take in an existing Group. The template argument is to specify the actual type of the Group as it is most likely a type that derives from Group that is we would like to create in the repository. Please see the doxygen documentation for a detailed description of each option.

## Getting Groups

The collection of functions to retrieve a Group and their descriptions are taken from source and shown here:

```

    /**
    * @name Sub-group retrieval methods.
    *
    * This collection of functions are used to get a sub-Group from the current group.
    ↪Various methods
    * for performing the lookup are provided (localIndex, string, KeyIndex), and each_
    ↪have their
    * advantages and costs. The lowest cost lookup is the "localIndex" lookup. The_
    ↪KeyIndex lookup
    * will add a cost for checking to make sure the index stored in KeyIndex is valid_
    ↪(a string
    * compare, and a hash if it is incorrect). The string lookup is the full cost hash_
    ↪lookup every

```

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```

    * time that it is called.
    *
    * The template parameter specifies the "type" that the caller expects to lookup,
    ↪ and thus attempts
    * to cast the pointer that is stored in m_subGroups to a pointer of the desired
    ↪ type. If this
    * cast fails, then a @p nullptr is returned. If no template parameter is specified,
    ↪ then a default
    * type of Group is assumed.
    */
    ///@{

/**
    * @brief Return a pointer to a sub-group of the current Group.
    * @tparam T The type of subgroup.
    * @tparam KEY The type of the lookup.
    * @param key The key used to perform the lookup.
    * @return A pointer to @p T that refers to the sub-group, if the Group does not
    ↪ exist or it
    * has an incompatible type a @c nullptr is returned.
    */
    template< typename T = Group, typename KEY = void >
    T * getGroupPointer( KEY const & key )
    { return dynamicCast< T * >( m_subGroups[ key ] ); }

/**
    * @copydoc getGroupPointer(KEY const &)
    */
    template< typename T = Group, typename KEY = void >
    T const * getGroupPointer( KEY const & key ) const
    { return dynamicCast< T const * >( m_subGroups[ key ] ); }

/**
    * @brief Return a reference to a sub-group of the current Group.
    * @tparam T The type of subgroup.
    * @tparam KEY The type of the lookup.
    * @param key The key used to perform the lookup.
    * @return A reference to @p T that refers to the sub-group.
    * @throw std::domain_error If the Group does not exist is thrown.
    */
    template< typename T = Group, typename KEY = void >
    T & getGroup( KEY const & key )
    {
        Group * const child = m_subGroups[ key ];
        GEOSX_THROW_IF( child == nullptr, "Group " << getPath() << " doesn't have a child
    ↪ " << key, std::domain_error );
        return dynamicCast< T & >( *child );
    }

/**
    * @copydoc getGroup( KEY const & )
    */
    template< typename T = Group, typename KEY = void >
    T const & getGroup( KEY const & key ) const
    {
        Group const * const child = m_subGroups[ key ];
        GEOSX_THROW_IF( child == nullptr, "Group " << getPath() << " doesn't have a child
    ↪ " << key, std::domain_error );

```

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```

    return dynamicCast< T const & >( *child );
}

/**
 * @brief Retrieve a group from the hierarchy using a path.
 * @tparam T type of subgroup
 * @param[in] path a unix-style string (absolute, relative paths valid)
 *             to lookup the Group to return. Absolute paths search
 *             from the tree root, while relative - from current group.
 * @return A reference to @p T that refers to the sub-group.
 * @throw std::domain_error If the Group doesn't exist.
 */
template< typename T = Group >
T & getGroupByPath( string const & path )
{ return dynamicCast< T & >( const_cast< Group & >( getBaseGroupByPath( path ) ) ); }

/**
 * @copydoc getGroupByPath(string const &)
 */
template< typename T = Group >
T const & getGroupByPath( string const & path ) const
{ return dynamicCast< T const & >( getBaseGroupByPath( path ) ); }

```

## Register Wrappers

```

/**
 * @name Wrapper registration interface
 */
//@{

/**
 * @brief Create and register a Wrapper around a new object.
 * @tparam T The type of the object allocated.
 * @tparam TBASE The type of the object that the Wrapper holds.
 * @param[in] name the name of the wrapper to use as a string key
 * @param[out] rkey a pointer to a index type that will be filled with the new
 *                 Wrapper index in this Group
 * @return A reference to the newly registered/created Wrapper
 */
template< typename T, typename TBASE=T >
Wrapper< TBASE > & registerWrapper( string const & name,
                                   wrapperMap::KeyIndex::index_type * const rkey =
↳ nullptr );

/**
 * @copybrief registerWrapper(string const &, wrapperMap::KeyIndex::index_type *
↳ const)
 * @tparam T the type of the wrapped object
 * @tparam TBASE the base type to cast the returned wrapper to
 * @param[in] viewKey The KeyIndex that contains the name of the new Wrapper.
 * @return A reference to the newly registered/created Wrapper
 */
template< typename T, typename TBASE=T >

```

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```

Wrapper< TBASE > & registerWrapper( Group::wrapperMap::KeyIndex const & viewKey );

/**
 * @brief Register a Wrapper around a given object and take ownership.
 * @tparam T the type of the wrapped object
 * @param[in] name the name of the wrapper to use as a string key
 * @param[in] newObject an owning pointer to the object that is being registered
 * @return A reference to the newly registered/created Wrapper
 * @note Not intended to register a @p WrapperBase instance. Use dedicated member_
↳function instead.
 */
template< typename T >
Wrapper< T > & registerWrapper( string const & name, std::unique_ptr< T > newObject_
↳);

/**
 * @brief Register a Wrapper around an existing object, does not take ownership of_
↳the object.
 * @tparam T the type of the wrapped object
 * @param[in] name the name of the wrapper to use as a string key
 * @param[in] newObject a pointer to the object that is being registered
 * @return A reference to the newly registered/created Wrapper
 * @note Not intended to register a @p WrapperBase instance. Use dedicated member_
↳function instead.
 */
template< typename T >
Wrapper< T > & registerWrapper( string const & name,
                               T * newObject );

/**
 * @brief Register and take ownership of an existing Wrapper.
 * @param wrapper A pointer to the an existing wrapper.
 * @return An un-typed pointer to the newly registered/created wrapper
 */
WrapperBase & registerWrapper( std::unique_ptr< WrapperBase > wrapper );

/**
 * @brief Removes a Wrapper from this group.
 * @param name the name of the Wrapper to remove from this group.
 */
void deregisterWrapper( string const & name );

///<@}

```

## Getting Wrappers/Wrapped Objects

```

/**
 * @name Untyped wrapper retrieval methods
 *
 * These functions query the collection of Wrapper objects for the given
 * index/name/KeyIndex and returns a WrapperBase pointer to the object if
 * it exists. If it is not found, nullptr is returned.
 */
///<@{

```

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```

/**
 * @brief Return a reference to a WrapperBase stored in this group.
 * @tparam KEY The lookup type.
 * @param key The value used to lookup the wrapper.
 * @return A reference to the WrapperBase that resulted from the lookup.
 * @throw std::domain_error if the wrapper doesn't exist.
 */
template< typename KEY >
WrapperBase const & getWrapperBase( KEY const & key ) const
{
    WrapperBase const * const wrapper = m_wrappers[ key ];
    GEOSX_THROW_IF( wrapper == nullptr, "Group " << getPath() << " doesn't have a_
↳child " << key, std::domain_error );
    return *wrapper;
}

/**
 * @copydoc getWrapperBase(KEY const &) const
 */
template< typename KEY >
WrapperBase & getWrapperBase( KEY const & key )
{
    WrapperBase * const wrapper = m_wrappers[ key ];
    GEOSX_THROW_IF( wrapper == nullptr, "Group " << getPath() << " doesn't have a_
↳child " << key, std::domain_error );
    return *wrapper;
}

/**
 * @brief
 * @param name
 * @return
 */
indexType getWrapperIndex( string const & name ) const
{ return m_wrappers.getIndex( name ); }

/**
 * @brief Get access to the internal wrapper storage.
 * @return a reference to wrapper map
 */
wrapperMap const & wrappers() const
{ return m_wrappers; }

/**
 * @copydoc wrappers() const
 */
wrapperMap & wrappers()
{ return m_wrappers; }

/**
 * @brief Return the number of wrappers.
 * @return The number of wrappers.
 */
indexType numWrappers() const
{ return m_wrappers.size(); }

```

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```

///@}

/**
 * @name Typed wrapper retrieval methods
 *
 * These functions query the collection of Wrapper objects for the given
 * index/key and returns a Wrapper<T> pointer to the object if
 * it exists. The template parameter @p T is used to perform a cast
 * on the WrapperBase pointer that is returned by the lookup, into
 * a Wrapper<T> pointer. If the wrapper is not found, or the
 * WrapperBase pointer cannot be cast to a Wrapper<T> pointer, then nullptr
 * is returned.
 */
///@{

/**
 * @brief Check if a wrapper exists
 * @tparam LOOKUP_TYPE the type of key used to perform the lookup.
 * @param[in] lookup a lookup value used to search the collection of wrappers
 * @return @p true if wrapper exists (regardless of type), @p false otherwise
 */
template< typename LOOKUP_TYPE >
bool hasWrapper( LOOKUP_TYPE const & lookup ) const
{ return m_wrappers[ lookup ] != nullptr; }

/**
 * @brief Retrieve a Wrapper stored in this group.
 * @tparam T the object type contained in the Wrapper
 * @tparam LOOKUP_TYPE the type of key used to perform the lookup
 * @param[in] index a lookup value used to search the collection of wrappers
 * @return A reference to the Wrapper<T> that resulted from the lookup.
 * @throw std::domain_error if the Wrapper doesn't exist.
 */
template< typename T, typename LOOKUP_TYPE >
Wrapper< T > const & getWrapper( LOOKUP_TYPE const & index ) const
{
    WrapperBase const & wrapper = getWrapperBase( index );
    return dynamicCast< Wrapper< T > const & >( wrapper );
}

/**
 * @copydoc getWrapper(LOOKUP_TYPE const &) const
 */
template< typename T, typename LOOKUP_TYPE >
Wrapper< T > & getWrapper( LOOKUP_TYPE const & index )
{
    WrapperBase & wrapper = getWrapperBase( index );
    return dynamicCast< Wrapper< T > & >( wrapper );
}

/**
 * @brief Retrieve a Wrapper stored in this group.
 * @tparam T the object type contained in the Wrapper
 * @tparam LOOKUP_TYPE the type of key used to perform the lookup
 * @param[in] index a lookup value used to search the collection of wrappers
 * @return A pointer to the Wrapper<T> that resulted from the lookup, if the Wrapper
 * doesn't exist or has a different type a @c nullptr is returned.

```

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```

*/
template< typename T, typename LOOKUP_TYPE >
Wrapper< T > * const * getWrapperPointer( LOOKUP_TYPE const & index ) const
{ return dynamicCast< Wrapper< T > const * >( m_wrappers[ index ] ); }

/**
 * @copydoc getWrapperPointer(LOOKUP_TYPE const &) const
 */
template< typename T, typename LOOKUP_TYPE >
Wrapper< T > * getWrapperPointer( LOOKUP_TYPE const & index )
{ return dynamicCast< Wrapper< T > * >( m_wrappers[ index ] ); }

///  

/**
 * @name Wrapper data access methods.
 *
 * These functions can be used to get referece/pointer access to the data
 * stored by wrappers in this group. They are essentially just shortcuts for
 * @p Group::getWrapper() and @p Wrapper<T>::getReference().
 * An additional template parameter can be provided to cast the return pointer
 * or reference to a base class pointer or reference (e.g. Array to ArrayView).
 */
///  

/**
 * @brief Look up a wrapper and get reference to wrapped object.
 * @tparam T return value type
 * @tparam WRAPPEDTYPE wrapped value type (by default, same as return)
 * @tparam LOOKUP_TYPE type of value used for wrapper lookup
 * @param lookup value for wrapper lookup
 * @return reference to @p T
 * @throw A std::domain_error if the Wrapper does not exist.
 */
template< typename T, typename LOOKUP_TYPE >
GEOSX_DECLTYPE_AUTO_RETURN
getReference( LOOKUP_TYPE const & lookup ) const
{ return getWrapper< T >( lookup ).reference(); }

/**
 * @copydoc getReference(LOOKUP_TYPE const &) const
 */
template< typename T, typename LOOKUP_TYPE >
T & getReference( LOOKUP_TYPE const & lookup )
{ return getWrapper< T >( lookup ).reference(); }

```

## Looping Interface

```

/**
 * @name Functor-based subgroup iteration
 *
 * These functions loop over sub-groups and executes a functor that uses the sub-
 * ↪group as an
 * argument. The functor is only executed if the group can be cast to a certain_
 * ↪type specified

```

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```

    * by the @p ROUPTYPE/S pack. The variadic list consisting of @p GROUPTYPE/S will
    ↳ be used recursively
    * to check if the group is able to be cast to the one of these types. The first
    ↳ type in the
    * @p GROUPTYPE/S list will be used to execute the functor, and the next sub-group
    ↳ will be processed.
    */
    ///@{

    /**
    * @brief Apply the given functor to subgroups that can be casted to one of
    ↳ specified types.
    * @tparam GROUPTYPE the first type that will be used in the attempted casting of
    ↳ group.
    * @tparam GROUPTYPES a variadic list of types that will be used in the attempted
    ↳ casting of group.
    * @tparam LAMBDA the type of functor to call
    * @param[in] lambda the functor to call on subgroups
    */
    template< typename GROUPTYPE = Group, typename ... GROUPTYPES, typename LAMBDA >
    void forSubGroups( LAMBDA && lambda )
    {
        for( auto & subGroupIter : m_subGroups )
        {
            applyLambdaToContainer< GROUPTYPE, GROUPTYPES... >( *subGroupIter.second, [&](
    ↳ auto & castedSubGroup )
            {
                lambda( castedSubGroup );
            } );
        }
    }

    /**
    * @copydoc forSubGroups(LAMBDA &&)
    */
    template< typename GROUPTYPE = Group, typename ... GROUPTYPES, typename LAMBDA >
    void forSubGroups( LAMBDA && lambda ) const
    {
        for( auto const & subGroupIter : m_subGroups )
        {
            applyLambdaToContainer< GROUPTYPE, GROUPTYPES... >( *subGroupIter.second, [&](
    ↳ auto const & castedSubGroup )
            {
                lambda( castedSubGroup );
            } );
        }
    }

    /**
    * @brief Apply the given functor to subgroups that can be casted to one of
    ↳ specified types.
    * @tparam GROUPTYPE the first type that will be used in the attempted casting of
    ↳ group.
    * @tparam GROUPTYPES a variadic list of types that will be used in the attempted
    ↳ casting of group.
    * @tparam LAMBDA the type of functor to call

```

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```

    * @param[in] lambda the functor to call on subgroups
    */
template< typename GROUPTYPE = Group, typename ... GROUPTYPES, typename LAMBDA >
void forSubGroupsIndex( LAMBDA && lambda )
{
    localIndex counter = 0;
    for( auto & subGroupIter : m_subGroups )
    {
        applyLambdaToContainer< GROUPTYPE, GROUPTYPES... >( *subGroupIter.second,
                                                                [&]( auto & castedSubGroup )
                                                                {
                                                                    lambda( counter, castedSubGroup );
                                                                } );
        ++counter;
    }
}

/**
 * @copydoc forSubGroupsIndex(LAMBDA &&)
 */
template< typename GROUPTYPE = Group, typename ... GROUPTYPES, typename LAMBDA >
void forSubGroupsIndex( LAMBDA && lambda ) const
{
    localIndex counter = 0;
    for( auto const & subGroupIter : m_subGroups )
    {
        applyLambdaToContainer< GROUPTYPE, GROUPTYPES... >( *subGroupIter.second,
                                                                [&]( auto const &
→castedSubGroup )
                                                                {
                                                                    lambda( counter, castedSubGroup );
                                                                } );
        ++counter;
    }
}

/**
 * @copybrief forSubGroups(LAMBDA &&)
 * @tparam GROUPTYPE the first type that will be used in the attempted_
→casting of group.
 * @tparam GROUPTYPES a variadic list of types that will be used in the_
→attempted casting of group.
 * @tparam LOOKUP_CONTAINER type of container of subgroup lookup keys (names or_
→indices), must support range-based for
 * loop
 * @tparam LAMBDA type of functor callable with an index in lookup_
→container and a reference to casted
 * subgroup
 * @param[in] subGroupKeys container with subgroup lookup keys (e.g. names or_
→indices) to apply the functor to
 * @param[in] lambda the functor to call
 */
template< typename GROUPTYPE = Group, typename ... GROUPTYPES, typename LOOKUP_
→CONTAINER, typename LAMBDA >
void forSubGroups( LOOKUP_CONTAINER const & subGroupKeys, LAMBDA && lambda )
{
    localIndex counter = 0;

```

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```

    for( auto const & subgroup : subGroupKeys )
    {
        applyLambdaToContainer< GROUPTYPE, GROUPTYPES... >( getGroup( subgroup ), [&](
↪auto & castedSubGroup )
        {
            lambda( counter, castedSubGroup );
        } );
        ++counter;
    }
}

/**
 * @copybrief forSubGroups(LAMBDA &&)
 * @tparam GROUPTYPE          the first type that will be used in the attempted
↪casting of group.
 * @tparam GROUPTYPES        a variadic list of types that will be used in the
↪attempted casting of group.
 * @tparam LOOKUP_CONTAINER type of container of subgroup lookup keys (names or
↪indices), must support range-based for
 * loop
 * @tparam LAMBDA            type of functor callable with an index in lookup
↪container and a reference to casted
 * subgroup
 * @param[in] subGroupKeys  container with subgroup lookup keys (e.g. names or
↪indices) to apply the functor to
 * @param[in] lambda        the functor to call
 */
template< typename GROUPTYPE = Group, typename ... GROUPTYPES, typename LOOKUP_
↪CONTAINER, typename LAMBDA >
void forSubGroups( LOOKUP_CONTAINER const & subGroupKeys, LAMBDA && lambda ) const
{
    localIndex counter = 0;
    for( auto const & subgroup : subGroupKeys )
    {
        applyLambdaToContainer< GROUPTYPE, GROUPTYPES... >( getGroup( subgroup ), [&](
↪auto const & castedSubGroup )
        {
            lambda( counter, castedSubGroup );
        } );
        ++counter;
    }
}
///  

/**
 * @name Functor-based wrapper iteration
 *
 * These functions loop over the wrappers contained in this group, and executes a
↪functor that
 * uses the Wrapper as an argument. The functor is only executed if the Wrapper can
↪be casted to
 * a certain type specified by the @p TYPE/S pack. The variadic list consisting of
 * @p TYPE/S will be used recursively to check if the Wrapper is able to be casted
↪to the
 * one of these types. The first type in the @p WRAPPERTYPE/S list will be used to
↪execute the
 * functor, and the next Wrapper will be processed.

```

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```

    */
    ///@{

    /**
     * @brief Apply the given functor to wrappers.
     * @tparam LAMBDA the type of functor to call
     * @param[in] lambda the functor to call
     */
    template< typename LAMBDA >
    void forWrappers( LAMBDA && lambda )
    {
        for( auto & wrapperIter : m_wrappers )
        {
            lambda( *wrapperIter.second );
        }
    }

    /**
     * @copydoc forWrappers( LAMBDA && )
     */
    template< typename LAMBDA >
    void forWrappers( LAMBDA && lambda ) const
    {
        for( auto const & wrapperIter : m_wrappers )
        {
            lambda( *wrapperIter.second );
        }
    }

    /**
     * @brief Apply the given functor to wrappers that can be cast to one of specified_
    ↪types.
     * @tparam TYPE the first type that will be used in the attempted casting of_
    ↪Wrapper
     * @tparam TYPES a variadic list of types that will be used in the attempted_
    ↪casting of Wrapper
     * @tparam LAMBDA the type of functor to call
     * @param[in] lambda the functor to call
     */
    template< typename TYPE, typename ... TYPES, typename LAMBDA >
    void forWrappers( LAMBDA && lambda )
    {
        for( auto & wrapperIter : m_wrappers )
        {
            applyLambdaToContainer< Wrapper< TYPE >, Wrapper< TYPES >... >( *wrapperIter.
    ↪second,
                                                                                               std::forward<_
    ↪LAMBDA >( lambda ));
        }
    }

    /**
     * @brief Apply the given functor to wrappers that can be cast to one of specified_
    ↪types.
     * @tparam TYPE the first type that will be used in the attempted casting of_
    ↪Wrapper
     * @tparam TYPES a variadic list of types that will be used in the attempted_
    ↪casting of Wrapper

```

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```

    * @tparam LAMBDA the type of functor to call
    * @param[in] lambda the functor to call
    */
    template< typename TYPE, typename ... TYPES, typename LAMBDA >
    void forWrappers( LAMBDA && lambda ) const
    {
        for( auto const & wrapperIter : m_wrappers )
        {
            applyLambdaToContainer< Wrapper< TYPE >, Wrapper< TYPES >... >( *wrapperIter.
↪second,
                                                                    std::forward<
↪LAMBDA >( lambda ));
        }
    }

    ///@}

```

## Wrapper

This class encapsulates an object for storage in a `Group` and provides an interface for performing some common operations on that object.

## Description

In the filesystem analogy, a `Wrapper` may be thought of as a file that stores actual data. Each `Wrapper` belong to a single `Group` much like a file belongs to a filesystem directory. In general, more than one wrapper in the tree may refer to the same wrapped object, just like symlinks in the file system may refer to the same file. However, only one wrapper should be *owning* the data (see below).

In the XML input file, `Wrapper` correspond to attribute of an XML element representing the containing `Group`. See [XML Input](#) for the relationship between XML input files and Data Repository.

`Wrapper<T>` is templated on the type of object it encapsulates, thus providing strong type safety when retrieving the objects. As each `Wrapper` class instantiation will be a distinct type, `Wrapper` derives from a non-templated `WrapperBase` class that defines a common interface. `WrapperBase` is the type of pointer that is stored in the `MappedVector` container within a `Group`.

`WrapperBase` provides several interface functions that delegate the work to the wrapped object if it supports the corresponding method signature. This allows a collection of heterogeneous wrappers (i.e. over different types) to be treated uniformly. Examples include:

- `size()`
- `resize(newSize)`
- `reserve(newCapacity)`
- `capacity()`
- `move(LvArray::MemorySpace)`

A `Wrapper` may be *owning* or *non-owning*, depending on how it's constructed. An *owning* `Wrapper` will typically either take a previously allocated object via `std::unique_ptr<T>` or no pointer at all and itself allocate the object. It will delete the wrapped object when destroyed. A *non-owning* `Wrapper` may be used to register with the data repository objects that are not directly heap-allocated, for example data members of other objects. It will take a raw pointer as input and not delete the wrapped object when destroyed.

## Attributes

Each instance of `Wrapper` has a set of attributes that control its function in the data repository. These attributes are:

- **InputFlags**

A strongly typed enum that defines the relationship between the `Wrapper` and the XML input. Possible values are:

Value	Explanation
FALSE	Data is not read from XML input (default).
OPTIONAL	Data is read from XML if an attribute matching <code>Wrapper</code> 's name is found.
REQUIRED	Data is read from XML and an error is raised if the attribute is not found.

Other values of `InputFlags` enumeration are reserved for `Group` objects.

---

**Note:** A runtime error will occur when attempting to read from XML a wrapped type `T` that does not have `operator>>` defined.

---

- **RestartFlags**

Enumeration that describes how the `Wrapper` interacts with restart files.

Value	Explanation
NO_WRITE	Data is not written into restart files.
WRITE	Data is written into restart files but not read upon restart.
WRITE_AND_READ	Data is both written and read upon restart (default).

---

**Note:** A runtime error will occur when attempting to write a wrapped type `T` that does not support buffer packing. Therefore, when registering custom types (i.e. not a basic C++ type or an `LvArray` container) we recommend setting the flag to `NO_WRITE`. A future documentation topic will explain how to extend buffer packing capabilities to custom user-defined types.

---

- **PlotLevel**

Enumeration that describes how the `Wrapper` interacts with plot (visualization) files.

Value	Explanation
LEVEL_0	Data always written to plot files.
LEVEL_1	Data written to plot when <code>plotLevel</code> $\geq 1$ is specified in input.
LEVEL_2	Data written to plot when <code>plotLevel</code> $\geq 2$ is specified in input.
LEVEL_3	Data written to plot when <code>plotLevel</code> $\geq 3$ is specified in input.
NO_PLOT	Data never written to plot files.

---

**Note:** Only data stored in `LvArray`'s `Array<T>` containers is currently written into plot files.

---

## Default Values

Wrapper supports setting a default value for its wrapped object. The default value is used if a wrapper with `InputFlags::OPTIONAL` attribute does not match an attribute in the input file. For *LvArray* containers it is also used as a default value for new elements upon resizing the container.

Default value can be set via one of the following two methods:

- `setDefaultValue` sets the default value but does not affect the actual value stored in the wrapper.
- `setApplyDefaultValue` sets the default value *and* applies it to the stored value.

---

**Note:** A runtime error is raised if a default value is not set for a wrapper with `InputFlags::OPTIONAL` attribute.

---

The type `DefaultValue<T>` is used to store the default value for the wrapper.

---

**Todo:** `DefaultValue` is actually not a type but an alias for another internal struct. As such, it cannot currently be specialized for a user's custom type.

---

## API documentation

### Wrapper

### ObjectCatalog

The “ObjectCatalog” is a collection of classes that acts as a statically initialized factory. It functions in a similar manner to a classic [factory method](#), except that there is no maintained list of derived objects that is required to create new objects. In other words, there is no case-switch/if-elseif block to maintain. Instead, the `ObjectCatalog` creates a “catalog” of derived objects using a `std::unordered_map`. The “catalog” is filled when new types are declared through the declaration of a helper class named `CatalogEntryConstructor`.

The key functional features of the “ObjectCatalog” concept may be summarized as:

- Anonymous addition of new objects to the catalog. Because we use a statically initialized singleton map object to store the catalog, no knowledge of the contents of the catalog is required in the main code. Therefore if a proprietary/sensitive catalog entry is desired, it is only required that the object definition be outside of the main repository and tied into the build system through some non-specific mechanism (i.e. a link in the `src/externalComponents` directory) and the catalog entry will be registered in the catalog without sharing any knowledge of its existence. Then a proprietary input file may refer to the object to call for its creation.
- Zero maintenance catalog. Again, because we use a singleton map to store the catalog, there is no updating of code required to add new entries into the catalog. The only modifications required are the actual source files of the catalog entry, as described in the [Usage](#) section below.

## Implementation Details

There are three key objects that are used to provide the `ObjectCatalog` functionality.

## CatalogInterface

The `CatalogInterface` class provides the base definitions and interface for the `ObjectCatalog` concept. It is templated on the common base class of all derived objects that are creatable by the “ObjectCatalog”. In addition, `CatalogInterface` is templated on a variadic parameter pack that allows for an arbitrary constructor argument list as shown in the declaration shown below:

```
template< typename BASETYPE, typename ... ARGS >
class CatalogInterface
```

The `CatalogInterface` also defines the actual catalog type using the template arguments:

```
typedef std::unordered_map< std::string,
                           std::unique_ptr< CatalogInterface< BASETYPE, ARGS... > >
    > CatalogType;
```

The `CatalogInterface::CatalogType` is a `std::unordered_map` with a string “key” and a value type that is a pointer to the `CatalogInterface` that represents a specific combination of `BASETYPE` and constructor arguments.

After from setting up and populating the catalog, which will be described in the “Usage” section, the only interface with the catalog will typically be when the `Factory()` method is called. The definition of the method is given as:

```
static std::unique_ptr< BASETYPE > factory( std::string const & objectTypeName,
                                           ARGS... args )
{
    // We stop the simulation if the product is not found
    if( !hasKeyName( objectTypeName ) )
    {
        std::list< typename CatalogType::key_type > keys = getKeys();
        string const tmp = stringutilities::join( keys.cbegin(), keys.cend(), ",\n" );

        string errorMsg = "Could not find keyword \"" + objectTypeName + "\" in this_
        context. ";
        errorMsg += "Please be sure that all your keywords are properly spelled or that_
        input file parameters have not changed.\n";
        errorMsg += "All available keys are: [\n" + tmp + "\n]";
        GEOSX_ERROR( errorMsg );
    }

    // We also stop the simulation if the builder is not here.
    CatalogInterface< BASETYPE, ARGS... > const * builder = getCatalog().at(
    objectTypeName ).get();
    if( builder == nullptr )
    {
        const string errorMsg = "\"" + objectTypeName + "\" could be found. But the_
        builder is invalid.\n";
        GEOSX_ERROR( errorMsg );
    }

    return builder->allocate( args ... );
}
```

It can be seen that the static `Factory` method is simply a wrapper that calls the virtual `Allocate` method on a the catalog which is returned by `getCatalog()`. The usage of the `Factory` method will be further discussed in the *Usage* section.

**Note:** The method for organizing constructing new objects relies on a common constructor list between the derived type and the `BASETYPE`. This means that there is a single catalog for each combination of `BASETYPE` and the variadic parameter pack representing the constructor arguments. In the future, we can investigate removing this restriction and allowing for construction of a hierarchy of objects with an arbitrary constructor parameter list.

## CatalogEntry

The `CatalogEntry` class derives from `CatalogInterface` and adds the a `TYPE` template argument to the arguments of the `CatalogInterface`.

```
template< typename BASETYPE, typename TYPE, typename ... ARGS >
class CatalogEntry final : public CatalogInterface< BASETYPE, ARGS... >
```

The `TYPE` template argument is the type of the object that you would like to be able to create with the “ObjectCatalog”. `TYPE` must be derived from `BASETYPE` and have a constructor that matches the variadic parameter pack specified in the template parameter list. The main purpose of the `CatalogEntry` is to override the `CatalogInterface::Allocate()` virtual function s.t. when key is retrieved from the catalog, then it is possible to create a new `TYPE`. The `CatalogEntry::Allocate()` function is a simple creation of the underlying `TYPE` as shown by its definition:

```
virtual std::unique_ptr< BASETYPE > allocate( ARGS... args ) const override
{
    #if OBJECTCATALOGVERBOSE > 0
        GEOSX_LOG( "Creating type " << LvArray::system::demangle( typeid(TYPE).name() )
                    << " from catalog of " << LvArray::system::demangle(
->typeid(BASETYPE).name() ) );
    #endif
    #if ( __cplusplus >= 201402L )
        return std::make_unique< TYPE >( args ... );
    #else
        return std::unique_ptr< BASETYPE >( new TYPE( args ... ) );
    #endif
}
```

## CatalogEntryConstructor

The `CatalogEntryConstructor` is a helper class that has a sole purpose of creating a new `CatalogEntry` and adding it to the catalog. When a new `CatalogEntryConstructor` is created, a new `CatalogEntry` entry is created and inserted into the catalog automatically.

## Usage

### Creating A New Catalog

When creating a new “ObjectCatalog”, it typically is done within the context of a specific `BASETYPE`. A simple example of a class hierarchy in which we would like to use the “ObjectCatalog” to use to generate new objects is given in the unit test located in `testObjectCatalog.cpp`.

The base class for this example is defined as:

```

class Base
{
public:
    Base( int & junk, double const & junk2 )
    {
        GEOSX_LOG( "calling Base constructor with arguments ("<<junk<<" "<<junk2<<")" );
    }

    virtual ~Base()
    {
        GEOSX_LOG( "calling Base destructor" );
    }

    using CatalogInterface = dataRepository::CatalogInterface< Base, int &, double_
    ↪const & >;
    static CatalogInterface::CatalogType & getCatalog()
    {
        static CatalogInterface::CatalogType catalog;
        return catalog;
    }

    virtual string getCatalogName() = 0;
};

```

There a couple of things to note in the definition of Base:

- Base has a convenience alias to use in place of the fully templated CatalogInterface name.
- Base defines a getCatalog() function that returns a static instantiation of a CatalogInterface::CatalogType. The CatalogInterface::getCatalog() function actually calls this function within the base class. This means that the base class actually owns the catalog, and the CatalogInterface is only operating on that Base::getCatalog(), and that the definition of this function is required.

## Adding A New Type To The Catalog

Once a Base class is defined with the required features, the next step is to add a new derived type to the catalog defined in Base. There are three requirements for the new type to be registered in the catalog:

- The derived type must have a constructor with the arguments specified by the variadic parameter pack specified in the catalog.
- There must be a static function static string catalogName() that returns the name of the type that will be used to as keyname when it is registered Base's catalog.
- The new type must be registered with the catalog held in Base. To accomplish this, a convenience macro REGISTER\_CATALOG\_ENTRY() is provided. The arguments to this macro are the name type of Base, the type of the derived class, and then the variadic pack of constructor arguments.

A pair of of simple derived class that have the required methods are used in the unit test.

```

class Derived1 : public Base
{
public:
    Derived1( int & junk, double const & junk2 ):
        Base( junk, junk2 )
    {

```

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```

    GEOSX_LOG( "calling Derived1 constructor with arguments ("<<junk<<" "<<junk2<<") " );
    ↪);
}

~Derived1()
{
    GEOSX_LOG( "calling Derived1 destructor" );
}
static string catalogName() { return "derived1"; }
string getCatalogName() { return catalogName(); }

};
REGISTER_CATALOG_ENTRY( Base, Derived1, int &, double const & )

```

```

class Derived2 : public Base
{
public:
    Derived2( int & junk, double const & junk2 ):
        Base( junk, junk2 )
    {
        GEOSX_LOG( "calling Derived2 constructor with arguments ("<<junk<<" "<<junk2<<") " );
        ↪);
    }

    ~Derived2()
    {
        GEOSX_LOG( "calling Derived2 destructor" );
    }
    static string catalogName() { return "derived2"; }
    string getCatalogName() { return catalogName(); }

};
REGISTER_CATALOG_ENTRY( Base, Derived2, int &, double const & )

```

## Allocating A New Object From The Catalog

The test function in the unit test shows how to allocate a new object of one of the derived types from Factory method. Note the call to Factory is scoped by Base::CatalogInterface, which is an alias to the full templated instantiation of CatalogInterface. The arguments for Factory

```

TEST( testObjectCatalog, testRegistration )
{
    GEOSX_LOG( "EXECUTING MAIN" );
    int junk = 1;
    double junk2 = 3.14;

    // allocate a new Derived1 object
    std::unique_ptr< Base >
    derived1 = Base::CatalogInterface::factory( "derived1", junk, junk2 );

    // allocate a new Derived2 object
    std::unique_ptr< Base >
    derived2 = Base::CatalogInterface::factory( "derived2", junk, junk2 );
}

```

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```

EXPECT_STREQ( derived1->getCatalogName().c_str(),
              Derived1::catalogName().c_str() );

EXPECT_STREQ( derived2->getCatalogName().c_str(),
              Derived2::catalogName().c_str() );
GEOSX_LOG( "EXITING MAIN" );
}

```

The unit test creates two new objects of type `Derived1` and `Derived2` using the catalogs `Factory` method. Then the test checks to see that the objects that were created are of the correct type. This unit test has some extra output to screen to help with understanding of the sequence of events. The result of running this test is:

```

$ tests/testObjectCatalog
Calling constructor for CatalogEntryConstructor< Derived1 , Base , ... >
Calling constructor for CatalogInterface< Base , ... >
Calling constructor for CatalogEntry< Derived1 , Base , ... >
Registered Base catalog component of derived type Derived1 where
↳Derived1::catalogName() = derived1
Calling constructor for CatalogEntryConstructor< Derived2 , Base , ... >
Calling constructor for CatalogInterface< Base , ... >
Calling constructor for CatalogEntry< Derived2 , Base , ... >
Registered Base catalog component of derived type Derived2 where
↳Derived2::catalogName() = derived2
Running main() from gtest_main.cc
[=====] Running 1 test from 1 test case.
[-----] Global test environment set-up.
[-----] 1 test from testObjectCatalog
[ RUN      ] testObjectCatalog.testRegistration
EXECUTING MAIN
Creating type Derived1 from catalog of Base
calling Base constructor with arguments (1 3.14)
calling Derived1 constructor with arguments (1 3.14)
Creating type Derived2 from catalog of Base
calling Base constructor with arguments (1 3.14)
calling Derived2 constructor with arguments (1 3.14)
EXITING MAIN
calling Derived2 destructor
calling Base destructor
calling Derived1 destructor
calling Base destructor
[ OK ] testObjectCatalog.testRegistration (0 ms)
[-----] 1 test from testObjectCatalog (0 ms total)

[-----] Global test environment tear-down
[=====] 1 test from 1 test case ran. (0 ms total)
[ PASSED ] 1 test.
Calling destructor for CatalogEntryConstructor< Derived2 , Base , ... >
Calling destructor for CatalogEntryConstructor< Derived1 , Base , ... >
Calling destructor for CatalogEntry< Derived2 , Base , ... >
Calling destructor for CatalogInterface< Base , ... >
Calling destructor for CatalogEntry< Derived1 , Base , ... >
Calling destructor for CatalogInterface< Base , ... >

```

In the preceding output, it is clear that the static catalog in `Base::getCatalog()` is initialized prior the execution of `main`, and destroyed after the completion of `main`. In practice, there have been no indicators of problems due to the use of a statically initialized/deinitialized catalog.

## 6.2.2 XML Input

In this document, you will learn how GEOSX classes interact with external information parsed from XML files, and how to add a new XML block that can be interpreted by GEOSX. Flow solvers and relative permeability are used as examples.

### GEOSX data structure overview

#### Group : the base class of GEOSX

All GEOSX classes derive from a base class called `dataRepository::Group`. The `Group` class provides a way to organize all GEOSX objects in a filesystem-like structure. One could think of `Group`s as *file folders* that can bear data (stored in `Wrapper`s), have a parent folder (another `Group`), and have possibly multiple subfolders (referred to as the subgroups). Below, we briefly review the data members of the `Group` class that are essential to understand the correspondence between the GEOSX data structure and the XML input. For more details, we refer the reader to the extensive documentation of the *Data Repository*, including the *Group* class documentation.

In the code listing below, we see that each `Group` object is at minimum equipped with the following member properties:

- A pointer to the parent `Group` called `m_parent` (member classes are prefixed by `m_`),
- The `Group`'s own data, stored for flexibility in an array of generic data `Wrapper`s called `m_wrappers`,
- A map of one or many children (also of type `Group`) called `m_subGroups`.
- The `m_size` and `m_capacity` members, that are used to set the size and capacity of any objects contained.
- The name of the `Group`, stored as a string in `m_name`. This name can be seen as the object unique ID.

```

/// The parent Group that contains "this" Group in its "sub-Group" collection.
Group * m_parent = nullptr;

/// Specification that this group will have the same m_size as m_parent.
integer m_sizedFromParent;

/// The container for the collection of all wrappers contained in "this" Group.
wrapperMap m_wrappers;

/// The container for the collection of all sub-groups contained in "this" Group.
subGroupMap m_subGroups;

/// The size/length of this Group...and all Wrapper<> that are are specified to_
↳ have the same size as their
/// owning group.
indexType m_size;

/// The capacity for wrappers in this group...and all Wrapper<> that are specified_
↳ to have the same size as their
/// owning group.
indexType m_capacity;

/// The name/key of this Group in its parent collection of sub-Groups.
string m_name;

/// Verbosity flag for group logs
integer m_logLevel;
//END_SPHINX_INCLUDE_02

```

(continues on next page)

```
/// Restart flag for this group... and subsequently all wrappers in this group.
```

[Source: `src/coreComponents/dataRepository/Group.hpp`]

## A few words about the ObjectCatalog

### What is an ObjectCatalog and why do we need it?

Some classes need external information (physical and/or algorithmic parameters for instance) provided by the user to be instantiated. This is the case when the `m_input_flags` data member of one of the `Group's Wrapper's` has an entry set to `REQUIRED` (we will illustrate this below). In this situation, the required information must be supplied in the XML input file, and if it is absent, an error is raised by GEOSX.

To connect the external (XML) and internal (C++) data structures, GEOSX uses an **ObjectCatalog** that maps keys (of type `string`) to the corresponding classes (one unique key per mapped class). These string keys, referred to as `catalogName's`, are essential to transfer the information from the XML file to the factory functions in charge of object instantiation (see below).

### What is a CatalogName?

The `catalogName` of an object is a *key* (of type `string`) associated with this object's class. On the one hand, in the XML file, the key is employed by the user as an XML tag to specify the type of object (e.g., the type of solver, constitutive model, etc) to create and use during the simulation. On the other hand, internally, the key provides a way to access the appropriate factory function to instantiate an object of the desired class.

Most of the time, the `catalogName` and the C++ class name are identical. This helps make the code easier to debug and allows the XML/C++ correspondence to be evident. But strictly speaking, the `catalogName` can be anything, as long as it refers uniquely to a specific class. The `catalogName` must not be confused with the object's *name* (`m_name` is a data member of the class that stores the object's unique ID, not its class key). You can have several objects of the same class and hence the same `catalogName`, but with different names (i.e. unique ID): several fluid models, several solvers, etc.

### How can I add my new externally-accessible class to the ObjectCatalog?

Let us consider a flow solver class derived from `FlowSolverBase`, that itself is derived from `SolverBase`. To instantiate and use this solver, the developer needs to make the derived flow solver class reachable from the XML file, via an XML tag. Internally, this requires adding the derived class information to `ObjectCatalog`, which is achieved with two main ingredients: 1) a `CatalogName()` method in the class that lets GEOSX know *what* to search for in the internal `ObjectCatalog` to instantiate an object of this class, 2) a macro that specifies *where* to search in the `ObjectCatalog`.

1. To let GEOSX know what to search for in the catalog to instantiate an object of the derived class, the developer must equip the class with a `CatalogName()` method that returns a `string`. In this document, we have referred to this returned `string` as the object's `catalogName`, but in fact, the method `CatalogName()` is what matters since the `ObjectCatalog` contains all the `CatalogName()` return values. Below, we illustrate this with the `CompositionalMultiphaseFlow` solver. The first code listing defines the class name, which in this case is the same as the `catalogName` shown in the second listing.

```
/**
 * @class CompositionalMultiphaseBase
 *
 * A compositional multiphase solver
 */
class CompositionalMultiphaseBase : public FlowSolverBase
{
```

[Source: *src/coreComponents/physicsSolvers/fluidFlow/CompositionalMultiphaseBase.hpp*]

2. To let GEOSX know where to search in the `ObjectCatalog`, a macro needs to be added at the end of the .cpp file implementing the class. This macro (illustrated below) must contain the type of the base class (in this case, `SolverBase`), and the name of the derived class (continuing with the example used above, this is `CompositionalMultiphaseFlow`). As a result of this construct, the `ObjectCatalog` is not a flat list of `strings` mapping the C++ classes. Instead, the `ObjectCatalog` forms a tree that reproduces locally the structure of the class diagram, from the base class to the derived classes.

```
REGISTER_CATALOG_ENTRY( SolverBase, CompositionalMultiphaseFVM, string const &, Group_
↳ * const )
```

[Source: *src/coreComponents/physicsSolvers/fluidFlow/CompositionalMultiphaseFVM.cpp*]

Summary: All GEOSX objects form a filesystem-like structure. If an object needs to be accessible externally, it must be registered in the `ObjectCatalog`. This is done by adding `CatalogName()` method that returns a `string` key to the object's class, and by adding the appropriate macro. The catalog has the same tree structure as the class diagram.

## Registration: parsing XML input files to instantiate GEOSX objects

In this section, we describe with more details the connection between **internal GEOSX objects** and **external XML tags** parsed from parameter files. We call this process *Registration*. The registration process works in three steps:

1. The XML document is parsed. Each time a new XML tag is found, the current local scope of the `ObjectCatalog` is inspected. The goal is to find a `catalogName` string that matches the XML tag.
2. If it is the case (the current local scope of the `ObjectCatalog` contains a `catalogName` identical to the XML tag), then the code creates a new instance of the class that the `catalogName` refers to. This new object is inserted in the `Group` tree structure at the appropriate location, as a subgroup.
3. By parsing the XML attributes of the tag, the new object properties are populated. Some checks are performed to ensure that the data supplied is conform, and that all the required information is present.

Let's look at this process in more details.

## Creating a new object and giving it a Catalog name

Consider again that we are registering a flow solver deriving from `FlowSolverBase`, and assume that this solver is called `CppNameOfMySolver`. This choice of name is not recommended (we want names that reflect what the solver does!), but for this particular example, we just need to know that this name is the class name inside the C++ code.

To specify parameters of this new solver from an XML file, we need to be sure that the XML tag and the `catalogName` of the class are identical. Therefore, we equip the `CppNameOfMySolver` class with a `CatalogName()` method that returns the solver `catalogName` (=XML name). Here, this method returns the string "`XmlNameOfMySolver`".

We have deliberately distinguished the class name from the catalog/XML name for the sake of clarity in this example. It is nevertheless a best practice to use the same name for the class and for the `catalogName`. This is the case below for the existing `CompositionalMultiphaseFVM` class.

```
class CompositionalMultiphaseFVM : public CompositionalMultiphaseBase
{
```

```

/**
 * @brief name of the solver in the object catalog
 * @return string that contains the catalog name to generate a new object through_
↳ the object catalog.
 */
static string catalogName() { return "CompositionalMultiphaseFVM"; }

```

[Source: src/coreComponents/physicsSolvers/fluidFlow/CompositionalMultiphaseFVM.hpp]

## Parsing XML and searching the ObjectCatalog in scope

Now that we have implemented a `CatalogName()` method returning a specific key (of type `string`), we can have a block in our XML input file with a tag that corresponds to the `catalogName` “`XmlNameOfMySolver`”. This is how the XML block would look like.

```

<Problem>
  <Solvers
    gravityVector="{ 0.0, 0.0, -9.81 }">
      <XmlNameOfMySolver name="nameOfThisSolverInstance"
        verboseLevel="1"
        gravityFlag="1"
        temperature="297.15" />
        <LinearSolverParameters newtonTol="1.0e-6"
          maxIterNewton="15"
          useDirectSolver="1"/>
      </XmlNameOfMySolver>
    </Solvers>
  </Problem>

```

Here, we see that the XML structure defines a parent node “Problem”, that has (among many others) a child node “Solvers”. In the “Solvers” block, we have placed the new solver block as a child node of the “Solvers” block with the XML tag corresponding to the `catalogName` of the new class. We will see in details next how the GEOSX internal structure constructed from this block mirrors the XML file structure.

## Instantiating the new solver

Above, we have specified an XML block with the tag “`XmlNameOfMySolver`”. Now, when reading the XML file and encountering an “`XmlNameOfMySolver`” solver block, we add a new instance of the class `CppNameOfMySolver` in the filesystem structure as explained below.

We saw that in the XML file, the new solver block appeared as child node of the XML block “Solvers”. The internal construction mirrors this XML structure. Specifically, the new object of class `CppNameOfMySolver` is registered as a subgroup (to continue the analogy used so far, as a subfolder) of its parent Group, the class `PhysicsSolverManager` (that has a `catalogName` “Solvers”). To do this, the method `CreateChild` of the `PhysicsSolverManager` class is used.

```

// Variable values in this example:
// -----
// childKey = "XmlNameOfMySolver" (string)
// childName = "nameOfThisSolverInstance" (string)
// SolverBase::CatalogInterface = the Catalog attached to the base Solver class
// hasKeyName = bool method to test if the childKey string is present in the Catalog
// registerGroup = method to create a new instance of the solver and add it to the_
↳ group tree

```

```

Group * PhysicsSolverManager::createChild( string const & childKey, string const &
↳ childName )
{
    Group * rval = nullptr;
    if( SolverBase::CatalogInterface::hasKeyName( childKey ) )
    {
        GEOSX_LOG_RANK_0( "Adding Solver of type " << childKey << ", named " << childName
↳ );
        rval = &registerGroup( childName,
                               SolverBase::CatalogInterface::factory( childKey, childName,
↳ this ) );
    }
    return rval;
}

```

[Source: *src/coreComponents/physicsSolvers/PhysicsSolverManager.cpp*]

In the code listing above, we see that in the `PhysicsSolverManager` class, the `ObjectCatalog` is searched to find the catalogName “`CompositionalMultiphaseFlow`” in the scope of the `SolverBase` class. Then, the factory function of the base class `SolverBase` is called. The catalogName (stored in `childKey`) is passed as an argument of the factory function to ensure that it instantiates an object of the desired derived class.

As explained above, this is working because 1) the XML tag matches the catalogName of the `CompositionalMultiphaseFlow` class and 2) a macro is placed at the end of the .cpp file implementing the `CompositionalMultiphaseFlow` class to let the `ObjectCatalog` know that `CompositionalMultiphaseFlow` is a derived class of `SolverBase`.

Note that several instances of the same type of solver can be created, as long as they each have a different name.

### Filling the objects with data (wrappers)

After finding and placing the new solver `Group` in the filesystem hierarchy, properties are read and stored. This is done by registering *data wrappers*. We refer to the documentation of the [Data Repository](#) for additional details about the `Wrapper`s. The method used to do that is called `registerWrapper` and is placed in the class constructor when the data is required in the XML file. Note that some properties are registered at the current (derived) class level, and other properties can also be registered at a base class level.

Here, the only data (=wrapper) that is defined at the level of our `CppNameOfMySolver` class is temperature, and everything else is registered at the base class level. We register a property of temperature, corresponding to the member class `m_temperature` of `CppNameOfMySolver`. The registration also checks if a property is required or optional (here, it is required), and provides a brief description that will be used in the auto-generated code documentation.

```

this->registerWrapper( viewKeyStruct::inputTemperatureString(), &m_inputTemperature
↳ ).
    setInputFlag( InputFlags::REQUIRED ).
    setDescription( "Temperature" );
//END_SPHINX_INCLUDE_00

```

[Source: *src/coreComponents/physicsSolvers/fluidFlow/CompositionalMultiphaseBase.cpp*]

This operation is done recursively if XML tags are nested.

### To summarize:

- Every class in GEOSX derive from a `Group` in a filesystem-like structure. A `Group` must have a parent `Group`, can have data (in `Wrapper` s), and can have one or many children (the subgroups). There is an

ObjectCatalog in which the classes derived from Group are identified by a key called the catalogName.

- When parsing XML input files, GEOSX inspects each object's scope in the ObjectCatalog to find classes with the same catalogName as the XML tag. Once it finds an XML tag in the ObjectCatalog, it registers it inside the filesystem structure.
- In the registration process, properties from the XML file are parsed and used to allocate member data Wrappers and fully instantiate the Group class.
- If XML tags are nested, subgroups are allocated and processed in a nested manner.

The correspondence between XML and class hierarchy is thus respected, and the internal object hierarchy mirrors the XML structure.

### Example: adding a new relative permeability model

This example is taken from the class BrooksCoreyRelativePermeability, derived from RelativePermeabilityBase.

#### Implement a CatalogName function (.hpp):

As explained above we add the class to the ObjectCatalog in two steps. First we implement the CatalogName function:

```
static string catalogName() { return "BrooksCoreyRelativePermeability"; }
```

[source: src/coreComponents/constitutive/relativePermeability/BrooksCoreyRelativePermeability.hpp]

Then in the .cpp file we add the macro to register the catalog entry:

```
REGISTER_CATALOG_ENTRY( ConstitutiveBase, BrooksCoreyRelativePermeability, string_  
↳const &, Group * const )
```

[source: src/coreComponents/constitutive/relativePermeability/BrooksCoreyRelativePermeability.cpp]

Now every time a “BrooksCoreyRelativePermeability” string is encountered inside a Relative Permeability catalog, we will instantiate a class BrooksCoreyRelativePermeability.

#### Declare the Wrapper s keys (.hpp):

When attaching properties (i.e. data Wrapper s) to a class, a similar registration process must be done. Every property is accessed through its ViewKey namespace. In this namespace, we define string s that correspond to the tags of XML attributes of the “BrooksCoreyRelativePermeability” block.

```
struct viewKeyStruct : RelativePermeabilityBase::viewKeyStruct  
{  
    static constexpr char const * phaseMinVolumeFractionString() { return  
↳"phaseMinVolumeFraction"; }  
    static constexpr char const * phaseRelPermExponentString() { return  
↳"phaseRelPermExponent"; }  
    static constexpr char const * phaseRelPermMaxValueString() { return  
↳"phaseRelPermMaxValue"; }  
    static constexpr char const * volFracScaleString() { return "volFracScale"; }  
} vieKeysBrooksCoreyRelativePermeability;
```

[source: src/coreComponents/constitutive/relativePermeability/BrooksCoreyRelativePermeability.hpp]



### Declare data members (.hpp):

The data members are defined in the class. They will ultimately contain the data read from the XML file (other data members not read from the XML file can also exist).

```
array1d< real64 > m_phaseMinVolumeFraction;
array1d< real64 > m_phaseRelPermExponent;
array1d< real64 > m_phaseRelPermMaxValue;

real64 m_volFracScale;
```

[source: *src/coreComponents/constitutive/relativePermeability/BrooksCoreyRelativePermeability.hpp*]

### Implement the data registration process (registerWrapper):

The registration process done in the class constructor puts everything together. It connects the attributes values in the XML file to class member data. For instance, in the listing below, the first registerWrapper call means that we want to read in the XML file the attribute value corresponding to the attribute tag “phaseMinVolumeFraction” defined in the .hpp file, and that we want to store the read values into the m\_phaseMinVolumeFraction data members. We see that this input is not required. If it is absent from the XML file, the default value is used instead. The short description that completes the registration will be added to the auto-generated documentation.

```
BrooksCoreyRelativePermeability::BrooksCoreyRelativePermeability( string const & name,
                                                                    Group * const
↳parent )
    : RelativePermeabilityBase( name, parent )
{
    registerWrapper( viewKeyStruct::phaseMinVolumeFractionString(), &m_
↳phaseMinVolumeFraction ).
        setApplyDefaultValue( 0.0 ).
        setInputFlag( InputFlags::OPTIONAL ).
        setDescription( "Minimum volume fraction value for each phase" );

    registerWrapper( viewKeyStruct::phaseRelPermExponentString(), &m_
↳phaseRelPermExponent ).
        setApplyDefaultValue( 1.0 ).
        setInputFlag( InputFlags::OPTIONAL ).
        setDescription( "Minimum relative permeability power law exponent for each phase"
↳);

    registerWrapper( viewKeyStruct::phaseRelPermMaxValueString(), &m_
↳phaseRelPermMaxValue ).
        setApplyDefaultValue( 0.0 ).
        setInputFlag( InputFlags::OPTIONAL ).
        setDescription( "Maximum relative permeability value for each phase" );

    registerWrapper( viewKeyStruct::volFracScaleString(), &m_volFracScale ).
        setApplyDefaultValue( 1.0 ).
        setDescription( "Factor used to scale the phase relative permeability, defined
↳as: one minus the sum of the phase minimum volume fractions." );
```

[source: *src/coreComponents/constitutive/relativePermeability/BrooksCoreyRelativePermeability.cpp*]

## The XML block

We are ready to use the relative permeability model in GEOSX. The corresponding XML block (child node of the “Constitutive” block) reads:

```
<Constitutive>
  <BrooksCoreyBakerRelativePermeability name="relperm"
    phaseNames="{oil, gas, water}"
    phaseMinVolumeFraction="{0.05, 0.05, 0.05}"
    waterOilRelPermExponent="{2.5, 1.5}"
    waterOilRelPermMaxValue="{0.8, 0.9}"
    gasOilRelPermExponent="{3, 3}"
    gasOilRelPermMaxValue="{0.4, 0.9}"/>
</Constitutive>
```

With this construct, we instruct the `ConstitutiveManager` class (whose `catalogName` is “Constitutive”) to instantiate a subgroup of type `BrooksCoreyRelativePermeability`. We also fill the data members of the values that we want to use for the simulation. For a simulation with multiple regions, we could define multiple relative permeability models in the “Constitutive” XML block (yielding multiple `relperm` subgroups in GEOSX), with a unique name attribute for each model.

*For more examples on how to contribute to GEOSX, please read [Adding a new Physics Solver](#)*

## Input Schema Generation

A schema file is a useful tool for validating input .xml files and constructing user-interfaces. Rather than manually maintaining the schema during development, GEOSX is designed to automatically generate one by traversing the documentation structure.

To generate the schema, run GEOSX with the input, schema, and the (optional) `schema_level` arguments, i.e.: `geosx -i input.xml -s schema.xsd`. There are two ways to limit the scope of the schema:

1. Setting the verbosity flag for an object in the documentation structure. If the schema-level argument is used, then only objects (and their children) and attributes with `(verbosity < schema-level)` will be output.
2. By supplying a limited input xml file. When GEOSX builds its data structure, it will only include objects that are listed within the xml (or those that are explicitly appended when those objects are initialized). The code will add all available *attributes* for these objects to the schema.

To take advantage of this design it is necessary to use the automatic xml parsing approach that relies upon the documentation node. If values are read in manually, then the schema can not be used to validate xml those inputs.

Note: the lightweight xml parser that is used in GEOSX cannot be used to validate inputs with the schema directly. As such, it is necessary to use an external tool for validation, such as the `geosx_tools` python module.

### 6.2.3 Working with data in GEOSX

In GEOSX, data is typically registered in the [Data Repository](#). This allows for the writing/reading of data to/from restart and plot files. Any object that derives from [Group](#) may have data registered on it through the methods described in [Group](#). Similarly, accessing data from outside the scope of an object is possible through one of the various `Group::get()`. Of course, for temporary data that does not need to persist between cycles, or across physics packages, you may simply define member or local variables which will not be registered with the [Data Repository](#).

## Working with data on the Mesh objects

the mesh objects in GEOSX such as the `FaceManager` or `NodeManager`, are derived from `ObjectManagerBase`, which in turn derives from `Group`. The important distinction is that `ObjectManagerBase` contains various members that are useful when defining mesh object managers. When considering data that is attached to a mesh object, we group the data into two categories:

- **Intrinsic** data is data that is required to describe the object. For instance, to define a `Node`, the `NodeManager` contains an array of positions corresponding to each `Node` it contains. Thus the `ReferencePosition` is Intrinsic data. Intrinsic data is almost always a member of the mesh object, and is registered on the mesh object in the constructor of mesh object itself.
- **Extrinsic** data is data that is not required to define the object. For instance, a physics package may request that a `Velocity` value be stored on the nodes. Appropriately the data will be registered on the `NodeManager`. However, this data is not required to define a `Node`, and is viewed as Extrinsic. Extrinsic data is never a member of the mesh object, and is typically registered on the mesh object outside of the definition of the mesh object (i.e. from a physics solver).

## Registering Intrinsic data on a Mesh Object

As mentioned above, Intrinsic data is typically a member of the mesh object, and is registered in the constructor of the mesh Object. Taking the `NodeManager` and the `referencePosition` as an example, we point out that the reference position is actually a member in the `NodeManager`.

```
/**
 * @brief Get the mutable reference position array. This table will contain all the
 * ↪node coordinates.
 * @return reference position array
 */
array2d< real64, nodes::REFERENCE_POSITION_PERM > & referencePosition() { return m_
↪referencePosition; }

/**
 * @brief Provide an immutable arrayView of the reference position. This table will
 * ↪contain all the node coordinates.
 * @return an immutable arrayView of the reference position.
 */

arrayView2d< real64 const, nodes::REFERENCE_POSITION_USD > referencePosition() const
{ return m_referencePosition; }
```

This member is registered in the constructor for the `NodeManager`.

```
NodeManager::NodeManager( string const & name,
                          Group * const parent ):
    ObjectManagerBase( name, parent ),
    m_referencePosition( 0, 3 )
{
    registerWrapper( viewKeyStruct::referencePositionString(), &m_referencePosition );
}
```

Finally in order to access this data, the `NodeManager` provides explicit accessors.

```
/**
 * @brief Get the mutable reference position array. This table will contain all the
 * ↪node coordinates.
 * @return reference position array
```

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```

    */
    array2d< real64, nodes::REFERENCE_POSITION_PERM > & referencePosition() { return m_
    ↳referencePosition; }

    /**
     * @brief Provide an immutable arrayView of the reference position. This table will
    ↳contain all the node coordinates.
     * @return an immutable arrayView of the reference position.
     */

    arrayView2d< real64 const, nodes::REFERENCE_POSITION_USD > referencePosition() const
    { return m_referencePosition; }

```

Thus the interface for Intrinsic data is set by the object that it is a part of, and the developer may only access the data through the accessors from outside of the mesh object class scope.

## Registering Extrinsic data on a Mesh Object

To register Extrinsic data, there are many ways a developer may proceed. We will use the example of registering a TotalDisplacement on the NodeManager from the SolidMechanics solver. The most general approach is to define a string key and call one of the `Group::registerWrapper()` functions from `SolverBase::RegisterMeshData()`. Then when you want to use the data, you can call `Group::getReference()`. For example this would look something like:

```

void SolidMechanicsLagrangianFEM::RegisterDataOnMesh( Group * const MeshBodies )
{
    for( auto & mesh : MeshBodies->GetSubGroups() )
    {
        NodeManager & nodes = mesh.second->groupCast< MeshBody * >()->getMeshLevel( 0 ).
        ↳getNodeManager();

        nodes.registerWrapper< array2d< real64, nodes::TOTAL_DISPLACEMENT_PERM > > (
        ↳keys::TotalDisplacement ).
            setPlotLevel( PlotLevel::LEVEL_0 ).
            setRegisteringObjects( this->getName() ).
            setDescription( "An array that holds the total displacements on the nodes." ).
            reference().resizeDimension< 1 >( 3 );
    }
}

```

and

```

arrayView2d< real64, nodes::TOTAL_DISPLACEMENT_USD > const & u = nodes.getReference
↳<array2d<real64, nodes::TOTAL_DISPLACEMENT_PERM >(keys::TotalDisplacement);
... do something with u

```

This approach is flexible and extendible, but is potentially error prone due to its verbosity. Therefore we also provide a more controlled/uniform method by which to register and extract commonly used data on the mesh. The trait approach requires the definition of a traits struct for each data object that will be supported. To apply the trait approach to the example use case shown above, there should be the following definition somewhere in a header file:

```

namespace extrinsicMeshData
{

```

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```

struct TotalDisplacement
{
    static constexpr auto key = "totalDisplacement";
    using DataType = real64;
    using Type = array2d< DataType, nodes::TOTAL_DISPLACEMENT_PERM >;
    static constexpr DataType defaultValue = 0;
    static constexpr auto plotLevel = dataRepository::PlotLevel::LEVEL_0;

    /// Description of the data associated with this trait.
    static constexpr auto description = "An array that holds the total displacements on_
↪the nodes.";
};
}

```

Then the registration is simplified as follows:

```

void SolidMechanicsLagrangianFEM::RegisterDataOnMesh( Group * const MeshBodies )
{
    for( auto & mesh : MeshBodies->GetSubGroups() )
    {
        NodeManager & nodes = mesh.second->groupCast< MeshBody * >()->getMeshLevel( 0 ).
↪getNodeManager();
        nodes.registerExtrinsicData< extrinsicMeshData::TotalDisplacement >( this->
↪getName() ).resizeDimension< 1 >( 3 );
    }
}

```

And to extract the data, the call would be:

```

arrayView2d< real64, nodes::TOTAL_DISPLACEMENT_USD > const & u = nodes.
↪getExtrinsicData< extrinsicMeshData::TotalDisplacement >();
... do something with u

```

The end result of the `trait` approach to this example is that the developer has defined a standard specification for `TotalDisplacement`, which may be used uniformly across the code.

## 6.2.4 Mesh Hierarchy

In GEOSX, the mesh structure consists of a hierarchy of classes intended to encapsulate data and functionality for each topological type. Each class in the mesh hierarchy represents a distinct topological object, such as a nodes, edges, faces, elements, etc. The mesh data structure is illustrated in an object instantiation hierarchy. The object instantiation hierarchy differs from a “class hierarchy” in that it shows how instantiations of each class relate to each other in the data hierarchy rather than how each class type relates to each other in an inheritance diagram.

To illustrate the mesh hierarchy, we propose to present it along with a model with two regions (Top and Bottom) (Fig. 6.2).

### DomainPartition

In `MeshObjectInstantiationHierarchy` the top level object `DomainPartition` represents a partition of the decomposed physical domain. At this time there is a unique `DomainPartition` for every MPI rank.

**Note:** Hypothetically, there may be more than one `DomainPartition` in cases where the ranks are overloaded. Currently GEOSX does not support overloading multiple `DomainPartition`’s onto a rank, although this may be a

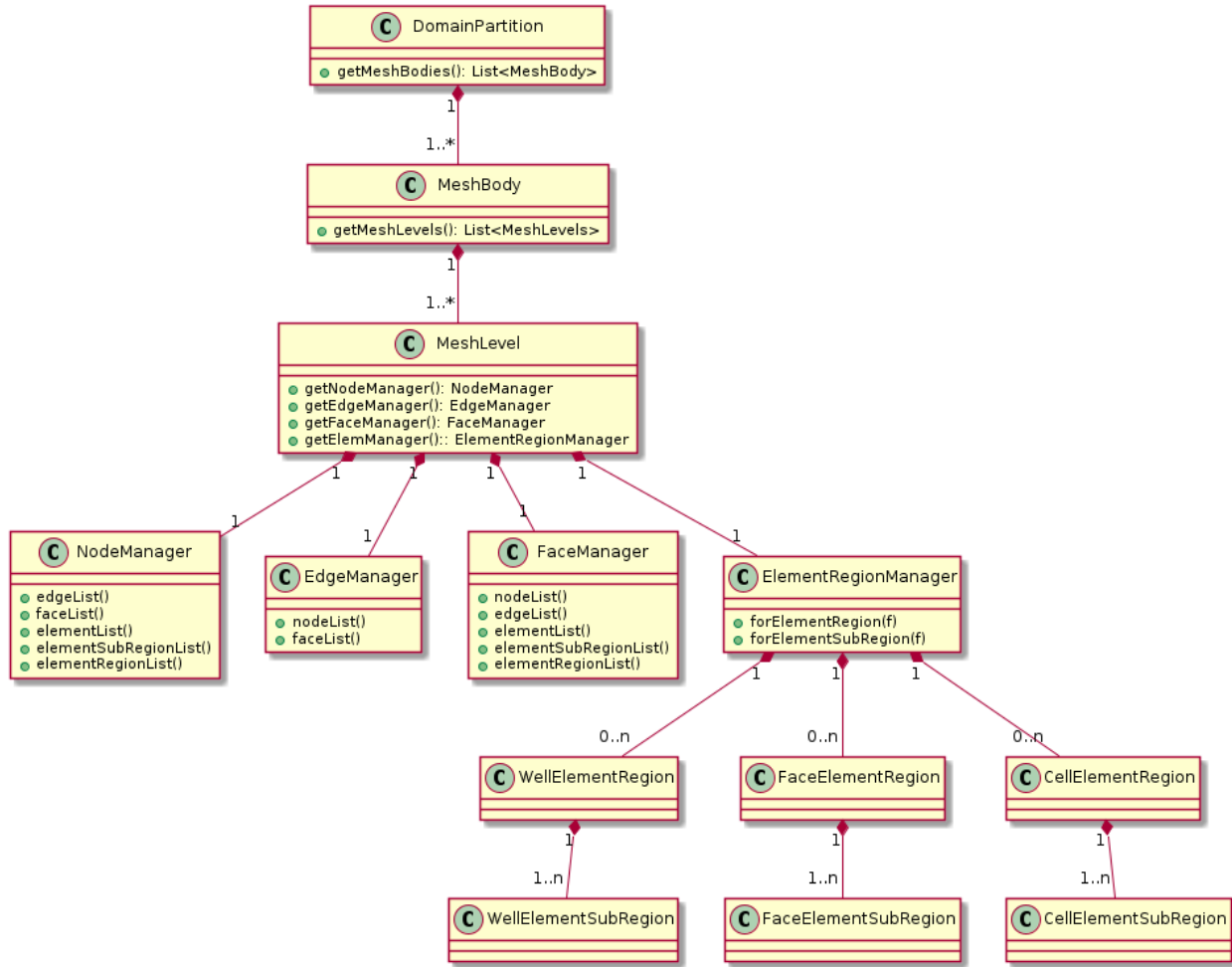


Fig. 6.1: Object instances describing the mesh domain. Cardinalities and relationships are indicated.

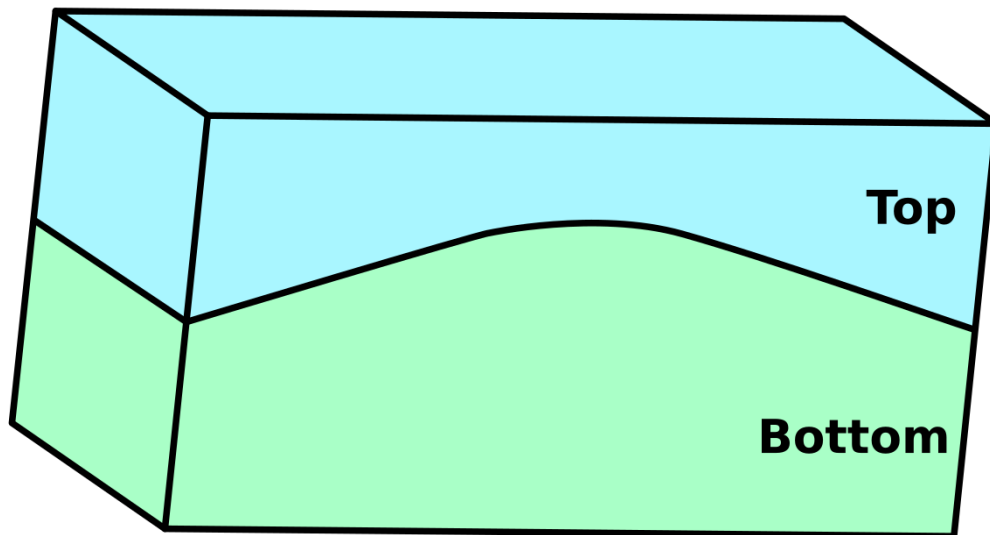


Fig. 6.2: Example of a model with two regions

future option if its use is properly motivated.

For instance, the model presented as example can be split into two different domains (Fig. 6.3).

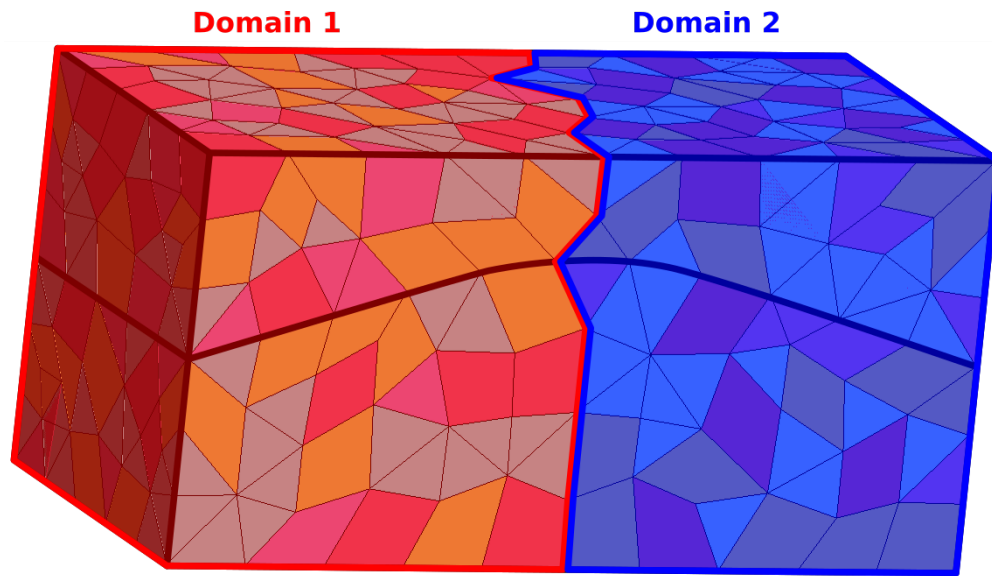


Fig. 6.3: Mesh partioned in two `DomainPartition`

### MeshBody

The `MeshBody` represents a topologically distinct mesh body. For instance if a simulation of two separate spheres was required, then one option would be to have both spheres as part of a single mesh body, while another option would be to have each sphere be a individual body.

---

**Note:** While not currently utilized in GEOSX, the intent is to have the ability to handle the bodies in a multi-body mesh on an individual basis. For instance, when conducting high resolution crush simulations of granular materials (i.e. sand), it may be advantageous to represent each particle as a `MeshBody`.

---

### MeshLevel

The `MeshLevel` is intended to facilitate the representation of a multi-level discretization of a `MeshBody`.

---

**Note:** In current practice, the code utilizes a single `MeshLevel` until such time as we implement a proper multi-level mesh capability. The `MeshLevel` contains the main components that compose a discretized mesh in GEOSX.

---

### Topological Mesh Objects

Each of the “Manager” objects are responsible for holding child objects, data, and providing functionality specific to a single topological object. Each topological object that is used to define a discretized mesh has a “Manager” to allow for simple traversal over the hierarchy, and to provide modular access to data. As such, the `NodeManager` manages data for the “nodes”, the `EdgeManager` manages data for the edges, the `FaceManager` holds data for the faces

and the `ElementRegionManager` manages the physical groups within the `MeshLevel` (regions, fractures, wells etc...). Additionally each manager contains index maps to the other types objects that are connected to the objects in that manager. For instance, the `FaceManager` contains a downward pointing map that gives the nodes that comprise each face in the mesh. Similarly the `FaceManager` contains an upward pointing map that gives the elements that are connected to a face.

## ElementRegionManager

The element data structure is significantly more complicated than the other Managers. While the other managers are “flat” across the `MeshLevel`, the element data structure seeks to provide a hierarchy in order to define groupings of the physical problem, as well as collecting discretization of similar topology. At the top of the element branch of the hierarchy is the `ElementRegionManager`. The `ElementRegionManager` holds a collection of instantiations of `ElementRegionBase` derived classes.

## ElementRegion

Conceptually the `ElementRegion` are used to defined regions of the problem domain where a `PhysicsSolver` will be applied.

- The `CellElementRegion` is related to all the polyhedra
- The `FaceElementRegion` is related to all the faces that have physical meaning in the domain, such as fractures and faults. This object should not be mistaken with the `FaceManager`. The `FaceManager` handles all the faces of the mesh, not only the faces of interest.
- The `WellElementRegion` is related to the well geometry.

An `ElementRegion` also has a list of materials allocated at each quadrature point across the entire region. One example of the utility of the `ElementRegion` is the case of the simulation of the mechanics and flow within sub-surface reservoir with an overburden. We could choose to have two `ElementRegion`, one being the reservoir, and one for the overburden. The mechanics solver would be applied to the entire problem, while the flow problem would be applied only to the reservoir region.

Each `ElementRegion` holds some number of `ElementSubRegion`. The `ElementSubRegion` is meant to hold all the element topologies present in an `ElementSubRegion` in their own groups. For instance, for a `CellElementRegion`, there may be one `CellElementSubRegion` for all tetrahedra, one for all hexahedra, one for all wedges and one for all the pyramids (Fig. 6.4).

Now that all the classes of the mesh hierarchy has been described, we propose to adapt the diagram presented in Fig. 6.1 to match with the example presented in Fig. 6.2.

## 6.2.5 DoF Manager

This will contains a description of the DoF manager in GEOSX.

### Brief description

The main aim of the Degrees-of-Freedom (DoF) Manager class is to handle all degrees of freedom associated with fields that exist on mesh elements, faces, edges and nodes. It creates a map between local mesh objects and global DoF indices. Additionally, `DofManager` simplifies construction of system matrix sparsity patterns.

Key concepts are locations and connectors. Locations, that can be elements, faces, edges or nodes, represent where the DoF is assigned. For example, a DoF for pressure in a two-point flux approximation will be on a cell (i.e. element), while a displacement DoF for structural equations will be on a node. The counterparts of locations are connectors, that



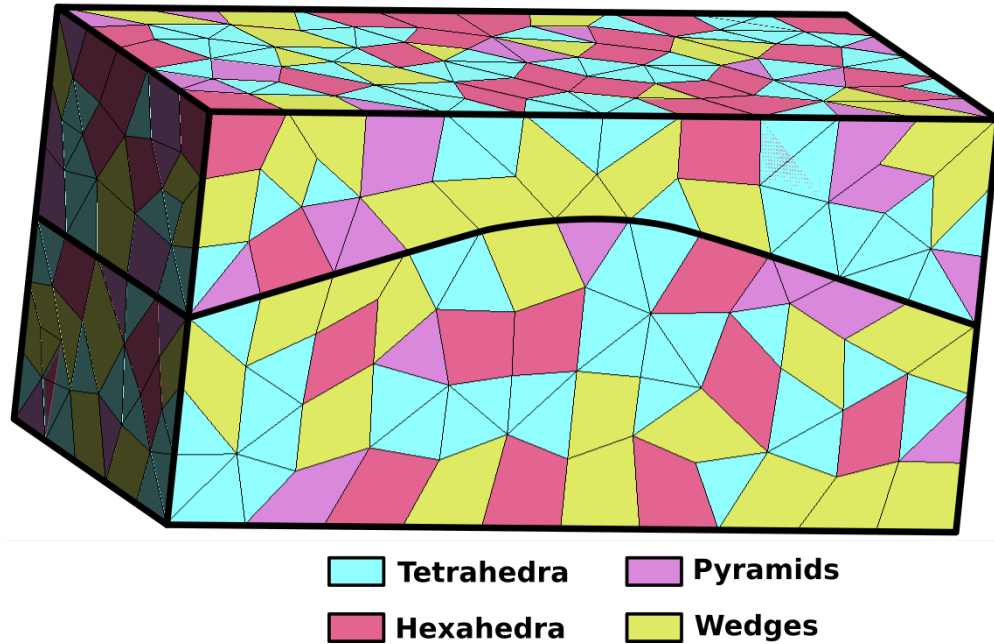


Fig. 6.4: Model meshed with different cell types

are the geometrical entities that link together different DoFs are create the sparsity pattern. Connectors can be elements, faces, edges, nodes or none. Using the same example as before, connectors will be faces and cells, respectively. The case of a mass matrix, where every element is linked only to itself, is an example when there are no connectors, i.e. these have to be set to none.

DoFs located on a mesh object are owned by the same rank that owns the object in parallel mesh partitioning. Two types of DoF numbering are supported, with the difference only showing in parallel runs of multi-field problems.

- Initially, each field is assigned an independent DoF numbering that starts at 0 and is contiguous across all MPI ranks. Within each rank, locally owned DoFs are numbered sequentially across mesh locations, and within each mesh location (e.g. node) - sequentially according to component number. With this numbering, sparsity patterns can be constructed for individual sub-matrices that represent diagonal/off-diagonal blocks of the global coupled system matrix.
- After all fields have been declared, the user can call `DofManager::reorderByRank()`, which constructs a globally contiguous DoF numbering across all fields. Specifically, all DoFs owned by rank 0 are numbered field-by-field starting from 0, then those on rank 1, etc. This makes global system sparsity pattern compatible with linear algebra packages that only support contiguous matrix rows on each rank. At this point, coupled system matrix sparsity pattern can be constructed.

Thus, each instance of `DofManager` only supports one type of numbering. If both types are required, the user is advised to maintain two separate instances of `DofManager`.

`DofManager` allocates a separate “DOF index” array for each field on the mesh. It is an array of global indices, where each value represents the first DoF index for that field and location (or equivalently, the row and column offset of that location’s equations and variables for the field in the matrix). For example, if index array for a field with 3 components contains the value  $N$ , global DoF numbers for that location will be  $N$ ,  $N+1$ ,  $N+2$ . DoF on ghosted locations have the same indices as on the owning rank. The array is stored under a generated key, which can be queried from the DoF manager, and is typically used in system assembly.

## Methods

The main methods of DoF Manager are:

- `setDomain`: sets the domain containing mesh bodies to operate on domain identifies the global domain

```
void setDomain( DomainPartition * const domain );
```

- `addField`: creates a new set of DoF, labeled field, with specific location. Default number of components is 1, like for pressure in flux. Default regions is the empty string, meaning all domain.

```
void addField( string const & fieldName,  
              Location const location,  
              localIndex const components,  
              arrayView1d< string const > const & regions );
```

- `addCoupling`: creates a coupling between two fields (`rowField` and `colField`) according to a given connectivity in the regions defined by `regions`. Both fields (row and column) must have already been defined on the regions where is required the coupling among them. Default value for `regions` is the whole intersection between the regions where the first and the second fields are defined. This method also creates the coupling between `colField` and `rowField`, i.e. the transpose of the rectangular sparsity pattern. This default behaviour can be disabled by passing `symmetric = false`.

```
void addCoupling( string const & rowField,  
                 string const & colField,  
                 Connectivity const connectivity,  
                 arrayView1d< string const > const & regions,  
                 bool const symmetric );
```

- `reorderByRank`: finish populating field and coupling information and apply DoF re-numbering

```
void reorderByRank();
```

- `getKey`: returns the “key” associated with the field, that can be used to access the index array on the mesh object manager corresponding to field’s location.

```
string const & getKey( string const & fieldName );
```

- `clear`: removes all fields, releases memory and re-opens the `DofManager`

```
void clear();
```

- `setSparsityPattern`: populates the sparsity for the given `rowField` and `colField` into matrix. Closes the matrix if `closePattern` is true.

```
void setSparsityPattern( MATRIX & matrix,  
                        string const & rowField,  
                        string const & colField,  
                        bool closePattern = true) const;
```

- `setSparsityPattern`: populates the sparsity for the full system matrix into matrix. Closes the matrix if `closePattern` is true.

```
void setSparsityPattern( MATRIX & matrix,  
                        bool closePattern = true ) const;
```

- `numGlobalDofs`: returns the total number of DoFs across all processors for the specified name field (if given) or all fields (if empty).

```
globalIndex numGlobalDofs( string const & field = "" ) const;
```

- `numLocalDofs`: returns the number of DoFs on this process for the specified name `field` (if given) or all fields (if empty).

```
localIndex numLocalDofs( string const & field = "" ) const;
```

- `printFieldInfo`: prints a short summary of declared fields and coupling to the output stream `os`.

```
void printFieldInfo( std::ostream & os = std::cout ) const;
```

## Example

Here we show how the sparsity pattern is computed for a simple 2D quadrilateral mesh with 6 elements. Unknowns are pressure, located on the element center, and displacements ( $x$  and  $y$  components), located on the nodes. For fluxes, a two-point flux approximation (TPFA) is used. The representation of the sparsity pattern of the  $C_L$  matrix (connectors/locations) for the simple mesh, shown in Fig. 6.5, is reported in Fig. 6.6. It can be noticed that the two unknowns for the displacements  $x$  and  $y$  are grouped together. Elements are the connectivity for DoF on nodes (Finite Element Method for displacements) and on elements (pressures). Faces are the connectivity for DoF on elements (Finite Volume Method for pressure), being the flux computation based on the pressure on the two adjacent elements.

Fig. 6.5: Small 2D quadrilateral mesh used for this examples. Nodes are labeled with black numbers, elements with light gray numbers and faces with italic dark gray numbers.

Fig. 6.6: Sparsity pattern of the binary matrix connections/locations.

The global sparsity pattern, shown in Fig. 6.7, is obtained through the symbolic multiplication of the transpose of the matrix  $C_L$  and the matrix itself, i.e.  $P = C_L^T C_L$ .

Fig. 6.7: Sparsity pattern of the global matrix, where red and green entries are related to the displacement field and to the pressure field, respectively. Blue entries represent coupling blocks.

## Real mesh and patterns

Now we build the pattern of the Jacobian matrix for a simple 3D mesh, shown in Fig. 6.8. Fields are:

- displacement (location: node, connectivity: element) defined on the blue, orange and red regions;
- pressure (location: element, connectivity: face) defined on the green, orange and red regions;
- mass matrix (location: element, connectivity: element) defined on the green region only.

Moreover, following coupling are imposed:

- displacement-pressure (connectivity: element) on the orange region only;
- pressure-mass matrix and transpose (connectivity: element) everywhere it is possible.

Fig. 6.9 shows the global pattern with the field-based ordering of unknowns. Different colors mean different fields. Red unknowns are associated with displacement, yellow ones with pressure and blue ones with mass matrix. Orange means the coupling among displacement and pressure, while green is the symmetric coupling among pressure and mass matrix.

Fig. 6.8: Real mesh used to compute the Jacobian pattern.

Fig. 6.9: Global pattern with field-based ordering. Red is associated with displacement unknowns, yellow with pressure ones and blue with those of mass matrix field. Orange means the coupling among displacement and pressure, while green is the symmetric coupling among pressure and mass matrix.

Fig. 6.10 shows the global pattern with the MPI rank-based ordering of unknowns. In this case, just two processes are used. Again, different colors indicate different ranks.

Fig. 6.10: Global pattern with MPI rank-based ordering. Red unknowns are owned by rank 0 and green ones by rank 1. Blue indicates the coupling among the two processes.

## 6.2.6 LvArray

### Use in GEOSX

LvArray containers are used in GEOSX as primary storage mechanism for mesh topology, field data and any other type of “large” data sets (i.e. ones that scale with the size of the problem). When allocating a new field, using one of LvArray containers is mandatory if the data is meant to be used in any computational kernel. The file `common/DataTypes.hpp` provides shorthand aliases for commonly used containers:

```
/**
 * @name Aliases for LvArray::Array class family.
 */
///<@{

/// Multidimensional array type. See LvArray:Array for details.
template< typename T,
          int NDIM,
          typename PERMUTATION=camp::make_idx_seq_t< NDIM > >
using Array = LvArray::Array< T, NDIM, PERMUTATION, localIndex, LvArray::ChaiBuffer >;

/// Multidimensional array view type. See LvArray:ArrayView for details.
template< typename T,
          int NDIM,
          int USD = NDIM - 1 >
using ArrayView = LvArray::ArrayView< T, NDIM, USD, localIndex, LvArray::ChaiBuffer >;

/// Multidimensional array slice type. See LvArray:ArraySlice for details.
template< typename T, int NDIM, int USD = NDIM - 1 >
using ArraySlice = LvArray::ArraySlice< T, NDIM, USD, localIndex >;

/// Multidimensional stack-based array type. See LvArray:StackArray for details.
template< typename T, int NDIM, int MAXSIZE, typename PERMUTATION=camp::make_idx_seq_t
  ↳< NDIM > >
using StackArray = LvArray::StackArray< T, NDIM, PERMUTATION, localIndex, MAXSIZE >;

///<@}

/**
 * @name Short-hand aliases for commonly used array types.
```

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```

*/
///<@{

/// Alias for a local (stack-based) rank-1 tensor type
using R1Tensor = Tensor< real64, 3 >;
/// Alias for a local (stack-based) rank-1 tensor type using 32 bits integers
using R1Tensor32 = Tensor< real32, 3 >;

/// Alias for a local (stack-based) rank-2 Voigt tensor type
using R2SymTensor = Tensor< real64, 6 >;

/// Alias for 1D array.
template< typename T >
using array1d = Array< T, 1 >;

/// Alias for 1D array view.
template< typename T >
using arrayView1d = ArrayView< T, 1 >;

/// Alias for 1D array slice.
template< typename T, int USD = 0 >
using arraySlice1d = ArraySlice< T, 1, USD >;

/// Alias for 1D stack array.
template< typename T, int MAXSIZE >
using stackArray1d = StackArray< T, 1, MAXSIZE >;

/// Alias for 2D array.
template< typename T, typename PERMUTATION=camp::make_idx_seq_t< 2 > >
using array2d = Array< T, 2, PERMUTATION >;

/// Alias for 2D array view.
template< typename T, int USD = 1 >
using arrayView2d = ArrayView< T, 2, USD >;

/// Alias for 2D array slice.
template< typename T, int USD = 1 >
using arraySlice2d = ArraySlice< T, 2, USD >;

/// Alias for 2D stack array.
template< typename T, int MAXSIZE >
using stackArray2d = StackArray< T, 2, MAXSIZE >;

/// Alias for 3D array.
template< typename T, typename PERMUTATION=camp::make_idx_seq_t< 3 > >
using array3d = Array< T, 3, PERMUTATION >;

/// Alias for 3D array view.
template< typename T, int USD=2 >
using arrayView3d = ArrayView< T, 3, USD >;

/// Alias for 3D array slice.
template< typename T, int USD=2 >
using arraySlice3d = ArraySlice< T, 3, USD >;

/// Alias for 3D stack array.

```

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```

template< typename T, int MAXSIZE >
using stackArray3d = StackArray< T, 3, MAXSIZE >;

/// Alias for 4D array.
template< typename T, typename PERMUTATION=camp::make_idx_seq_t< 4 > >
using array4d = Array< T, 4, PERMUTATION >;

/// Alias for 4D array view.
template< typename T, int USD=3 >
using arrayView4d = ArrayView< T, 4, USD >;

/// Alias for 4D array slice.
template< typename T, int USD=3 >
using arraySlice4d = ArraySlice< T, 4, USD >;

/// Alias for 4D stack array.
template< typename T, int MAXSIZE >
using stackArray4d = StackArray< T, 4, MAXSIZE >;

/// Alias for 5D array.
template< typename T, typename PERMUTATION=camp::make_idx_seq_t< 5 > >
using array5d = Array< T, 5, PERMUTATION >;

/// Alias for 5D array view.
template< typename T, int USD=4 >
using arrayView5d = ArrayView< T, 5, USD >;

/// Alias for 5D array slice.
template< typename T, int USD=4 >
using arraySlice5d = ArraySlice< T, 5, 4 >;

/// Alias for 5D stack array.
template< typename T, int MAXSIZE >
using stackArray5d = StackArray< T, 5, MAXSIZE >;

///  

/**
 * @name Aliases for sorted arrays and set types.
 */
///  

///  

/// A set of local indices.
template< typename T >
using set = std::set< T >;

/// A sorted array of local indices.
template< typename T >
using SortedArray = LvArray::SortedArray< T, localIndex, LvArray::ChaiBuffer >;

/// A sorted array view of local indices.
template< typename T >
using SortedArrayView = LvArray::SortedArrayView< T, localIndex, LvArray::ChaiBuffer >
↪;

///  


```

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```

/**
 * @name Aliases for LvArray::ArrayOfArrays class family.
 */
///<@{

/// Array of variable-sized arrays. See LvArray::ArrayOfArrays for details.
template< typename T, typename INDEX_TYPE=localIndex >
using ArrayOfArrays = LvArray::ArrayOfArrays< T, INDEX_TYPE, LvArray::ChaiBuffer >;

/// View of array of variable-sized arrays. See LvArray::ArrayOfArraysView for
↳ details.
template< typename T, typename INDEX_TYPE=localIndex, bool CONST_SIZES=std::is_const<
↳ T >::value >
using ArrayOfArraysView = LvArray::ArrayOfArraysView< T, INDEX_TYPE const, CONST_
↳ SIZES, LvArray::ChaiBuffer >;

/// Array of variable-sized sets. See LvArray::ArrayOfSets for details.
template< typename T, typename INDEX_TYPE=localIndex >
using ArrayOfSets = LvArray::ArrayOfSets< T, INDEX_TYPE, LvArray::ChaiBuffer >;

/// View of array of variable-sized sets. See LvArray::ArrayOfSetsView for details.
template< typename T, typename INDEX_TYPE=localIndex >
using ArrayOfSetsView = LvArray::ArrayOfSetsView< T, INDEX_TYPE const,
↳ LvArray::ChaiBuffer >;

/// Alias for Sparsity pattern class.
template< typename COL_INDEX, typename INDEX_TYPE=localIndex >
using SparsityPattern = LvArray::SparsityPattern< COL_INDEX, INDEX_TYPE,
↳ LvArray::ChaiBuffer >;

/// Alias for Sparsity pattern View.
template< typename COL_INDEX, typename INDEX_TYPE=localIndex >
using SparsityPatternView = LvArray::SparsityPatternView< COL_INDEX, INDEX_TYPE const,
↳ LvArray::ChaiBuffer >;

/// Alias for CRS Matrix class.
template< typename T, typename COL_INDEX=globalIndex >
using CRSMatrix = LvArray::CRSMatrix< T, COL_INDEX, localIndex, LvArray::ChaiBuffer >;

/// Alias for CRS Matrix View.
template< typename T, typename COL_INDEX=globalIndex >
using CRSMatrixView = LvArray::CRSMatrixView< T, COL_INDEX, localIndex const,
↳ LvArray::ChaiBuffer >;

///<@}

```

## LvArray documentation

Please refer to the full `LvArray` documentation for details on each of the classes.

## 6.2.7 Kernel interface

## Finite Element Method Kernel Interface

The finite element method kernel interface (FEMKI) specifies an API for the launching of computational kernels for solving physics discretized using the finite element method. Using this approach, a set of generic element looping patterns and kernel launching functions may be implemented, and reused by various physics solvers that contain kernels conforming to the FEMKI.

There are several main components of the FEMKI:

1. A collection of element looping functions that provide various looping patterns, and call the `launch` function.
2. The kernel interface, which is specified by the `finiteElement::KernelBase` class. Each physics solver will define a class that contains its kernels functions, most likely deriving, or conforming to the API specified by the `KernelBase` class. Also part of this class will typically be a nested `StackVariables` class that defines a collection of stack variables for use in the various kernel interface functions.
3. A `launch` function, which launches the kernel, and calls the kernel interface functions conforming to the interface defined by `KernelBase`. This function is actually a member function of the `Kernel` class, so it may be overridden by a specific physics kernel, allowing complete customization of the interface, while maintaining the usage of the looping patterns.

## A Generic Element Looping Pattern

One example of a looping pattern is the `regionBasedKernelApplication` function.

The contents of the looping function are displayed here:

```
/**
 * @brief Performs a loop over specific regions (by type and name) and calls a kernel_
 * launch on the subregions
 * with compile time knowledge of sub-loop bounds such as number of nodes and_
 * quadrature points per element.
 * @tparam POLICY The RAJA launch policy to pass to the kernel launch.
 * @tparam CONSTITUTIVE_BASE The common base class for constitutive pass-thru/
 * dispatch which gives the kernel
 * launch compile time knowledge of the constitutive model. This is achieved_
 * through a call to the
 * ConstitutivePassThru function which should have a specialization for_
 * CONSTITUTIVE_BASE implemented in
 * order to perform the compile time dispatch.
 * @tparam SUBREGION_TYPE The type of subregion to loop over. TODO make this a_
 * parameter pack?
 * @tparam KERNEL_FACTORY The type of @p kernelFactory, typically an instantiation of_
 * @c KernelFactory, and
 * must adhere to that interface.
 * @param mesh The MeshLevel object.
 * @param targetRegions The names of the target regions(of type @p SUBREGION_TYPE) to_
 * apply the @p KERNEL_TEMPLATE.
 * @param finiteElementName The name of the finite element.
 * @param constitutiveStringName The key to the constitutive model name found on the_
 * Region.
 * @param kernelFactory The object used to construct the kernel.
 * @return The maximum contribution to the residual, which may be used to scale the_
 * residual.
 *
 * @details Loops over all regions Applies/Launches a kernel specified by the @p_
 * KERNEL_TEMPLATE through
```

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```

* #::geosx::finiteElement::KernelBase::kernelLaunch().
*/
template< typename POLICY,
          typename CONSTITUTIVE_BASE,
          typename SUBREGION_TYPE,
          typename KERNEL_FACTORY >
static
real64 regionBasedKernelApplication( MeshLevel & mesh,
                                     arrayViewId< string const > const &_
↳targetRegions,
                                     string const & finiteElementName,
                                     string const & constitutiveStringName,
                                     KERNEL_FACTORY & kernelFactory )
{
    GEOSX_MARK_FUNCTION;
    // save the maximum residual contribution for scaling residuals for convergence_
↳criteria.
    real64 maxResidualContribution = 0;

    NodeManager & nodeManager = mesh.getNodeManager();
    EdgeManager & edgeManager = mesh.getEdgeManager();
    FaceManager & faceManager = mesh.getFaceManager();
    ElementRegionManager & elementRegionManager = mesh.getElemManager();

    // Loop over all sub-regions in regions of type SUBREGION_TYPE, that are listed in_
↳the targetRegions array.
    elementRegionManager.forElementSubRegions< SUBREGION_TYPE >( targetRegions,
                                                                    [&
↳constitutiveStringName,
                                                                    &
↳maxResidualContribution,
                                                                    &nodeManager,
                                                                    &edgeManager,
                                                                    &faceManager,
                                                                    &kernelFactory,
                                                                    &finiteElementName]
                                                                    ( localIndex const_
↳targetRegionIndex, auto & elementSubRegion )
    {
        localIndex const numElems = elementSubRegion.size();

        // Get the constitutive model...and allocate a null constitutive model if_
↳required.

        constitutive::ConstitutiveBase * constitutiveRelation = nullptr;
        constitutive::NullModel * nullConstitutiveModel = nullptr;
        if( elementSubRegion.template hasWrapper< string >( constitutiveStringName ) )
        {
            string const & constitutiveName = elementSubRegion.template getReference<_
↳string >( constitutiveStringName );
            constitutiveRelation = &elementSubRegion.template getConstitutiveModel(_
↳constitutiveName );
        }
        else
        {
            nullConstitutiveModel = &elementSubRegion.template registerGroup<_
↳constitutive::NullModel >( "nullModelGroup" );

```

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```

        constitutiveRelation = nullConstitutiveModel;
    }

    // Call the constitutive dispatch which converts the type of constitutive model
    ↪ into a compile time constant.
    constitutive::ConstitutivePassThru< CONSTITUTIVE_BASE >::execute(
    ↪ *constitutiveRelation,
                                                                    [&
                                                                    &nodeManager,
                                                                    &edgeManager,
                                                                    &faceManager,
                                                                    ↪
                                                                    &kernelFactory,
                                                                    &
                                                                    &
                                                                    numElems]
                                                                    ( auto & ↪
    ↪ castedConstitutiveRelation )
    {
        FiniteElementBase &
        subRegionFE = elementSubRegion.template getReference< FiniteElementBase >(
    ↪ finiteElementName );

        finiteElement::dispatch3D( subRegionFE,
                                    [&maxResidualContribution,
                                    &nodeManager,
                                    &edgeManager,
                                    &faceManager,
                                    targetRegionIndex,
                                    &kernelFactory,
                                    &elementSubRegion,
                                    numElems,
                                    &castedConstitutiveRelation] ( auto const ↪
    ↪ finiteElement )
        {
            auto kernel = kernelFactory.createKernel( nodeManager,
                                                        edgeManager,
                                                        faceManager,
                                                        targetRegionIndex,
                                                        elementSubRegion,
                                                        finiteElement,
                                                        castedConstitutiveRelation );

            using KERNEL_TYPE = decltype( kernel );

            // Call the kernelLaunch function, and store the maximum contribution to the
    ↪ residual.
            maxResidualContribution =
                std::max( maxResidualContribution,
                          KERNEL_TYPE::template kernelLaunch< POLICY, KERNEL_TYPE >(
    ↪ numElems, kernel ) );
        } );
    } );

```

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```

// Remove the null constitutive model (not required, but cleaner)
if( nullConstitutiveModel )
{
    elementSubRegion.deregisterGroup( "nullModelGroup" );
}

} );

return maxResidualContribution;
}

```

This pattern may be used with any kernel class that either:

1. Conforms to the `KernelBase` interface by defining each of the kernel functions in `KernelBase`.
2. Defines its own `kernelLaunch` function that conforms the the signature of `KernelBase::kernelLaunch`. This option essentially allows for a custom kernel that does not conform to the interface defined by `KernelBase` and `KernelBase::kernelLaunch`.

### The `KernelBase::kernelLaunch` Interface

The `kernelLaunch` function is a member of the kernel class itself. As mentioned above, a physics implementation may use the existing `KernelBase` interface, or define its own. The `KernelBase::kernelLaunch` function defines a launching policy, and an internal looping pattern over the quadrature points, and calls the functions defined by the `KernelBase` as shown here:

```

template< typename POLICY,
          typename KERNEL_TYPE >
static
real64
kernelLaunch( localIndex const numElems,
              KERNEL_TYPE const & kernelComponent )
{
    GEOSX_MARK_FUNCTION;

    // Define a RAJA reduction variable to get the maximum residual contribution.
    RAJA::ReduceMax< ReducePolicy< POLICY >, real64 > maxResidual( 0 );

    forAll< POLICY >( numElems,
                    [=] GEOSX_HOST_DEVICE ( localIndex const k )
    {
        typename KERNEL_TYPE::StackVariables stack;

        kernelComponent.setup( k, stack );
        for( integer q=0; q<numQuadraturePointsPerElem; ++q )
        {
            kernelComponent.quadraturePointKernel( k, q, stack );
        }
        maxResidual.max( kernelComponent.complete( k, stack ) );
    } );
    return maxResidual.get();
}

```

Each of the `KernelBase` functions called in the `KernelBase::kernelLaunch` function are intended to provide a certain amount of modularity and flexibility for the physics implementations. The general purpose of each function

is described by the function name, but may be further described by the function documentation found [here](#).

## 6.2.8 Constitutive models

In GEOSX, all constitutive models defining fluid and rock properties are implemented in the namespace `constitutive` and derived from a common base class, `ConstitutiveBase`. All objects are owned and handled by the `ConstitutiveManager`.

### Standalone models

Standalone constitutive models implement constitutive laws such as:

- mechanical material models (linear elasticity, plasticity, etc.),
- PVT fluid behaviors,
- relative permeability relationships,
- porosity and permeability dependencies on state variables,
- contact laws.

### Storage, allocation, and update of properties

Each constitutive model owns, as member variables, `LvArray : Array` containers that hold the properties (or fields) and their derivatives with respect to the other fields needed to update each property. Each property is stored as an array with the first dimension representing the `elementIndex` and the second dimension storing the index of the integration point. These dimensions are determined by the number of elements of the subregion on which each constitutive model is registered, and by the chosen discretization method. Vector and tensor fields have an additional dimension to identify their components. Similarly, an additional dimension is necessary for multiphase fluid models with properties defined for each component in each phase. For example, a single-phase fluid model where density and viscosity are functions of the fluid pressure has the following members:

```
array2d< real64 > m_density;  
array2d< real64 > m_dDensity_dPressure;  
array2d< real64 > m_dDensity_dTemperature;  
  
array2d< real64 > m_initialDensity;  
array2d< real64 > m_density_n;  
  
array2d< real64 > m_viscosity;  
array2d< real64 > m_dViscosity_dPressure;  
array2d< real64 > m_dViscosity_dTemperature;  
  
array2d< real64 > m_internalEnergy;  
array2d< real64 > m_internalEnergy_n;  
array2d< real64 > m_dInternalEnergy_dPressure;  
array2d< real64 > m_dInternalEnergy_dTemperature;  
  
array2d< real64 > m_enthalpy;  
array2d< real64 > m_dEnthalpy_dPressure;  
array2d< real64 > m_dEnthalpy_dTemperature;
```

Resizing all fields of the constitutive models happens during the initialization phase by the `ConstitutiveManager` through a call to `ConstitutiveManager::hangConstitutiveRelation`, which sets the appropriate `subRegion` as the parent `Group` of each constitutive model object. This function also resizes

all fields based on the size of the subregion and the number of quadrature points on it, by calling `CONSTITUTIVE_MODEL::allocateConstitutiveData`. For the single phase fluid example used before, this call is:

```
void SingleFluidBase::allocateConstitutiveData( Group & parent,
                                              localIndex const_
↳numConstitutivePointsPerParentIndex )
{
    ConstitutiveBase::allocateConstitutiveData( parent,
↳numConstitutivePointsPerParentIndex );

    resize( parent.size() );

    m_density.resize( parent.size(), numConstitutivePointsPerParentIndex );
    m_dDensity_dPressure.resize( parent.size(), numConstitutivePointsPerParentIndex );
    m_dDensity_dTemperature.resize( parent.size(), numConstitutivePointsPerParentIndex_
↳);
    m_initialDensity.resize( parent.size(), numConstitutivePointsPerParentIndex );
    m_density_n.resize( parent.size(), numConstitutivePointsPerParentIndex );

    m_viscosity.resize( parent.size(), numConstitutivePointsPerParentIndex );
    m_dViscosity_dPressure.resize( parent.size(), numConstitutivePointsPerParentIndex );
    m_dViscosity_dTemperature.resize( parent.size(),
↳numConstitutivePointsPerParentIndex );

    m_internalEnergy.resize( parent.size(), numConstitutivePointsPerParentIndex );
    m_internalEnergy_n.resize( parent.size(), numConstitutivePointsPerParentIndex );
    m_dInternalEnergy_dPressure.resize( parent.size(),
↳numConstitutivePointsPerParentIndex );
    m_dInternalEnergy_dTemperature.resize( parent.size(),
↳numConstitutivePointsPerParentIndex );

    m_enthalpy.resize( parent.size(), numConstitutivePointsPerParentIndex );
    m_dEnthalpy_dPressure.resize( parent.size(), numConstitutivePointsPerParentIndex );
    m_dEnthalpy_dTemperature.resize( parent.size(), numConstitutivePointsPerParentIndex_
↳);
}
```

Any property or field stored on a constitutive model must be updated within a computational kernel to ensure that *host* and *device* memory in GPUs are properly synced, and that any updates are performed on *device*. Some properties are updated within finite element kernels of specific physics (such as stress in a mechanics kernel). Consequently, for each constitutive model class, a corresponding *nameOfTheModelUpdates*, which only contains `LvArray::arrayView` containers to the data, can be captured by value inside computational kernels. For example, for the single phase fluid model *Updates* are:

```
/**
 * @brief Base class for single-phase fluid model kernel wrappers.
 */
class SingleFluidBaseUpdate
{
public:

    /**
     * @brief Get number of elements in this wrapper.
     * @return number of elements
     */
    GEOSX_HOST_DEVICE
```

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```

localIndex numElems() const { return m_density.size( 0 ); }

/**
 * @brief Get number of gauss points per element.
 * @return number of gauss points per element
 */
GEOSX_HOST_DEVICE
localIndex numGauss() const { return m_density.size( 1 ); };

protected:

/**
 * @brief Constructor.
 * @param density      fluid density
 * @param dDens_dPres  derivative of density w.r.t. pressure
 * @param viscosity    fluid viscosity
 * @param dVisc_dPres  derivative of viscosity w.r.t. pressure
 */
SingleFluidBaseUpdate( arrayView2d< real64 > const & density,
                      arrayView2d< real64 > const & dDens_dPres,
                      arrayView2d< real64 > const & viscosity,
                      arrayView2d< real64 > const & dVisc_dPres )
    : m_density( density ),
      m_dDens_dPres( dDens_dPres ),
      m_viscosity( viscosity ),
      m_dVisc_dPres( dVisc_dPres )
{}

/**
 * @brief Copy constructor.
 */
SingleFluidBaseUpdate( SingleFluidBaseUpdate const & ) = default;

/**
 * @brief Move constructor.
 */
SingleFluidBaseUpdate( SingleFluidBaseUpdate && ) = default;

/**
 * @brief Deleted copy assignment operator
 * @return reference to this object
 */
SingleFluidBaseUpdate & operator=( SingleFluidBaseUpdate const & ) = delete;

/**
 * @brief Deleted move assignment operator
 * @return reference to this object
 */
SingleFluidBaseUpdate & operator=( SingleFluidBaseUpdate && ) = delete;

/// Fluid density
arrayView2d< real64 > m_density;

/// Derivative of density w.r.t. pressure
arrayView2d< real64 > m_dDens_dPres;

```

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```

/// Fluid viscosity
arrayView2d< real64 > m_viscosity;

/// Derivative of viscosity w.r.t. pressure
arrayView2d< real64 > m_dVisc_dPres;

```

Because *Updates* classes are responsible for updating the fields owned by the constitutive models, they also implement all functions needed to perform property updates, such as:

```

private:

/**
 * @brief Compute fluid properties at a single point.
 * @param[in]  pressure the target pressure value
 * @param[out] density fluid density
 * @param[out] viscosity fluid viscosity
 */
GEOSX_HOST_DEVICE
virtual void compute( real64 const pressure,
                     real64 & density,
                     real64 & viscosity ) const = 0;

/**
 * @brief Compute fluid properties and derivatives at a single point.
 * @param[in]  pressure the target pressure value
 * @param[out] density fluid density
 * @param[out] dDensity_dPressure fluid density derivative w.r.t. pressure
 * @param[out] viscosity fluid viscosity
 * @param[out] dViscosity_dPressure fluid viscosity derivative w.r.t. pressure
 */
GEOSX_HOST_DEVICE
virtual void compute( real64 const pressure,
                     real64 & density,
                     real64 & dDensity_dPressure,
                     real64 & viscosity,
                     real64 & dViscosity_dPressure ) const = 0;

/**
 * @brief Compute fluid properties and derivatives at a single point.
 * @param[in]  pressure the target pressure value
 * @param[in]  temperature the target temperature value
 * @param[out] density fluid density
 * @param[out] dDensity_dPressure fluid density derivative w.r.t. pressure
 * @param[out] dDensity_dTemperature fluid density derivative w.r.t. temperature
 * @param[out] viscosity fluid viscosity
 * @param[out] dViscosity_dPressure fluid viscosity derivative w.r.t. pressure
 * @param[out] dViscosity_dTemperature fluid viscosity derivative w.r.t. temperature
 * @param[out] internalEnergy fluid internal energy
 * @param[out] dInternalEnergy_dPressure fluid internal energy derivative w.r.t.
↪pressure
 * @param[out] dInternalEnergy_dTemperature fluid internal energy derivative w.r.t.
↪temperature
 * @param[out] enthalpy fluid enthalpy
 * @param[out] dEnthalpy_dPressure fluid enthalpy derivative w.r.t. pressure
 * @param[out] dEnthalpy_dTemperature fluid enthalpy derivative w.r.t. temperature
 */

```

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```

GEOSX_HOST_DEVICE
virtual void compute( real64 const pressure,
                     real64 const temperature,
                     real64 & density,
                     real64 & dDensity_dPressure,
                     real64 & dDensity_dTemperature,
                     real64 & viscosity,
                     real64 & dViscosity_dPressure,
                     real64 & dViscosity_dTemperature,
                     real64 & internalEnergy,
                     real64 & dInternalEnergy_dPressure,
                     real64 & dInternalEnergy_dTemperature,
                     real64 & enthalpy,
                     real64 & dEnthalpy_dPressure,
                     real64 & dEnthalpy_dTemperature ) const = 0;

/**
 * @brief Update fluid state at a single point.
 * @param[in] k      element index
 * @param[in] q      gauss point index
 * @param[in] pressure the target pressure value
 */
GEOSX_HOST_DEVICE
virtual void update( localIndex const k,
                    localIndex const q,
                    real64 const pressure ) const = 0;

/**
 * @brief Update fluid state at a single point.
 * @param[in] k      element index
 * @param[in] q      gauss point index
 * @param[in] pressure the target pressure value
 * @param[in] temperature the target temperature value
 */
GEOSX_HOST_DEVICE
virtual void update( localIndex const k,
                    localIndex const q,
                    real64 const pressure,
                    real64 const temperature ) const = 0;

};

```

## Compound models

Compound constitutive models are employed to mimic the behavior of a material that requires a combination of constitutive models linked together. These compound models do not hold any data. They serve only as an interface with the individual models that they couple.

## Coupled Solids

CoupledSolid models are employed to represent porous materials that require both a mechanical behavior and constitutive laws that describe the dependency of porosity and permeability on the primary unknowns.

The base class CoupledSolidBase implements some basic behaviors and is used to access a generic



CoupledSolid in a physics solver:

```
CoupledSolidBase const & solid =
    getConstitutiveModel< CoupledSolidBase >( subRegion, subRegion.template_
    ↪getReference< string >( viewKeyStruct::solidNamesString() ) );
```

Additionally, a *template class* defines a base CoupledSolid model templated on the types of solid, porosity, and permeability models:

```
template< typename SOLID_TYPE,
          typename PORO_TYPE,
          typename PERM_TYPE >
class CoupledSolid : public CoupledSolidBase
```

While physics solvers that need a porous material only interface with a compound model, this one has access to the standalone models needed:

```
protected:
    SOLID_TYPE const & getSolidModel() const
    { return this->getParent().template getGroup< SOLID_TYPE >( m_solidModelName ); }

    PORO_TYPE const & getPorosityModel() const
    { return this->getParent().template getGroup< PORO_TYPE >( m_porosityModelName ); }

    PERM_TYPE const & getPermModel() const
    { return this->getParent().template getGroup< PERM_TYPE >( m_permeabilityModelName_
    ↪); }
```

There are two specializations of a CoupledSolid:

- **CompressibleSolid:** this model is used whenever there is no need to define a full mechanical model, but only simple correlations that compute material properties (like porosity or permeability). This model assumes that the solid model is of type *NullModel* and is only templated on the types of porosity and permeability models.
- **PorousSolid:** this model is used to represent a full porous material where the porosity and permeability models need to be aware of the mechanical response of the material.

## 6.2.9 PVT Package Hierarchy

The architecture of the PVT package is as follows

The color scheme is:

- Green is for computational flash classes
- Purple is for data classes
- Orange is for fluid models (black oil, free water...)
- Light blue are for computational system (algorithms and data combined)

Only classes exposed in the public interface of the PVT package is meant to be used outside the PVT package.

- `PHASE_TYPE`, `EOS_TYPE` and `COMPOSITIONAL_FLASH_TYPE` enums are meant to be used to select the models one wants to use.
- `ScalarPropertyAndDerivatives` and `VectorPropertyAndDerivatives` are utility classes used to return the results. Those classes should eventually be replaced by `LvArray`.
- `MultiphaseSystemProperties` agglomerates the result of the computation.



- `MultiphaseSystem` is responsible for performing the computation and serving the results.
- `MultiphaseSystemBuilder` builds the system.

### 6.2.10 Adding a new Physics Solver

In this tutorial, you will learn how to construct a new GEOSX Physics Solver class. We will use *LaplaceFEM* solver, computing the solution of the Laplace problem in a specified material, as a starting point.

$$\begin{aligned} D^* \Delta X &= f \quad \text{in } \Omega \\ X &= X^g \quad \text{on } \Gamma_g \end{aligned}$$

It is advised to read *XML Input* preliminary to this tutorial. The goal of this document is to explain how to develop a new solver that solves Laplace's equation with a constant diffusion coefficient that is specified by users in the XML input.

For readability, member functions in the text will be referenced by their names but their arguments will be omitted.

#### *LaplaceFEM* overview

The *LaplaceFEM* solver can be found in `./src/coreComponents/physicsSolvers/simplePDE/`. Let us inspect declarations in `LaplaceFEM.hpp` and implementations in `LaplaceFEM.cpp` before diving into specifying a new solver class that meets our needs.

#### Declaration file (reference)

The included header is `physicsSolvers/simplePDE/LaplaceBaseH1.hpp` which declares the base class `LaplaceBaseH1`, shared by all Laplace solvers. Moreover, `physicsSolver/simplePDE/LaplaceBaseH1.hpp` includes the following headers:

- `common/EnumStrings.hpp` which includes facilities for enum-string conversion (useful for reading enum values from input);
- `physicsSolver/SolverBase.hpp` which declares the abstraction class shared by all physics solvers.
- `managers/FieldSpecification/FieldSpecificationManager.hpp` which declares a manager used to access and to set field on the discretized domain.

Let us jump forward to the class enum and variable as they contain the data used specifically in the implementation of *LaplaceFEM*.

#### class enums and variables (reference)

The class exhibits two member variables:

- `m_fieldName` which stores the name of the diffused variable (e.g. the temperature) as a *string*;
- `m_timeIntegrationOption` an *enum* value allowing to dispatch with respect to the transient treatment.

`TimeIntegrationOption` is an *enum* specifying the transient treatment which can be chosen respectively between *SteadyState* and *ImplicitTransient* depending on whether we are interested in the transient state.

```
enum class TimeIntegrationOption : integer
{
    SteadyState,
    ImplicitTransient
};
```

In order to register an enumeration type with the Data Repository and have its value read from input, we must define stream insertion/extraction operators. This is a common task, so GEOSX provides a facility for automating it. Upon including `common/EnumStrings.hpp`, we can call the following macro at the namespace scope (in this case, right after the `LaplaceBaseH1` class definition is complete):

```
ENUM_STRINGS( LaplaceBaseH1::TimeIntegrationOption,
              "SteadyState",
              "ImplicitTransient" );
```

Once explained the main variables and enum, let us start reading through the different member functions:

```
class LaplaceFEM : public LaplaceBaseH1
{
public:
    /// The default nullary constructor is disabled to avoid compiler auto-generation:
    LaplaceFEM() = delete;

    /// The constructor needs a user-defined "name" and a parent Group (to place this_
    ↪instance in the
    /// tree structure of classes)
    LaplaceFEM( const string & name,
                Group * const parent );

    /// Destructor
    virtual ~LaplaceFEM() override;

    /// "CatalogName()" return the string used as XML tag in the input file. It ties_
    ↪the XML tag with
    /// this C++ classes. This is important.
    static string catalogName() { return "LaplaceFEM"; }
```

Start looking at the class *LaplaceFEM* constructor and destructor declarations shows the usual *string* name and *Group\** pointer to parent that are required to build the global file-system like structure of GEOSX (see *Group : the base class of GEOSX* for details). It can also be noted that the nullary constructor is deleted on purpose to avoid compiler automatic generation and user misuse.

The next method `catalogName()` is static and returns the key to be added to the *Catalog* for this type of solver (see *A few words about the ObjectCatalog* for details). It has to be paired with the following macro in the implementation file.

```
REGISTER_CATALOG_ENTRY( SolverBase, LaplaceFEM, string const &, Group * const )
```

Finally, the member function `registerDataOnMesh()` is declared in the `LaplaceBaseH1` class as

```
/// This method ties properties with their supporting mesh
virtual void registerDataOnMesh( Group & meshBodies ) override final;
```

It is used to assign fields onto the discretized mesh object and will be further discussed in the *Implementation File (reference)* section.

The next block consists in solver interface functions. These member functions set up and specialize every time step from the system matrix assembly to the solver stage.

```

virtual void
setupSystem( DomainPartition & domain,
             DofManager & dofManager,
             CRSMatrix< real64, globalIndex > & localMatrix,
             ParallelVector & rhs,
             ParallelVector & solution,
             bool const setSparsity = false ) override;

virtual void
assembleSystem( real64 const time,
               real64 const dt,
               DomainPartition & domain,
               DofManager const & dofManager,
               CRSMatrixView< real64, globalIndex const > const & localMatrix,
               arrayViewld< real64 > const & localRhs ) override;

```

Furthermore, the following functions are inherited from the base class.

```

virtual real64 solverStep( real64 const & time_n,
                        real64 const & dt,
                        integer const cycleNumber,
                        DomainPartition & domain ) override;

virtual void
implicitStepSetup( real64 const & time_n,
                 real64 const & dt,
                 DomainPartition & domain ) override;

virtual void
setupDofs( DomainPartition const & domain,
          DofManager & dofManager ) const override;

virtual void
applyBoundaryConditions( real64 const time,
                       real64 const dt,
                       DomainPartition & domain,
                       DofManager const & dofManager,
                       CRSMatrixView< real64, globalIndex const > const &_
↳localMatrix,
                       arrayViewld< real64 > const & localRhs ) override;

virtual void
applySystemSolution( DofManager const & dofManager,
                   arrayViewld< real64 const > const & localSolution,
                   real64 const scalingFactor,
                   DomainPartition & domain ) override;

virtual void updateState( DomainPartition & domain ) override final;

virtual void
  resetStateToBeginningOfStep( DomainPartition & GEOSX_UNUSED_PARAM( domain ) )_
↳override;

virtual void
implicitStepComplete( real64 const & time,
                    real64 const & dt,
                    DomainPartition & domain ) override;

```

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```

/// This method is specific to this Laplace solver.
/// It is used to apply Dirichlet boundary condition
/// and called when the base class applyBoundaryConditions() is called.
virtual void applyDirichletBCImplicit( real64 const time,
                                      DofManager const & dofManager,
                                      DomainPartition & domain,
                                      CRSMatrixView< real64, globalIndex const >
↳const & localMatrix,
                                      arrayView1d< real64 > const & localRhs );

```

Eventually, `applyDirichletBCImplicit()` is the working specialized member functions called when `applyBoundaryConditions()` is called in this particular class override.

Browsing the base class `SolverBase`, it can be noted that most of the solver interface functions are called during either `SolverBase::linearImplicitStep()` or `SolverBase::nonlinearImplicitStep()` depending on the solver strategy chosen.

Switching to protected members, `postProcessInput()` is a central member function and will be called by Group object after input is read from XML entry file. It will set and dispatch solver variables from the base class `SolverBase` to the most derived class. For `LaplaceFEM`, it will allow us to set the right time integration scheme based on the XML value as will be further explored in the next [Implementation File \(reference\)](#) section.

Let us focus on a struct that plays an important role: the `viewKeyStruct` structure.

### `viewKeyStruct` structure (reference)

This embedded instantiated structure is a common pattern shared by all solvers. It stores `dataRepository::ViewKey` type objects that are used as binding data between the input XML file and the source code.

```

struct viewKeyStruct : public SolverBase::viewKeyStruct
{
    static constexpr char const * timeIntegrationOption() { return
↳"timeIntegrationOption"; }
    static constexpr char const * fieldVarName() { return "fieldName"; }
};

```

We can check that in the `LaplaceFEM` companion `integratedTest`

```

<LaplaceFEM
  name="laplace"
  discretization="FE1"
  timeIntegrationOption="SteadyState"
  fieldName="Temperature"
  targetRegions="{ Domain }">
  <LinearSolverParameters
    directParallel="0"/>
</LaplaceFEM>

```

In the following section, we will see where this binding takes place.

## Implementation File (reference)

Switching to implementation, we will focus on few implementations, leaving details to other tutorials. The LaplaceFEM constructor is implemented as follows.

```
LaplaceFEM::LaplaceFEM( const string & name,
                        Group * const parent ):
    LaplaceBaseH1( name, parent )
{ }
```

As we see, it calls the LaplaceBaseH1 constructor, that is implemented as follows.

```
LaplaceBaseH1::LaplaceBaseH1( const string & name,
                               Group * const parent ):
    SolverBase( name, parent ),
    m_fieldName( "primaryField" ),
    m_timeIntegrationOption( TimeIntegrationOption::ImplicitTransient )
{
    this->registerWrapper( viewKeyStruct::timeIntegrationOption(), &m_
    ↪timeIntegrationOption ).
        setInputFlag( InputFlags::REQUIRED ).
        setDescription( "Time integration method. Options are:\n* " + EnumStrings<
    ↪TimeIntegrationOption >::concat( "\n* " ) );

    this->registerWrapper( viewKeyStruct::fieldVarName(), &m_fieldName ).
        setInputFlag( InputFlags::REQUIRED ).
        setDescription( "Name of field variable" );
}
```

Checking out the constructor, we can see that the use of a registerWrapper<T>(...) allows us to register the key value from the *enum* viewKeyStruct defining them as:

- InputFlags::OPTIONAL if they are optional and can be provided;
- InputFlags::REQUIRED if they are required and will throw error if not;

and their associated descriptions for auto-generated docs.

```
void LaplaceBaseH1::registerDataOnMesh( Group & meshBodies )
{
    meshBodies.forSubGroups< MeshBody >( [&] ( MeshBody & meshBody )
    {
        NodeManager & nodes = meshBody.getBaseDiscretization().getNodeManager();

        nodes.registerWrapper< real64_array >( m_fieldName ).
            setApplyDefaultValue( 0.0 ).
            setPlotLevel( PlotLevel::LEVEL_0 ).
            setDescription( "Primary field variable" );
    } );
}
```

registerDataOnMesh() is browsing all subgroups in the mesh Group object and for all nodes in the sub group:

- register the observed field under the chosen m\_fieldName key;
- apply a default value;
- set the output verbosity level (here PlotLevel::LEVEL\_0);
- set the field associated description for auto generated docs.

```

void LaplaceFEM::assembleSystem( real64 const GEOSX_UNUSED_PARAM( time_n ),
                                real64 const GEOSX_UNUSED_PARAM( dt ),
                                DomainPartition & domain,
                                DofManager const & dofManager,
                                CRSMatrixView< real64, globalIndex const > const &
    ↪ localMatrix,
                                arrayViewId< real64 > const & localRhs )
{
    forDiscretizationOnMeshTargets( domain.getMeshBodies(), [&] ( string const &,
                                                                MeshLevel & mesh,
                                                                arrayViewId< string &
    ↪ const > const & regionNames )
    {
        NodeManager & nodeManager = mesh.getNodeManager();
        string const dofKey = dofManager.getKey( m_fieldName );
        arrayViewId< globalIndex const > const &
        dofIndex = nodeManager.getReference< arrayId< globalIndex > >( dofKey );

        LaplaceFEMKernelFactory kernelFactory( dofIndex, dofManager.rankOffset(),
    ↪ localMatrix, localRhs, m_fieldName );

        string const dummyString = "dummy";
        finiteElement::
            regionBasedKernelApplication< parallelDevicePolicy< 32 >,
                                         constitutive::NullModel,
                                         CellElementSubRegion >( mesh,
                                                                regionNames,
                                                                this->
    ↪ getDiscretizationName(),
                                                                dummyString,
                                                                kernelFactory );

    } );
}

```

`assembleSystem()` will be our core focus as we want to change the diffusion coefficient from its hard coded value to a XML read user-defined value. One can see that this method is in charge of constructing in a parallel fashion the FEM system matrix. Bringing `nodeManager` and `ElementRegionManager` from domain local `MeshLevel` object together with `FiniteElementDiscretizationManager` from the `NumericalMethodManager`, it uses nodes embedded loops on degrees of freedom in a local index embedded loops to fill a matrix and a rhs container.

As we spotted the place to change in a code to get a user-defined diffusion coefficient into the game, let us jump to writing our new *LaplaceDiffFEM* solver.

---

**Note:** We might want to remove final keyword from `postProcessInput()` as it will prevent you from overriding it.

---

## Start doing your own Physic solver

As we will extend *LaplaceFEM* capabilities, we will derive publicly from it.



## Declaration File

As there is only few places where we have to change, the whole declaration file is reported below and commented afterwards.

```
#include "physicsSolvers/simplePDE/LaplaceFEM.hpp"

namespace geosx
{

class LaplaceDiffFEM : public LaplaceFEM
{
public:

    LaplaceDiffFEM() = delete;

    LaplaceDiffFEM( const string& name,
                    Group * const parent );

    virtual ~LaplaceDiffFEM() override;

    static string catalogName() { return "LaplaceDiffFEM"; }

    virtual void
    assembleSystem( real64 const time,
                    real64 const dt,
                    DomainPartition * const domain,
                    DofManager const & dofManager,
                    ParallelMatrix & matrix,
                    ParallelVector & rhs ) override;

    struct viewKeyStruct : public LaplaceFEM::viewKeyStruct
    {
        dataRepository::ViewKey diffusionCoeff = { "diffusionCoeff" };
    } laplaceDiffFEMViewKeys;

protected:
    virtual void postProcessInput() override final;

private:
    real64 m_diffusion;

};
```

We intend to have a user-defined diffusion coefficient, we then need a *real64* class variable `m_diffusion` to store it.

Consistently with *LaplaceFEM*, we will also delete the nullary constructor and declare a constructor with the same arguments for forwarding to *Group* master class. Another mandatory step is to override the static `CatalogName()` method to properly register any data from the new solver class.

Then as mentioned in *Implementation File (reference)*, the diffusion coefficient is used when assembling the matrix coefficient. Hence we will have to override the `assembleSystem()` function as detailed below.

Moreover, if we want to introduce a new binding between the input XML and the code we will have to work on the three `struct viewKeyStruct`, `postProcessInput()` and the constructor.

Our new solver `viewKeyStruct` will have its own structure inheriting from the *LaplaceFEM* one to have the

timeIntegrationOption and fieldName field. It will also create a diffusionCoeff field to be bound to the user defined homogeneous coefficient on one hand and to our m\_diffusion class variable on the other.

## Implementation File

As we have seen in *Implementation File (reference)*, the first place where to implement a new register from XML input is in the constructor. The diffusionCoeff entry we have defined in the laplaceDiffFEMViewKeys will then be asked as a required input. If not provided, the error thrown will ask for it described as an “input uniform diffusion coefficient for the Laplace equation”.

```
LaplaceDiffFEM::LaplaceDiffFEM( const string& name,
                                Group * const parent ):
LaplaceFEM( name, parent ), m_diffusion(0.0)
{
    registerWrapper<string>(laplaceDiffFEMViewKeys.diffusionCoeff.Key()).
        setInputFlag(InputFlags::REQUIRED).
        setDescription("input uniform diffusion coeff for the laplace equation");
}
```

Another important spot for binding the value of the XML read parameter to our m\_diffusion is in postProcessInput().

```
void LaplaceDiffFEM::postProcessInput()
{
    LaplaceFEM::postProcessInput();

    string sDiffCoeff = this->getReference<string>(laplaceDiffFEMViewKeys.
->diffusionCoeff);
    this->m_diffusion = std::stof(sDiffCoeff);
}
```

Now that we have required, read and bind the user-defined diffusion value to a variable, we can use it in the construction of our matrix into the overridden assembleSystem().

```
// begin element loop, skipping ghost elements
for( localIndex k=0 ; k<elementSubRegion->size() ; ++k )
{
    if(elemGhostRank[k] < 0)
    {
        element_rhs = 0.0;
        element_matrix = 0.0;
        for( localIndex q=0 ; q<n_q_points ; ++q)
        {
            for( localIndex a=0 ; a<numNodesPerElement ; ++a)
            {
                elemDofIndex[a] = dofIndex[ elemNodes( k, a ) ];

                for( localIndex b=0 ; b<numNodesPerElement ; ++b)
                {
                    element_matrix(a,b) += detJ[k][q] *
                                            m_diffusion *
                                            + Dot( dNdX[k][q][a], dNdX[k][q][b] );
                }
            }
        }
    }
}
```

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```

    matrix.add( elemDofIndex, elemDofIndex, element_matrix );
    rhs.add( elemDofIndex, element_rhs );
}
}

```

This completes the implementation of our new solver *LaplaceDiffFEM*.

Nonetheless, the compiler should complain that `m_fieldName` is privately as inherited from *LaplaceFEM*. One should then either promote `m_fieldName` to protected or add a getter in *LaplaceFEM* class to correct the error. The getter option has been chosen and the fix in our solver is then:

```

arrayId<globalIndex> const & dofIndex =
    nodeManager->getReference< arrayId<globalIndex> >( dofManager.getKey(
    ↪getFieldName() ) );

```

Note: For consistency do not forget to change LaplaceFEM to LaplaceDiffFEM in the guards comments

## Last steps

After assembling both declarations and implementations for our new solver, the final steps go as:

- add declarations to parent CMakeLists.txt (here add to `physicsSolvers_headers`);
- add implementations to parent CMakeLists.txt (here add to `physicsSolvers_sources`);
- check that Doxygen comments are properly set in our solver class;
- uncrustify it to match the code style;
- write unit tests for each new features in the solver class;
- write an integratedTests for the solver class.



## CHAPTER 7

---

### Doxygen

---

The c++ source in GEOSX is annotated using doxygen. Our doxygen pages are linked below.

Developers may find it helpful to review the key code components described in the Developer Guide before diving into the doxygen.



## 8.1 Python Tools Setup

The preferred method to setup the GEOSX python tools is to run the following command in the build directory:

```
make geosx_xml_tools
```

This will attempt to install the required packages into one of the following locations (in order of preference):

1. The python distribution indicated via the *PYTHON\_POST\_EXECUTABLE* cmake variable
2. The python distribution indicated via the *Python3\_EXECUTABLE* cmake variable (also used by pygeosx)
3. The python distribution that was used to configure GEOSX

If the user does not have write access for the target python distribution, the installation will attempt to create a new virtual python environment (Note: this requires that the virtualenv package be installed). If any package dependencies are missing, then the install script will attempt to fetch them from the internet using pip. After installation, these packages will be available for import within the associated python distribution, and a set of console scripts will be available within the GEOSX build bin directory.

Alternatively, these packages can be installed manually into a python environment using pip:

```
cd GEOSX/src/coreComponents/python/modules/geosx_mesh_tools_package
pip install --upgrade .

cd ../geosx_xml_tools_package
pip install --upgrade .

# Etc.
```

## 8.2 Packages

### 8.2.1 HDF5 Wrapper

The *hdf5\_wrapper* python package adds a wrapper to *h5py* that greatly simplifies reading/writing to/from hdf5-format files.

#### Usage

Once loaded, the contents of a file can be navigated in the same way as a native python dictionary.

```
import hdf5_wrapper

data = hdf5_wrapper.hdf5_wrapper('data.hdf5')

test = data['test']
for k, v in data.items():
    print('key: %s, value: %s' % (k, str(v)))
```

If the user indicates that a file should be opened in write-mode (*w*) or read/write-mode (*a*), then the file can be created or modified. Note: for these changes to be written to the disk, the wrapper may need to be closed or deleted.

```
import hdf5_wrapper
import numpy as np

data = hdf5_wrapper.hdf5_wrapper('data.hdf5', mode='w')
data['string'] = 'string'
data['integer'] = 123
data['array'] = np.random.randn(3, 4, 5)
data['child'] = {'float': 1.234}
```

Existing dictionaries can be placed on the current level:

```
existing_dict = {'some': 'value'}
data.insert(existing_dict)
```

And external hdf5 format files can be linked together:

```
for k in ['child_a', 'child_b']:
    data.link(k, '%s.hdf5' % (k))
```

#### API

**class** `hdf5_wrapper.wrapper.hdf5_wrapper` (*fname: str = "", target: Optional[h5py.\_hl.files.File] = None, mode: str = 'r'*)

A class for reading/writing hdf5 files, which behaves similar to a native dict

**close** () → None

Closes the database

**copy** () → Dict[str, Any]

Copy the entire database into memory

**Returns** a dictionary holding the database contents



**Return type** dict

**get\_copy** () → Dict[str, Any]

Copy the entire database into memory

**Returns** a dictionary holding the database contents

**Return type** dict

**insert** (*x*: Union[Dict[str, Any], hdf5\_wrapper]) → None

Insert the contents of the target object to the current location

**Parameters** **x** (*dict*, *hdf5\_wrapper*) – the dictionary to insert

**keys** () → Iterable[str]

Get a list of groups and arrays located at the current level

**Returns** a list of key names pointing to objects at the current level

**Return type** list

**link** (*k*: str, *target*: str) → None

Link an external hdf5 file to this location in the database

**Parameters**

- **k** (*str*) – the name of the new link in the database
- **target** (*str*) – the path to the external database

**values** () → Iterable[Union[hdf5\_wrapper.wrapper.hdf5\_wrapper, Any]]

Get a list of values located on the current level

## 8.2.2 GEOSX Mesh Tools

The *geosx\_mesh\_tools* python package includes tools for converting meshes from common formats (abaqus, etc.) to those that can be read by GEOSX (gmsh, vtk). See [Python Tools Setup](#) for details on setup instructions, and [Using an External Mesh](#) for a detailed description of how to use external meshes in GEOSX. The available console scripts for this package and its API are described below.

### convert\_abaqus

Compile an xml file with advanced features into a single file that can be read by GEOSX.

```
usage: convert_abaqus [-h] [-v] input output
```

### Positional Arguments

<b>input</b>	Input abaqus mesh file name
<b>output</b>	Output gmsh/vtu mesh file name

### Named Arguments

<b>-v, --verbose</b>	Increase verbosity level
	Default: False

**Note:** For vtk format meshes, the user also needs to determine the region ID numbers and names of nodesets to import into GEOSX. The following shows how these could look in an input XML file for a mesh with three regions (*REGIONA*, *REGIONB*, and *REGIONC*) and six nodesets (*xneg*, *xpos*, *yneg*, *ypos*, *zneg*, and *zpos*):

```
<Problem>
  <Mesh>
    <VTKMesh
      name="external_mesh"
      file="mesh.vtu"
      regionAttribute="REGIONA-REGIONB-REGIONC"
      nodesetNames="{ xneg, xpos, yneg, ypos, zneg, zpos }"/>
    </Mesh>

    <ElementRegions>
      <CellElementRegion
        name="ALL"
        cellBlocks="{ 0_tetrahedra, 1_tetrahedra, 2_tetrahedra }"
        materialList="{ water, porousRock }"
        meshBody="external_mesh"/>
      </ElementRegions>
    </Problem>
```

## API

`geosx_mesh_tools.abaqus_converter.convert_abaqus_to_gmsh` (*input\_mesh*: *str*, *output\_mesh*: *str*, *logger*: *logging.Logger* = *None*)  
→ int

Convert an abaqus mesh to gmsh 2 format, preserving nodeset information.

If the code encounters any issues with region/element indices, the conversion will attempt to continue, with errors indicated by -1 values in the output file.

### Parameters

- **input\_mesh** (*str*) – path of the input abaqus file
- **output\_mesh** (*str*) – path of the output gmsh file
- **logger** (*logging.Logger*) – an instance of *logging.Logger*

**Returns** Number of potential warnings encountered during conversion

**Return type** int

`geosx_mesh_tools.abaqus_converter.convert_abaqus_to_vtu` (*input\_mesh*: *str*, *output\_mesh*: *str*, *logger*: *logging.Logger* = *None*)  
→ int

Convert an abaqus mesh to vtu format, preserving nodeset information.

If the code encounters any issues with region/element indices, the conversion will attempt to continue, with errors indicated by -1 values in the output file.

### Parameters

- **input\_mesh** (*str*) – path of the input abaqus file
- **output\_mesh** (*str*) – path of the output vtu file

- **logger** (*logging.Logger*) – a logger instance

**Returns** Number of potential warnings encountered during conversion

**Return type** int

### 8.2.3 GEOSX XML Tools

The *geosx\_xml\_tools* python package adds a set of advanced features to the GEOSX xml format: units, parameters, and symbolic expressions. See [Python Tools Setup](#) for details on setup instructions, and [Advanced XML Features](#) for a detailed description of the input format. The available console scripts for this package and its API are described below.

#### convert\_abaqus

Convert an abaqus format mesh file to gmsh or vtk format.

```
usage: preprocess_xml [-h] [-i INPUT] [-c COMPILED_NAME] [-s SCHEMA]
                    [-v VERBOSE] [-p PARAMETERS [PARAMETERS ...]]
```

#### Named Arguments

<b>-i, --input</b>	Input file name (multiple allowed)
<b>-c, --compiled-name</b>	Compiled xml file name (otherwise, it is randomly genrated) Default: ""
<b>-s, --schema</b>	GEOSX schema to use for validation Default: ""
<b>-v, --verbose</b>	Verbosity of outputs Default: 0
<b>-p, --parameters</b>	Parameter overrides (name value, multiple allowed) Default: []

#### format\_xml

Formats an xml file.

```
usage: format_xml [-h] [-i INDENT] [-s STYLE] [-d DEPTH] [-a ALPHEBITIZE]
                 [-c CLOSE] [-n NAMESPACE]
                 input
```

#### Positional Arguments

<b>input</b>	Input file name
--------------	-----------------

## Named Arguments

<b>-i, --indent</b>	Indent size Default: 2
<b>-s, --style</b>	Indent style Default: 0
<b>-d, --depth</b>	Block separation depth Default: 2
<b>-a, --alphabetize</b>	Alphabetize attributes Default: 0
<b>-c, --close</b>	Close tag style Default: 0
<b>-n, --namespace</b>	Include namespace Default: 0

## check\_xml\_attribute\_coverage

Checks xml attribute coverage for files in the GEOSX repository.

```
usage: check_xml_attribute_coverage [-h] [-r ROOT] [-o OUTPUT]
```

## Named Arguments

<b>-r, --root</b>	GEOSX root Default: ""
<b>-o, --output</b>	Output file name Default: "attribute_test.xml"

## check\_xml\_redundancy

Checks for redundant attribute definitions in an xml file, such as those that duplicate the default value.

```
usage: check_xml_redundancy [-h] [-r ROOT]
```

## Named Arguments

<b>-r, --root</b>	GEOSX root Default: ""
-------------------	---------------------------

## API

Command line tools for `geosx_xml_tools`

`geosx_xml_tools.main.check_mpi_rank()` → int

Check the MPI rank

**Returns** MPI rank

**Return type** int

`geosx_xml_tools.main.format_geosx_arguments(compiled_name: str, unknown_args: Iterable[str])` → Iterable[str]

Format GEOSX arguments

**Parameters**

- **compiled\_name** (*str*) – Name of the compiled xml file
- **unknown\_args** (*list*) – List of unprocessed arguments

**Returns** List of arguments to pass to GEOSX

**Return type** list

`geosx_xml_tools.main.preprocess_parallel()` → Iterable[str]

MPI aware xml preprocessing

`geosx_xml_tools.main.preprocess_serial()` → None

Entry point for the `geosx_xml_tools` console script

`geosx_xml_tools.main.wait_for_file_write_rank_0(target_file_argument: Union[int, str] = 0, max_wait_time: float = 100, max_startup_delay: float = 1)`  
→ Callable[[Callable[[...], Any]], Callable[[...], Any]]

Constructor for a function decorator that waits for a target file to be written on rank 0

**Parameters**

- **target\_file\_argument** (*int*, *str*) – Index or keyword of the filename argument in the decorated function
- **max\_wait\_time** (*float*) – Maximum amount of time to wait (seconds)
- **max\_startup\_delay** (*float*) – Maximum delay allowed for thread startup (seconds)

**Returns** Wrapped function

Tools for processing xml files in GEOSX

`geosx_xml_tools.xml_processor.apply_regex_to_node(node: <sphinx.ext.autodoc.importer._MockObject object at 0x7f95ebf66910>)` → None

Apply regexes that handle parameters, units, and symbolic math to each xml attribute in the structure.

**Parameters** **node** (*lxml.etree.Element*) – The target node in the xml structure.

`geosx_xml_tools.xml_processor.generate_random_name(prefix: str = "", suffix: str = '.xml')`  
→ str

If the target name is not specified, generate a random name for the compiled xml

**Parameters**

- **prefix** (*str*) – The file prefix (default = '').

- **suffix** (*str*) – The file suffix (default = ‘.xml’)

**Returns** Random file name

**Return type** *str*

`geosx_xml_tools.xml_processor.merge_included_xml_files` (*root*:  
    <*sphinx.ext.autodoc.importer.\_MockObject*  
    *object at 0x7f95ebf66910*>,  
    *fname*: *str*, *includeCount*:  
    *int*, *maxInclude*: *int* = 100)  
    → None

Recursively merge included files into the current structure.

#### Parameters

- **root** (*lxml.etree.Element*) – The root node of the base xml structure.
- **fname** (*str*) – The name of the target xml file to merge.
- **includeCount** (*int*) – The current recursion depth.
- **maxInclude** (*int*) – The maximum number of xml files to include (default = 100)

`geosx_xml_tools.xml_processor.merge_xml_nodes` (*existingNode*:  
    <*sphinx.ext.autodoc.importer.\_MockObject*  
    *object at 0x7f95ebf66910*>, *targetNode*:  
    <*sphinx.ext.autodoc.importer.\_MockObject*  
    *object at 0x7f95ebf66910*>, *level*: *int*) →  
    None

Merge nodes in an included file into the current structure level by level.

#### Parameters

- **existingNode** (*lxml.etree.Element*) – The current node in the base xml structure.
- **targetNode** (*lxml.etree.Element*) – The node to insert.
- **level** (*int*) – The xml file depth.

`geosx_xml_tools.xml_processor.process` (*inputFiles*: *Iterable[str]*, *outputFile*: *str* = "", *schema*:  
    *str* = "", *verbose*: *int* = 0, *parameter\_override*:  
    *List[Tuple[str, str]]* = [], *keep\_parameters*: *bool* =  
    *True*, *keep\_includes*: *bool* = *True*) → *str*

Process an xml file

#### Parameters

- **inputFiles** (*list*) – Input file names.
- **outputFile** (*str*) – Output file name (if not specified, then generate randomly).
- **schema** (*str*) – Schema file name to validate the final xml (if not specified, then do not validate).
- **verbose** (*int*) – Verbosity level.
- **parameter\_override** (*list*) – Parameter value overrides
- **keep\_parameters** (*bool*) – If True, then keep parameters in the compiled file (default = True)
- **keep\_includes** (*bool*) – If True, then keep includes in the compiled file (default = True)

**Returns** Output file name

**Return type** `str`

`geosx_xml_tools.xml_processor.validate_xml` (*fname: str, schema: str, verbose: int*) → `None`  
 Validate an xml file, and parse the warnings.

**Parameters**

- **fname** (*str*) – Target xml file name.
- **schema** (*str*) – Schema file name.
- **verbose** (*int*) – Verbosity level.

`geosx_xml_tools.xml_formatter.format_attribute` (*attribute\_indent: str, ka: str, attribute\_value: str*) → `str`

Format xml attribute strings

**Parameters**

- **attribute\_indent** (*str*) – Attribute indent string
- **ka** (*str*) – Attribute name
- **attribute\_value** (*str*) – Attribute value

**Returns** Formatted attribute value

**Return type** `str`

`geosx_xml_tools.xml_formatter.format_file` (*input\_fname: str, indent\_size: int = 2, indent\_style: bool = False, block\_separation\_max\_depth: int = 2, alphabitize\_attributes: bool = False, close\_style: bool = False, namespace: bool = False*) → `None`

Script to format xml files

**Parameters**

- **input\_fname** (*str*) – Input file name
- **indent\_size** (*int*) – Indent size
- **indent\_style** (*bool*) – Style of indentation (0=fixed, 1=hanging)
- **block\_separation\_max\_depth** (*int*) – Max depth to separate xml blocks
- **alphabitize\_attributes** (*bool*) – Alphabitize attributes
- **close\_style** (*bool*) – Style of close tag (0=same line, 1=new line)
- **namespace** (*bool*) – Insert this namespace in the xml description

`geosx_xml_tools.xml_formatter.format_xml_level` (*output: TextIO, node: <sphinx.ext.autodoc.importer.\_MockObject object at 0x7f95ea7b2690>, level: int, indent: str = ' ', block\_separation\_max\_depth: int = 2, modify\_attribute\_indent: bool = False, sort\_attributes: bool = False, close\_tag\_newline: bool = False, include\_namespace: bool = False*) → `None`

Iteratively format the xml file

**Parameters**

- **output** (*file*) – the output text file handle
- **node** (*lxml.etree.Element*) – the current xml element
- **level** (*int*) – the xml depth
- **indent** (*str*) – the xml indent style
- **block\_separation\_max\_depth** (*int*) – the maximum depth to separate adjacent elements
- **modify\_attribute\_indent** (*bool*) – option to have flexible attribute indentation
- **sort\_attributes** (*bool*) – option to sort attributes alphabetically
- **close\_tag\_newline** (*bool*) – option to place close tag on a separate line
- **include\_namespace** (*bool*) – option to include the xml namespace in the output

`geosx_xml_tools.xml_formatter.main()` → None

Script to format xml files

#### Parameters

- **input** (*str*) – Input file name
- **-i/--indent** (*int*) – Indent size
- **-s/--style** (*int*) – Indent style
- **-d/--depth** (*int*) – Block separation depth
- **-a/--alphabetize** (*int*) – Alphabetize attributes
- **-c/--close** (*int*) – Close tag style
- **-n/--namespace** (*int*) – Include namespace

Tools for managing units in GEOSX

**class** `geosx_xml_tools.unit_manager.UnitManager`

This class is used to manage unit definitions.

**buildUnits** () → None

Build the unit definitions.

**regexHandler** (*match: re.Match*) → str

Split the matched string into a scale and unit definition.

**Parameters** **match** (*re.match*) – The matching string from the regex.

**Returns** The string with evaluated unit definitions

**Return type** str

Tools for managing regular expressions in `geosx_xml_tools`

**class** `geosx_xml_tools.regex_tools.DictRegexHandler`

This class is used to substitute matched values with those stored in a dict.

`geosx_xml_tools.regex_tools.SymbolicMathRegexHandler` (*match: re.Match*) → str

Evaluate symbolic expressions that are identified using the `regex_tools.patterns['symbolic']`.

**Parameters** **match** (*re.match*) – A matching string identified by the regex.



`geosx_xml_tools.xml_redundancy_check.check_redundancy_level` (*local\_schema*:  
*Dict[str, Any]*, *node*:  
*<sphinx.ext.autodoc.importer.\_MockObject*  
*object* *at*  
*0x7f95eba76050>*,  
*whitelist*: *Iter-*  
*able[str] = ['compo-*  
*nent']*) → *int*

Check xml redundancy at the current level

#### Parameters

- **local\_schema** (*dict*) – Schema definitions
- **node** (*lxml.etree.Element*) – current xml node
- **whitelist** (*list*) – always match nodes containing these attributes

**Returns** Number of required attributes in the node and its children

**Return type** *int*

`geosx_xml_tools.xml_redundancy_check.check_xml_redundancy` (*schema*: *Dict[str, Any]*,  
*fname*: *str*) → *None*

Check redundancy in an xml file

#### Parameters

- **schema** (*dict*) – Schema definitions
- **fname** (*str*) – Name of the target file

`geosx_xml_tools.xml_redundancy_check.main` () → *None*  
 Entry point for the xml attribute usage test script

**Parameters** **-r/--root** (*str*) – GEOSX root directory

`geosx_xml_tools.xml_redundancy_check.process_xml_files` (*geosx\_root*: *str*) → *None*  
 Test for xml redundancy

**Parameters** **geosx\_root** (*str*) – GEOSX root directory

`geosx_xml_tools.attribute_coverage.collect_xml_attributes` (*xml\_types*: *Dict[str,*  
*Dict[str, Any]]*, *fname*:  
*str*, *folder*: *str*) →  
*None*

Collect xml attribute usage in a file

#### Parameters

- **xml\_types** (*dict*) – dictionary containing attribute usage
- **fname** (*str*) – name of the target file
- **folder** (*str*) – the source folder for the current file

`geosx_xml_tools.attribute_coverage.collect_xml_attributes_level` (*local\_types*:  
*Dict[str,*  
*Dict[str,*  
*Any]]*, *node*:  
*<sphinx.ext.autodoc.importer.\_MockOb*  
*ject* *at*  
*0x7f95eaa68d50>*,  
*folder*: *str*) →  
*None*

Collect xml attribute usage at the current level

**Parameters**

- **local\_types** (*dict*) – dictionary containing attribute usage
- **node** (*lxml.etree.Element*) – current xml node
- **folder** (*str*) – the source folder for the current file

`geosx_xml_tools.attribute_coverage.main()` → None

Entry point for the xml attribute usage test script

**Parameters**

- **-r/--root** (*str*) – GEOSX root directory
- **-o/--output** (*str*) – output file name

`geosx_xml_tools.attribute_coverage.parse_schema(fname: str)` → Dict[str, Dict[str, Any]]

Parse the schema file into the xml attribute usage dict

**Parameters** **fname** (*str*) – schema name

**Returns** Dictionary of attributes and children for the entire schema

**Return type** dict

`geosx_xml_tools.attribute_coverage.parse_schema_element` (*root:*  
*<sphinx.ext.autodoc.importer.\_MockObject*  
*object* *at*  
*0x7f95eaa68d50>*, *node:*  
*<sphinx.ext.autodoc.importer.\_MockObject*  
*object* *at*  
*0x7f95eaa68d50>*,  
*xsd:* *str* =  
*'{http://www.w3.org/2001/XMLSchema}'*,  
*recursive\_types:* *It-*  
*erable[str]* = [*'Peri-*  
*odicEvent'*, *'SoloEvent'*,  
*'HaltEvent'*], *folders:*  
*Iterable[str]* = [*'src'*,  
*'examples'*]) → Dict[str, Dict[str, Any]]

Parse the xml schema at the current level

**Parameters**

- **root** (*lxml.etree.Element*) – the root schema node
- **node** (*lxml.etree.Element*) – current schema node
- **xsd** (*str*) – the file namespace
- **recursive\_types** (*list*) – node tags that allow recursive nesting
- **folders** (*list*) – folders to sort xml attribute usage into

**Returns** Dictionary of attributes and children for the current node

**Return type** dict

`geosx_xml_tools.attribute_coverage.process_xml_files` (*geosx\_root: str*, *output\_name:*  
*str*) → None

Test for xml attribute usage

**Parameters**

- **geosx\_root** (*str*) – GEOSX root directory
- **output\_name** (*str*) – output file name

`geosx_xml_tools.attribute_coverage.write_attribute_usage_xml` (*xml\_types*:  
*Dict[str, Dict[str, Any]], fname: str*)  
 → None

Write xml attribute usage file

**Parameters**

- **xml\_types** (*dict*) – dictionary containing attribute usage by xml type
- **fname** (*str*) – output file name

`geosx_xml_tools.attribute_coverage.write_attribute_usage_xml_level` (*local\_types*:  
*Dict[str, Dict[str, Any]], node:*  
*<sphinx.ext.autodoc.importer.\_Module*  
*object at*  
*0x7f95eaa68d50>, folders:*  
*Iterable[str]*  
*= ['src', 'examples']*) →  
 None

Write xml attribute usage file at a given level

**Parameters**

- **local\_types** (*dict*) – dict containing attribute usage at the current level
- **node** (*lxml.etree.Element*) – current xml node

Tools for reading/writing GEOSX ascii tables

`geosx_xml_tools.table_generator.read_GEOS_table` (*axes\_files: Iterable[str], prop-*  
*erty\_files: Iterable[str]*) → *Tu-*  
*ple[Iterable[numpy.ndarray], Dict[str,*  
*numpy.ndarray]]*

Read an GEOS-compatible ascii table.

**Parameters**

- **axes\_files** (*list*) – List of the axes file names in order.
- **property\_files** (*list*) – List of property file names

**Returns** List of axis definitions, dict of property values

**Return type** tuple

```
geosx_xml_tools.table_generator.write_GEOS_table(axes_values: Iterable
                                                    able[numpy.ndarray],
                                                    properties: Dict[str, numpy.ndarray],
                                                    axes_names: Iterable[str] = ['x',
                                                    'y', 'z', 't'], string_format: str =
                                                    '%1.5e') → None
```

Write an GEOS-compatible ascii table.

#### Parameters

- **axes\_values** (*list*) – List of arrays containing the coordinates for each axis of the table.
- **properties** (*dict*) – Dict of arrays with dimensionality/size defined by the axes\_values
- **axes\_names** (*list*) – Names for each axis (default = ['x', 'y', 'z', 't'])
- **string\_format** (*str*) – Format for output values (default = %1.5e)

```
geosx_xml_tools.table_generator.write_read_GEOS_table_example() → None
```

Table read / write example.

## 8.2.4 PyGEOSX Tools

The *pygeosx\_tools* python package adds a variety of tools for working with pygeosx objects. These include common operations such as setting the value of geosx wrappers with python functions, parallel communication, and file IO. Examples using these tools can be found here: [pygeosx Examples](#).

### API

```
pygeosx_tools.wrapper.allgather_wrapper(problem, key, ghost_key="")
```

Get a global copy of a wrapper as a numpy ndarray on all ranks

#### Parameters

- **problem** (*pygeosx.Group*) – GEOSX problem handle
- **target\_key** (*str*) – Key for the target wrapper

**Returns** The wrapper as a numpy ndarray

**Return type** np.ndarray

```
pygeosx_tools.wrapper.gather_wrapper(problem, key, ghost_key="")
```

Get a global copy of a wrapper as a numpy ndarray on rank 0

#### Parameters

- **problem** (*pygeosx.Group*) – GEOSX problem handle
- **target\_key** (*str*) – Key for the target wrapper

**Returns** The wrapper as a numpy ndarray

**Return type** np.ndarray

```
pygeosx_tools.wrapper.get_global_value_range(problem, key)
```

Get the range of a target value across all processes

#### Parameters

- **problem** (*pygeosx.Group*) – GEOSX problem handle

- **target\_key** (*str*) – Key for the target wrapper

**Returns** The global min/max of the target

**Return type** tuple

`pygeosx_tools.wrapper.get_matching_wrapper_path(problem, filters)`

Recursively search the group and its children for wrappers that match the filters. A successful match is identified if the wrapper path contains all of the strings in the filter. For example, if `filters=['a', 'b', 'c']`, the following could match any of the following: 'a/b/c', 'c/b/a', 'd/e/c/f/b/a/a'

**Parameters**

- **problem** (`pygeosx.Group`) – GEOSX problem handle
- **filters** (*list*) – a list of strings

**Returns** Key of the matching wrapper

**Return type** str

`pygeosx_tools.wrapper.get_wrapper(problem, target_key, write_flag=False)`

Get a local copy of a wrapper as a numpy ndarray

**Parameters**

- **filename** (*str*) – Catalog file name
- **problem** (`pygeosx.Group`) – GEOSX problem handle
- **target\_key** (*str*) – Key for the target wrapper
- **write\_flag** (*bool*) – Sets write mode (default=False)

**Returns** The wrapper as a numpy ndarray

**Return type** np.ndarray

`pygeosx_tools.wrapper.get_wrapper_par(problem, target_key, allgather=False, ghost_key="")`

Get a global copy of a wrapper as a numpy ndarray. Note: if `ghost_key` is set, it will try to remove any ghost elements

**Parameters**

- **problem** (`pygeosx.Group`) – GEOSX problem handle
- **target\_key** (*str*) – Key for the target wrapper
- **allgather** (*bool*) – Flag to trigger allgather across ranks (False)
- **ghost\_key** (*str*) – Key for the corresponding ghost wrapper (default="")

**Returns** The wrapper as a numpy ndarray

**Return type** np.ndarray

`pygeosx_tools.wrapper.plot_history(records, output_root='.', save_figures=True, show_figures=True)`

Plot the time-histories for the records structure. Note: If figures are shown, the GEOSX process will be blocked until they are closed

**Parameters**

- **records** (*dict*) – A dict of dicts containing the queries
- **output\_root** (*str*) – Path to save figures (default = '.')
- **save\_figures** (*bool*) – Flag to indicate whether figures should be saved (default = True)

- **show\_figures** (*bool*) – Flag to indicate whether figures should be drawn (default = False)

`pygeosx_tools.wrapper.print_global_value_range` (*problem, key, header, scale=1.0, precision='%1.4f'*)

Print the range of a target value across all processes

#### Parameters

- **problem** (`pygeosx.Group`) – GEOSX problem handle
- **target\_key** (*str*) – Key for the target wrapper
- **header** (*str*) – Header to print with the range
- **scale** (*float*) – Multiply the range with this value before printing (default = 1.0)
- **precision** (*str*) – Format for printing the range (default = %1.4f)

**Returns** The global min/max of the target

**Return type** tuple

`pygeosx_tools.wrapper.run_queries` (*problem, records*)

Query the current GEOSX datastructure Note: The expected record request format is as follows. For now, the only supported query is to find the min/max values of the target record = { 'path/of/wrapper': { 'label': 'aperture (m)', # A label to include with plots 'scale': 1.0, # Value to scale results by 'history': [], # A list to store values over time 'fhandle': plt.figure() # A figure handle }}

#### Parameters

- **problem** (`pygeosx.Group`) – GEOSX problem handle
- **records** (*dict*) – A dict of dicts that specifies the queries to run

`pygeosx_tools.wrapper.search_datastructure_wrappers_recursive` (*group, filters, matching\_paths, level=0, group\_path=[]*)

Recursively search the group and its children for wrappers that match the filters

#### Parameters

- **problem** (`pygeosx.Group`) – GEOSX problem handle
- **filters** (*list*) – a list of strings
- **matching\_paths** (*list*) – a list of matching values

`pygeosx_tools.wrapper.set_wrapper_to_value` (*problem, key, value*)

Set the value of a wrapper

#### Parameters

- **problem** (`pygeosx.Group`) – GEOSX problem handle
- **target\_key** (*str*) – Key for the target wrapper
- **value** (*float*) – Value to set the wrapper

`pygeosx_tools.wrapper.set_wrapper_with_function` (*problem, target\_key, input\_keys, fn, target\_index=-1*)

Set the value of a wrapper using a function

#### Parameters

- **problem** (`pygeosx.Group`) – GEOSX problem handle

- **target\_key** (*str*) – Key for the target wrapper
- **input\_keys** (*str*, *list*) – The input key(s)
- **fn** (*function*) – Vectorized function used to calculate target values
- **target\_index** (*int*) – Target index to write the output (default = all)

`pygeosx_tools.file_io.load_tables` (*axes\_names: Iterable[str], property\_names: Iterable[str], table\_root: str = '/tables', extension: str = 'csv'*) → `Tuple[Iterable[numpy.ndarray], Dict[str, numpy.ndarray]]`

Load a set of tables in GEOSX format

#### Parameters

- **axes\_names** (*list*) – Axis file names in the target directory (with no extension)
- **property\_names** (*list*) – Property file names in the target directory (with not extension)
- **table\_root** (*str*) – Root path for the table directory
- **extension** (*str*) – Table file extension (default = 'csv')

**Returns** List of axes values, and dictionary of table values

**Return type** tuple

`pygeosx_tools.file_io.save_tables` (*axes: Iterable[numpy.ndarray], properties: Dict[str, numpy.ndarray], table\_root: str = '/tables', axes\_names: List[str] = []*) → `None`

Saves a set of tables in GEOSX format

The shape of these arrays should match the length of each axis in the specified order. The output directory will be created if it does not exist yet. If `axes_names` are not supplied, then they will be selected based on the dimensionality of the grid: 1D=[t]; 3D=[x, y, z]; 4D=[x, y, z, t].

#### Parameters

- **axes** (*list*) – A list of numpy ndarrays defining the table axes
- **properties** (*dict*) – A dict of numpy ndarrays defining the table values
- **table\_root** (*str*) – The root path for the output directory
- **axes\_names** (*list*) – A list of names for each potential axis (optional)

`pygeosx_tools.mesh_interpolation.apply_to_bins` (*fn: Callable[[Union[float, numpy.ndarray]], float], position: numpy.ndarray, value: numpy.ndarray, bins: numpy.ndarray, collapse\_edges: bool = True*)

Apply a function to values that are located within a series of bins Note: if a bin is empty, this function will fill a nan value

#### Parameters

- **fn** (*function*) – Function that takes a single scalar or array input
- **position** (*np.ndarray*) – A 1D list/array describing the location of each sample
- **value** (*np.ndarray*) – A 1D list/array of values at each location
- **bins** (*np.ndarray*) – The bin edges for the position data
- **collapse\_edges** (*bool*) – Controls the behavior of edge-data (default=True)

**Returns** an array of function results for each bin

**Return type** `np.ndarray`

`pygeosx_tools.mesh_interpolation.extrapolate_nan_values` (*x*, *y*, *slope\_scale*=0.0)  
Fill in any nan values in two 1D arrays by extrapolating

**Parameters**

- **x** (`np.ndarray`) – 1D list/array of positions
- **y** (`np.ndarray`) – 1D list/array of values
- **slope\_scale** (`float`) – value to scale the extrapolation slope (default=0.0)

**Returns** The input array with nan values replaced by extrapolated data

**Return type** `np.ndarray`

`pygeosx_tools.mesh_interpolation.get_random_realization` (*x*, *bins*, *value*, *rand\_fill*=0,  
*rand\_scale*=0,  
*slope\_scale*=0)

Get a random realization for a noisy signal with a set of bins

**Parameters**

- **x** (`np.ndarray`) – 1D list/array of positions
- **bins** (`np.ndarray`) – 1D list/array of bin edges
- **value** (`np.ndarray`) – 1D list/array of values
- **rand\_fill** (`float`) – The standard deviation to use where data is not defined (default=0)
- **rand\_scale** (`float`) – Value to scale the standard deviation for the realization (default=0)
- **slope\_scale** (`float`) – Value to scale the extrapolation slope (default=0.0)

**Returns** An array containing the random realization

**Return type** `np.ndarray`

`pygeosx_tools.mesh_interpolation.get_realizations` (*x*, *bins*, *targets*)  
Get random realizations for noisy signals on target bins

**Parameters**

- **x** (`np.ndarray`) – 1D list/array of positions
- **bins** (`np.ndarray`) – 1D list/array of bin edges
- **targets** (`dict`) – Dict of geosx target keys, inputs to `get_random_realization`

**Returns** Dictionary of random realizations

**Return type** `dict`

`pygeosx_tools.well_log.convert_E_nu_to_K_G` (*E*, *nu*)  
Convert young's modulus and poisson's ratio to bulk and shear modulus

**Parameters**

- **E** (`float`, `np.ndarray`) – Young's modulus
- **nu** (`float`, `np.ndarray`) – Poisson's ratio

**Returns** bulk modulus, shear modulus with same size as inputs

**Return type** `tuple`



`pygeosx_tools.well_log.estimate_shmin(z, rho, nu)`

Estimate the minimum horizontal stress using the poisson's ratio

**Parameters**

- **z** (*float, np.ndarray*) – Depth
- **rho** (*float, np.ndarray*) – Density
- **nu** (*float, np.ndarray*) – Poisson's ratio

**Returns** minimum horizontal stress

**Return type** float

`pygeosx_tools.well_log.parse_las(fname, variable_start='~C', body_start='~A')`

Parse an las format log file

**Parameters**

- **fname** (*str*) – Path to the log file
- **variable\_start** (*str*) – A string that indicates the start of variable header information (default = '~CURVE INFORMATION')
- **body\_start** (*str*) – a string that indicates the start of the log body (default = '~A')

**Returns** a dict containing the values and unit definitions for each variable in the log

**Return type** np.ndarray

## 8.2.5 Time History Tools

`timehistory.plot_time_history.getHistorySeries(database, variable, setname, indices=None, components=None)`

Retrieve a series of time history structures suitable for plotting in addition to the specific set index and component for the time series

**Parameters**

- **database** (*hdf5\_wrapper.hdf5\_wrapper*) – database to retrieve time history data from
- **variable** (*str*) – the name of the time history variable for which to retrieve time-series data
- **setname** (*str*) – the name of the index set as specified in the geosx input xml for which to query time-series data
- **indices** (*int, list*) – the indices in the named set to query for, if None, defaults to all
- **components** (*int, list*) – the components in the flattened data types to retrieve, defaults to all

**Returns** list of (time, data, idx, comp) timeseries tuples for each time history data component

**Return type** list



Welcome to the GEOSX build guide.

## 9.1 System prerequisites

To configure and build GEOSX you will need the following tools available on your system.

### 9.1.1 List of prerequisites

Minimal requirements:

- [CMake](#) build system generator (3.17+).
- build tools ([GNU make](#) or [ninja](#) on Linux, XCode on MacOS).
- a C++ compiler with full c++14 standard support ([gcc](#) 8.3+ or [clang](#) 8.0+ are recommended).
- [python](#) (2.7+ or 3.6+).
- [zlib](#), [blas](#) and [lapack](#) libraries
- any compatible MPI runtime and compilers (if building with MPI)

If you want to build from a repository check out (instead of a release tarball):

- [git](#) (2.20+ is tested, but most versions should work fine)

If you plan on building bundled third-party library (TPLs) dependencies yourself:

- Compatible C and Fortran compilers

If you will be checking out and running integrated tests (a submodule of GEOSX, currently not publicly available):

- [git-lfs](#) (Git Large File Storage extension)
- [h5py](#) and [mpi4py](#) python modules

If you are interested in building Doxygen documentation:

- GNU bison
- LaTeX
- ghostscript
- Graphviz

In order for XML validation to work (executed as an optional build step):

- xmlint

### 9.1.2 Installing prerequisites

On a local development machine with sudo/root privileges, most of these dependencies can be installed with a system package manager. For example, on a Debian-based system (check your package manager for specific package names):

```
sudo apt install build-essential git git-lfs gcc g++ gfortran cmake libopenmpi-dev_  
↳ libblas-dev liblapack-dev zlib1g-dev python3 python3-h5py python3-mpi4py libxml2-  
↳ utils
```

On HPC systems it is typical for these tools to be installed by system administrators and provided via [modules](#). To list available modules, type:

```
module avail
```

Then load the appropriate modules using `module load` command. Please contact your system administrator if you need help choosing or installing appropriate modules.

## 9.2 Third-party dependencies

GEOSX makes use of multiple third-party libraries (TPLs) and tools, some of which are mandatory and some optional. We only test against specific versions, and sometimes even require development snapshots (specific git commits). Not all of these guarantee backwards compatibility, so we strongly recommend building with these specific versions.

### 9.2.1 List of third-party libraries and tools

The two tables below lists the dependencies with their specific versions and relevant CMake variables. Some of these libraries may have their own system prerequisites.

#### Libraries

The following libraries are linked to by GEOSX:

Name	Version	Enable option	Path variable	Description
Adiak	0.2.0	ENABLE_CALIPER	ADIAK_DIR	Library for collecting metadata from HPC application runs, and distributing that metadata to subscriber tools.
Caliper	2.4.0	ENABLE_CALIPER	CALIPER_DIR	Instrumentation and performance profiling library.
conduit	0.5.0	<i>mandatory</i>	CONDUIT_DIR	Simplified Data Exchange for HPC Simulations.
CHAI	2.2.2	<i>mandatory</i>	CHAI_DIR	Copy-hiding array abstraction to automatically migrate data between memory spaces.
RAJA	0.12.1	<i>mandatory</i>	RAJA_DIR	Collection of C++ software abstractions that enable architecture portability for HPC applications.
hdf5	1.10.5	<i>mandatory</i>	HDF5_DIR	High-performance data management and storage suite.
math-presso	2015-12-15	ENABLE_MATHPRESSO	MATHPRESSO_DIR	Mathematical Expression Parser and JIT Compiler.
pugixml	1.8.0	<i>mandatory</i>	PUGIXML_DIR	Light-weight, simple and fast XML parser for C++ with XPath support.
parmetis	4.0.3	<i>mandatory</i> (with MPI)	PARMETIS_DIR	Parallel Graph Partitioning library. Should be built with 64-bit <code>idx_t</code> type.
suites-parse	5.8.1	ENABLE_SUITESPARSE	SUITESPARSE_DIR	A suite of sparse matrix software.
superlu-dist	0f6efc3	ENABLE_SUPERLU	SUPERLU_DIST_DIR	General purpose library for the direct solution of large, sparse, nonsymmetric systems of linear equations.
hypre	2186a8f	ENABLE_HYPRE	HYPRE_DIR	Library of high performance preconditioners and solvers for large sparse linear systems on massively parallel computers.
PETSc	3.13.0	ENABLE_PETSC	PETSC_DIR	Suite of data structures and routines for the scalable (parallel) solution of scientific applications.
Trilinos	12.18.1	ENABLE_TRILINOS	TRILINOS_DIR	Collection of reusable scientific software libraries, known in particular for linear solvers.
siilo	4.10.3	<i>mandatory</i>	SILO_DIR	A Mesh and Field I/O Library and Scientific Database.
VTK	9.0.0-rc3	ENABLE_VTK	VTK_DIR	Open source software for manipulating and displaying scientific data.

## Tools

The following tools are used as part of the build process to support GEOSX development:

Name	Version	Enable option	Path variable	Description
doxygen	1.8.20	ENABLE_DOXYGEN	DOXYGEN_EXECUTABLE	De facto standard tool for generating documentation from annotated C++ sources.
sphinx	1.8.5	ENABLE_SPHINX	SPHINX_EXECUTABLE	A tool that makes it easy to create intelligent and beautiful documentation.
uncrustify	401a409	ENABLE_UNCRUSTIFY	UNCrustify_EXECUTABLE	A source code beautifier for C, C++, C#, ObjectiveC, D, Java, Pawn and VALA.

Some other dependencies ([GoogleTest](#), [GoogleBenchmark](#)) are provided through [BLT](#) build system which is embedded in GEOSX source. No actions are needed to build them.

If you would like to create a Docker image with all dependencies, take a look at [Dockerfiles](#) that are used in our CI process.

## 9.2.2 Building bundled dependencies

To simplify the process of building TPLs, we provide a git repository [thirdPartyLibs](#). It contains source copies of exact TPL versions required and is updated periodically. It also contains a CMake script for building all TPLs in a single command.

The recommended steps to build TPLs are:

- Create a host-config file that sets all system-specific CMake variables (compiler and library paths, configuration flags, etc.) Take a look at [host-config examples](#).
- Configure via `config-build.py` script:

```
cd thirdPartyLibs
python scripts/config-build.py --hostconfig=/path/to/host-config.cmake --
    ↪ buildtype=Release --installpath=/path/to/install/dir -DNUM_PROC=8
```

where

- `--buildpath` or `-bp` is the build directory (by default, created under current).
  - `--installpath` or `-ip` is the installation directory (wraps `CMAKE_INSTALL_PREFIX`).
  - `--buildtype` or `-bt` is a wrapper to the `CMAKE_BUILD_TYPE` option.
  - `--hostconfig` or `-hc` is a path to host-config file.
  - all other command-line options are passed to CMake.
- Run the build:

```
cd <buildpath>
make
```

**Warning:** Do not provide `-j` argument to `make` here, since the top-level `make` only launches sub-project builds. Instead use `-DNUM_PROC` option above, which is passed to each sub-project's `make` command.

You may also run the CMake configure step manually instead of relying on `config-build.py`. The full TPL build may take anywhere between 15 minutes and 2 hours, depending on your machine, number of threads and libraries enabled.

---

**Note:** An exception from the above pattern, `sphinx` is currently not a part of the TPL bundle and must be installed with your Python or package manager.

---

---

**Note:** PETSc build currently downloads [pt-scotch](#) from the internet. If you do not have access to internet, modify the `./configure` step of `petsc` in `CMakeLists.txt` and change the `--download-ptscotch` option accordingly. `pt-scotch` also relies on `bison` and `flex`.

---

## 9.2.3 Installing dependencies individually

You may also install each individual TPL separately, either manually or through a package manager. This is a more difficult route, since you are responsible for configuring dependencies in a compatible manner. Again, we strongly recommend using the exact versions listed above, to avoid possible build problems.

You may look at [our TPL CMake script](#) to see how we configure TPL builds.

## 9.3 Building GEOSX

### 9.3.1 Build steps

- Create a host-config file that sets all system-specific CMake variables. Take a look at [host-config examples](#). We recommend the same host-config is used for both TPL and GEOSX builds. In particular, certain options (such as `ENABLE_MPI` or `ENABLE_CUDA`) need to match between the two.
- Provide paths to all enabled TPLs. This can be done in one of two ways:
  - Provide each path via a separate CMake variable (see [Third-party dependencies](#) for path variable names).
  - If you built TPLs from the `tplMirror` repository, you can set `GEOSX_TPL_DIR` variable in your host-config to point to the TPL installation path, and

```
include("/path/to/GEOSX/host-configs/tpls.cmake")
```

which will set all the individual TPL paths for you.

- Configure via `config-build.py` script:

```
cd GEOSX
python scripts/config-build.py --hostconfig=/path/to/host-config.cmake --
↪buildtype=Release --installpath=/path/to/install/dir
```

where

- `--buildpath` or `-bp` is the build directory (by default, created under current working dir).
- `--installpath` or `-ip` is the installation directory (wraps `CMAKE_INSTALL_PREFIX`).
- `--buildtype` or `-bt` is a wrapper to the `CMAKE_BUILD_TYPE` option.
- `--hostconfig` or `-hc` is a path to host-config file.
- all unrecognized options are passed to CMake.

If `--buildpath` is not used, build directory is automatically named `build-<config-filename-without-extension>-<buildtype>`. It is possible to keep automatic naming and change the build root directory with `--buildrootdir`. In that case, build path will be set to `<buildrootdir>/<config-filename-without-extension>-<buildtype>`. Both `--buildpath` and `--buildrootdir` are incompatible and cannot be used in the same time. Same pattern is applicable to install path, with `--installpath` and `--installrootdir` options.

- Run the build:

```
cd <buildpath>
make -j $(nproc)
```

You may also run the CMake configure step manually instead of relying on `config-build.py`. A full build typically takes between 10 and 30 minutes, depending on chosen compilers, options and number of cores.

### 9.3.2 Configuration options

Below is a list of CMake configuration options, in addition to TPL options above. Some options, when enabled, require additional settings (e.g. `ENABLE_CUDA`). Please see [host-config examples](#).

Option	Default	Explanation
ENABLE_MPI	ON	Build with MPI (also applies to TPLs)
ENABLE_OPENMP	OFF	Build with OpenMP (also applies to TPLs)
ENABLE_CUDA	OFF	Build with CUDA (also applies to TPLs)
ENABLE_CUDA_NVTX	OFF	Enable CUDA NVTX user instrumentation (via GEOSX_MARK_SCOPE or GEOSX_MARK_FUNCTION macros)
ENABLE_DOCS	ON	Build documentation (Sphinx and Doxygen)
ENABLE_WARNINGS_AS_ERRORS	OFF	Treat all warnings as errors
ENABLE_PAMELA	ON	Enable PAMELA library (required for external mesh import)
ENABLE_PVTPackage	ON	Enable PVTPackage library (required for compositional flow runs)
ENABLE_TOTALVIEW_OUTPUT	OFF	Enables TotalView debugger custom view of GEOSX data structures
GEOSX_ENABLE_FPE	ON	Enable floating point exception trapping
GEOSX_LA_INTERFACE	HyPre	Choice of Linear Algebra backend (HyPre/Petsc/Trilinos)
GEOSX_BUILD_OBJ_LIBS	ON	Use CMake Object Libraries build
GEOSX_BUILD_SHARED_LIBS	OFF	Build geosx_core as a shared library instead of static
GEOSX_PARALLEL_COMPILE_JOBS	1	Max. number of compile jobs (when using Ninja), in addition to -j flag
GEOSX_PARALLEL_LINK_JOBS	1	Max. number of link jobs (when using Ninja), in addition to -j flag

## 9.4 Spack and Uberenv

GEOSX is transitioning to a new [Spack](#) and [Uberenv](#) system for building our dependencies. We refer the reader to the [Spack documentation](#) and [Uberenv documentation](#), in particular the Spack documentation for [specs](#) and [dependencies](#), [manual compiler configuration](#) and [external packages](#) are worth reading.

Building the dependencies can be as simple as running

```
./scripts/uberenv/uberenv.py
```

This will create a directory `uberenv_libs` in the current working directory, clone [Spack](#) into `uberenv_libs/spack` and install the dependencies into `uberenv_libs/system_dependent_path`. It will then spit out a host-config file in the current directory which you can use to build GEOSX. While the above command **should** work on every system, it should never be used. Invoked as such, Spack will ignore any system libraries you have installed and will go down a rabbit hole building dependencies. Furthermore this does not allow you to choose the compiler to build. Both of these are easily solved by creating a directory with a `packages.yaml` and a `compilers.yaml`.

To prevent this from happening you'll need to create a directory with a `packages.yaml` file and a `compilers.yaml` file. You can find working examples for commonly used systems in [scripts/uberenv/spack\\_configs](#). It is worth noting that each LC system type has two such directories, for example there is a `toss_3_x85_54_ib` and `toss_3_x85_54_ib_python` directory. This is because when building `pygeosx` Python needs to be built from scratch, and as such cannot be listed in `packages.yaml`. However, when not building `pygeosx` other dependencies depend on python, but an existing system version works just fine, so it can be put in `packages.yaml` to prevent Spack from building it.

Once you have these files setup you can run Uberenv again and instruct it to use them with. If for instance you added Clang 10.0.1 to the `compilers.yaml` file the your command would look something like this:

```
./scripts/uberenv/uberenv.py --spack-config-dir=/path/to/your/config/directory/ --
↪ spec="%clang@10.0.1"
```

**Note:** When building `pygeosx`, Spack will build various python packages, however by default they are not installed in python. There are various ways of accomplishing [this](#), but the recommended approach is to use `spack`



activate. The command would look something like this `./uberenv_libs/spack/bin/spack activate py-numpy py-scipy py-pip py-mpi4py`

## 9.4.1 Build Configuration

The GEOSX Spack package has a lot of options for controlling which dependencies you would like to build and how you'd like them built. The GEOSX Spack package file is at `scripts/uberenv/packages/geosx/package.py` <<https://github.com/GEOSX/GEOSX/tree/develop/scripts/uberenv/packages/geosx/package.py>>`\_. The variants for the package are as follows

```
variant('shared', default=True, description='Build Shared Libs.')
variant('caliper', default=True, description='Build Caliper support.')
variant('mkl', default=False, description='Use the Intel MKL library.')
variant('essl', default=False, description='Use the IBM ESSL library.')
variant('suite-sparse', default=True, description='Build SuiteSparse support.')
variant('trilinos', default=True, description='Build Trilinos support.')
variant('hypre', default=True, description='Build HYPRE support.')
variant('hypre-cuda', default=False, description='Build HYPRE with CUDA support.')
variant('petsc', default=True, description='Build PETSc support.')
variant('scotch', default=True, description='Build Scotch support.')
variant('lai',
      default='trilinos',
      description='Linear algebra interface.',
      values=('trilinos', 'hypre', 'petsc'),
      multi=False)
variant('pygeosx', default=False, description='Build the GEOSX python interface.')
```

For example if you wanted to build with GCC 8.3.1, without Caliper and with PETSC as the Linear Algebra Interface, your spec would be `%gcc@8.3.1 ~caliper lai=petsc`.

The GEOSX Spack package lists out the libraries that GEOSX depends on. Currently these dependencies are

```
depends_on('cmake@3.8:', type='build')
depends_on('cmake@3.9:', when='+cuda', type='build')

#
# Virtual packages
#
depends_on('mpi')
depends_on('blas')
depends_on('lapack')

#
# Performance portability
#
depends_on('raja@0.12.1 +openmp +shared ~examples ~exercises')
depends_on('raja +cuda', when='+cuda')

depends_on('umpire@4.1.2 ~c +shared +openmp ~examples')
depends_on('umpire +cuda', when='+cuda')

depends_on('chai@2.2.2 +shared +raja ~benchmarks ~examples')
```

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```

depends_on('chai@2.2.2 +cuda', when='+cuda')

#
# IO
#
depends_on('hdf5@1.10.5: +shared +pic +mpi', when='~vtk')

depends_on('conduit@0.5.0 +shared ~test ~fortran +mpi +hdf5 ~hdf5_compat')

depends_on('silo@4.10: ~fortran +shared ~silex +pic +mpi ~zlib')

depends_on('adiak@0.2: +mpi +shared', when='+caliper')
depends_on('caliper@2.4: +shared +adiak +mpi ~callpath ~libpfm ~gotcha ~sampler',
↳when='+caliper')

depends_on('pugixml@1.8: +shared')

depends_on('fmt@8.0: cxxstd=14 +pic')

#
# Math
#
depends_on('intel-mkl +shared ~ilp64', when='+mkl')

# depends_on('essl ~ilp64 threads=openmp +lapack +cuda', when='+essl')

depends_on('parmetis@4.0.3: +shared +int64')

depends_on('scotch@6.0.9: +mpi +int64', when='+scotch')

depends_on('superlu-dist +int64 +openmp +shared', when='~petsc')
depends_on('superlu-dist@6.3.0 +int64 +openmp +shared', when='+petsc')

depends_on('suite-sparse@5.8.1: +pic +openmp +amd +camd +colamd +ccolamd +cholmod_
↳+umfpack', when='+suite-sparse')
depends_on('suite-sparse +blas-no-underscore', when='%gcc +suite-sparse +essl')

trilinos_build_options = '~fortran +openmp +shared'
trilinos_tpls = '~boost ~glm ~gtest ~hdf5 ~hypr ~matio ~metis +mpi ~mumps ~
↳netcdf ~suite-sparse'
trilinos_packages = '+amesos +aztec +epetra +epetraext +ifpack +kokkos +ml +stk_
↳+stratimikos +teuchos +tpetra ~amesos2 ~anasazi ~belos ~exodus ~ifpack2 ~muelu ~
↳sacado ~zoltan ~zoltan2'
depends_on('trilinos@12.18.1 ' + trilinos_build_options + trilinos_tpls +
↳trilinos_packages, when='+trilinos')
depends_on('trilinos +blas_lowercase_no_underscore', when='+trilinos +essl')
# depends_on('trilinos +force-new-lapack', when='+trilinos +essl')

depends_on('hypr@2.20.300 +shared +superlu-dist +mixedint +mpi +openmp', when=
↳'+hypr')
depends_on('hypr@2.20.300 +cuda +shared +superlu-dist +mpi +openmp +unified-
↳memory +cusparse', when='+hypr-cuda')

petsc_build_options = '+shared +mpi'
petsc_tpls = '+metis ~hdf5 ~hypr +superlu-dist +int64'
depends_on('petsc@3.13.0: ' + petsc_build_options + petsc_tpls, when='+petsc')

```

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```

#
# Python
#
depends_on('python +shared +pic', when='+pygeosx')
depends_on('py-numpy@1.19: +blas +lapack +force-parallel-build', when='+pygeosx')
depends_on('py-scipy@1.5.2: +force-parallel-build', when='+pygeosx')
depends_on('py-mpi4py@3.0.3:', when='+pygeosx')
depends_on('py-pip', when='+pygeosx')

#
# Dev tools
#
depends_on('uncrustify@0.71:')

#
# Documentation
#
depends_on('doxygen@1.8.13:', when='+docs', type='build')
depends_on('py-sphinx@1.6.3:', when='+docs', type='build')

```

Using the Spack spec syntax you can inturn specify variants for each of the dependencies of GEOSX. So for example if you could modify the spec above to build RAJA in debug by using `%gcc@8.3.1 ~caliper lai=petsc ^raja build_type=Debug`. When building with Uberenv Spack should print out a table containing the full spec for every dependency it will build. If you would like to look at the variants for say RAJA in more detail you can find the package file at `uberenv_libs/spack/var/spack/repos/builtin/packages/raja/package.py`.

## 9.4.2 Adding a Dependency (Advanced)

Adding a dependency to GEOSX is straight forward if the dependency already builds with Spack. If that is the case then all you need to do is add a `depends_on('cool-new-library')` to the GEOSX `package.py` file. If however the dependency doesn't have a Spack package, you will have to add one by creating a `cool-new-library/package.yaml` file in the `scripts/uberenv/packages` directory and adding the logic to build it there.

Oftentimes (unfortunately), even when a package already exists, it might not work out of the box for your system. In this case copy over the existing `package.py` file from the Spack repository into `scripts/uberenv/packages/cool-new-library/package.py`, as if you were adding a new package, and perform your modifications there. Once you have the package working, copy the package back into the Spack repository (running Uberenv should do this for you) and commit+push your changes to Spack.

## 9.5 Continuous Integration process

To save building time, the third party libraries (that do not change so often) and GEOSX are build separately.

Everytime a pull is requested in the TPL repository, docker images are generated and deployed on [dockerhub](#). The repository names (`ubuntu18.04-gcc8`, `centos7.7.1908-clang9.0.0`, `centos7.6.1810-gcc8.3.1-cuda10.1.243` etc.) obviously reflect the OS and the compiler flavour used. For each image, the unique `${TRAVIS_PULL_REQUEST}-${TRAVIS_BUILD_NUMBER}` tag is used so we can connect the related code source in a rather convenient way. Each docker contains the `org.opencontainers.image.created` and `org.opencontainers.image.revision` labels to provide additional information.

For the OSX builds, we construct a tarball of the TPLs and save them in a remote cloud storage. There is currently only one mac osx tested environment (xcode 11.2) and the same

`${TRAVIS_PULL_REQUEST}-${TRAVIS_BUILD_NUMBER}` pattern is used as an identifier for the build. An important counterpart to using a tarball and not a docker image is that the tarball does not provide the whole system the precompiled binaries rely on. Problems may arise since we use the rolling release [Homebrew](#) (to install open-mpi in particular). To circumvent this potential issue, the brew version is fixed to a specific commit (see `BREW_HASH` variable in [third party's .travis.yml](#)) and stored as a metainformation of the tarball blob inside the cloud storage. It is therefore possible for GEOSX to recover this information and build against the same revision of brew packages. Note that the `TRAVIS_PULL_REQUEST`, `TRAVIS_BUILD_NUMBER` and `TRAVIS_COMMIT` are also stored as metainformation in the same way (have a look at the OSX build section of [GEOSX's .travis.yml](#) to see how to retrieve these informations).

There thus is only one unique identifier for both dockers and mac osx builds for one TPL code base. It is necessary to define the global environment `GEOSX_TPL_TAG` (e.g. something like 82-254) to build against one selected version of the TPL.

It must be mentioned that one and only one version of the compiled TPL tarball is stored per pull request (older ones are removed automatically). Therefore, a client building against a work in progress PR may experience a 404 error sooner or later.

## 9.5.1 Building docker images

Our continuous integration process builds the TPL and GEOSX against two operating systems (ubuntu and centos) and two compilers (clang and gcc). The docker files use [multi-stage builds](#) in order to minimise the sizes of the images.

- First stage installs and defines all the elements that are commons to both TPL and GEOSX (for example, MPI and c++ compiler, BLAS, LAPACK, path to the installation directory...).
- As a second stage, we install everything needed to build (*not run*) the TPLs. We keep nothing from this second step for GEOSX, except the compiled TPL themselves. For example, a fortran compiler is needed by the TPL but not by GEOSX: it shall be installed during this step, so GEOSX won't access a fortran compiler (it does not have to).
- Last stage copies the compiled TPL from second stage and installs the elements only required by GEOSX (there are few).

## 9.5.2 Docker images contract

GEOSX will find a compiled version of the third party libraries.

As part of the contract provided by the TPL, the docker images also defines several environment variables. The

```
GEOSX_TPL_DIR
```

variable contains the absolute path of the installation root directory of the third party libraries. GEOSX must use it when building.

Other variables are classical absolute path compiler variables.

```
CC
CXX
MPICC
MPICXX
```

And the absolute path the `mpirun` (or equivalent) command.

```
MPIEXEC
```

The following `openmpi` environment variables allow it to work properly in the docker container. But there should be no reason to access or use them explicitly.

```
OMPI_CC=$CC  
OMPI_CXX=$CXX
```



### 10.1 Input Schema Definitions

XML Schema

## 10.2 Element: AcousticSEM

Name	Type	De- fault	Description
cflFactor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
discretization	string	required	Name of discretization object (defined in the <a href="#">Numerical Methods</a> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <a href="#">Finite Element Discretization</a> should be specified. If this is a Finite Volume Method, the name of a <a href="#">Finite Volume Discretization</a> discretization should be specified.
dtSeismo-Trace	real64	0	Time step for output pressure at receivers
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
logLevel	integer	0	Log level
name	string	required	A name is required for any non-unique nodes
output-Seismo-Trace	integer	0	Flag that indicates if we write the seismo trace in a file .txt, 0 no output, 1 otherwise
receiver-Coordinates	real64_array	2, required	Coordinates (x,y,z) of the receivers
rickerOrder	integer	2	Flag that indicates the order of the Ricker to be used 0, 1 or 2. Order 2 by default
source-Coordinates	real64_array	2, required	Coordinates (x,y,z) of the sources
targetRegions	string_array	required	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
time-Source-Frequency	real32	required	Central frequency for the time source
Linear-Solver-Parameters	node	unique	<a href="#">Element: LinearSolverParameters</a>
Non-linear-Solver-Parameters	node	unique	<a href="#">Element: NonlinearSolverParameters</a>





## 10.3 Element: Aquifer

Name	Type	Default	Description
allowAllPhasesIntoAquifer	integer	0	<p>Flag to allow all phases to flow into the aquifer. This flag only matters for the configuration in which flow is from reservoir to aquifer.</p> <ul style="list-style-type: none"> <li>- If the flag is equal to 1, then all phases, including non-aqueous phases, are allowed to flow into the aquifer.</li> <li>- If the flag is equal to 0, then only the water phase is allowed to flow into the aquifer.</li> </ul> <p>If you are in a configuration in which flow is from reservoir to aquifer and you expect non-aqueous phases to saturate the reservoir cells next to the aquifer, set this flag to 1.</p> <p>This keyword is ignored for single-phase flow simulations</p>
aquiferAngle	real64	required	Angle subtended by the aquifer boundary from the center of the reservoir [degress]
aquiferElevation	real64	required	Aquifer elevation (positive going upward) [m]
aquiferInitialPressure	real64	required	Aquifer initial pressure [Pa]
aquiferInnerRadius	real64	required	Aquifer inner radius [m]
aquiferPermeability	real64	required	Aquifer permeability [m <sup>2</sup> ]
aquiferPorosity	real64	required	Aquifer porosity
aquiferThickness	real64	required	Aquifer thickness [m]
aquiferTotalCompressibility	real64	required	Aquifer total compressibility (rock and fluid) [Pa <sup>-1</sup> ]
aquiferWaterDensity	real64	required	Aquifer water density
aquiferWaterPhaseComponentFraction	real64 array	{0}	Aquifer water phase component fraction. This keyword is ignored for single-

## 10.4 Element: Benchmarks

Name	Type	Default	Description
lassen	node	unique	<i>Element: lassen</i>
quartz	node	unique	<i>Element: quartz</i>

## 10.5 Element: BiotPorosity

Name	Type	Default	Description
defaultReferencePorosity	real64	required	Default value of the reference porosity
grainBulkModulus	real64	required	Grain bulk modulus
name	string	required	A name is required for any non-unique nodes

## 10.6 Element: BlackOilFluid

Name	Type	Default	Description
componentMolarWeight	real64_array	required	Component molar weights
componentNames	string_array	{}	List of component names
hydrocarbonFormationVolumeFactorTableNames	string_array	{}	<p>List of formation volume factor TableFunction names from the Functions block.</p> <p>The user must provide one TableFunction per hydrocarbon phase, in the order provided in “phaseNames”.</p> <p>For instance, if “oil” is before “gas” in “phaseNames”, the table order should be: oilTableName, gasTableName</p>
hydrocarbonViscosityTableNames	string_array	{}	<p>List of viscosity TableFunction names from the Functions block.</p> <p>The user must provide one TableFunction per hydrocarbon phase, in the order provided in “phaseNames”.</p> <p>For instance, if “oil” is before “gas” in “phaseNames”, the table order should be: oilTableName, gasTableName</p>
name	string	required	A name is required for any non-unique nodes
phaseNames	string_array	required	List of fluid phases
surfaceDensities	real64_array	required	List of surface mass densities for each phase
tableFiles	path_array	{}	List of filenames with input PVT tables (one per phase)
waterCompressibility	real64	0	Water compressibility
waterFormationVolumeFactor	real64	0	Water formation volume factor
waterReferencePressure	real64	0	Water reference pressure
waterViscosity	real64	0	Water viscosity

## 10.7 Element: Blueprint

Name	Type	De- fault	Description
childDirectory	string		Child directory path
name	string	re- quired	A name is required for any non-unique nodes
outputFullQuadratureData	integer	0	If true writes out data associated with every quadrature point.
parallelThreads	integer	1	Number of plot files.
plotLevel	geosx_dataRepository_PlotLevel		Determines which fields to write.

## 10.8 Element: BoundedPlane

Name	Type	De- fault	Description
dimensions	real64_array	re- quired	Length and width of the bounded plane
length-Vector	R1Tensor	re- quired	Tangent vector defining the orthonormal basis along with the normal.
name	string	re- quired	A name is required for any non-unique nodes
normal	R1Tensor	re- quired	Normal (n_x,n_y,n_z) to the plane (will be normalized automatically)
origin	R1Tensor	re- quired	Origin point (x,y,z) of the plane (basically, any point on the plane)
tolerance	real64	1e-05	Tolerance to determine if a point sits on the plane or not. It is relative to the maximum dimension of the plane.
widthVec- tor	R1Tensor	re- quired	Tangent vector defining the orthonormal basis along with the normal.

## 10.9 Element: Box

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes
strike	real64	-90	The strike angle of the box
xMax	R1Tensor	required	Maximum (x,y,z) coordinates of the box
xMin	R1Tensor	required	Minimum (x,y,z) coordinates of the box

## 10.10 Element: BrooksCoreyBakerRelativePermeability

Name	Type	Default	Description
gasOilRelPermExponent	real64_array	{1}	Rel perm power law exponent for the pair (gas phase, oil phase) at residual water saturation The expected format is “{ gasExp, oilExp }”, in that order
gasOilRelPermMaxValue	real64_array	{0}	Maximum rel perm value for the pair (gas phase, oil phase) at residual water saturation The expected format is “{ gasMax, oilMax }”, in that order
name	string	required	A name is required for any non-unique nodes
phaseMinVolumeFraction	real64_array	{0}	Minimum volume fraction value for each phase
phaseNames	string_array	required	List of fluid phases
waterOilRelPermExponent	real64_array	{1}	Rel perm power law exponent for the pair (water phase, oil phase) at residual gas saturation The expected format is “{ waterExp, oilExp }”, in that order
waterOilRelPermMaxValue	real64_array	{0}	Maximum rel perm value for the pair (water phase, oil phase) at residual gas saturation The expected format is “{ waterMax, oilMax }”, in that order

## 10.11 Element: BrooksCoreyCapillaryPressure

Name	Type	De- fault	Description
capPressureEpsilon	real64	1e-06	Wetting-phase saturation at which the max cap. pressure is attained; used to avoid infinite cap. pressure values for saturations close to zero
name	string	re- quired	A name is required for any non-unique nodes
phaseCapPressureExponentInv	real64_array	{1}	Inverse of capillary power law exponent for each phase
phaseEntryPressure	real64_array	{1}	Entry pressure value for each phase
phaseMinVolumeFraction	real64_array	{0}	Minimum volume fraction value for each phase
phaseNames	string_array	re- quired	List of fluid phases

## 10.12 Element: BrooksCoreyRelativePermeability

Name	Type	De- fault	Description
name	string	re- quired	A name is required for any non-unique nodes
phaseMinVolumeFraction	real64_array	{0}	Minimum volume fraction value for each phase
phaseNames	string_array	re- quired	List of fluid phases
phaseRelPermExponent	real64_array	{1}	Minimum relative permeability power law exponent for each phase
phaseRelPermMaxValue	real64_array	{0}	Maximum relative permeability value for each phase

## 10.13 Element: CO2BrineEzrokhiFluid

Name	Type	De- fault	Description
componentMolarWeight	real64_array	{0}	Component molar weights
componentNames	string_array	{ }	List of component names
flashModelParaFile	path	re- quired	Name of the file defining the parameters of the flash model
name	string	re- quired	A name is required for any non-unique nodes
phaseNames	string_array	{ }	List of fluid phases
phasePVTParaFiles	path_array	re- quired	Names of the files defining the parameters of the viscosity and density models

## 10.14 Element: CO2BrineEzrokhiThermalFluid

Name	Type	De- fault	Description
componentMolar-Weight	real64_array	{0}	Component molar weights
componentNames	string_array	{ }	List of component names
flashModelParaFile	path	re- quired	Name of the file defining the parameters of the flash model
name	string	re- quired	A name is required for any non-unique nodes
phaseNames	string_array	{ }	List of fluid phases
phasePVTParaFiles	path_array	re- quired	Names of the files defining the parameters of the viscosity and density models

## 10.15 Element: CO2BrinePhillipsFluid

Name	Type	De- fault	Description
componentMolar-Weight	real64_array	{0}	Component molar weights
componentNames	string_array	{ }	List of component names
flashModelParaFile	path	re- quired	Name of the file defining the parameters of the flash model
name	string	re- quired	A name is required for any non-unique nodes
phaseNames	string_array	{ }	List of fluid phases
phasePVTParaFiles	path_array	re- quired	Names of the files defining the parameters of the viscosity and density models

## 10.16 Element: CO2BrinePhillipsThermalFluid

Name	Type	De- fault	Description
componentMolar-Weight	real64_array	{0}	Component molar weights
componentNames	string_array	{ }	List of component names
flashModelParaFile	path	re- quired	Name of the file defining the parameters of the flash model
name	string	re- quired	A name is required for any non-unique nodes
phaseNames	string_array	{ }	List of fluid phases
phasePVTParaFiles	path_array	re- quired	Names of the files defining the parameters of the viscosity and density models



## 10.17 Element: CarmanKozenyPermeability

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes
particleDiameter	real64	required	Diameter of the spherical particles.
sphericity	real64	required	Sphericity of the particles.

## 10.18 Element: CellElementRegion

Name	Type	Default	Description
cellBlocks	string_array	{ }	(no description available)
coarseningRatio	real64	0	(no description available)
materialList	string_array	required	List of materials present in this region
meshBody	string		Mesh body that contains this region
name	string	required	A name is required for any non-unique nodes

## 10.19 Element: ChomboLO

Name	Type	Default	Description
beginCycle	real64	required	Cycle at which the coupling will commence.
childDirectory	string		Child directory path
inputPath	string	/IN- VALID_INPUT_PATH	Path at which the chombo to geosx file will be written.
name	string	required	A name is required for any non-unique nodes
outputPath	string	required	Path at which the geosx to chombo file will be written.
parallelThreads	integer	1	Number of plot files.
useChombo-Pressures	integer	0	True iff geosx should use the pressures chombo writes out.
waitForInput	integer	required	True iff geosx should wait for chombo to write out a file. When true the inputPath must be set.

## 10.20 Element: CompositeFunction

Name	Type	De- fault	Description
expression	string		Composite math expression
function-Names	string_array	{ }	List of source functions. The order must match the variableNames argument.
inputVar-Names	string_array	{ }	Name of fields are input to function.
name	string	re- quired	A name is required for any non-unique nodes
variable-Names	string_array	{ }	List of variables in expression



## 10.21 Element: CompositionalMultiphaseFVM

Name	Type	Default	Description
allowLocal-CompDen-sityChop-ping	in-te-ger	1	Flag indicating whether local (cell-wise) chopping of negative compositions is allowed
cflFactor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
discretiza-tion	string	re-quired	Name of discretization object (defined in the <a href="#">Numerical Methods</a> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <a href="#">Finite Element Discretization</a> should be specified. If this is a Finite Volume Method, the name of a <a href="#">Finite Volume Discretization</a> discretization should be specified.
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
inputFlux-Estimate	real64	1	Initial estimate of the input flux used only for residual scaling. This should be essentially equivalent to the input flux * dt.
isThermal	in-te-ger	0	Flag indicating whether the problem is thermal or not.
logLevel	in-te-ger	0	Log level
max-CompFrac-tionChange	real64	0.5	Maximum (absolute) change in a component fraction in a Newton iteration
maxRela-tivePres-sureChange	real64	0.5	Maximum (relative) change in pressure in a Newton iteration (expected value between 0 and 1)
maxRela-tiveTem-pera-tureChange	real64	0.5	Maximum (relative) change in temperature in a Newton iteration (expected value between 0 and 1)
name	string	re-quired	A name is required for any non-unique nodes
solution-ChangeScal-ingFactor	real64	0.5	Damping factor for solution change targets
target-PhaseVol-Fraction-ChangeIn-TimeStep	real64	0.2	Target (absolute) change in phase volume fraction in a time step
targetRe-gions	string array	re-quired	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
targetRel-ativePres-sureChangeIn-TimeStep	real64	0.2	Target (relative) change in pressure in a time step (expected value between 0 and 1)
targetRel-ativeTem-pera-tureChangeIn-TimeStep	real64	0.2	Target (relative) change in temperature in a time step (expected value between 0 and 1)
temperature	real64	re-quired	Temperature

## 10.22 Element: CompositionalMultiphaseFluid

Name	Type	Default	Description
componentAcentricFactor	real64_array	required	Component acentric factors
componentBinaryCoeff	real64_array2d	{{0}}	Table of binary interaction coefficients
componentCriticalPressure	real64_array	required	Component critical pressures
componentCriticalTemperature	real64_array	required	Component critical temperatures
componentMolarWeight	real64_array	required	Component molar weights
componentNames	string_array	required	List of component names
componentVolumeShift	real64_array	{0}	Component volume shifts
equationsOfState	string_array	required	List of equation of state types for each phase
name	string	required	A name is required for any non-unique nodes
phaseNames	string_array	required	List of fluid phases



## 10.23 Element: CompositionalMultiphaseHybridFVM

Name	Type	Default	Description
allowLocal-CompDen-sityChop-ping	in-te-ger	1	Flag indicating whether local (cell-wise) chopping of negative compositions is allowed
cflFactor	real64	0.5	Factor to apply to the CFL condition when calculating the maximum allowable time step. Values should be in the interval (0,1]
discretiza-tion	string	re-quired	Name of discretization object (defined in the <i>Numerical Methods</i> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <i>Finite Element Discretization</i> should be specified. If this is a Finite Volume Method, the name of a <i>Finite Volume Discretization</i> discretization should be specified.
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
inputFlux-Estimate	real64	1	Initial estimate of the input flux used only for residual scaling. This should be essentially equivalent to the input flux * dt.
isThermal	in-te-ger	0	Flag indicating whether the problem is thermal or not.
logLevel	in-te-ger	0	Log level
max-CompFrac-tionChange	real64	0.5	Maximum (absolute) change in a component fraction in a Newton iteration
maxRela-tivePres-sureChange	real64	0.5	Maximum (relative) change in pressure in a Newton iteration (expected value between 0 and 1)
maxRela-tiveTem-pera-tureChange	real64	0.5	Maximum (relative) change in temperature in a Newton iteration (expected value between 0 and 1)
name	string	re-quired	A name is required for any non-unique nodes
solution-ChangeScal-ingFactor	real64	0.5	Damping factor for solution change targets
target-PhaseVol-Fraction-ChangeIn-TimeStep	real64	0.2	Target (absolute) change in phase volume fraction in a time step
targetRe-gions	string array	re-quired	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
targetRel-ativePres-sureChangeIn-TimeStep	real64	0.2	Target (relative) change in pressure in a time step (expected value between 0 and 1)
targetRel-ativeTem-pera-tureChangeIn-TimeStep	real64	0.2	Target (relative) change in temperature in a time step (expected value between 0 and 1)
temperature	real64	re-quired	Temperature

## 10.24 Element: CompositionalMultiphaseReservoir

Name	Type	De- fault	Description
cflFactor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
flow-Solver-Name	string	re- quired	Name of the flow solver used by the coupled solver
ini- tialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
logLevel	in- te- ger	0	Log level
name	string	re- quired	A name is required for any non-unique nodes
targetRe- gions	string_array	re- quired	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
well-Solver-Name	string	re- quired	Name of the well solver used by the coupled solver
Linear-Solver-Parameters	node	unique	<i>Element: LinearSolverParameters</i>
Non-linear-Solver-Parameters	node	unique	<i>Element: NonlinearSolverParameters</i>

## 10.25 Element: CompositionalMultiphaseStatistics

Name	Type	Default	Description
computeCFLNumbers	integer	0	Flag to decide whether CFL numbers are computed or not
computeRegionStatistics	integer	1	Flag to decide whether region statistics are computed or not
flowSolverName	string	required	Name of the flow solver
logLevel	integer	0	Log level
name	string	required	A name is required for any non-unique nodes



## 10.26 Element: CompositionalMultiphaseWell

Name	Type	De- fault	Description
allowLocalCompDensityChopping	integer	1	Flag indicating whether local (cell-wise) chopping of negative compositions is allowed
cflFactor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
logLevel	integer	0	Log level
maxCompFractionChange	real64	1	Maximum (absolute) change in a component fraction between two Newton iterations
maxRelativePressureChange	real64	1	Maximum (relative) change in pressure between two Newton iterations (recommended with rate control)
name	string	required	A name is required for any non-unique nodes
targetRegions	string array	required	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
useMass	integer	0	Use mass formulation instead of molar
Linear-SolverParameters	node	unique	<a href="#">Element: LinearSolverParameters</a>
Non-linear-SolverParameters	node	unique	<a href="#">Element: NonlinearSolverParameters</a>
WellControls	node		<a href="#">Element: WellControls</a>

## 10.27 Element: CompressibleSinglePhaseFluid

Name	Type	Default	Description
compressibility	real64	0	Fluid compressibility
defaultDensity	real64	required	Default value for density.
defaultViscosity	real64	required	Default value for viscosity.
densityModelType	geosx_constitutive_ExponentialApproximationType		Type of density model. Valid options: * exponential * linear * quadratic
name	string	required	A name is required for any non-unique nodes
referenceDensity	real64	1000	Reference fluid density
referencePressure	real64	0	Reference pressure
referenceViscosity	real64	0.001	Reference fluid viscosity
viscosibility	real64	0	Fluid viscosity exponential coefficient
viscosityModelType	geosx_constitutive_ExponentialApproximationType		Type of viscosity model. Valid options: * exponential * linear * quadratic

## 10.28 Element: CompressibleSolidCarmanKozenyPermeability

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes
permeabilityModelName	string	required	Name of the permeability model.
porosityModelName	string	required	Name of the porosity model.
solidInternalEnergyModelName	string		Name of the solid internal energy model.
solidModelName	string	required	Name of the solid model.

## 10.29 Element: CompressibleSolidConstantPermeability

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes
permeabilityModelName	string	required	Name of the permeability model.
porosityModelName	string	required	Name of the porosity model.
solidInternalEnergyModelName	string		Name of the solid internal energy model.
solidModelName	string	required	Name of the solid model.

## 10.30 Element: CompressibleSolidParallelPlatesPermeability

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes
permeabilityModelName	string	required	Name of the permeability model.
porosityModelName	string	required	Name of the porosity model.
solidInternalEnergyModelName	string		Name of the solid internal energy model.
solidModelName	string	required	Name of the solid model.

## 10.31 Element: CompressibleSolidSlipDependentPermeability

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes
permeabilityModelName	string	required	Name of the permeability model.
porosityModelName	string	required	Name of the porosity model.
solidInternalEnergyModelName	string		Name of the solid internal energy model.
solidModelName	string	required	Name of the solid model.

## 10.32 Element: ConstantPermeability

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes
permeabilityComponents	RTensor	required	xx, yy and zz components of a diagonal permeability tensor.

## 10.33 Element: Constitutive

Name	Type	Default	Description
BiotPorosity	node		<i>Element: BiotPorosity</i>
BlackOilFluid	node		<i>Element: BlackOilFluid</i>
BrooksCoreyBakerRelativePermeability	node		<i>Element: BrooksCoreyBakerRelativePermeability</i>
BrooksCoreyCapillaryPressure	node		<i>Element: BrooksCoreyCapillaryPressure</i>

Continued on next page

Table 10.1 – continued from previous page

Name	Type	Default	Description
BrooksCoreyRelativePermeability	node		<i>Element: BrooksCoreyRelativePermeability</i>
CO2BrineEzrokhiFluid	node		<i>Element: CO2BrineEzrokhiFluid</i>
CO2BrineEzrokhiThermalFluid	node		<i>Element: CO2BrineEzrokhiThermalFluid</i>
CO2BrinePhillipsFluid	node		<i>Element: CO2BrinePhillipsFluid</i>
CO2BrinePhillipsThermalFluid	node		<i>Element: CO2BrinePhillipsThermalFluid</i>
CarmanKozenyPermeability	node		<i>Element: CarmanKozenyPermeability</i>
CompositionalMultiphaseFluid	node		<i>Element: CompositionalMultiphaseFluid</i>
CompressibleSinglePhaseFluid	node		<i>Element: CompressibleSinglePhaseFluid</i>
CompressibleSolidCarmanKozenyPermeability	node		<i>Element: CompressibleSolidCarmanKozenyPermeability</i>
CompressibleSolidConstantPermeability	node		<i>Element: CompressibleSolidConstantPermeability</i>
CompressibleSolidParallelPlatesPermeability	node		<i>Element: CompressibleSolidParallelPlatesPermeability</i>
CompressibleSolidSlipDependentPermeability	node		<i>Element: CompressibleSolidSlipDependentPermeability</i>
ConstantPermeability	node		<i>Element: ConstantPermeability</i>
Coulomb	node		<i>Element: Coulomb</i>
DamageElasticIsotropic	node		<i>Element: DamageElasticIsotropic</i>
DamageSpectralElasticIsotropic	node		<i>Element: DamageSpectralElasticIsotropic</i>
DamageVolDevElasticIsotropic	node		<i>Element: DamageVolDevElasticIsotropic</i>
DeadOilFluid	node		<i>Element: DeadOilFluid</i>
DelftEgg	node		<i>Element: DelftEgg</i>
DruckerPrager	node		<i>Element: DruckerPrager</i>
ElasticIsotropic	node		<i>Element: ElasticIsotropic</i>
ElasticIsotropicPressureDependent	node		<i>Element: ElasticIsotropicPressureDependent</i>
ElasticOrthotropic	node		<i>Element: ElasticOrthotropic</i>
ElasticTransverseIsotropic	node		<i>Element: ElasticTransverseIsotropic</i>
ExtendedDruckerPrager	node		<i>Element: ExtendedDruckerPrager</i>
FrictionlessContact	node		<i>Element: FrictionlessContact</i>
JFunctionCapillaryPressure	node		<i>Element: JFunctionCapillaryPressure</i>
ModifiedCamClay	node		<i>Element: ModifiedCamClay</i>
MultiPhaseConstantThermalConductivity	node		<i>Element: MultiPhaseConstantThermalConductivity</i>
MultiPhaseVolumeWeightedThermalConductivity	node		<i>Element: MultiPhaseVolumeWeightedThermalConductivity</i>
NullModel	node		<i>Element: NullModel</i>
ParallelPlatesPermeability	node		<i>Element: ParallelPlatesPermeability</i>
ParticleFluid	node		<i>Element: ParticleFluid</i>
PermeabilityBase	node		<i>Element: PermeabilityBase</i>
PorousDelftEgg	node		<i>Element: PorousDelftEgg</i>
PorousDruckerPrager	node		<i>Element: PorousDruckerPrager</i>
PorousElasticIsotropic	node		<i>Element: PorousElasticIsotropic</i>
PorousElasticOrthotropic	node		<i>Element: PorousElasticOrthotropic</i>
PorousElasticTransverseIsotropic	node		<i>Element: PorousElasticTransverseIsotropic</i>
PorousExtendedDruckerPrager	node		<i>Element: PorousExtendedDruckerPrager</i>
PorousModifiedCamClay	node		<i>Element: PorousModifiedCamClay</i>
PressurePorosity	node		<i>Element: PressurePorosity</i>
ProppantPermeability	node		<i>Element: ProppantPermeability</i>
ProppantPorosity	node		<i>Element: ProppantPorosity</i>
ProppantSlurryFluid	node		<i>Element: ProppantSlurryFluid</i>
ProppantSolidProppantPermeability	node		<i>Element: ProppantSolidProppantPermeability</i>
SinglePhaseConstantThermalConductivity	node		<i>Element: SinglePhaseConstantThermalConductivity</i>
SlipDependentPermeability	node		<i>Element: SlipDependentPermeability</i>
SolidInternalEnergy	node		<i>Element: SolidInternalEnergy</i>

Continued on next page

Table 10.1 – continued from previous page

Name	Type	Default	Description
TableCapillaryPressure	node		<i>Element: TableCapillaryPressure</i>
TableRelativePermeability	node		<i>Element: TableRelativePermeability</i>
TableRelativePermeabilityHysteresis	node		<i>Element: TableRelativePermeabilityHysteresis</i>
ThermalCompressibleSinglePhaseFluid	node		<i>Element: ThermalCompressibleSinglePhaseFluid</i>
VanGenuchtenBakerRelativePermeability	node		<i>Element: VanGenuchtenBakerRelativePermeability</i>
VanGenuchtenCapillaryPressure	node		<i>Element: VanGenuchtenCapillaryPressure</i>

## 10.34 Element: Coulomb

Name	Type	De- fault	Description
aper- tureTable- Name	string	re- quired	Name of the aperture table
aper- ture- Toler- ance	real64	1e- 09	Value to be used to avoid floating point errors in expressions involving aperture. For example in the case of dividing by the actual aperture (not the effective aperture that results from the aperture function) this value may be used to avoid the 1/0 error. Note that this value may have some physical significance in its usage, as it may be used to smooth out highly nonlinear behavior associated with 1/0 in addition to avoiding the 1/0 error.
cohe- sion	real64	re- quired	Cohesion
dis- place- men- tJumpThresh- old	real64	2.2204e- 16	Threshold valued to determine whether a fracture is open or not.
fric- tion- Co- effi- cient	real64	re- quired	Friction coefficient
name	string	re- quired	A name is required for any non-unique nodes
penal- tyS- tiff- ness	real64	0	Value of the penetration penalty stiffness. Units of Pressure/length
shearS- tiff- ness	real64	0	Value of the shear elastic stiffness. Units of Pressure/length

## 10.35 Element: Cylinder

Name	Type	Default	Description
innerRadius	real64	-1	Inner radius of the anulus
name	string	required	A name is required for any non-unique nodes
point1	R1Tensor	required	Center point of one (upper or lower) face of the cylinder
point2	R1Tensor	required	Center point of the other face of the cylinder
radius	real64	required	Radius of the cylinder

## 10.36 Element: DamageElasticIsotropic

Name	Type	Default	Description
compressiveStrength	real64	0	Compressive strength from the uniaxial compression test
criticalFractureEnergy	real64	required	Critical fracture energy
criticalStrainEnergy	real64	required	Critical stress in a 1d tension test
defaultBulkModulus	real64	-1	Default Bulk Modulus Parameter
defaultDensity	real64	required	Default Material Density
defaultPoissonRatio	real64	-1	Default Poisson's Ratio
defaultShearModulus	real64	-1	Default Shear Modulus Parameter
defaultYoungModulus	real64	-1	Default Young's Modulus
degradationLowerLimit	real64	0	The lower limit of the degradation function
deltaCoefficient	real64	-1	Coefficient in the calculation of the external driving force
extDrivingForceFlag	integer	0	Whether to have external driving force. Can be 0 or 1
lengthScale	real64	required	Length scale l in the phase-field equation
name	string	required	A name is required for any non-unique nodes
tensileStrength	real64	0	Tensile strength from the uniaxial tension test

## 10.37 Element: DamageSpectralElasticIsotropic

Name	Type	Default	Description
compressiveStrength	real64	0	Compressive strength from the uniaxial compression test
criticalFractureEnergy	real64	required	Critical fracture energy
criticalStrainEnergy	real64	required	Critical stress in a 1d tension test
defaultBulkModulus	real64	-1	Default Bulk Modulus Parameter
defaultDensity	real64	required	Default Material Density
defaultPoissonRatio	real64	-1	Default Poisson's Ratio
defaultShearModulus	real64	-1	Default Shear Modulus Parameter
defaultYoungModulus	real64	-1	Default Young's Modulus
degradationLowerLimit	real64	0	The lower limit of the degradation function
deltaCoefficient	real64	-1	Coefficient in the calculation of the external driving force
extDrivingForceFlag	integer	0	Whether to have external driving force. Can be 0 or 1
lengthScale	real64	required	Length scale l in the phase-field equation
name	string	required	A name is required for any non-unique nodes
tensileStrength	real64	0	Tensile strength from the uniaxial tension test

## 10.38 Element: DamageVolDevElasticIsotropic

Name	Type	Default	Description
compressiveStrength	real64	0	Compressive strength from the uniaxial compression test
criticalFractureEnergy	real64	required	Critical fracture energy
criticalStrainEnergy	real64	required	Critical stress in a 1d tension test
defaultBulkModulus	real64	-1	Default Bulk Modulus Parameter
defaultDensity	real64	required	Default Material Density
defaultPoissonRatio	real64	-1	Default Poisson's Ratio
defaultShearModulus	real64	-1	Default Shear Modulus Parameter
defaultYoungModulus	real64	-1	Default Young's Modulus
degradationLowerLimit	real64	0	The lower limit of the degradation function
deltaCoefficient	real64	-1	Coefficient in the calculation of the external driving force
extDrivingForceFlag	integer	0	Whether to have external driving force. Can be 0 or 1
lengthScale	real64	required	Length scale $l$ in the phase-field equation
name	string	required	A name is required for any non-unique nodes
tensileStrength	real64	0	Tensile strength from the uniaxial tension test

## 10.39 Element: DeadOilFluid

Name	Type	Default	Description
componentMolarWeight	real64_array	required	Component molar weights
componentNames	string_array	{}	List of component names
hydrocarbonFormationVolumeFactorTableNames	string_array	{}	<p>List of formation volume factor TableFunction names from the Functions block.</p> <p>The user must provide one TableFunction per hydrocarbon phase, in the order provided in “phaseNames”.</p> <p>For instance, if “oil” is before “gas” in “phaseNames”, the table order should be: oilTableName, gasTableName</p>
hydrocarbonViscosityTableNames	string_array	{}	<p>List of viscosity TableFunction names from the Functions block.</p> <p>The user must provide one TableFunction per hydrocarbon phase, in the order provided in “phaseNames”.</p> <p>For instance, if “oil” is before “gas” in “phaseNames”, the table order should be: oilTableName, gasTableName</p>
name	string	required	A name is required for any non-unique nodes
phaseNames	string_array	required	List of fluid phases
surfaceDensities	real64_array	required	List of surface mass densities for each phase
tableFiles	path_array	{}	List of filenames with input PVT tables (one per phase)
waterCompressibility	real64	0	Water compressibility
waterFormationVolumeFactor	real64	0	Water formation volume factor
waterReferencePressure	real64	0	Water reference pressure
waterViscosity	real64	0	Water viscosity



## 10.40 Element: DelftEgg

Name	Type	Default	Description
defaultBulkModulus	real64	-1	Default Bulk Modulus Parameter
defaultCslSlope	real64	1	Slope of the critical state line
defaultDensity	real64	required	Default Material Density
defaultPoissonRatio	real64	-1	Default Poisson's Ratio
defaultPreConsolidationPressure	real64	-1.5	Initial preconsolidation pressure
defaultRecompressionIndex	real64	0.002	Recompression Index
defaultShapeParameter	real64	1	Shape parameter for the yield surface
defaultShearModulus	real64	-1	Default Shear Modulus Parameter
defaultVirginCompressionIndex	real64	0.005	Virgin compression index
defaultYoungModulus	real64	-1	Default Young's Modulus
name	string	required	A name is required for any non-unique nodes

## 10.41 Element: Dirichlet

Name	Type	De- fault	Description
bcApplicationTable- Name	string		Name of table that specifies the on/off application of the boundary condition.
beginTime	real64	- 1e+99	Time at which the boundary condition will start being applied.
component	integer	-1	Component of field (if tensor) to apply boundary condition to.
direction	R1Tensor	{0,0,0}	Direction to apply boundary condition to.
endTime	real64	1e+99	Time at which the boundary condition will stop being applied.
fieldName	string		Name of field that boundary condition is applied to.
functionName	string		Name of function that specifies variation of the boundary condition.
initialCondition	integer	0	Boundary condition is applied as an initial condition.
logLevel	integer	0	Log level
name	string	re- quired	A name is required for any non-unique nodes
objectPath	string		Path to the target field
scale	real64	0	Scale factor for value of the boundary condition.
setNames	string_array	re- quired	Name of sets that boundary condition is applied to.

## 10.42 Element: DruckerPrager

Name	Type	Default	Description
defaultBulkModulus	real64	-1	Default Bulk Modulus Parameter
defaultCohesion	real64	0	Initial cohesion
defaultDensity	real64	required	Default Material Density
defaultDilationAngle	real64	30	Dilation angle (degrees)
defaultFrictionAngle	real64	30	Friction angle (degrees)
defaultHardeningRate	real64	0	Cohesion hardening/softening rate
defaultPoissonRatio	real64	-1	Default Poisson's Ratio
defaultShearModulus	real64	-1	Default Shear Modulus Parameter
defaultYoungModulus	real64	-1	Default Young's Modulus
name	string	required	A name is required for any non-unique nodes

## 10.43 Element: ElasticIsotropic

Name	Type	Default	Description
defaultBulkModulus	real64	-1	Default Bulk Modulus Parameter
defaultDensity	real64	required	Default Material Density
defaultPoissonRatio	real64	-1	Default Poisson's Ratio
defaultShearModulus	real64	-1	Default Shear Modulus Parameter
defaultYoungModulus	real64	-1	Default Young's Modulus
name	string	required	A name is required for any non-unique nodes

## 10.44 Element: ElasticIsotropicPressureDependent

Name	Type	Default	Description
defaultDensity	real64	required	Default Material Density
defaultRecompressionIndex	real64	0.002	Recompression Index
defaultRefPressure	real64	-1	Reference Pressure
defaultRefStrainVol	real64	0	Reference Volumetric Strain
defaultShearModulus	real64	-1	Elastic Shear Modulus Parameter
name	string	required	A name is required for any non-unique nodes

## 10.45 Element: ElasticOrthotropic

Name	Type	Default	Description
defaultC11	real64	-1	Default C11 Component of Voigt Stiffness Tensor
defaultC12	real64	-1	Default C12 Component of Voigt Stiffness Tensor
defaultC13	real64	-1	Default C13 Component of Voigt Stiffness Tensor
defaultC22	real64	-1	Default C22 Component of Voigt Stiffness Tensor
defaultC23	real64	-1	Default C23 Component of Voigt Stiffness Tensor
defaultC33	real64	-1	Default C33 Component of Voigt Stiffness Tensor
defaultC44	real64	-1	Default C44 Component of Voigt Stiffness Tensor
defaultC55	real64	-1	Default C55 Component of Voigt Stiffness Tensor
defaultC66	real64	-1	Default C66 Component of Voigt Stiffness Tensor
defaultDensity	real64	required	Default Material Density
defaultE1	real64	-1	Default Young's Modulus E1
defaultE2	real64	-1	Default Young's Modulus E2
defaultE3	real64	-1	Default Young's Modulus E3
defaultG12	real64	-1	Default Shear Modulus G12
defaultG13	real64	-1	Default Shear Modulus G13
defaultG23	real64	-1	Default Shear Modulus G23
defaultNu12	real64	-1	Default Poission's Ratio Nu12
defaultNu13	real64	-1	Default Poission's Ratio Nu13
defaultNu23	real64	-1	Default Poission's Ratio Nu23
name	string	required	A name is required for any non-unique nodes

## 10.46 Element: ElasticSEM

Name	Type	De- fault	Description
cflFactor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
discretization	string	required	Name of discretization object (defined in the <i>Numerical Methods</i> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <i>Finite Element Discretization</i> should be specified. If this is a Finite Volume Method, the name of a <i>Finite Volume Discretization</i> discretization should be specified.
dtSeismo-Trace	real64	0	Time step for output pressure at receivers
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
logLevel	integer	0	Log level
name	string	required	A name is required for any non-unique nodes
output-Seismo-Trace	integer	0	Flag that indicates if we write the seismo trace in a file .txt, 0 no output, 1 otherwise
receiver-Coordinates	real64_array	2, required	Coordinates (x,y,z) of the receivers
rickerOrder	integer	2	Flag that indicates the order of the Ricker to be used 0, 1 or 2. Order 2 by default
source-Coordinates	real64_array	2, required	Coordinates (x,y,z) of the sources
targetRegions	string_array	required	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
time-Source-Frequency	real32	required	Central frequency for the time source
Linear-Solver-Parameters	node	unique	<i>Element: LinearSolverParameters</i>
Non-linear-Solver-Parameters	node	unique	<i>Element: NonlinearSolverParameters</i>

## 10.47 Element: ElasticTransverselsotropic

Name	Type	Default	Description
defaultC11	real64	-1	Default Stiffness Parameter C11
defaultC13	real64	-1	Default Stiffness Parameter C13
defaultC33	real64	-1	Default Stiffness Parameter C33
defaultC44	real64	-1	Default Stiffness Parameter C44
defaultC66	real64	-1	Default Stiffness Parameter C66
defaultDensity	real64	required	Default Material Density
defaultPoissonRatioAxialTransverse	real64	-1	Default Axial-Transverse Poisson's Ratio
defaultPoissonRatioTransverse	real64	-1	Default Transverse Poisson's Ratio
defaultShearModulusAxialTransverse	real64	-1	Default Axial-Transverse Shear Modulus
defaultYoungModulusAxial	real64	-1	Default Axial Young's Modulus
defaultYoungModulusTransverse	real64	-1	Default Transverse Young's Modulus
name	string	required	A name is required for any non-unique nodes

## 10.48 Element: ElementRegions

Name	Type	Default	Description
CellElementRegion	node		<i>Element: CellElementRegion</i>
SurfaceElementRegion	node		<i>Element: SurfaceElementRegion</i>
WellElementRegion	node		<i>Element: WellElementRegion</i>

## 10.49 Element: EmbeddedSurfaceGenerator

Name	Type	Default	Description
cflFactor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
discretization	string	required	Name of discretization object (defined in the <i>Numerical Methods</i> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <i>Finite Element Discretization</i> should be specified. If this is a Finite Volume Method, the name of a <i>Finite Volume Discretization</i> discretization should be specified.
fractureRegion	string	FractureRegion	(no description available)
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
logLevel	integer	0	Log level
mpiCommOrder	integer	0	Flag to enable MPI consistent communication ordering
name	string	required	A name is required for any non-unique nodes
targetRegions	string array	required	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
LinearSolverParameters	node	unique	<i>Element: LinearSolverParameters</i>
NonlinearSolverParameters	node	unique	<i>Element: NonlinearSolverParameters</i>

## 10.50 Element: Events

Name	Type	Default	Description
logLevel	integer	0	Log level
maxCycle	integer	2147483647	Maximum simulation cycle for the global event loop.
maxTime	real64	1.79769e+308	Maximum simulation time for the global event loop.
minTime	real64	0	Start simulation time for the global event loop.
HaltEvent	node		<i>Element: HaltEvent</i>
PeriodicEvent	node		<i>Element: PeriodicEvent</i>
SoloEvent	node		<i>Element: SoloEvent</i>

## 10.51 Element: ExtendedDruckerPrager

Name	Type	Default	Description
defaultBulkModulus	real64	-1	Default Bulk Modulus Parameter
defaultCohesion	real64	0	Initial cohesion
defaultDensity	real64	re- quired	Default Material Density
defaultDilationRatio	real64	1	Dilation ratio [0,1] (ratio = $\tan \text{dilationAngle} / \tan \text{frictionAngle}$ )
defaultHardening	real64	0	Hardening parameter (hardening rate is faster for smaller values)
defaultInitialFrictionAngle	real64	30	Initial friction angle (degrees)
defaultPoissonRatio	real64	-1	Default Poisson's Ratio
defaultResidualFrictionAngle	real64	30	Residual friction angle (degrees)
defaultShearModulus	real64	-1	Default Shear Modulus Parameter
defaultYoungModulus	real64	-1	Default Young's Modulus
name	string	re- quired	A name is required for any non-unique nodes

## 10.52 Element: FieldSpecification

Name	Type	Default	Description
bcApplicationTableName	string		Name of table that specifies the on/off application of the boundary condition.
beginTime	real64	- 1e+99	Time at which the boundary condition will start being applied.
component	integer	-1	Component of field (if tensor) to apply boundary condition to.
direction	R1Tensor	{0,0,0}	Direction to apply boundary condition to.
endTime	real64	1e+99	Time at which the boundary condition will stop being applied.
fieldName	string		Name of field that boundary condition is applied to.
functionName	string		Name of function that specifies variation of the boundary condition.
initialCondition	integer	0	Boundary condition is applied as an initial condition.
logLevel	integer	0	Log level
name	string	re- quired	A name is required for any non-unique nodes
objectPath	string		Path to the target field
scale	real64	0	Scale factor for value of the boundary condition.
setNames	string_array	re- quired	Name of sets that boundary condition is applied to.

## 10.53 Element: FieldSpecifications

Name	Type	Default	Description
Aquifer	node		<i>Element: Aquifer</i>
Dirichlet	node		<i>Element: Dirichlet</i>
FieldSpecification	node		<i>Element: FieldSpecification</i>
HydrostaticEquilibrium	node		<i>Element: HydrostaticEquilibrium</i>
PML	node		<i>Element: PML</i>
SourceFlux	node		<i>Element: SourceFlux</i>
Traction	node		<i>Element: Traction</i>

## 10.54 Element: File

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes

## 10.55 Element: FiniteElementSpace

Name	Type	De- fault	Description
formula- tion	string	de- fault	Specifier to indicate any specialized formulations. For instance, one of the many enhanced assumed strain methods of the Hexahedron parent shape would be indicated here
name	string	re- quired	A name is required for any non-unique nodes
order	in- te- ger	re- quired	The order of the finite element basis.
useVir- tualEle- ments	in- te- ger	0	Specifier to indicate whether to force the use of VEM

## 10.56 Element: FiniteElements

Name	Type	Default	Description
FiniteElementSpace	node		<i>Element: FiniteElementSpace</i>
LinearSolverParameters	node	unique	<i>Element: LinearSolverParameters</i>
NonlinearSolverParameters	node	unique	<i>Element: NonlinearSolverParameters</i>



## 10.57 Element: FiniteVolume

Name	Type	Default	Description
HybridMimeticDiscretization	node		<i>Element: HybridMimeticDiscretization</i>
TwoPointFluxApproximation	node		<i>Element: TwoPointFluxApproximation</i>

## 10.58 Element: FlowProppantTransport

Name	Type	De- fault	Description
cflFac- tor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
flow- Solver- Name	string	re- quired	Name of the flow solver used by the coupled solver
ini- tialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
logLevel	in- te- ger	0	Log level
name	string	re- quired	A name is required for any non-unique nodes
prop- pant- Solver- Name	string	re- quired	Name of the proppant solver used by the coupled solver
targe- tRe- gions	string array	re- quired	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
Linear- Solver- Param- eters	node	unique	<i>Element: LinearSolverParameters</i>
Non- linear- Solver- Param- eters	node	unique	<i>Element: NonlinearSolverParameters</i>

## 10.59 Element: FrictionlessContact

Name	Type	Default	Description
apertureTable-Name	string	required	Name of the aperture table
aperture-Tolerance	real64	1e-09	Value to be used to avoid floating point errors in expressions involving aperture. For example in the case of dividing by the actual aperture (not the effective aperture that results from the aperture function) this value may be used to avoid the 1/0 error. Note that this value may have some physical significance in its usage, as it may be used to smooth out highly nonlinear behavior associated with 1/0 in addition to avoiding the 1/0 error.
displacement-JumpThreshold	real64	2.2204e-16	Threshold valued to determine whether a fracture is open or not.
name	string	required	A name is required for any non-unique nodes
penaltyStiffness	real64	0	Value of the penetration penalty stiffness. Units of Pressure/length
shearStiffness	real64	0	Value of the shear elastic stiffness. Units of Pressure/length

## 10.60 Element: Functions

Name	Type	Default	Description
CompositeFunction	node		<i>Element: CompositeFunction</i>
MultivariableTableFunction	node		<i>Element: MultivariableTableFunction</i>
SymbolicFunction	node		<i>Element: SymbolicFunction</i>
TableFunction	node		<i>Element: TableFunction</i>

## 10.61 Element: Geometry

Name	Type	Default	Description
BoundedPlane	node		<i>Element: BoundedPlane</i>
Box	node		<i>Element: Box</i>
Cylinder	node		<i>Element: Cylinder</i>
ThickPlane	node		<i>Element: ThickPlane</i>

## 10.62 Element: HaltEvent

Name	Type	De- fault	Description
beginTime	real64	0	Start time of this event.
endTime	real64	1e+100	End time of this event.
finalDtStretch	real64	0.001	Allow the final dt request for this event to grow by this percentage to match the endTime exactly.
forceDt	real64	-1	While active, this event will request this timestep value (ignoring any children/targets requests).
logLevel	integer	0	Log level
maxEventDt	real64	-1	While active, this event will request a timestep $\leq$ this value (depending upon any child/target requests).
maxRuntime	real64	required	The maximum allowable runtime for the job.
name	string	required	A name is required for any non-unique nodes
target	string		Name of the object to be executed when the event criteria are met.
targetExact-StartStop	integer	1	If this option is set, the event will reduce its timestep requests to match any specified beginTime/endTimes exactly.
HaltEvent	node		<i>Element: HaltEvent</i>
PeriodicEvent	node		<i>Element: PeriodicEvent</i>
SoloEvent	node		<i>Element: SoloEvent</i>

## 10.63 Element: HybridMimeticDiscretization

Name	Type	Default	Description
innerProductType	string	required	Type of inner product used in the hybrid FVM solver
name	string	required	A name is required for any non-unique nodes



## 10.64 Element: Hydrofracture

Name	Type	Default	Description
cflFactor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
contactRelationName	string	required	Name of contact relation to enforce constraints on fracture boundary.
couplingTypeOption	geosx_HydrofractureSolver_CouplingTypeOption		Coupling method. Valid options: * FIM * SIM_FixedStress
flowSolverName	string	required	Name of the flow solver used by the coupled solver
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
logLevel	integer	0	Log level
maxNumResolves	integer	10	Value to indicate how many resolves may be executed to perform surface generation after the execution of flow and mechanics solver.
name	string	required	A name is required for any non-unique nodes
solidSolverName	string	required	Name of the solid solver used by the coupled solver
surfaceGeneratorName	string	required	Name of the surface generator to use in the hydrofracture solver
targetRegions	string_array	required	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
LinearSolverParameters	node	unique	<a href="#">Element: LinearSolverParameters</a>
NonlinearSolverParameters	node	unique	<a href="#">Element: Nonlinear-SolverParameters</a>

## 10.65 Element: HydrostaticEquilibrium

Name	Type	De- fault	Description
bcApplicationTableName	string		Name of table that specifies the on/off application of the boundary condition.
beginTime	real64	- 1e+99	Time at which the boundary condition will start being applied.
componentFractionVsElevationTableNames	string_array		Names of the tables specifying the (component fraction vs elevation) relationship for each component
componentNames	string_array		Names of the fluid components
datumElevation	real64	re- quired	Datum elevation [m]
datumPressure	real64	re- quired	Datum pressure [Pa]
direction	R1Tensor	{0,0,0}	Direction to apply boundary condition to.
elevationIncrementInHydrostaticPressureTable	real64	0.6096	Elevation increment [m] in the hydrostatic pressure table constructed internally
endTime	real64	1e+99	Time at which the boundary condition will stop being applied.
equilibrationTolerance	real64	0.001	Tolerance in the fixed-point iteration scheme used for hydrostatic initialization
functionName	string		Name of function that specifies variation of the boundary condition.
initialPhaseName	string		Name of the phase initially saturating the reservoir
logLevel	integer	0	Log level
maxNumberOfEquilibrationIterations	integer	5	Maximum number of equilibration iterations
name	string	re- quired	A name is required for any non-unique nodes
objectPath	string		Path to the target field
scale	real64	0	Scale factor for value of the boundary condition.
temperatureVsElevationTableName	string		Name of the table specifying the (temperature [K] vs elevation) relationship

## 10.66 Element: Included

Name	Type	Default	Description
File	node		<i>Element: File</i>

## 10.67 Element: InternalMesh

Name	Type	De- fault	Description
cellBlock-Names	string_array	re- quired	Names of each mesh block
element-Types	string_array	re- quired	Element types of each mesh block
name	string	re- quired	A name is required for any non-unique nodes
nx	inte- ger_array	re- quired	Number of elements in the x-direction within each mesh block
ny	inte- ger_array	re- quired	Number of elements in the y-direction within each mesh block
nz	inte- ger_array	re- quired	Number of elements in the z-direction within each mesh block
positionTol- erance	real64	1e-10	A position tolerance to verify if a node belong to a nodeset
trianglePat- tern	integer	0	Pattern by which to decompose the hex mesh into wedges
xBias	real64_array	{1}	Bias of element sizes in the x-direction within each mesh block ( $dx\_left=(1+b)*L/N$ , $dx\_right=(1-b)*L/N$ )
xCoords	real64_array	re- quired	x-coordinates of each mesh block vertex
yBias	real64_array	{1}	Bias of element sizes in the y-direction within each mesh block ( $dy\_left=(1+b)*L/N$ , $dy\_right=(1-b)*L/N$ )
yCoords	real64_array	re- quired	y-coordinates of each mesh block vertex
zBias	real64_array	{1}	Bias of element sizes in the z-direction within each mesh block ( $dz\_left=(1+b)*L/N$ , $dz\_right=(1-b)*L/N$ )
zCoords	real64_array	re- quired	z-coordinates of each mesh block vertex

## 10.68 Element: InternalWell

Name	Type	De- fault	Description
logLevel	integer	0	Log level
meshName	string	re- quired	Name of the reservoir mesh associated with this well
minEle- mentLength	real64	0.001	Minimum length of a well element, computed as (segment length / number of elements per segment ) [m]
minSeg- mentLength	real64	0.01	Minimum length of a well segment [m]
name	string	re- quired	A name is required for any non-unique nodes
numEle- mentsPerSeg- ment	integer	re- quired	Number of well elements per polyline segment
polylineNode- Coords	real64_array2d	re- quired	Physical coordinates of the well polyline nodes
polylineSeg- mentConn	globalIn- dex_array2d	re- quired	Connectivity of the polyline segments
radius	real64	re- quired	Radius of the well [m]
wellControl- sName	string	re- quired	Name of the set of constraints associated with this well
wellRegion- Name	string	re- quired	Name of the well element region
Perforation	node		<i>Element: Perforation</i>



## 10.69 Element: InternalWellbore

Name	Type	De- fault	Description
autoSpaceRadialElems	real64_array	1}	Automatically set number and spacing of elements in the radial direction. This overrides the values of nr! Value in each block indicates factor to scale the radial increment. Larger numbers indicate larger radial elements.
cartesianMappingInnerRadius	real64	1e+99	If using a Cartesian aligned outer boundary, this is inner radius at which to start the mapping.
cellBlockNames	string_array	required	Names of each mesh block
elementTypes	string_array	required	Element types of each mesh block
hardRadialCoords	real64_array	required	Sets the radial spacing to specified values
name	string	required	A name is required for any non-unique nodes
nr	integer_array	required	Number of elements in the radial direction
nt	integer_array	required	Number of elements in the tangent direction
nz	integer_array	required	Number of elements in the z-direction within each mesh block
positionTolerance	real64	1e-10	A position tolerance to verify if a node belong to a nodeset
rBias	real64_array	0.8}	Bias of element sizes in the radial direction
radius	real64_array	required	Wellbore radius
theta	real64_array	required	Tangent angle defining geometry size: 90 for quarter, 180 for half and 360 for full wellbore geometry
trajectory	real64_array	required	Coordinates defining the wellbore trajectory
trianglePattern	integer	0	Pattern by which to decompose the hex mesh into wedges
useCartesianOuterBoundary	integer	1000000	Enforce a Cartesian aligned outer boundary on the outer block starting with the radial block specified in this value
xBias	real64_array	required	Bias of element sizes in the x-direction within each mesh block ( $dx\_left=(1+b)*L/N$ , $dx\_right=(1-b)*L/N$ )
yBias	real64_array	required	Bias of element sizes in the y-direction within each mesh block ( $dy\_left=(1+b)*L/N$ , $dx\_right=(1-b)*L/N$ )
zBias	real64_array	required	Bias of element sizes in the z-direction within each mesh block ( $dz\_left=(1+b)*L/N$ , $dz\_right=(1-b)*L/N$ )
zCoords	real64_array	required	z-coordinates of each mesh block vertex



## 10.70 Element: JFunctionCapillaryPressure

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes
nonWettingIntermediateJFunctionTableName	string		<p>J-function table (dimensionless) for the pair (non-wetting phase, intermediate phase)</p> <p>Note that this input is only used for three-phase flow.</p> <p>If you want to do a two-phase simulation, please use instead wettingNonWettingJFunctionTableName to specify the table names.</p>
nonWettingIntermediateSurfaceTension	real64	0	<p>Surface tension [N/m] for the pair (non-wetting phase, intermediate phase)</p> <p>If you have a value in [dyne/cm], divide it by 1000 to obtain the value in [N/m]</p> <p>Note that this input is only used for three-phase flow.</p> <p>If you want to do a two-phase simulation, please use instead wettingNonWettingSurfaceTension to specify the surface tensions.</p>
permeabilityDirection	geosx_constitutive_JFunctionCapillaryPressure_PermeabilityDirection		<p>Permeability direction. Options are:</p> <p>XY - use the average of the permeabilities in the x and y directions,</p> <p>X - only use the permeability in the x direction,</p> <p>Y - only use the permeability in the y direction,</p>
<b>10.70. Element: JFunctionCapillaryPressure</b>			<p>Z - only use the permeability in the z direction.</p>
permeabilityExponent	real64	0.5	Permeability exponent

## 10.71 Element: LagrangianContact

Name	Type	De- fault	Description
cflFactor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
contactRelationName	string	required	Name of contact relation to enforce constraints on fracture boundary.
discretization	string	required	Name of discretization object (defined in the <a href="#">Numerical Methods</a> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <a href="#">Finite Element Discretization</a> should be specified. If this is a Finite Volume Method, the name of a <a href="#">Finite Volume Discretization</a> discretization should be specified.
fractureRegionName	string	required	Name of the fracture region.
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
logLevel	integer	0	Log level
name	string	required	A name is required for any non-unique nodes
solidSolverName	string	required	Name of the solid mechanics solver in the rock matrix
stabilizationName	string	required	Name of the stabilization to use in the lagrangian contact solver
targetRegions	string_array	required	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
LinearSolverParameters	node	unique	<a href="#">Element: LinearSolverParameters</a>
NonlinearSolverParameters	node	unique	<a href="#">Element: NonlinearSolverParameters</a>



## 10.72 Element: LaplaceFEM

Name	Type	Default	Description
cflFactor	real64	0.5	Factor to apply to the CFL condition when calculating the maximum allowable time step. Values should be in the interval (0,1]
discretization	string	required	Name of discretization object (defined in the <i>Numerical Methods</i> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <i>Finite Element Discretization</i> should be specified. If this is a Finite Volume Method, the name of a <i>Finite Volume Discretization</i> should be specified.
fieldName	string	required	Name of field variable
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
logLevel	integer	0	Log level
name	string	required	A name is required for any non-unique nodes
targetRegions	string_array	required	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
timeIntegrationOption	geosx_LaplaceBaseH1_TimeIntegrationOption	required	Time integration method. Options are: * SteadyState * ImplicitTransient
LinearSolverParameters	node	unique	<i>Element: LinearSolverParameters</i>
NonlinearSolverParameters	node	unique	<i>Element: NonlinearSolverParameters</i>



## 10.73 Element: LinearSolverParameters

Name	Type	Default	Description
amgAggressiveCoarseningLevel	integer	0	AMG number levels for aggressive coarsening Available options are: TODO
amgCoarseSolver	geosx_LinearSolverParameters AMG_CoarseType	direct	AMG coarsest level solver/smoother type. Available options are: default\ jacobi\ llt\ jacobi\ fgsv\
amgCoarseningType	string	HMIS	AMG coarsening algorithm Available options are: TODO
amgInterpolationType	integer	6	AMG interpolation algorithm Available options are: TODO
amgNullSpaceType	geosx_LinearSolverParameters AMG_NullSpaceType	constantModes	AMG near null space approximation. Available options are: constantModes\ rigidBodyModes\
amgNumFunctions	integer	1	AMG number of functions Available options are: TODO
amgNumSweeps	integer	2	AMG smoother sweeps
amgSmootherType	geosx_LinearSolverParameters AMG_SmootherType	fgsv	AMG smoother type. Available options are: default\ jacobi\ llt\ jacobi\ fgsv\
amgThreshold	real64	0	AMG strength-of-connection threshold
directCheckResidual	integer	0	Whether to check the linear system solution residual
directColPerm	geosx_LinearSolverParameters Direct_ColPerm	none	How to permute the columns. Available options are: none\ MMD_AtplusA\ MMD_AtA\ col
directEquil	integer	1	Whether to scale the rows and columns of the matrix
directIterRef	integer	1	Whether to perform iterative refinement
directParallel	integer	1	Whether to use a parallel solver (instead of a serial one)
directRepTinyPivot	integer	1	Whether to raise a warning when a tiny pivot is encountered



## 10.74 Element: Mesh

Name	Type	Default	Description
InternalMesh	node		<i>Element: InternalMesh</i>
InternalWell	node		<i>Element: InternalWell</i>
InternalWellbore	node		<i>Element: InternalWellbore</i>
PAMELAMesh	node		<i>Element: PAMELAMesh</i>
VTKMesh	node		<i>Element: VTKMesh</i>

## 10.75 Element: ModifiedCamClay

Name	Type	Default	Description
defaultCsISlope	real64	1	Slope of the critical state line
defaultDensity	real64	required	Default Material Density
defaultPreConsolidationPressure	real64	-1.5	Initial preconsolidation pressure
defaultRecompressionIndex	real64	0.002	Recompression Index
defaultRefPressure	real64	-1	Reference Pressure
defaultRefStrainVol	real64	0	Reference Volumetric Strain
defaultShearModulus	real64	-1	Elastic Shear Modulus Parameter
defaultVirginCompressionIndex	real64	0.005	Virgin compression index
name	string	required	A name is required for any non-unique nodes

## 10.76 Element: MultiPhaseConstantThermalConductivity

Name	Type	De- fault	Description
name	string	re- quired	A name is required for any non-unique nodes
phaseNames	string_array	re- quired	List of fluid phases
thermalConductivity- Components	RTensor	re- quired	xx, yy, and zz components of a diagonal thermal conductivity tensor [J/(s.m.K)]

## 10.77 Element: MultiPhaseVolumeWeightedThermalConductivity

Name	Type	De- fault	Description
name	string	re- quired	A name is required for any non-unique nodes
phaseNames	string_array	re- quired	List of fluid phases
phaseThermalConductivity	real64_array	re- quired	Phase thermal conductivity [W/(m.K)]
rockThermalConductivity- Components	R1Tensor	re- quired	xx, yy, and zz components of a diagonal rock thermal conductivity tensor [W/(m.K)]

## 10.78 Element: MultiphasePoromechanics

Name	Type	De- fault	Description
cflFactor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
flow-Solver-Name	string	re- quired	Name of the flow solver used by the coupled solver
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
logLevel	integer	0	Log level
name	string	re- quired	A name is required for any non-unique nodes
solid-Solver-Name	string	re- quired	Name of the solid solver used by the coupled solver
targetRegions	string_array	re- quired	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
Linear-Solver-Parameters	node	unique	<i>Element: LinearSolverParameters</i>
Non-linear-Solver-Parameters	node	unique	<i>Element: NonlinearSolverParameters</i>

## 10.79 Element: MultiphasePoromechanicsReservoir

Name	Type	Default	Description
cflFactor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
logLevel	integer	0	Log level
name	string	required	A name is required for any non-unique nodes
poromechanics-Solver-Name	string	required	Name of the poromechanics solver used by the coupled solver
targetRegions	string_array	required	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
well-Solver-Name	string	required	Name of the well solver used by the coupled solver
Linear-Solver-Parameters	node	unique	<i>Element: LinearSolverParameters</i>
Non-linear-Solver-Parameters	node	unique	<i>Element: NonlinearSolverParameters</i>

## 10.80 Element: MultivariableTableFunction

Name	Type	Default	Description
inputVarNames	string_array	{ }	Name of fields are input to function.
name	string	required	A name is required for any non-unique nodes



## 10.81 Element: NonlinearSolverParameters

Name	Type	Default	Description
allowNonConverged	integer	0	Allow non-converged solution to be accepted. (i.e. exit from the Newton loop without achieving the desired tolerance)
lineSearchAction	geosx_NonlinearSolverParameters.lineSearchAction	Attempt	How the line search is to be used. Options are: <ul style="list-style-type: none"> <li>* None - Do not use line search.</li> <li>* Attempt - Use line search. Allow exit from line search without achieving smaller residual than starting residual.</li> <li>* Require - Use line search. If smaller residual than starting residual is not achieved, cut time step.</li> </ul>
lineSearchCutFactor	real64	0.5	Line search cut factor. For instance, a value of 0.5 will result in the effective application of the last solution by a factor of (0.5, 0.25, 0.125, ...)
lineSearchInterpolationType	geosx_NonlinearSolverParameters.lineSearchInterpolationType	Linear	Strategy to cut the solution update during the line search. Options are: <ul style="list-style-type: none"> <li>* Linear</li> <li>* Parabolic</li> </ul>
lineSearchMaxCuts	integer	4	Maximum number of line search cuts.
logLevel	integer	0	Log level
maxAllowedResidualNorm	real64	1e+09	Maximum value of residual norm that is allowed in a Newton loop
maxNumConfigurationAttempts	integer	10	Max number of times that the configuration can be changed
maxSubSteps	integer	10	Maximum number of time sub-steps allowed for the solver
maxTimeStepCuts	integer	2	Max number of time step cuts
newtonMaxIter	integer	5	Maximum number of iterations that are allowed in a Newton loop.
newtonMinIter	integer	1	Minimum number of iterations that are required before exiting the Newton loop.

## 10.82 Element: NullModel

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes

## 10.83 Element: NumericalMethods

Name	Type	Default	Description
FiniteElements	node	unique	<i>Element: FiniteElements</i>
FiniteVolume	node	unique	<i>Element: FiniteVolume</i>

## 10.84 Element: Outputs

Name	Type	Default	Description
Blueprint	node		<i>Element: Blueprint</i>
ChomboIO	node		<i>Element: ChomboIO</i>
Python	node		<i>Element: Python</i>
Restart	node		<i>Element: Restart</i>
Silo	node		<i>Element: Silo</i>
TimeHistory	node		<i>Element: TimeHistory</i>
VTK	node		<i>Element: VTK</i>

## 10.85 Element: PAMELAMesh

Name	Type	De- fault	Description
fieldNamesIn- GEOSX	string_array	{ }	Names of fields in GEOSX to import into
fieldsToImport	string_array	{ }	Fields to be imported from the external mesh file
file	path	re- quired	Path to the mesh file
logLevel	integer	0	Log level
name	string	re- quired	A name is required for any non-unique nodes
scale	R1Tensor	{1,1,1}	Scale the coordinates of the vertices by given scale factors (after translation)
translate	R1Tensor	{0,0,0}	Translate the coordinates of the vertices by a given vector (prior to scaling)

## 10.86 Element: PML

Name	Type	Default	Description
bcApplicationTable-Name	string		Name of table that specifies the on/off application of the boundary condition.
beginTime	real64	-1e+99	Time at which the boundary condition will start being applied.
component	integer	-1	Component of field (if tensor) to apply boundary condition to.
direction	R1Tensor	{0,0,0}	Direction to apply boundary condition to.
endTime	real64	1e+99	Time at which the boundary condition will stop being applied.
functionName	string		Name of function that specifies variation of the boundary condition.
logLevel	integer	0	Log level
name	string	required	A name is required for any non-unique nodes
objectPath	string		Path to the target field
reflectivity	real32	0.001	Desired reflectivity of the PML region, used to compute the damping profile
scale	real64	0	Scale factor for value of the boundary condition.
setNames	string_array	required	Name of sets that boundary condition is applied to.
thickness-MaxXYZ	R1Tensor	{21,-1,-1}	Thickness of the PML region, at right, back, and bottom sides, used to compute the damping profile
thickness-MinXYZ	R1Tensor	{21,-1,-1}	Thickness of the PML region, at left, front, and top sides, used to compute the damping profile
waveSpeed-MaxXYZ	R1Tensor	{21,-1,-1}	Wave speed in the PML, at right, back, and bottom sides, used to compute the damping profile
waveSpeed-MinXYZ	R1Tensor	{21,-1,-1}	Wave speed in the PML, at left, front, and top sides, used to compute the damping profile
xMax	R1Tensor	{3.40282e+38,3.40282e+38,3.40282e+38}	Maximum (x,y,z) coordinates of the inner PML boundaries
xMin	R1Tensor	{-3.40282e+38,-3.40282e+38,-3.40282e+38}	Minimum (x,y,z) coordinates of the inner PML boundaries

## 10.87 Element: PVTDriver

Name	Type	Default	Description
baseline	path	none	Baseline file
feedComposition	real64_array	required	Feed composition array [mol fraction]
fluid	string	required	Fluid to test
logLevel	integer	0	Log level
name	string	required	A name is required for any non-unique nodes
output	string	none	Output file
pressureControl	string	required	Function controlling pressure time history
steps	integer	required	Number of load steps to take
temperatureControl	string	required	Function controlling temperature time history

## 10.88 Element: PackCollection

Name	Type	De- fault	Description
fieldName	string	re- quired	The name of the (packable) field associated with the specified object to retrieve data from
name	string	re- quired	A name is required for any non-unique nodes
objectPath	string	re- quired	The name of the object from which to retrieve field values.
onlyOn- SetChange	integer	0	Whether or not to only collect when the collected sets of indices change in any way.
setNames	string_array	{ }	The set(s) for which to retrieve data.

## 10.89 Element: ParallelPlatesPermeability

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes

## 10.90 Element: Parameter

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes
value	string	required	Input parameter definition for the preprocessor

## 10.91 Element: Parameters

Name	Type	Default	Description
Parameter	node		<i>Element: Parameter</i>



## 10.92 Element: ParticleFluid

Name	Type	Default	Description
collisionAlpha	real64	1.27	Collision alpha coefficient
collisionBeta	real64	1.5	Collision beta coefficient
fluidViscosity	real64	0.001	Fluid viscosity
hinderedSettlingCoefficient	real64	5.9	Hindered settling coefficient
isCollisionalSlip	integer	0	Whether the collisional component of the slip velocity is considered
maxProppantConcentration	real64	0.6	Max proppant concentration
name	string	required	A name is required for any non-unique nodes
particleSettlingModel	geosx_constitutive_ParticleSettlingModel	required	Particle settling velocity model. Valid options: * Stokes * Intermediate * Turbulence
proppantDensity	real64	1400	Proppant density
proppantDiameter	real64	0.0002	Proppant diameter
slipConcentration	real64	0.1	Slip concentration
sphericity	real64	1	Sphericity

## 10.93 Element: Perforation

Name	Type	Default	Description
distanceFromHead	real64	required	Linear distance from well head to the perforation
name	string	required	A name is required for any non-unique nodes
transmissibility	real64	-1	Perforation transmissibility

## 10.94 Element: PeriodicEvent

Name	Type	De- fault	Description
begin-Time	real64	0	Start time of this event.
cycleFrequency	integer	1	Event application frequency (cycle, default)
endTime	real64	1e+100	End time of this event.
finalDt-Stretch	real64	0.001	Allow the final dt request for this event to grow by this percentage to match the endTime exactly.
forceDt	real64	-1	While active, this event will request this timestep value (ignoring any children/targets requests).
function	string		Name of an optional function to evaluate when the time/cycle criteria are met. If the result is greater than the specified eventThreshold, the function will continue to execute.
logLevel	integer	0	Log level
max-EventDt	real64	-1	While active, this event will request a timestep $\leq$ this value (depending upon any child/target requests).
name	string	required	A name is required for any non-unique nodes
object	string		If the optional function requires an object as an input, specify its path here.
set	string		If the optional function is applied to an object, specify the setname to evaluate (default = everything).
stat	integer	0	If the optional function is applied to an object, specify the statistic to compare to the eventThreshold. The current options include: min, avg, and max.
target	string		Name of the object to be executed when the event criteria are met.
targetExactStart-Stop	integer	1	If this option is set, the event will reduce its timestep requests to match any specified beginTime/endTimes exactly.
targetExact-Timestep	integer	1	If this option is set, the event will reduce its timestep requests to match the specified timeFrequency perfectly: $dt\_request = \min(dt\_request, t\_last + time\_frequency - time)$ .
threshold	real64	0	If the optional function is used, the event will execute if the value returned by the function exceeds this threshold.
timeFrequency	real64	-1	Event application frequency (time). Note: if this value is specified, it will override any cycle-based behavior.
HaltEvent	node		<i>Element: HaltEvent</i>
PeriodicEvent	node		<i>Element: PeriodicEvent</i>
SoloEvent	node		<i>Element: SoloEvent</i>

## 10.95 Element: PermeabilityBase

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes

## 10.96 Element: PhaseFieldDamageFEM

Name	Type	De- fault	Description
cflFactor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
damage-Upper-Bound	real64	1.5	The upper bound of the damage
discretization	string	required	Name of discretization object (defined in the <a href="#">Numerical Methods</a> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <a href="#">Finite Element Discretization</a> should be specified. If this is a Finite Volume Method, the name of a <a href="#">Finite Volume Discretization</a> discretization should be specified.
field-Name	string	required	name of field variable
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
irreversibilityFlag	integer	0	The flag to indicate whether to apply the irreversibility constraint
localDissipation	string	required	Type of local dissipation function. Can be Linear or Quadratic
logLevel	integer	0	Log level
name	string	required	A name is required for any non-unique nodes
targetRegions	string array	required	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
timeIntegrationOption	string	required	option for default time integration method
Linear-Solver-Parameters	node	unique	<a href="#">Element: LinearSolverParameters</a>
Non-linear-Solver-Parameters	node	unique	<a href="#">Element: NonlinearSolverParameters</a>



## 10.97 Element: PhaseFieldFracture

Name	Type	Default	Description
cflFactor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
couplingTypeOption	geosx_PhaseFieldFractureSolverRequiredCouplingTypeOption	required	Coupling option. Valid options: * FixedStress * TightlyCoupled
damageSolverName	string	required	Name of the damage mechanics solver to use in the PhaseFieldFracture solver
discretization	string	required	Name of discretization object (defined in the <a href="#">Numerical Methods</a> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <a href="#">Finite Element Discretization</a> should be specified. If this is a Finite Volume Method, the name of a <a href="#">Finite Volume Discretization</a> discretization should be specified.
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
logLevel	integer	0	Log level
name	string	required	A name is required for any non-unique nodes
solidSolverName	string	required	Name of the solid mechanics solver to use in the PhaseFieldFracture solver
subcycling	integer	required	turn on subcycling on each load step
targetRegions	string_array	required	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions
<b>10.97. Element: PhaseFieldFracture</b>			this solver will be applied to rests in the EventManager. <b>623</b>
LinearSolverParameters	node	unique	<a href="#">Element: LinearSolverParameters</a>

## 10.98 Element: PorousDelftEgg

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes
permeabilityModelName	string	required	Name of the permeability model.
porosityModelName	string	required	Name of the porosity model.
solidInternalEnergyModelName	string		Name of the solid internal energy model.
solidModelName	string	required	Name of the solid model.

## 10.99 Element: PorousDruckerPrager

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes
permeabilityModelName	string	required	Name of the permeability model.
porosityModelName	string	required	Name of the porosity model.
solidInternalEnergyModelName	string		Name of the solid internal energy model.
solidModelName	string	required	Name of the solid model.

## 10.100 Element: PorousElasticIsotropic

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes
permeabilityModelName	string	required	Name of the permeability model.
porosityModelName	string	required	Name of the porosity model.
solidInternalEnergyModelName	string		Name of the solid internal energy model.
solidModelName	string	required	Name of the solid model.

## 10.101 Element: PorousElasticOrthotropic

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes
permeabilityModelName	string	required	Name of the permeability model.
porosityModelName	string	required	Name of the porosity model.
solidInternalEnergyModelName	string		Name of the solid internal energy model.
solidModelName	string	required	Name of the solid model.

## 10.102 Element: PorousElasticTransverselsotropic

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes
permeabilityModelName	string	required	Name of the permeability model.
porosityModelName	string	required	Name of the porosity model.
solidInternalEnergyModelName	string		Name of the solid internal energy model.
solidModelName	string	required	Name of the solid model.

## 10.103 Element: PorousExtendedDruckerPrager

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes
permeabilityModelName	string	required	Name of the permeability model.
porosityModelName	string	required	Name of the porosity model.
solidInternalEnergyModelName	string		Name of the solid internal energy model.
solidModelName	string	required	Name of the solid model.

## 10.104 Element: PorousModifiedCamClay

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes
permeabilityModelName	string	required	Name of the permeability model.
porosityModelName	string	required	Name of the porosity model.
solidInternalEnergyModelName	string		Name of the solid internal energy model.
solidModelName	string	required	Name of the solid model.

## 10.105 Element: PressurePorosity

Name	Type	Default	Description
compressibility	real64	required	Solid compressibility
defaultReferencePorosity	real64	required	Default value of the reference porosity
name	string	required	A name is required for any non-unique nodes
referencePressure	real64	required	Reference pressure for solid compressibility

## 10.106 Element: Problem

Name	Type	Default	Description
Benchmarks	node	unique	<i>Element: Benchmarks</i>
Constitutive	node	unique	<i>Element: Constitutive</i>
ElementRegions	node	unique	<i>Element: ElementRegions</i>
Events	node	unique, required	<i>Element: Events</i>
FieldSpecifications	node	unique	<i>Element: FieldSpecifications</i>
Functions	node	unique	<i>Element: Functions</i>
Geometry	node	unique	<i>Element: Geometry</i>
Included	node	unique	<i>Element: Included</i>
Mesh	node	unique, required	<i>Element: Mesh</i>
NumericalMethods	node	unique	<i>Element: NumericalMethods</i>
Outputs	node	unique, required	<i>Element: Outputs</i>
Parameters	node	unique	<i>Element: Parameters</i>
Solvers	node	unique, required	<i>Element: Solvers</i>
Tasks	node	unique	<i>Element: Tasks</i>

## 10.107 Element: ProppantPermeability

Name	Type	Default	Description
maxProppantConcentration	real64	required	Maximum proppant concentration.
name	string	required	A name is required for any non-unique nodes
proppantDiameter	real64	required	Proppant diameter.

## 10.108 Element: ProppantPorosity

Name	Type	Default	Description
defaultReferencePorosity	real64	required	Default value of the reference porosity
maxProppantConcentration	real64	required	Maximum proppant concentration
name	string	required	A name is required for any non-unique nodes



## 10.109 Element: ProppantSlurryFluid

Name	Type	Default	Description
componentNames	string_array	{ }	List of fluid component names
compressibility	real64	0	Fluid compressibility
defaultComponentDensity	real64_array	{0}	Default value for the component density.
defaultComponentViscosity	real64_array	{0}	Default value for the component viscosity.
defaultCompressibility	real64_array	{0}	Default value for the component compressibility.
flowBehaviorIndex	real64_array	{0}	Flow behavior index
flowConsistencyIndex	real64_array	{0}	Flow consistency index
maxProppantConcentration	real64	0.6	Maximum proppant concentration
name	string	required	A name is required for any non-unique nodes
referenceDensity	real64	1000	Reference fluid density
referencePressure	real64	100000	Reference pressure
referenceProppantDensity	real64	1400	Reference proppant density
referenceViscosity	real64	0.001	Reference fluid viscosity

## 10.110 Element: ProppantSolidProppantPermeability

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes
permeabilityModelName	string	required	Name of the permeability model.
porosityModelName	string	required	Name of the porosity model.
solidInternalEnergyModelName	string		Name of the solid internal energy model.
solidModelName	string	required	Name of the solid model.



## 10.111 Element: ProppantTransport

Name	Type	De- fault	Description
bridg- ingFac- tor	real64	0	Bridging factor used for bridging/screen-out calculation
cflFac- tor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
critical- Shield- sNum- ber	real64	0	Critical Shields number
dis- cretiza- tion	string	re- quired	Name of discretization object (defined in the <a href="#">Numerical Methods</a> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <a href="#">Finite Element Discretiza- tion</a> should be specified. If this is a Finite Volume Method, the name of a <a href="#">Finite Volume Discretization</a> discretization should be specified.
fric- tion- Coeffi- cient	real64	0.03	Friction coefficient
ini- tialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
input- FluxEs- timate	real64	1	Initial estimate of the input flux used only for residual scaling. This should be essentially equivalent to the input flux * dt.
isTher- mal	in- te- ger	0	Flag indicating whether the problem is thermal or not.
logLevel	in- te- ger	0	Log level
max- Prop- ant- Con- centra- tion	real64	0.6	Maximum proppant concentration
name	string	re- quired	A name is required for any non-unique nodes
prop- ant- Density	real64	2500	Proppant density
prop- antDi- ameter	real64	0.0004	Proppant diameter
targe- tRe- gions	string array	re- quired	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
up- dateProp- ant-Pack-	in- te- ger	0	Flag that enables/disables proppant-packing update

### 10.111. Element: ProppantTransport

629

Linear-  
Solver-  
Param-  
eters

node unique

[Element: LinearSolverParameters](#)

## 10.112 Element: Python

Name	Type	Default	Description
childDirectory	string		Child directory path
name	string	required	A name is required for any non-unique nodes
parallelThreads	integer	1	Number of plot files.



## 10.113 Element: ReactiveCompositionalMultiphaseOBL

Name	Type	De- fault	Description
OBL- Opera- torsTable- File	path	re- quired	File containing OBL operator values
al- lowLocalOBL- Chop- ping	in- te- ger	1	Allow keeping solution within OBL limits
cflFac- tor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
compo- nent- Names	string_4array		List of component names
dis- cretiza- tion	string	re- quired	Name of discretization object (defined in the <a href="#">Numerical Methods</a> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <a href="#">Finite Element Discretization</a> should be specified. If this is a Finite Volume Method, the name of a <a href="#">Finite Volume Discretization</a> discretization should be specified.
en- ableEn- ergy- Bal- ance	in- te- ger	re- quired	Enable energy balance calculation and temperature degree of freedom
ini- tialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
input- FluxEs- timate	real64	1	Initial estimate of the input flux used only for residual scaling. This should be essentially equivalent to the input flux * dt.
isTher- mal	in- te- ger	0	Flag indicating whether the problem is thermal or not.
logLevel	in- te- ger	0	Log level
max- CompFrac- tion- Change	real64	1	Maximum (absolute) change in a component fraction between two Newton iterations
name	string	re- quired	A name is required for any non-unique nodes
num- Com- ponents	in- te- ger	re- quired	Number of components
numPhases	in- te- ger	re- quired	Number of phases
phase- Names	string_4array		List of fluid phases
targetRe- gions	string_array	re- quired	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
trans- Mult- Exp	real64	1	Exponent of dynamic transmissibility multiplier

## 10.114 Element: Restart

Name	Type	Default	Description
childDirectory	string		Child directory path
name	string	required	A name is required for any non-unique nodes
parallelThreads	integer	1	Number of plot files.

## 10.115 Element: Run

Name	Type	De- fault	Description
args	string		Any extra command line arguments to pass to GEOSX.
autoParti- tion	string		May be 'Off' or 'On', if 'On' partitioning arguments are created automati- cally. Default is Off.
name	string	re- quired	The name of this benchmark.
nodes	integer	re- quired	The number of nodes needed to run the benchmark.
strongScal- ing	inte- ger_array	{0}	Repeat the benchmark N times, scaling the number of nodes in the bench- mark by these values.
tasksPerN- ode	integer	re- quired	The number of tasks per node to run the benchmark with.
threadsPer- Task	integer	0	The number of threads per task to run the benchmark with.
timeLimit	integer	0	The time limit of the benchmark.

## 10.116 Element: Silo

Name	Type	De- fault	Description
childDirectory	string		Child directory path
fieldNames	string array		Names of the fields to output. If this attribute is specified, GEOSX outputs all (and only) the fields specified by the user, regardless of their <i>plotLevel</i>
name	string	re- quired	A name is required for any non-unique nodes
onlyPlotSpecifiedFieldNames	integer	0	If this flag is equal to 1, then we only plot the fields listed in <i>fieldNames</i> . Otherwise, we plot all the fields with the required <i>plotLevel</i> , plus the fields listed in <i>fieldNames</i>
parallelThreads	integer	1	Number of plot files.
plotFileRoot	string	plot	(no description available)
plotLevel	integer	1	(no description available)
writeCellElementMesh	integer	1	(no description available)
writeEdgeMesh	integer	0	(no description available)
writeFEM-Faces	integer	0	(no description available)
writeFaceElementMesh	integer	1	(no description available)

## 10.117 Element: SinglePhaseConstantThermalConductivity

Name	Type	De- fault	Description
name	string	re- quired	A name is required for any non-unique nodes
thermalConductivity-Components	R1Tensor	re- quired	xx, yy, and zz components of a diagonal thermal conductivity tensor [J/(s.m.K)]



## 10.118 Element: SinglePhaseFVM

Name	Type	De- fault	Description
cflFactor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
discretization	string	required	Name of discretization object (defined in the <a href="#">Numerical Methods</a> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <a href="#">Finite Element Discretization</a> should be specified. If this is a Finite Volume Method, the name of a <a href="#">Finite Volume Discretization</a> discretization should be specified.
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
input-FluxEstimate	real64	1	Initial estimate of the input flux used only for residual scaling. This should be essentially equivalent to the input flux * dt.
isThermal	integer	0	Flag indicating whether the problem is thermal or not.
logLevel	integer	0	Log level
name	string	required	A name is required for any non-unique nodes
targetRegions	string array	required	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
temperature	real64	0	Temperature
Linear-Solver-Parameters	node	unique	<a href="#">Element: LinearSolverParameters</a>
Non-linear-Solver-Parameters	node	unique	<a href="#">Element: NonlinearSolverParameters</a>

## 10.119 Element: SinglePhaseHybridFVM

Name	Type	De- fault	Description
cflFactor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
discretization	string	required	Name of discretization object (defined in the <a href="#">Numerical Methods</a> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <a href="#">Finite Element Discretization</a> should be specified. If this is a Finite Volume Method, the name of a <a href="#">Finite Volume Discretization</a> discretization should be specified.
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
input-FluxEstimate	real64	1	Initial estimate of the input flux used only for residual scaling. This should be essentially equivalent to the input flux * dt.
isThermal	integer	0	Flag indicating whether the problem is thermal or not.
logLevel	integer	0	Log level
name	string	required	A name is required for any non-unique nodes
targetRegions	string_array	required	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
temperature	real64	0	Temperature
Linear-Solver-Parameters	node	unique	<a href="#">Element: LinearSolverParameters</a>
Non-linear-Solver-Parameters	node	unique	<a href="#">Element: NonlinearSolverParameters</a>

## 10.120 Element: SinglePhasePoromechanics

Name	Type	De- fault	Description
cflFactor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
flow-Solver-Name	string	re- quired	Name of the flow solver used by the coupled solver
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
logLevel	integer	0	Log level
name	string	re- quired	A name is required for any non-unique nodes
solid-Solver-Name	string	re- quired	Name of the solid solver used by the coupled solver
targetRegions	string array	re- quired	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
Linear-Solver-Parameters	node	unique	<i>Element: LinearSolverParameters</i>
Non-linear-Solver-Parameters	node	unique	<i>Element: NonlinearSolverParameters</i>

## 10.121 Element: SinglePhasePoromechanicsEmbeddedFractures

Name	Type	De- fault	Description
cflFactor	real64	0.5	Factor to apply to the CFL condition when calculating the maximum allowable time step. Values should be in the interval (0,1]
flow-Solver-Name	string	required	Name of the flow solver used by the coupled solver
fractures-Solver-Name	string	required	Name of the fractures solver to use in the fractured poroelastic solver
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
logLevel	integer	0	Log level
name	string	required	A name is required for any non-unique nodes
solid-Solver-Name	string	required	Name of the solid solver used by the coupled solver
targetRegions	string_array	required	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
Linear-Solver-Parameters	node	unique	<i>Element: LinearSolverParameters</i>
Non-linear-Solver-Parameters	node	unique	<i>Element: NonlinearSolverParameters</i>

## 10.122 Element: SinglePhasePoromechanicsReservoir

Name	Type	De- fault	Description
cflFac- tor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
ini- tialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
logLevel	in- te- ger	0	Log level
name	string	re- quired	A name is required for any non-unique nodes
porome- chan- ics- Solver- Name	string	re- quired	Name of the poromechanics solver used by the coupled solver
targe- tRe- gions	string array	re- quired	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
well- Solver- Name	string	re- quired	Name of the well solver used by the coupled solver
Linear- Solver- Param- eters	node	unique	<i>Element: LinearSolverParameters</i>
Non- linear- Solver- Param- eters	node	unique	<i>Element: NonlinearSolverParameters</i>

## 10.123 Element: SinglePhaseProppantFVM

Name	Type	De- fault	Description
cflFactor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
discretization	string	required	Name of discretization object (defined in the <a href="#">Numerical Methods</a> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <a href="#">Finite Element Discretization</a> should be specified. If this is a Finite Volume Method, the name of a <a href="#">Finite Volume Discretization</a> discretization should be specified.
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
input-FluxEstimate	real64	1	Initial estimate of the input flux used only for residual scaling. This should be essentially equivalent to the input flux * dt.
isThermal	integer	0	Flag indicating whether the problem is thermal or not.
logLevel	integer	0	Log level
name	string	required	A name is required for any non-unique nodes
targetRegions	string_array	required	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
temperature	real64	0	Temperature
Linear-Solver-Parameters	node	unique	<a href="#">Element: LinearSolverParameters</a>
Non-linear-Solver-Parameters	node	unique	<a href="#">Element: NonlinearSolverParameters</a>

## 10.124 Element: SinglePhaseReservoir

Name	Type	De- fault	Description
cflFactor	real64	0.5	Factor to apply to the <a href="#">CFL condition</a> when calculating the maximum allowable time step. Values should be in the interval (0,1]
flow-Solver-Name	string	re- quired	Name of the flow solver used by the coupled solver
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
logLevel	in- te- ger	0	Log level
name	string	re- quired	A name is required for any non-unique nodes
targetRegions	string_array	re- quired	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
well-Solver-Name	string	re- quired	Name of the well solver used by the coupled solver
Linear-Solver-Parameters	node	unique	<i>Element: LinearSolverParameters</i>
Non-linear-Solver-Parameters	node	unique	<i>Element: NonlinearSolverParameters</i>

## 10.125 Element: SinglePhaseStatistics

Name	Type	Default	Description
flowSolverName	string	required	Name of the flow solver
logLevel	integer	0	Log level
name	string	required	A name is required for any non-unique nodes

## 10.126 Element: SinglePhaseWell

Name	Type	Default	Description
cflFactor	real64	0.5	Factor to apply to the CFL condition when calculating the maximum allowable time step. Values should be in the interval (0,1]
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
logLevel	integer	0	Log level
name	string	required	A name is required for any non-unique nodes
targetRegions	string array	required	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
Linear-Solver-Parameters	node	unique	<i>Element: LinearSolverParameters</i>
Non-linear-Solver-Parameters	node	unique	<i>Element: NonlinearSolverParameters</i>
Well-Controls	node		<i>Element: WellControls</i>

## 10.127 Element: SlipDependentPermeability

Name	Type	Default	Description
initialPermeability	R1Tensor	required	initial permeability of the fracture.
maxPermMultiplier	real64	required	Maximum permeability multiplier.
name	string	required	A name is required for any non-unique nodes
shearDispThreshold	real64	required	Threshold of shear displacement.

## 10.128 Element: SolidInternalEnergy

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes
referenceInternalEnergy	real64	required	Internal energy at the reference temperature [J/kg]
referenceTemperature	real64	required	Reference temperature [K]
volumetricHeatCapacity	real64	required	Solid volumetric heat capacity [J/(kg.K)]



## 10.129 Element: SolidMechanicsEmbeddedFractures

Name	Type	De- fault	Description
cflFactor	real64	0.5	Factor to apply to the CFL condition when calculating the maximum allowable time step. Values should be in the interval (0,1]
contactRelation-Name	string	re- quired	Name of contact relation to enforce constraints on fracture boundary.
fractureRegion-Name	string	re- quired	Name of the fracture region.
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
logLevel	integer	0	Log level
name	string	re- quired	A name is required for any non-unique nodes
solid-Solver-Name	string	re- quired	Name of the solid mechanics solver in the rock matrix
targetRegions	string array	re- quired	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
useStatic-Condensation	integer	0	Defines whether to use static condensation or not.
Linear-Solver-Parameters	node	unique	<i>Element: LinearSolverParameters</i>
Non-linear-Solver-Parameters	node	unique	<i>Element: NonlinearSolverParameters</i>



## 10.130 Element: SolidMechanicsLagrangianSSLE

Name	Type	Default	Description
cflFactor	real64	0.5	Factor to apply to the CFL condition when calculating the maximum allowable time step. Values should be in the interval (0,1]
contactRelationName	string	NOCONTACT	Name of contact relation to enforce constraints on fracture boundary.
discretization	string	required	Name of discretization object (defined in the <i>Numerical Methods</i> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <i>Finite Element Discretization</i> should be specified. If this is a Finite Volume Method, the name of a <i>Finite Volume Discretization</i> discretization should be specified.
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
logLevel	integer	0	Log level
massDamping	real64	0	Value of mass based damping coefficient.
maxNumResolves	integer	10	Value to indicate how many resolves may be executed after some other event is executed. For example, if a SurfaceGenerator is specified, it will be executed after the mechanics solve. However if a new surface is generated, then the mechanics solve must be executed again due to the change in topology.
name	string	required	A name is required for any non-unique nodes
newmarkBeta	real64	0.25	Value of $\beta$ in the Newmark Method for Implicit Dynamic time integration option. This should be $\text{pow}(\text{newmarkGamma}+0.5, 2.0)/4.0$ unless you know what you are doing.
newmarkGamma	real64	0.5	Value of $\gamma$ in the Newmark Method for Implicit Dynamic time integration option
stiffnessDamping	real64	0	Value of stiffness based

### 10.131 Element: SolidMechanicsStateReset

Name	Type	Default	Description
disableInelasticity	integer	0	Flag to enable/disable inelastic behavior
logLevel	integer	0	Log level
name	string	required	A name is required for any non-unique nodes
resetDisplacements	integer	1	Flag to reset displacements (and velocities)
solidSolverName	string	required	Name of the solid mechanics solver

### 10.132 Element: SolidMechanicsStatistics

Name	Type	Default	Description
logLevel	integer	0	Log level
name	string	required	A name is required for any non-unique nodes
solidSolverName	string	required	Name of the solid solver



## 10.133 Element: SolidMechanics\_LagrangianFEM

Name	Type	Default	Description
cflFactor	real64	0.5	Factor to apply to the CFL condition when calculating the maximum allowable time step. Values should be in the interval (0,1]
contactRelationName	string	NOCONTACT	Name of contact relation to enforce constraints on fracture boundary.
discretization	string	required	Name of discretization object (defined in the <i>Numerical Methods</i> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <i>Finite Element Discretization</i> should be specified. If this is a Finite Volume Method, the name of a <i>Finite Volume Discretization</i> should be specified.
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
logLevel	integer	0	Log level
massDamping	real64	0	Value of mass based damping coefficient.
maxNumResolves	integer	10	Value to indicate how many resolves may be executed after some other event is executed. For example, if a SurfaceGenerator is specified, it will be executed after the mechanics solve. However if a new surface is generated, then the mechanics solve must be executed again due to the change in topology.
name	string	required	A name is required for any non-unique nodes
newmarkBeta	real64	0.25	Value of $\beta$ in the Newmark Method for Implicit Dynamic time integration option. This should be $\text{pow}(\text{newmarkGamma}+0.5, 2.0)/4.0$ unless you know what you are doing.
newmarkGamma	real64	0.5	Value of $\gamma$ in the Newmark Method for Implicit Dynamic time integration option
stiffnessDamping	real64	0	Value of stiffness based

## 10.134 Element: SoloEvent

Name	Type	De- fault	Description
beginTime	real64	0	Start time of this event.
endTime	real64	1e+100	End time of this event.
finalDt-Stretch	real64	0.001	Allow the final dt request for this event to grow by this percentage to match the end-Time exactly.
forceDt	real64	-1	While active, this event will request this timestep value (ignoring any children/targets requests).
logLevel	in- te- ger	0	Log level
max-EventDt	real64	-1	While active, this event will request a timestep $\leq$ this value (depending upon any child/target requests).
name	string	re- quired	A name is required for any non-unique nodes
target	string		Name of the object to be executed when the event criteria are met.
targetCy- cle	in- te- ger	-1	Targeted cycle to execute the event.
targetEx- actStart- Stop	in- te- ger	1	If this option is set, the event will reduce its timestep requests to match any specified beginTime/endTimes exactly.
targe- tExact- Timestep	in- te- ger	1	If this option is set, the event will reduce its timestep requests to match the specified execution time exactly: $dt\_request = \min(dt\_request, t\_target - time)$ .
targetTime	real64	-1	Targeted time to execute the event.
HaltEvent	node		<i>Element: HaltEvent</i>
Peri- odicEvent	node		<i>Element: PeriodicEvent</i>
SoloEvent	node		<i>Element: SoloEvent</i>

## 10.135 Element: Solvers

Name	Type	Default	Description
gravityVector	RI Tensor	{0,0,-9.81}	Gravity vector used in the physics solvers
AcousticSEM	node		<i>Element: AcousticSEM</i>
CompositionalMultiphaseFVM	node		<i>Element: CompositionalMultiphaseFVM</i>
CompositionalMultiphaseHybridFVM	node		<i>Element: CompositionalMultiphaseHybridFVM</i>
CompositionalMultiphaseReservoir	node		<i>Element: CompositionalMultiphaseReservoir</i>
CompositionalMultiphaseWell	node		<i>Element: CompositionalMultiphaseWell</i>
ElasticSEM	node		<i>Element: ElasticSEM</i>
EmbeddedSurfaceGenerator	node		<i>Element: EmbeddedSurfaceGenerator</i>
FlowProppantTransport	node		<i>Element: FlowProppantTransport</i>
Hydrofracture	node		<i>Element: Hydrofracture</i>
LagrangianContact	node		<i>Element: LagrangianContact</i>
LaplaceFEM	node		<i>Element: LaplaceFEM</i>

Continued on next page

Table 10.2 – continued from previous page

Name	Type	Default	Description
MultiphasePoromechanics	node		<i>Element: MultiphasePoromechanics</i>
MultiphasePoromechanicsReservoir	node		<i>Element: MultiphasePoromechanicsReservoir</i>
PhaseFieldDamageFEM	node		<i>Element: PhaseFieldDamageFEM</i>
PhaseFieldFracture	node		<i>Element: PhaseFieldFracture</i>
ProppantTransport	node		<i>Element: ProppantTransport</i>
ReactiveCompositionalMultiphaseOBL	node		<i>Element: ReactiveCompositionalMultiphaseOBL</i>
SinglePhaseFVM	node		<i>Element: SinglePhaseFVM</i>
SinglePhaseHybridFVM	node		<i>Element: SinglePhaseHybridFVM</i>
SinglePhasePoromechanics	node		<i>Element: SinglePhasePoromechanics</i>
SinglePhasePoromechanicsEmbeddedFractures	node		<i>Element: SinglePhasePoromechanicsEmbeddedFractures</i>
SinglePhasePoromechanicsReservoir	node		<i>Element: SinglePhasePoromechanicsReservoir</i>
SinglePhaseProppantFVM	node		<i>Element: SinglePhaseProppantFVM</i>
SinglePhaseReservoir	node		<i>Element: SinglePhaseReservoir</i>
SinglePhaseWell	node		<i>Element: SinglePhaseWell</i>
SolidMechanicsEmbeddedFractures	node		<i>Element: SolidMechanicsEmbeddedFractures</i>
SolidMechanicsLagrangianSSLE	node		<i>Element: SolidMechanicsLagrangianSSLE</i>
SolidMechanics_LagrangianFEM	node		<i>Element: SolidMechanics_LagrangianFEM</i>
SurfaceGenerator	node		<i>Element: SurfaceGenerator</i>

## 10.136 Element: SourceFlux

Name	Type	De- fault	Description
bcApplicationTable- Name	string		Name of table that specifies the on/off application of the boundary condition.
beginTime	real64	- 1e+99	Time at which the boundary condition will start being applied.
component	integer	-1	Component of field (if tensor) to apply boundary condition to.
direction	R1Tensor	{0,0,0}	Direction to apply boundary condition to.
endTime	real64	1e+99	Time at which the boundary condition will stop being applied.
fieldName	string		Name of field that boundary condition is applied to.
functionName	string		Name of function that specifies variation of the boundary condition.
initialCondition	integer	0	Boundary condition is applied as an initial condition.
logLevel	integer	0	Log level
name	string	re- quired	A name is required for any non-unique nodes
objectPath	string		Path to the target field
scale	real64	0	Scale factor for value of the boundary condition.
setNames	string_array	re- quired	Name of sets that boundary condition is applied to.



## 10.137 Element: SurfaceElementRegion

Name	Type	Default	Description
defaultAperture	real64	required	The default aperture of newly formed surface elements.
faceBlock	string	FractureSubRegion	The name of the face block in the mesh, or the embedded surface.
materialList	string_array	required	List of materials present in this region
meshBody	string		Mesh body that contains this region
name	string	required	A name is required for any non-unique nodes
subRegionType	geosx_SurfaceElementRegion or faceSubRegionType	faceSubRegionType	Type of surface element subregion. Valid options: * faceElement * embeddedElement

## 10.138 Element: SurfaceGenerator

Name	Type	Default	Description
cflFactor	real64	0.5	Factor to apply to the CFL condition when calculating the maximum allowable time step. Values should be in the interval (0,1]
fractureRegion	string	Fracture	(no description available)
initialDt	real64	1e+99	Initial time-step value required by the solver to the event manager.
logLevel	integer	0	Log level
mpiCommOrder	integer	0	Flag to enable MPI consistent communication ordering
name	string	required	A name is required for any non-unique nodes
nodeBasedSIF	integer	0	Flag for choosing between node or edge based criteria: 1 for node based criterion
rockToughness	real64	required	Rock toughness of the solid material
targetRegions	string array	required	Allowable regions that the solver may be applied to. Note that this does not indicate that the solver will be applied to these regions, only that allocation will occur such that the solver may be applied to these regions. The decision about what regions this solver will be applied to rests in the EventManager.
LinearSolverParameters	node	unique	<i>Element: LinearSolverParameters</i>
NonlinearSolverParameters	node	unique	<i>Element: NonlinearSolverParameters</i>

## 10.139 Element: SymbolicFunction

Name	Type	De- fault	Description
expression	string	re- quired	Symbolic math expression
inputVar- Names	string_array	{ }	Name of fields are input to function.
name	string	re- quired	A name is required for any non-unique nodes
variable- Names	string_array	re- quired	List of variables in expression. The order must match the evaluate argument



## 10.140 Element: TableCapillaryPressure

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes
nonWettingIntermediateCapPressureTableName	string		<p>Capillary pressure table [Pa] for the pair (non-wetting phase, intermediate phase)</p> <p>Note that this input is only used for three-phase flow.</p> <p>If you want to do a two-phase simulation, please use instead wettingNonWettingCapPressureTableName to specify the table names</p>
phaseNames	string_array	required	List of fluid phases
wettingIntermediateCapPressureTableName	string		<p>Capillary pressure table [Pa] for the pair (wetting phase, intermediate phase)</p> <p>Note that this input is only used for three-phase flow.</p> <p>If you want to do a two-phase simulation, please use instead wettingNonWettingCapPressureTableName to specify the table names</p>
wettingNonWettingCapPressureTableName	string		<p>Capillary pressure table [Pa] for the pair (wetting phase, non-wetting phase)</p> <p>Note that this input is only used for two-phase flow.</p> <p>If you want to do a three-phase simulation, please use instead wettingIntermediateCapPressureTableName and nonWettingIntermediateCapPressureTableName to specify the table names</p>

## 10.141 Element: TableFunction

Name	Type	Default	Description
coordinateFiles	path_array	{}	List of coordinate file names for ND Table
coordinates	real64_array	{0}	Coordinates inputs for 1D tables
inputVarNames	string_array	{}	Name of fields are input to function.
interpolation	geosx_TableFunction_InterpolationType		Interpolation method. Valid options: * linear * nearest * upper * lower
name	string	required	A name is required for any non-unique nodes
values	real64_array	{0}	Values for 1D tables
voxelFile	path		Voxel file name for ND Table



## 10.142 Element: TableRelativePermeability

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes
nonWettingIntermediateRelPermTableNames	string_array	{}	<p>List of relative permeability tables for the pair (non-wetting phase, intermediate phase)</p> <p>The expected format is “{ nonWetting-PhaseRelPermTableName, intermediatePhaseRelPermTableName }”, in that order</p> <p>Note that this input is only used for three-phase flow.</p> <p>If you want to do a two-phase simulation, please use instead wettingNonWettingRelPermTableNames to specify the table names</p>
phaseNames	string_array	required	List of fluid phases
wettingIntermediateRelPermTableNames	string_array	{}	<p>List of relative permeability tables for the pair (wetting phase, intermediate phase)</p> <p>The expected format is “{ wetting-PhaseRelPermTableName, intermediatePhaseRelPermTableName }”, in that order</p> <p>Note that this input is only used for three-phase flow.</p> <p>If you want to do a two-phase simulation, please use instead wettingNonWettingRelPermTableNames to specify the table names</p>
wettingNonWettingRelPermTableNames	string_array	{}	<p>List of relative permeability tables for the pair (wetting phase,</p>





## 10.143 Element: TableRelativePermeabilityHysteresis

Name	Type	Default	Description
drainageNonWettingIntermediatePhaseRelPermTableNames	string	{}	<p>List of drainage relative permeability tables for the pair (non-wetting phase, intermediate phase)</p> <p>The expected format is “{ nonWetting-PhaseRelPermTableName, intermediatePhaseRelPermTableName }”, in that order</p> <p>Note that this input is only used for three-phase flow.</p> <p>If you want to do a two-phase simulation, please use instead drainageWettingNonWettingRelPermTableNames to specify the table names</p>
drainageWettingIntermediatePhaseRelPermTableNames	string	{}	<p>List of drainage relative permeability tables for the pair (wetting phase, intermediate phase)</p> <p>The expected format is “{ wetting-PhaseRelPermTableName, intermediatePhaseRelPermTableName }”, in that order</p> <p>Note that this input is only used for three-phase flow.</p> <p>If you want to do a two-phase simulation, please use instead drainageWettingNonWettingRelPermTableNames to specify the table names</p>
drainageWettingNonWettingRelPermTableNames	string	{}	<p>List of drainage relative permeability tables for the pair (wetting phase, non-wetting phase)</p> <p>The expected format is “{ wetting-</p>

## 10.144 Element: Tasks

Name	Type	Default	Description
CompositionalMultiphaseStatistics	node		<i>Element: CompositionalMultiphaseStatistics</i>
PVTDriver	node		<i>Element: PVTDriver</i>
PackCollection	node		<i>Element: PackCollection</i>
SinglePhaseStatistics	node		<i>Element: SinglePhaseStatistics</i>
SolidMechanicsStateReset	node		<i>Element: SolidMechanicsStateReset</i>
SolidMechanicsStatistics	node		<i>Element: SolidMechanicsStatistics</i>
TriaxialDriver	node		<i>Element: TriaxialDriver</i>

## 10.145 Element: ThermalCompressibleSinglePhaseFluid

Name	Type	Default	Description
compressibility	real64	0	Fluid compressibility
defaultDensity	real64	required	Default value for density.
defaultViscosity	real64	required	Default value for viscosity.
densityModelType	geosx_constitutive_ExponentialApproximationType		Type of density model. Valid options: * exponential * linear * quadratic
internalEnergyModelType	geosx_constitutive_ExponentialApproximationType		Type of internal energy model. Valid options: * exponential * linear * quadratic
name	string	required	A name is required for any non-unique nodes
referenceDensity	real64	1000	Reference fluid density
referenceInternalEnergy	real64	0.001	Reference fluid internal energy
referencePressure	real64	0	Reference pressure
referenceTemperature	real64	0	Reference temperature
referenceViscosity	real64	0.001	Reference fluid viscosity
thermalExpansionCoeff	real64	0	Fluid thermal expansion coefficient. Unit: 1/K
viscosibility	real64	0	Fluid viscosity exponential coefficient
viscosityModelType	geosx_constitutive_ExponentialApproximationType		Type of viscosity model. Valid options: * exponential * linear * quadratic
volumetricHeatCapacity	real64	0	Fluid volumetric heat capacity. Unit: J/kg/K

## 10.146 Element: ThickPlane

Name	Type	Default	Description
name	string	required	A name is required for any non-unique nodes
normal	R1Tensor	required	Normal (n_x,n_y,n_z) to the plane (will be normalized automatically)
origin	R1Tensor	required	Origin point (x,y,z) of the plane (basically, any point on the plane)
thickness	real64	required	The total thickness of the plane (with half to each side)

## 10.147 Element: TimeHistory

Name	Type	Default	Description
childDirectory	string		Child directory path
filename	string	TimeHistory	The filename to which to write time history output.
format	string	hdf	The output file format for time history output.
name	string	required	A name is required for any non-unique nodes
parallelThreads	integer	1	Number of plot files.
sources	string_array	required	A list of collectors from which to collect and output time history information.

## 10.148 Element: Traction

Name	Type	Default	Description
bcApplicationTableName	string		Name of table that specifies the on/off application of the boundary condition.
beginTime	real64	-1e+99	Time at which the boundary condition will start being applied.
direction	R1Tensor	{0,0,0}	Direction to apply boundary condition to.
endTime	real64	1e+99	Time at which the boundary condition will stop being applied.
functionName	string		Name of function that specifies variation of the boundary condition.
initialCondition	integer	0	Boundary condition is applied as an initial condition.
inputStress	R2SymTensor	{0,0,0,0,0,0}	Input stress for traction-Type = stress
logLevel	integer	0	Log level
name	string	required	A name is required for any non-unique nodes
objectPath	string		Path to the target field
scale	real64	0	Scale factor for value of the boundary condition.
setNames	string_array	required	Name of sets that boundary condition is applied to.
tractionType	geosx_TractionBoundaryCondition	vector_TractionType	Type of traction boundary condition. Options are: vector - traction is applied to the faces as specified from the scale and direction, normal - traction is applied to the faces as a pressure specified from the product of scale and the outward face normal, stress - traction is applied to the faces as specified by the inner product of input stress and face normal.

## 10.149 Element: TriaxialDriver

Name	Type	Default	Description
axialControl	string	required	Function controlling axial stress or strain (depending on test mode)
baseline	path	none	Baseline file
initialStress	real64	required	Initial stress (scalar used to set an isotropic stress state)
logLevel	integer	0	Log level
material	string	required	Solid material to test
mode	string	required	Test mode [stressControl, strainControl, mixedControl]
name	string	required	A name is required for any non-unique nodes
output	string	none	Output file
radialControl	string	required	Function controlling radial stress or strain (depending on test mode)
steps	integer	required	Number of load steps to take

## 10.150 Element: TwoPointFluxApproximation

Name	Type	Default	Description
areaRelTol	real64	1e-08	Relative tolerance for area calculations.
meanPermCoefficient	real64	1	(no description available)
name	string	required	A name is required for any non-unique nodes
usePEDFM	integer	0	(no description available)

## 10.151 Element: VTK

Name	Type	Default	Description
childDirectory	string		Child directory path
fieldNames	string_array	{ }	Names of the fields to output. If this attribute is specified, GEOSX outputs all the fields specified by the user, regardless of their <i>plotLevel</i>
format	geosx_vtk_VTKOutputFormat	binary	Output data format. Valid options: binary, ascii
name	string	required	A name is required for any non-unique nodes
onlyPlot-Specified-FieldNames	integer	0	If this flag is equal to 1, then we only plot the fields listed in <i>fieldNames</i> . Otherwise, we plot all the fields with the required <i>plotLevel</i> , plus the fields listed in <i>fieldNames</i>
outputRegionType	geosx_vtk_VTKRegionType	cell	Output region types. Valid options: cell, well, surface, all
parallelThreads	integer	1	Number of plot files.
plotFileRoot	string	VTK	Name of the root file for this output.
plotLevel	integer	1	Level detail plot. Only fields with lower of equal plot level will be output.
writeFEM-Faces	integer	0	(no description available)

## 10.152 Element: VTKMesh

Name	Type	Default	Description
field-NamesIn-GEOSX	string_array{ }		Names of fields in GEOSX to import into
field-sToImport	string_array{ }		Fields to be imported from the external mesh file
file	path	required	Path to the mesh file
logLevel	integer	0	Log level
name	string	required	A name is required for any non-unique nodes
node-set-Names	string_array{ }		Names of the VTK nodesets to import
partition-Method	geosx_VTKMeshMethod (PartitionMethod)		Method to partition the mesh
partition-Re-fine-ment	integer	1	Number of partitioning refinement iterations (defaults to 1, recommended value). A value of 0 disables graph partitioning and keeps simple kd-tree partitions (not recommended). Values higher than 1 may lead to slightly improved partitioning, but yield diminishing returns.
region-Attribute	string	attribute	Name of the VTK cell attribute to use as region marker
scale	R1Tensor	{1,1,1}	Scale the coordinates of the vertices by given scale factors (after translation)
translate	R1Tensor	{0,0,0}	Translate the coordinates of the vertices by a given vector (prior to scaling)
use-GlobalIds	integer	0	Controls the use of global IDs in the input file for cells and points. If set to 0 (default value), the GlobalId arrays in the input mesh are used if available, and generated otherwise. If set to a negative value, the GlobalId arrays in the input mesh are not used, and generated global Ids are automatically generated. If set to a positive value, the GlobalId arrays in the input mesh are used and required, and the simulation aborts if they are not available



## 10.153 Element: VanGenuchtenBakerRelativePermeability

Name	Type	Default	Description
gasOilRelPermExponentInv	real64_array	{0.5}	Rel perm power law exponent inverse for the pair (gas phase, oil phase) at residual water saturation The expected format is “{ gasExp, oilExp }”, in that order
gasOilRelPermMaxValue	real64_array	{0}	Maximum rel perm value for the pair (gas phase, oil phase) at residual water saturation The expected format is “{ gasMax, oilMax }”, in that order
name	string	required	A name is required for any non-unique nodes
phaseMinVolumeFraction	real64_array	{0}	Minimum volume fraction value for each phase
phaseNames	string_array	required	List of fluid phases
waterOilRelPermExponentInv	real64_array	{0.5}	Rel perm power law exponent inverse for the pair (water phase, oil phase) at residual gas saturation The expected format is “{ waterExp, oilExp }”, in that order
waterOilRelPermMaxValue	real64_array	{0}	Maximum rel perm value for the pair (water phase, oil phase) at residual gas saturation The expected format is “{ waterMax, oilMax }”, in that order

## 10.154 Element: VanGenuchtenCapillaryPressure

Name	Type	De- fault	Description
capPressureEpsilon	real64	1e-06	Saturation at which the extremum capillary pressure is attained; used to avoid infinite capillary pressure values for saturations close to 0 and 1
name	string	required	A name is required for any non-unique nodes
phaseCapPressureExponentInv	real64_array	{0.5}	Inverse of capillary power law exponent for each phase
phaseCapPressureMultiplier	real64_array	{1}	Entry pressure value for each phase
phaseMinVolumeFraction	real64_array	{0}	Minimum volume fraction value for each phase
phaseNames	string_array	required	List of fluid phases



## 10.155 Element: WellControls

Name	Type	Default	Description
control	geosx_WellControls_Control	required	Well control. Valid options: * BHP * phaseVolRate * totalVolRate * uninitialized
enableCrossflow	integer	1	Flag to enable crossflow. Currently only supported for injectors: - If the flag is set to 1, both reservoir-to-well flow and well-to-reservoir flow are allowed at the perforations. - If the flag is set to 0, we only allow well-to-reservoir flow at the perforations.
initialPressureCoefficient	real64	0.1	Tuning coefficient for the initial well pressure of rate-controlled wells: - Injector pressure at reference depth initialized as: $(1 + \text{initialPressureCoefficient}) * \text{reservoirPressureAtClosestPerforation} + \text{density} * g * (z_{\text{Ref}} - z_{\text{Perf}})$ - Producer pressure at reference depth initialized as: $(1 - \text{initialPressureCoefficient}) * \text{reservoirPressureAtClosestPerforation} + \text{density} * g * (z_{\text{Ref}} - z_{\text{Perf}})$
injectionStream	real64_array	{-1}	Global component densities of the injection stream [moles/m <sup>3</sup> or kg/m <sup>3</sup> ]
injectionTemperature	real64	-1	Temperature of the injection stream [K]
logLevel	integer	0	Log level
name	string	required	A name is required for any non-unique nodes
referenceElevation	real64	required	Reference elevation where

## 10.156 Element: WellElementRegion

Name	Type	Default	Description
materialList	string_array	required	List of materials present in this region
meshBody	string		Mesh body that contains this region
name	string	required	A name is required for any non-unique nodes

## 10.157 Element: lassen

Name	Type	Default	Description
Run	node	unique	<i>Element: Run</i>

## 10.158 Element: quartz

Name	Type	Default	Description
Run	node	unique	<i>Element: Run</i>

## 10.158.1 Datastructure Definitions

## 10.159 Datastructure: AcousticSEM

Name	Type	Description
indexSeismoTrace	integer	Count for output pressure at receivers
maxStableDt	real64	Value of the Maximum Stable Timestep for this solver.
meshTargets	geosx_mapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >	MeshBody/Region combinations that the solver will be applied to.
pressureNp1AtReceivers	real32_array2d	Pressure value at each receiver for each timestep
receiverConstants	real64_array2d	Constant part of the receiver for the nodes listed in m_receiverNodeIds
receiverIsLocal	integer_array	Flag that indicates whether the receiver is local to this MPI rank
receiverNodeIds	integer_array2d	Indices of the nodes (in the right order) for each receiver point
sourceConstants	real64_array2d	Constant part of the source for the nodes listed in m_sourceNodeIds
sourceIsAccessible	integer_array	Flag that indicates whether the source is accessible to this MPI rank
sourceNodeIds	integer_array2d	Indices of the nodes (in the right order) for each source point
sourceValue	real32_array2d	Source Value of the sources
usePML	integer	Flag to apply PML
LinearSolverParameters	node	<i>Datastructure: LinearSolverParameters</i>
NonlinearSolverParameters	node	<i>Datastructure: NonlinearSolverParameters</i>
SolverStatistics	node	<i>Datastructure: SolverStatistics</i>

## 10.160 Datastructure: Aquifer

Name	Type	Description
component	integer	Component of field (if tensor) to apply boundary condition to.
cumulativeFlux	real64	(no description available)
fieldName	string	Name of field that boundary condition is applied to.
objectPath	string	Path to the target field

## 10.161 Datastructure: Benchmarks

Name	Type	Description
lassen	node	<i>Datastructure: lassen</i>
quartz	node	<i>Datastructure: quartz</i>

## 10.162 Datastructure: BiotPorosity

Name	Type	Description
biotCoefficient	real64_array	Biot coefficient.
dPorosity_dPressure	real64_array2d	Derivative of rock porosity with respect to pressure
initialPorosity	real64_array2d	Initial porosity
porosity	real64_array2d	Rock porosity
porosity_n	real64_array2d	Rock porosity at the previous converged time step
referencePorosity	real64_array	Reference porosity





## 10.163 Datastructure: BlackOilFluid

Name	Type	Description
PVTO	geosx_constitutive_PVTOData	(no description available)
dPhaseCompFraction	LvArray_Array< double, 5, comp_int_seq< long, 0l, 1l, 2l, 3l, 4l >, int, LvArray_ChaiBuffer >	Derivative of phase component fraction with respect to pressure, temperature, and global component fractions
dPhaseDensity	real64_array4d	Derivative of phase density with respect to pressure, temperature, and global component fractions
dPhaseEnthalpy	real64_array4d	Derivative of phase enthalpy with respect to pressure, temperature, and global component fractions
dPhaseFraction	real64_array4d	Derivative of phase fraction with respect to pressure, temperature, and global component fractions
dPhaseInternalEnergy	real64_array4d	Derivative of phase internal energy with respect to pressure, temperature, and global component fractions
dPhaseMassDensity	real64_array4d	Derivative of phase mass density with respect to pressure, temperature, and global component fractions
dPhaseViscosity	real64_array4d	Derivative of phase viscosity with respect to pressure, temperature, and global component fractions
dTotalDensity	real64_array3d	Derivative of total density with respect to pressure, temperature, and global component fractions
formation-VolFactorTableWrappers	LvArray_Array< geosx_TableFunction_KernelWrapper, 1, comp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >	(no description available)
hydrocarbon-PhaseOrder	integer_array	(no description available)
initialTotalMassDensity	real64_array2d	Initial total mass density
phaseCompFraction	real64_array4d	Phase component fraction
phaseCompFraction_n	real64_array4d	Phase component fraction at the previous converged time step
phaseDensity	real64_array3d	Phase density
phaseDensity_n	real64_array3d	Phase density at the previous converged time step
phaseEnthalpy	real64_array3d	Phase enthalpy
phaseEnthalpy_n	real64_array3d	Phase enthalpy at the previous converged time step
phaseFraction	real64_array3d	Phase fraction
phaseInternalEnergy	real64_array3d	Phase internal energy
phaseInternalEnergy_n	real64_array3d	Phase internal energy at the previous converged time step
phaseMassDensity	real64_array3d	Phase mass density
phaseOrder	integer_array	(no description available)
phaseTypes	integer_array	(no description available)
phaseViscosity	real64_array3d	Phase viscosity

## 10.164 Datastructure: Blueprint

Name	Type	Description

## 10.165 Datastructure: BoundedPlane

Name	Type	Description

## 10.166 Datastructure: Box

Name	Type	Description
center	R1Tensor	(no description available)
cosStrike	real64	(no description available)
sinStrike	real64	(no description available)

## 10.167 Datastructure: BrooksCoreyBakerRelativePermeability

Name	Type	Description
dPhaseRelPerm_dPhaseVolFrac	real64_array	Derivative of phase relative permeability with respect to phase volume fraction
phaseOrder	integer_array	(no description available)
phaseRelPerm	real64_array	Phase relative permeability
phaseTrappedVol-Fraction	real64_array	Phase trapped volume fraction
phaseTypes	integer_array	(no description available)
volFracScale	real64	Factor used to scale the phase capillary pressure, defined as: one minus the sum of the phase minimum volume fractions.

## 10.168 Datastructure: BrooksCoreyCapillaryPressure

Name	Type	Description
dPhaseCapPressure_dPhaseVolFraction	real64_array	Derivative of phase capillary pressure with respect to phase volume fraction
phaseCapPressure	real64_array	Phase capillary pressure
phaseOrder	integer_array	(no description available)
phaseTypes	integer_array	(no description available)
volFracScale	real64	Factor used to scale the phase capillary pressure, defined as: one minus the sum of the phase minimum volume fractions.

## 10.169 Datastructure: BrooksCoreyRelativePermeability

Name	Type	Description
dPhaseRelPerm_dPhaseVolFraction	real64_array	Derivative of phase relative permeability with respect to phase volume fraction
phaseOrder	integer_array	(no description available)
phaseRelPerm	real64_array	Phase relative permeability
phaseTrappedVol-Fraction	real64_array	Phase trapped volume fraction
phaseTypes	integer_array	(no description available)
volFracScale	real64	Factor used to scale the phase relative permeability, defined as: one minus the sum of the phase minimum volume fractions.



## 10.170 Datastructure: CO2BrineEzrokhiFluid

Name	Type	Description
dPhaseCompFraction	LvArray_Array< double, 5, comp_int_seq< long, 0l, 1l, 2l, 3l, 4l >, int, LvArray_ChaiBuffer >	Derivative of phase component fraction with respect to pressure, temperature, and global component fractions
dPhaseDensity	real64_array4d	Derivative of phase density with respect to pressure, temperature, and global component fractions
dPhaseEnthalpy	real64_array4d	Derivative of phase enthalpy with respect to pressure, temperature, and global component fractions
dPhaseFraction	real64_array4d	Derivative of phase fraction with respect to pressure, temperature, and global component fractions
dPhaseInternalEnergy	real64_array4d	Derivative of phase internal energy with respect to pressure, temperature, and global component fractions
dPhaseMassDensity	real64_array4d	Derivative of phase mass density with respect to pressure, temperature, and global component fractions
dPhaseViscosity	real64_array4d	Derivative of phase viscosity with respect to pressure, temperature, and global component fractions
dTotalDensity	real64_array3d	Derivative of total density with respect to pressure, temperature, and global component fractions
initialTotalMassDensity	real64_array2d	Initial total mass density
phaseCompFraction	real64_array4d	Phase component fraction
phaseCompFraction_n	real64_array4d	Phase component fraction at the previous converged time step
phaseDensity	real64_array3d	Phase density
phaseDensity_n	real64_array3d	Phase density at the previous converged time step
phaseEnthalpy	real64_array3d	Phase enthalpy
phaseEnthalpy_n	real64_array3d	Phase enthalpy at the previous converged time step
phaseFraction	real64_array3d	Phase fraction
phaseInternalEnergy	real64_array3d	Phase internal energy
phaseInternalEnergy_n	real64_array3d	Phase internal energy at the previous converged time step
phaseMassDensity	real64_array3d	Phase mass density
phaseViscosity	real64_array3d	Phase viscosity
totalDensity	real64_array2d	Total density
totalDensity_n	real64_array2d	Total density at the previous converged time step
<b>10.170. Datastructure: CO2BrineEzrokhiFluid</b>		<b>679</b>
useMass	integer	(no description available)



## 10.171 Datastructure: CO2BrineEzrokhiThermalFluid

Name	Type	Description
dPhaseCompFraction	LvArray_Array< double, 5, comp_int_seq< long, 0l, 1l, 2l, 3l, 4l >, int, LvArray_ChaiBuffer >	Derivative of phase component fraction with respect to pressure, temperature, and global component fractions
dPhaseDensity	real64_array4d	Derivative of phase density with respect to pressure, temperature, and global component fractions
dPhaseEnthalpy	real64_array4d	Derivative of phase enthalpy with respect to pressure, temperature, and global component fractions
dPhaseFraction	real64_array4d	Derivative of phase fraction with respect to pressure, temperature, and global component fractions
dPhaseInternalEnergy	real64_array4d	Derivative of phase internal energy with respect to pressure, temperature, and global component fractions
dPhaseMassDensity	real64_array4d	Derivative of phase mass density with respect to pressure, temperature, and global component fractions
dPhaseViscosity	real64_array4d	Derivative of phase viscosity with respect to pressure, temperature, and global component fractions
dTotalDensity	real64_array3d	Derivative of total density with respect to pressure, temperature, and global component fractions
initialTotalMassDensity	real64_array2d	Initial total mass density
phaseCompFraction	real64_array4d	Phase component fraction
phaseCompFraction_n	real64_array4d	Phase component fraction at the previous converged time step
phaseDensity	real64_array3d	Phase density
phaseDensity_n	real64_array3d	Phase density at the previous converged time step
phaseEnthalpy	real64_array3d	Phase enthalpy
phaseEnthalpy_n	real64_array3d	Phase enthalpy at the previous converged time step
phaseFraction	real64_array3d	Phase fraction
phaseInternalEnergy	real64_array3d	Phase internal energy
phaseInternalEnergy_n	real64_array3d	Phase internal energy at the previous converged time step
phaseMassDensity	real64_array3d	Phase mass density
phaseViscosity	real64_array3d	Phase viscosity
totalDensity	real64_array2d	Total density
totalDensity_n	real64_array2d	Total density at the previous converged time step
useMass	integer	(no description available)





## 10.172 Datastructure: CO2BrinePhillipsFluid

Name	Type	Description
dPhaseCompFraction	LvArray_Array< double, 5, comp_int_seq< long, 0l, 1l, 2l, 3l, 4l >, int, LvArray_ChaiBuffer >	Derivative of phase component fraction with respect to pressure, temperature, and global component fractions
dPhaseDensity	real64_array4d	Derivative of phase density with respect to pressure, temperature, and global component fractions
dPhaseEnthalpy	real64_array4d	Derivative of phase enthalpy with respect to pressure, temperature, and global component fractions
dPhaseFraction	real64_array4d	Derivative of phase fraction with respect to pressure, temperature, and global component fractions
dPhaseInternalEnergy	real64_array4d	Derivative of phase internal energy with respect to pressure, temperature, and global component fractions
dPhaseMassDensity	real64_array4d	Derivative of phase mass density with respect to pressure, temperature, and global component fractions
dPhaseViscosity	real64_array4d	Derivative of phase viscosity with respect to pressure, temperature, and global component fractions
dTotalDensity	real64_array3d	Derivative of total density with respect to pressure, temperature, and global component fractions
initialTotalMassDensity	real64_array2d	Initial total mass density
phaseCompFraction	real64_array4d	Phase component fraction
phaseCompFraction_n	real64_array4d	Phase component fraction at the previous converged time step
phaseDensity	real64_array3d	Phase density
phaseDensity_n	real64_array3d	Phase density at the previous converged time step
phaseEnthalpy	real64_array3d	Phase enthalpy
phaseEnthalpy_n	real64_array3d	Phase enthalpy at the previous converged time step
phaseFraction	real64_array3d	Phase fraction
phaseInternalEnergy	real64_array3d	Phase internal energy
phaseInternalEnergy_n	real64_array3d	Phase internal energy at the previous converged time step
phaseMassDensity	real64_array3d	Phase mass density
phaseViscosity	real64_array3d	Phase viscosity
totalDensity	real64_array2d	Total density
totalDensity_n	real64_array2d	Total density at the previous converged time step
<b>10.172. Datastructure: CO2BrinePhillipsFluid</b>		<b>683</b>
useMass	integer	(no description available)



## 10.173 Datastructure: CO2BrinePhillipsThermalFluid

Name	Type	Description
dPhaseCompFraction	LvArray_Array< double, 5, comp_int_seq< long, 0l, 1l, 2l, 3l, 4l >, int, LvArray_ChaiBuffer >	Derivative of phase component fraction with respect to pressure, temperature, and global component fractions
dPhaseDensity	real64_array4d	Derivative of phase density with respect to pressure, temperature, and global component fractions
dPhaseEnthalpy	real64_array4d	Derivative of phase enthalpy with respect to pressure, temperature, and global component fractions
dPhaseFraction	real64_array4d	Derivative of phase fraction with respect to pressure, temperature, and global component fractions
dPhaseInternalEnergy	real64_array4d	Derivative of phase internal energy with respect to pressure, temperature, and global component fractions
dPhaseMassDensity	real64_array4d	Derivative of phase mass density with respect to pressure, temperature, and global component fractions
dPhaseViscosity	real64_array4d	Derivative of phase viscosity with respect to pressure, temperature, and global component fractions
dTotalDensity	real64_array3d	Derivative of total density with respect to pressure, temperature, and global component fractions
initialTotalMassDensity	real64_array2d	Initial total mass density
phaseCompFraction	real64_array4d	Phase component fraction
phaseCompFraction_n	real64_array4d	Phase component fraction at the previous converged time step
phaseDensity	real64_array3d	Phase density
phaseDensity_n	real64_array3d	Phase density at the previous converged time step
phaseEnthalpy	real64_array3d	Phase enthalpy
phaseEnthalpy_n	real64_array3d	Phase enthalpy at the previous converged time step
phaseFraction	real64_array3d	Phase fraction
phaseInternalEnergy	real64_array3d	Phase internal energy
phaseInternalEnergy_n	real64_array3d	Phase internal energy at the previous converged time step
phaseMassDensity	real64_array3d	Phase mass density
phaseViscosity	real64_array3d	Phase viscosity
totalDensity	real64_array2d	Total density
totalDensity_n	real64_array2d	Total density at the previous converged time step
useMass	integer	(no description available)

## 10.174 Datastructure: CarmanKozenyPermeability

Name	Type	Description
dPerm_dPorosity	real64_array3d	(no description available)
dPerm_dPressure	real64_array3d	Derivative of rock permeability with respect to pressure
permeability	real64_array3d	Rock permeability

## 10.175 Datastructure: CellElementRegion

Name	Type	Description
domainBoundaryIndicator	integer_array	(no description available)
ghostRank	integer_array	(no description available)
globalToLocalMap	geosx_mapBase< long long, int, std_integral_constant< bool, false > >	(no description available)
isExternal	integer_array	(no description available)
localToGlobalMap	globalIndex_array	Array that contains a map from localIndex to globalIndex.
elementSubRegions	node	<i>Datastructure: elementSubRegions</i>
neighborData	node	<i>Datastructure: neighborData</i>
sets	node	<i>Datastructure: sets</i>

## 10.176 Datastructure: ChomboIO

Name	Type	Description

## 10.177 Datastructure: CompositeFunction

Name	Type	Description

## 10.178 Datastructure: CompositionalMultiphaseFVM

Name	Type	Description
maxStableDt	real64	Value of the Maximum Stable Timestep for this solver.
meshTargets	geosx_mapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >	MeshBody/Region combinations that the solver will be applied to.
Linear-SolverParameters	node	<i>Datastructure: Linear-SolverParameters</i>
Nonlinear-SolverParameters	node	<i>Datastructure: Nonlinear-SolverParameters</i>
Solver-Statistics	node	<i>Datastructure: Solver-Statistics</i>



## 10.179 Datastructure: CompositionalMultiphaseFluid

Name	Type	Description
dPhaseCompFraction	LvArray_Array< double, 5, comp_int_seq< long, 0l, 1l, 2l, 3l, 4l >, int, LvArray_ChaiBuffer >	Derivative of phase component fraction with respect to pressure, temperature, and global component fractions
dPhaseDensity	real64_array4d	Derivative of phase density with respect to pressure, temperature, and global component fractions
dPhaseEnthalpy	real64_array4d	Derivative of phase enthalpy with respect to pressure, temperature, and global component fractions
dPhaseFraction	real64_array4d	Derivative of phase fraction with respect to pressure, temperature, and global component fractions
dPhaseInternalEnergy	real64_array4d	Derivative of phase internal energy with respect to pressure, temperature, and global component fractions
dPhaseMassDensity	real64_array4d	Derivative of phase mass density with respect to pressure, temperature, and global component fractions
dPhaseViscosity	real64_array4d	Derivative of phase viscosity with respect to pressure, temperature, and global component fractions
dTotalDensity	real64_array3d	Derivative of total density with respect to pressure, temperature, and global component fractions
initialTotalMassDensity	real64_array2d	Initial total mass density
phaseCompFraction	real64_array4d	Phase component fraction
phaseCompFraction_n	real64_array4d	Phase component fraction at the previous converged time step
phaseDensity	real64_array3d	Phase density
phaseDensity_n	real64_array3d	Phase density at the previous converged time step
phaseEnthalpy	real64_array3d	Phase enthalpy
phaseEnthalpy_n	real64_array3d	Phase enthalpy at the previous converged time step
phaseFraction	real64_array3d	Phase fraction
phaseInternalEnergy	real64_array3d	Phase internal energy
phaseInternalEnergy_n	real64_array3d	Phase internal energy at the previous converged time step
phaseMassDensity	real64_array3d	Phase mass density
phaseViscosity	real64_array3d	Phase viscosity
totalDensity	real64_array2d	Total density
totalDensity_n	real64_array2d	Total density at the previous converged time step
useMass	integer	(no description available)

## 10.180 Datastructure: CompositionalMultiphaseHybridFVM

Name	Type	Registered On	Description
maxStableDt	real64		Value of the Maximum Stable Timestep for this solver.
meshTargets	geosx_mapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >		MeshBody/Region combinations that the solver will be applied to.
facePressure_n	real64_array	<i>Datastructure: faceManager</i>	Face pressure at the previous converged time step
mim-Gravity-Coefficient	real64_array	<i>Datastructure: faceManager</i>	Mimetic gravity coefficient
Linear-SolverParameters	node		<i>Datastructure: Linear-SolverParameters</i>
Non-linear-SolverParameters	node		<i>Datastructure: NonlinearSolverParameters</i>
Solver-Statistics	node		<i>Datastructure: Solver-Statistics</i>



## 10.181 Datastructure: CompositionalMultiphaseReservoir

Name	Type	Description
discretization	string	Name of discretization object (defined in the <i>Numerical Methods</i> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <i>Finite Element Discretization</i> should be specified. If this is a Finite Volume Method, the name of a <i>Finite Volume Discretization</i> discretization should be specified.
maxStableTime	double	Value of the Maximum Stable Timestep for this solver.
meshTargets	std::map< string, std::pair< string, long, 0l >, int, LvArray_ChainBuffer >, std::integral_constant< bool, true > >	MeshBody/Region combinations that the solver will be applied to.
LinearSolverParameters	node	<i>Datastructure: LinearSolverParameters</i>
NonlinearSolverParameters	node	<i>Datastructure: NonlinearSolverParameters</i>
SolverStatistics	node	<i>Datastructure: SolverStatistics</i>

## 10.182 Datastructure: CompositionalMultiphaseStatistics

Name	Type	Description

## 10.183 Datastructure: CompositionalMultiphaseWell

Name	Type	Description
discretization	string	Name of discretization object (defined in the <i>Numerical Methods</i> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <i>Finite Element Discretization</i> should be specified. If this is a Finite Volume Method, the name of a <i>Finite Volume Discretization</i> discretization should be specified.
maxStableTime	double	Value of the Maximum Stable Timestep for this solver.
meshTargets	std::map< std::pair< string, string >, LvArray< string, 1, camp_int_seq< long, 0l >, int, LvArray< char, 1, camp_int_seq< long, 0l >, std::integral_constant< bool, true > >>	MeshBody/Region combinations that the solver will be applied to.
LinearSolverParameters	node	<i>Datastructure: LinearSolverParameters</i>
NonlinearSolverParameters	node	<i>Datastructure: NonlinearSolverParameters</i>
SolverStatistics	node	<i>Datastructure: SolverStatistics</i>
WellControls	node	<i>Datastructure: WellControls</i>

## 10.184 Datastructure: CompressibleSinglePhaseFluid

Name	Type	Description
dDensity_dPressure	real64_array2d	Derivative of density with respect to pressure
dDensity_dTemperature	real64_array2d	Derivative of density with respect to temperature
dEnthalpy_dPressure	real64_array2d	Derivative of enthalpy with respect to pressure
dEnthalpy_dTemperature	real64_array2d	Derivative of enthalpy with respect to temperature
dInternalEnergy_dPressure	real64_array2d	Derivative of internal energy with respect to pressure
dInternalEnergy_dTemperature	real64_array2d	Derivative of internal energy with respect to temperature
dViscosity_dPressure	real64_array2d	Derivative of viscosity with respect to pressure
dViscosity_dTemperature	real64_array2d	Derivative of viscosity with respect to temperature
density	real64_array2d	Density
density_n	real64_array2d	Density at the previous converged time step
enthalpy	real64_array2d	Enthalpy
initialDensity	real64_array2d	Initial density
internalEnergy	real64_array2d	Internal energy
internalEnergy_n	real64_array2d	Fluid internal energy at the previous converged step
viscosity	real64_array2d	Viscosity

## 10.185 Datastructure: CompressibleSolidCarmanKozenyPermeability

Name	Type	Description

## 10.186 Datastructure: CompressibleSolidConstantPermeability

Name	Type	Description

## 10.187 Datastructure: CompressibleSolidParallelPlatesPermeability

Name	Type	Description

## 10.188 Datastructure: CompressibleSolidSlipDependentPermeability

Name	Type	Description

## 10.189 Datastructure: ConstantPermeability

Name	Type	Description
dPerm_dPressure	real64_array3d	Derivative of rock permeability with respect to pressure
permeability	real64_array3d	Rock permeability

## 10.190 Datastructure: Constitutive

Name	Type	Description
BiotPorosity	node	<i>Datastructure: BiotPorosity</i>
BlackOilFluid	node	<i>Datastructure: BlackOilFluid</i>
BrooksCoreyBakerRelativePermeability	node	<i>Datastructure: BrooksCoreyBakerRelativePermeability</i>
BrooksCoreyCapillaryPressure	node	<i>Datastructure: BrooksCoreyCapillaryPressure</i>
BrooksCoreyRelativePermeability	node	<i>Datastructure: BrooksCoreyRelativePermeability</i>
CO2BrineEzrokhiFluid	node	<i>Datastructure: CO2BrineEzrokhiFluid</i>
CO2BrineEzrokhiThermalFluid	node	<i>Datastructure: CO2BrineEzrokhiThermalFluid</i>
CO2BrinePhillipsFluid	node	<i>Datastructure: CO2BrinePhillipsFluid</i>
CO2BrinePhillipsThermalFluid	node	<i>Datastructure: CO2BrinePhillipsThermalFluid</i>
CarmanKozenyPermeability	node	<i>Datastructure: CarmanKozenyPermeability</i>
CompositionalMultiphaseFluid	node	<i>Datastructure: CompositionalMultiphaseFluid</i>
CompressibleSinglePhaseFluid	node	<i>Datastructure: CompressibleSinglePhaseFluid</i>
CompressibleSolidCarmanKozenyPermeability	node	<i>Datastructure: CompressibleSolidCarmanKozenyPermeability</i>
CompressibleSolidConstantPermeability	node	<i>Datastructure: CompressibleSolidConstantPermeability</i>
CompressibleSolidParallelPlatesPermeability	node	<i>Datastructure: CompressibleSolidParallelPlatesPermeability</i>
CompressibleSolidSlipDependentPermeability	node	<i>Datastructure: CompressibleSolidSlipDependentPermeability</i>
ConstantPermeability	node	<i>Datastructure: ConstantPermeability</i>
Coulomb	node	<i>Datastructure: Coulomb</i>
DamageElasticIsotropic	node	<i>Datastructure: DamageElasticIsotropic</i>
DamageSpectralElasticIsotropic	node	<i>Datastructure: DamageSpectralElasticIsotropic</i>
DamageVolDevElasticIsotropic	node	<i>Datastructure: DamageVolDevElasticIsotropic</i>
DeadOilFluid	node	<i>Datastructure: DeadOilFluid</i>
DelftEgg	node	<i>Datastructure: DelftEgg</i>
DruckerPrager	node	<i>Datastructure: DruckerPrager</i>
ElasticIsotropic	node	<i>Datastructure: ElasticIsotropic</i>
ElasticIsotropicPressureDependent	node	<i>Datastructure: ElasticIsotropicPressureDependent</i>
ElasticOrthotropic	node	<i>Datastructure: ElasticOrthotropic</i>
ElasticTransverseIsotropic	node	<i>Datastructure: ElasticTransverseIsotropic</i>
ExtendedDruckerPrager	node	<i>Datastructure: ExtendedDruckerPrager</i>
FrictionlessContact	node	<i>Datastructure: FrictionlessContact</i>
JFunctionCapillaryPressure	node	<i>Datastructure: JFunctionCapillaryPressure</i>
ModifiedCamClay	node	<i>Datastructure: ModifiedCamClay</i>
MultiPhaseConstantThermalConductivity	node	<i>Datastructure: MultiPhaseConstantThermalConductivity</i>
MultiPhaseVolumeWeightedThermalConductivity	node	<i>Datastructure: MultiPhaseVolumeWeightedThermalConductivity</i>
NullModel	node	<i>Datastructure: NullModel</i>
ParallelPlatesPermeability	node	<i>Datastructure: ParallelPlatesPermeability</i>
ParticleFluid	node	<i>Datastructure: ParticleFluid</i>
PermeabilityBase	node	<i>Datastructure: PermeabilityBase</i>
PorousDelftEgg	node	<i>Datastructure: PorousDelftEgg</i>

Continued on next page

Table 10.3 – continued from previous page

Name	Type	Description
PorousDruckerPrager	node	<i>Datastructure: PorousDruckerPrager</i>
PorousElasticIsotropic	node	<i>Datastructure: PorousElasticIsotropic</i>
PorousElasticOrthotropic	node	<i>Datastructure: PorousElasticOrthotropic</i>
PorousElasticTransverseIsotropic	node	<i>Datastructure: PorousElasticTransverseIsotropic</i>
PorousExtendedDruckerPrager	node	<i>Datastructure: PorousExtendedDruckerPrager</i>
PorousModifiedCamClay	node	<i>Datastructure: PorousModifiedCamClay</i>
PressurePorosity	node	<i>Datastructure: PressurePorosity</i>
ProppantPermeability	node	<i>Datastructure: ProppantPermeability</i>
ProppantPorosity	node	<i>Datastructure: ProppantPorosity</i>
ProppantSlurryFluid	node	<i>Datastructure: ProppantSlurryFluid</i>
ProppantSolidProppantPermeability	node	<i>Datastructure: ProppantSolidProppantPermeability</i>
SinglePhaseConstantThermalConductivity	node	<i>Datastructure: SinglePhaseConstantThermalConductivity</i>
SlipDependentPermeability	node	<i>Datastructure: SlipDependentPermeability</i>
SolidInternalEnergy	node	<i>Datastructure: SolidInternalEnergy</i>
TableCapillaryPressure	node	<i>Datastructure: TableCapillaryPressure</i>
TableRelativePermeability	node	<i>Datastructure: TableRelativePermeability</i>
TableRelativePermeabilityHysteresis	node	<i>Datastructure: TableRelativePermeabilityHysteresis</i>
ThermalCompressibleSinglePhaseFluid	node	<i>Datastructure: ThermalCompressibleSinglePhaseFluid</i>
VanGenuchtenBakerRelativePermeability	node	<i>Datastructure: VanGenuchtenBakerRelativePermeability</i>
VanGenuchtenCapillaryPressure	node	<i>Datastructure: VanGenuchtenCapillaryPressure</i>

## 10.191 Datastructure: ConstitutiveModels

Name	Type	Description

## 10.192 Datastructure: Coulomb

Name	Type	Description
elasticSlip	real64_array2d	Elastic Slip

## 10.193 Datastructure: Cylinder

Name	Type	Description

## 10.194 Datastructure: DamageElasticIsotropic

Name	Type	Description
bulkModulus	real64_array	Elastic Bulk Modulus Field
damage	real64_array2d	Material Damage Variable
density	real64_array2d	Material Density
extDrivingForce	real64_array2d	External Driving Force
oldStress	real64_array3d	Previous Material Stress
shearModulus	real64_array	Elastic Shear Modulus Field
strainEnergyDensity	real64_array2d	Strain Energy Density
stress	real64_array3d	Current Material Stress

## 10.195 Datastructure: DamageSpectralElasticIsotropic

Name	Type	Description
bulkModulus	real64_array	Elastic Bulk Modulus Field
damage	real64_array2d	Material Damage Variable
density	real64_array2d	Material Density
extDrivingForce	real64_array2d	External Driving Force
oldStress	real64_array3d	Previous Material Stress
shearModulus	real64_array	Elastic Shear Modulus Field
strainEnergyDensity	real64_array2d	Strain Energy Density
stress	real64_array3d	Current Material Stress

## 10.196 Datastructure: DamageVolDevElasticIsotropic

Name	Type	Description
bulkModulus	real64_array	Elastic Bulk Modulus Field
damage	real64_array2d	Material Damage Variable
density	real64_array2d	Material Density
extDrivingForce	real64_array2d	External Driving Force
oldStress	real64_array3d	Previous Material Stress
shearModulus	real64_array	Elastic Shear Modulus Field
strainEnergyDensity	real64_array2d	Strain Energy Density
stress	real64_array3d	Current Material Stress



## 10.197 Datastructure: DeadOilFluid

Name	Type	Description
dPhaseC-ompFraction	LvArray_Array< double, 5, camp_int_seq< long, 0l, 1l, 2l, 3l, 4l >, int, LvArray_ChaiBuffer >	Derivative of phase component fraction with respect to pressure, temperature, and global component fractions
dPhaseDensity	real64_array4d	Derivative of phase density with respect to pressure, temperature, and global component fractions
dPhaseEnthalpy	real64_array4d	Derivative of phase enthalpy with respect to pressure, temperature, and global component fractions
dPhaseFraction	real64_array4d	Derivative of phase fraction with respect to pressure, temperature, and global component fractions
dPhaseInternalEnergy	real64_array4d	Derivative of phase internal energy with respect to pressure, temperature, and global component fractions
dPhaseMassDensity	real64_array4d	Derivative of phase mass density with respect to pressure, temperature, and global component fractions
dPhaseViscosity	real64_array4d	Derivative of phase viscosity with respect to pressure, temperature, and global component fractions
dTotalDensity	real64_array3d	Derivative of total density with respect to pressure, temperature, and global component fractions
formation-VolFactorTableWrappers	LvArray_Array< geosx_TableFunction_KernelWrapper, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >	(no description available)
hydrocarbon-PhaseOrder	integer_array	(no description available)
initialTotalMassDensity	real64_array2d	Initial total mass density
phaseC-ompFraction	real64_array4d	Phase component fraction
phaseC-ompFraction_n	real64_array4d	Phase component fraction at the previous converged time step
phaseDensity	real64_array3d	Phase density
phaseDensity_n	real64_array3d	Phase density at the previous converged time step
phaseEnthalpy	real64_array3d	Phase enthalpy
phaseEnthalpy_n	real64_array3d	Phase enthalpy at the previous converged time step
phaseFraction	real64_array3d	Phase fraction
phaseInternalEnergy	real64_array3d	Phase internal energy
phaseInternalEnergy_n	real64_array3d	Phase internal energy at the previous converged time step
phaseMassDensity	real64_array3d	Phase mass density
phaseOrder	integer_array	(no description available)
phaseTypes	integer_array	(no description available)
phaseViscosity	real64_array3d	Phase viscosity



## 10.198 Datastructure: DelftEgg

Name	Type	Description
bulkModulus	real64_array	Elastic Bulk Modulus Field
csiSlope	real64_array	Slope of the critical state line
density	real64_array2d	Material Density
oldPreConsolidationPressure	real64_array2d	Old preconsolidation pressure
oldStress	real64_array3d	Previous Material Stress
preConsolidationPressure	real64_array2d	New preconsolidation pressure
recompressionIndex	real64_array	Recompression index
shapeParameter	real64_array	Shape parameter for the yield surface
shearModulus	real64_array	Elastic Shear Modulus Field
stress	real64_array3d	Current Material Stress
virginCompressionIndex	real64_array	Virgin compression index

## 10.199 Datastructure: Dirichlet

Name	Type	Description

## 10.200 Datastructure: DruckerPrager

Name	Type	Description
bulkModulus	real64_array	Elastic Bulk Modulus Field
cohesion	real64_array2d	New cohesion state
density	real64_array2d	Material Density
dilation	real64_array	Plastic potential slope
friction	real64_array	Yield surface slope
hardening	real64_array	Hardening rate
oldCohesion	real64_array2d	Old cohesion state
oldStress	real64_array3d	Previous Material Stress
shearModulus	real64_array	Elastic Shear Modulus Field
stress	real64_array3d	Current Material Stress

## 10.201 Datastructure: ElasticIsotropic

Name	Type	Description
bulkModulus	real64_array	Elastic Bulk Modulus Field
density	real64_array2d	Material Density
oldStress	real64_array3d	Previous Material Stress
shearModulus	real64_array	Elastic Shear Modulus Field
stress	real64_array3d	Current Material Stress

## 10.202 Datastructure: ElasticIsotropicPressureDependent

Name	Type	Description
density	real64_array2d	Material Density
oldStress	real64_array3d	Previous Material Stress
recompressionIndex	real64_array	Recompression Index Field
refPressure	real64	Reference Pressure Field
refStrainVol	real64	Reference Volumetric Strain
shearModulus	real64_array	Elastic Shear Modulus
stress	real64_array3d	Current Material Stress

## 10.203 Datastructure: ElasticOrthotropic

Name	Type	Description
c11	real64_array	Elastic Stiffness Field C11
c12	real64_array	Elastic Stiffness Field C12
c13	real64_array	Elastic Stiffness Field C13
c22	real64_array	Elastic Stiffness Field C22
c23	real64_array	Elastic Stiffness Field C23
c33	real64_array	Elastic Stiffness Field C33
c44	real64_array	Elastic Stiffness Field C44
c55	real64_array	Elastic Stiffness Field C55
c66	real64_array	Elastic Stiffness Field C66
density	real64_array2d	Material Density
oldStress	real64_array3d	Previous Material Stress
stress	real64_array3d	Current Material Stress

## 10.204 Datastructure: ElasticSEM

Name	Type	Description
displacementXNp1AtReceivers	real32_array2d	Displacement value at each receiver for each timestep (x-component)
displacementYNp1AtReceivers	real32_array2d	Displacement value at each receiver for each timestep (y-component)
displacementZNp1AtReceivers	real32_array2d	Displacement value at each receiver for each timestep (z-component)
indexSeismoTrace	integer	Count for output pressure at receivers
maxStableDt	real64	Value of the Maximum Stable Timestep for this solver.
meshTargets	geosx_mapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >	MeshBody/Region combinations that the solver will be applied to.
receiverIsLocal	integer_array	Flag that indicates whether the receiver is local to this MPI rank
receiverNodeIds	integer_array2d	Indices of the nodes (in the right order) for each receiver point
sourceConstants	real64_array2d	Constant part of the receiver for the nodes listed in m_receiverNodeIds
sourceIsAccessible	integer_array	Flag that indicates whether the source is accessible to this MPI rank
sourceNodeIds	integer_array2d	Indices of the nodes (in the right order) for each source point
sourceValue	real32_array2d	Source Value of the sources
usePML	integer	Flag to apply PML
Linear-SolverParameters	node	<i>Datastructure: Linear-SolverParameters</i>
Nonlinear-SolverParameters	node	<i>Datastructure: Nonlinear-SolverParameters</i>
Solver-Statistics	node	<i>Datastructure: SolverStatistics</i>

## 10.205 Datastructure: ElasticTransverselsotropic

Name	Type	Description
c11	real64_array	Elastic Stiffness Field C11
c13	real64_array	Elastic Stiffness Field C13
c33	real64_array	Elastic Stiffness Field C33
c44	real64_array	Elastic Stiffness Field C44
c66	real64_array	Elastic Stiffness Field C66
density	real64_array2d	Material Density
oldStress	real64_array3d	Previous Material Stress
stress	real64_array3d	Current Material Stress

## 10.206 Datastructure: ElementRegions

Name	Type	Description
domainBound- aryIndicator	integer_array	(no description available)
ghostRank	integer_array	(no description available)
globalToLo- calMap	geosx_mapBase< long long, int, std_integral_constant< bool, false > >	(no description available)
isExternal	integer_array	(no description available)
localToGlob- alMap	globalIndex_array	Array that contains a map from localIn- dex to globalIndex.
CellElementRe- gion	node	<i>Datastructure: CellElementRegion</i>
SurfaceElemen- tRegion	node	<i>Datastructure: SurfaceElementRegion</i>
WellElementRe- gion	node	<i>Datastructure: WellElementRegion</i>
elementRegions- Group	node	<i>Datastructure: elementRegionsGroup</i>
neighborData	node	<i>Datastructure: neighborData</i>
sets	node	<i>Datastructure: sets</i>

## 10.207 Datastructure: EmbeddedSurfaceGenerator

Name	Type	Registered On	Description
maxStableDt	real64		Value of the Maximum Stable Timestep for this solver.
meshTargets	geosx_mapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >		MeshBody/Region combinations that the solver will be applied to.
parent-EdgeIndex	integer_array	<i>Datastructure: embeddedSurfacesNodeManager</i>	Index of parent edge within the mesh object it is registered on.
Linear-Solver-Parameters	node		<i>Datastructure: LinearSolverParameters</i>
Non-linear-Solver-Parameters	node		<i>Datastructure: Non-linearSolverParameters</i>
Solver-Statistics	node		<i>Datastructure: Solver-Statistics</i>

## 10.208 Datastructure: Events

Name	Type	Description
currentSubEvent	integer	Index of the current subevent.
cycle	integer	Current simulation cycle number.
dt	real64	Current simulation timestep.
time	real64	Current simulation time.
HaltEvent	node	<i>Datastructure: HaltEvent</i>
PeriodicEvent	node	<i>Datastructure: PeriodicEvent</i>
SoloEvent	node	<i>Datastructure: SoloEvent</i>

## 10.209 Datastructure: ExtendedDruckerPrager

Name	Type	Description
bulkModulus	real64_array	Elastic Bulk Modulus Field
density	real64_array2d	Material Density
dilationRatio	real64_array	Plastic potential slope ratio
hardening	real64_array	Hardening parameter
initialFriction	real64_array	Initial yield surface slope
oldStateVariable	real64_array2d	Old equivalent plastic shear strain
oldStress	real64_array3d	Previous Material Stress
pressureIntercept	real64_array	Pressure point at cone vertex
residualFriction	real64_array	Residual yield surface slope
shearModulus	real64_array	Elastic Shear Modulus Field
stateVariable	real64_array2d	New equivalent plastic shear strain
stress	real64_array3d	Current Material Stress

## 10.210 Datastructure: FieldSpecification

Name	Type	Description

## 10.211 Datastructure: FieldSpecifications

Name	Type	Description
Aquifer	node	<i>Datastructure: Aquifer</i>
Dirichlet	node	<i>Datastructure: Dirichlet</i>
FieldSpecification	node	<i>Datastructure: FieldSpecification</i>
HydrostaticEquilibrium	node	<i>Datastructure: HydrostaticEquilibrium</i>
PML	node	<i>Datastructure: PML</i>
SourceFlux	node	<i>Datastructure: SourceFlux</i>
Traction	node	<i>Datastructure: Traction</i>

## 10.212 Datastructure: File

Name	Type	Description

## 10.213 Datastructure: FiniteElementSpace

Name	Type	Description

## 10.214 Datastructure: FiniteElements

Name	Type	Description
FiniteElementSpace	node	<i>Datastructure: FiniteElementSpace</i>
LinearSolverParameters	node	<i>Datastructure: LinearSolverParameters</i>
NonlinearSolverParameters	node	<i>Datastructure: NonlinearSolverParameters</i>

## 10.215 Datastructure: FiniteVolume

Name	Type	Description
HybridMimeticDiscretization	node	<i>Datastructure: HybridMimeticDiscretization</i>
TwoPointFluxApproximation	node	<i>Datastructure: TwoPointFluxApproximation</i>

## 10.216 Datastructure: FlowProppantTransport

Name	Type	Description
discretization	string	Name of discretization object (defined in the <i>Numerical Methods</i> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <i>Finite Element Discretization</i> should be specified. If this is a Finite Volume Method, the name of a <i>Finite Volume Discretization</i> discretization should be specified.
maxStableDeltaT	double	Value of the Maximum Stable Timestep for this solver.
meshTargets	geosx_mapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >	MeshBody/Region combinations that the solver will be applied to.
LinearSolverParameters	node	<i>Datastructure: LinearSolverParameters</i>
NonlinearSolverParameters	node	<i>Datastructure: NonlinearSolverParameters</i>
SolverStatistics	node	<i>Datastructure: SolverStatistics</i>

## 10.217 Datastructure: FrictionlessContact

Name	Type	Description

## 10.218 Datastructure: Functions

Name	Type	Description
CompositeFunction	node	<i>Datastructure: CompositeFunction</i>
MultivariableTableFunction	node	<i>Datastructure: MultivariableTableFunction</i>
SymbolicFunction	node	<i>Datastructure: SymbolicFunction</i>
TableFunction	node	<i>Datastructure: TableFunction</i>

## 10.219 Datastructure: Geometry

Name	Type	Description
BoundedPlane	node	<i>Datastructure: BoundedPlane</i>
Box	node	<i>Datastructure: Box</i>
Cylinder	node	<i>Datastructure: Cylinder</i>
ThickPlane	node	<i>Datastructure: ThickPlane</i>

## 10.220 Datastructure: HaltEvent

Name	Type	Description
currentSubEvent	integer	Index of the current subevent
eventForecast	integer	Indicates when the event is expected to execute
isTargetExecuting	integer	Index of the current subevent
lastCycle	integer	Last event occurrence (cycle)
lastTime	real64	Last event occurrence (time)
HaltEvent	node	<i>Datastructure: HaltEvent</i>
PeriodicEvent	node	<i>Datastructure: PeriodicEvent</i>
SoloEvent	node	<i>Datastructure: SoloEvent</i>

## 10.221 Datastructure: HybridMimeticDiscretization

Name	Type	Description



## 10.222 Datastructure: Hydrofracture

Name	Type	Description
discretization	string	Name of discretization object (defined in the <i>Numerical Methods</i> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <i>Finite Element Discretization</i> should be specified. If this is a Finite Volume Method, the name of a <i>Finite Volume Discretization</i> discretization should be specified.
maxStableTime	double	Value of the Maximum Stable Timestep for this solver.
meshTargets	TargetMapBase< std::pair< string, string >, LvArray< string, 1, camp_int_seq< long, 0l >, int, LvArray< char, 1, camp_int_seq< long, 0l >, std::integral_constant< bool, true > >>	MeshBody/Region combinations that the solver will be applied to.
LinearSolverParameters	node	<i>Datastructure: LinearSolverParameters</i>
NonlinearSolverParameters	node	<i>Datastructure: NonlinearSolverParameters</i>
SolverStatistics	node	<i>Datastructure: SolverStatistics</i>

## 10.223 Datastructure: HydrostaticEquilibrium

Name	Type	Description
component	integer	Component of field (if tensor) to apply boundary condition to.
fieldName	string	Name of field that boundary condition is applied to.
initialCondition	integer	Boundary condition is applied as an initial condition.
setNames	string_array	Name of sets that boundary condition is applied to.

## 10.224 Datastructure: Included

Name	Type	Description
File	node	<i>Datastructure: File</i>

## 10.225 Datastructure: InternalMesh

Name	Type	Description
meshLevels	node	<i>Datastructure: meshLevels</i>

## 10.226 Datastructure: InternalWell

Name	Type	Description
Perforation	node	<i>Datastructure: Perforation</i>

## 10.227 Datastructure: InternalWellbore

Name	Type	Description
nx	integer_array	Number of elements in the x-direction within each mesh block
ny	integer_array	Number of elements in the y-direction within each mesh block
xCoords	real64_array	x-coordinates of each mesh block vertex
yCoords	real64_array	y-coordinates of each mesh block vertex
meshLevels	node	<i>Datastructure: meshLevels</i>

## 10.228 Datastructure: JFunctionCapillaryPressure

Name	Type	Description
dPhaseCapPressure_dPhaseVolFraction	real64_array4d	Derivative of phase capillary pressure with respect to phase volume fraction
jFuncMultiplier	real64_array2d	Multiplier for the Leverett J-function
jFunctionWrappers	LvArray_Array< geosx_TableFunction_KernelWrapper, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >	(no description available)
phaseCapPressure	real64_array3d	Phase capillary pressure
phaseOrder	integer_array	(no description available)
phaseTypes	integer_array	(no description available)

## 10.229 Datastructure: LagrangianContact

Name	Type	Description
maxStableDt	real64	Value of the Maximum Stable Timestep for this solver.
meshTargets	geosx_mapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >	MeshBody/Region combinations that the solver will be applied to.
Linear-SolverParameters	node	<i>Datastructure: Linear-SolverParameters</i>
Nonlinear-SolverParameters	node	<i>Datastructure: Nonlinear-SolverParameters</i>
Solver-Statistics	node	<i>Datastructure: Solver-Statistics</i>

## 10.230 Datastructure: LaplaceFEM

Name	Type	Description
maxStableDt	real64	Value of the Maximum Stable Timestep for this solver.
meshTargets	geosx_mapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >	MeshBody/Region combinations that the solver will be applied to.
Linear-SolverParameters	node	<i>Datastructure: Linear-SolverParameters</i>
Nonlinear-SolverParameters	node	<i>Datastructure: Nonlinear-SolverParameters</i>
Solver-Statistics	node	<i>Datastructure: Solver-Statistics</i>

## 10.231 Datastructure: Level0

Name	Type	Description
meshLevel	integer	(no description available)
ElementRegions	node	<i>Datastructure: ElementRegions</i>
edgeManager	node	<i>Datastructure: edgeManager</i>
embeddedSurfacesEdgeManager	node	<i>Datastructure: embeddedSurfacesEdgeManager</i>
embeddedSurfacesNodeManager	node	<i>Datastructure: embeddedSurfacesNodeManager</i>
faceManager	node	<i>Datastructure: faceManager</i>
nodeManager	node	<i>Datastructure: nodeManager</i>

## 10.232 Datastructure: LinearSolverParameters

Name	Type	Description

## 10.233 Datastructure: Mesh

Name	Type	Description
InternalMesh	node	<i>Datastructure: InternalMesh</i>
InternalWell	node	<i>Datastructure: InternalWell</i>
InternalWellbore	node	<i>Datastructure: InternalWellbore</i>
PAMELAMesh	node	<i>Datastructure: PAMELAMesh</i>
VTKMesh	node	<i>Datastructure: VTKMesh</i>

## 10.234 Datastructure: MeshBodies

Name	Type	Description
InternalMesh	node	<i>Datastructure: InternalMesh</i>
InternalWellbore	node	<i>Datastructure: InternalWellbore</i>
PAMELAMesh	node	<i>Datastructure: PAMELAMesh</i>
VTKMesh	node	<i>Datastructure: VTKMesh</i>

## 10.235 Datastructure: ModifiedCamClay

Name	Type	Description
csISlope	real64_array	Slope of the critical state line
density	real64_array2d	Material Density
oldPreConsolidationPressure	real64_array2d	Old preconsolidation pressure
oldStress	real64_array3d	Previous Material Stress
preConsolidationPressure	real64_array2d	New preconsolidation pressure
recompressionIndex	real64_array	Recompression Index Field
refPressure	real64	Reference Pressure Field
refStrainVol	real64	Reference Volumetric Strain
shearModulus	real64_array	Elastic Shear Modulus
stress	real64_array3d	Current Material Stress
virginCompressionIndex	real64_array	Virgin compression index

## 10.236 Datastructure: MultiPhaseConstantThermalConductivity

Name	Type	Description
dEffectiveConductivity_dPhaseVolFraction	real64_array4d	Derivative of effective conductivity with respect to phase volume fraction
effectiveConductivity	real64_array3d	Effective conductivity

## 10.237 Datastructure: MultiPhaseVolumeWeightedThermalConductivity

Name	Type	Description
dEffectiveConductivity_dPhaseVolFraction	real64_array4d	Derivative of effective conductivity with respect to phase volume fraction
effectiveConductivity	real64_array3d	Effective conductivity
rockThermalConductivity	real64_array3d	Rock thermal conductivity

## 10.238 Datastructure: MultiphasePoromechanics

Name	Type	Description
discretization	string	Name of discretization object (defined in the <i>Numerical Methods</i> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <i>Finite Element Discretization</i> should be specified. If this is a Finite Volume Method, the name of a <i>Finite Volume Discretization</i> discretization should be specified.
maxStableDt	real64	Value of the Maximum Stable Timestep for this solver.
meshTargets	geosx_mapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >	MeshBody/Region combinations that the solver will be applied to.
LinearSolverParameters	node	<i>Datastructure: LinearSolverParameters</i>
NonlinearSolverParameters	node	<i>Datastructure: NonlinearSolverParameters</i>
SolverStatistics	node	<i>Datastructure: SolverStatistics</i>

## 10.239 Datastructure: MultiphasePoromechanicsReservoir

Name	Type	Description
discretization	string	Name of discretization object (defined in the <i>Numerical Methods</i> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <i>Finite Element Discretization</i> should be specified. If this is a Finite Volume Method, the name of a <i>Finite Volume Discretization</i> discretization should be specified.
maxStableTime	double	Value of the Maximum Stable Timestep for this solver.
meshTargets	geosx_mapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >	MeshBody/Region combinations that the solver will be applied to.
LinearSolverParameters	node	<i>Datastructure: LinearSolverParameters</i>
NonlinearSolverParameters	node	<i>Datastructure: NonlinearSolverParameters</i>
SolverStatistics	node	<i>Datastructure: SolverStatistics</i>

## 10.240 Datastructure: MultivariableTableFunction

Name	Type	Description

## 10.241 Datastructure: NonlinearSolverParameters

Name	Type	Description
newtonNumberOfIterations	integer	Number of Newton's iterations.

## 10.242 Datastructure: NullModel

Name	Type	Description

## 10.243 Datastructure: NumericalMethods

Name	Type	Description
FiniteElements	node	<i>Datastructure: FiniteElements</i>
FiniteVolume	node	<i>Datastructure: FiniteVolume</i>

## 10.244 Datastructure: Outputs

Name	Type	Description
Blueprint	node	<i>Datastructure: Blueprint</i>
ChomboIO	node	<i>Datastructure: ChomboIO</i>
Python	node	<i>Datastructure: Python</i>
Restart	node	<i>Datastructure: Restart</i>
Silo	node	<i>Datastructure: Silo</i>
TimeHistory	node	<i>Datastructure: TimeHistory</i>
VTK	node	<i>Datastructure: VTK</i>

## 10.245 Datastructure: PAMELAMesh

Name	Type	Description
meshLevels	node	<i>Datastructure: meshLevels</i>

## 10.246 Datastructure: PML

Name	Type	Description
fieldName	string	Name of field that boundary condition is applied to.
initialCondition	integer	Boundary condition is applied as an initial condition.

## 10.247 Datastructure: PVTDriver

Name	Type	Description

## 10.248 Datastructure: PackCollection

Name	Type	Description

## 10.249 Datastructure: ParallelPlatesPermeability

Name	Type	Description
dPerm_dDispJump	real64_array4d	Derivative of rock permeability with respect to displacement jump
dPerm_dPressure	real64_array3d	Derivative of rock permeability with respect to pressure
permeability	real64_array3d	Rock permeability

## 10.250 Datastructure: Parameter

Name	Type	Description

## 10.251 Datastructure: Parameters

Name	Type	Description
Parameter	node	<i>Datastructure: Parameter</i>

## 10.252 Datastructure: ParticleFluid

Name	Type	Description
collisionFactor	real64_array	Collision factor
dCollisionFactor_dProppantConcentration	real64_array	Derivative of collision factor with respect to proppant concentration
dSettlingFactor_dComponentConcentration	real64_array2d	Derivative of settling factor with respect to component concentration
dSettlingFactor_dPressure	real64_array	Derivative of settling factor with respect to pressure
dSettlingFactor_dProppantConcentration	real64_array	Derivative of settling factor with respect to proppant concentration
proppantPackPermeability	real64_array	Proppant pack permeability
settlingFactor	real64_array	Settling factor

## 10.253 Datastructure: Perforation

Name	Type	Description



## 10.254 Datastructure: PeriodicEvent

Name	Type	Description
currentSubEvent	integer	Index of the current subevent
eventForecast	integer	Indicates when the event is expected to execute
isTargetExecuting	integer	Index of the current subevent
lastCycle	integer	Last event occurrence (cycle)
lastTime	real64	Last event occurrence (time)
HaltEvent	node	<i>Datastructure: HaltEvent</i>
PeriodicEvent	node	<i>Datastructure: PeriodicEvent</i>
SoloEvent	node	<i>Datastructure: SoloEvent</i>

## 10.255 Datastructure: PermeabilityBase

Name	Type	Description
dPerm_dPressure	real64_array3d	Derivative of rock permeability with respect to pressure
permeability	real64_array3d	Rock permeability

## 10.256 Datastructure: PhaseFieldDamageFEM

Name	Type	Description
maxStableDt	real64	Value of the Maximum Stable Timestep for this solver.
meshTargets	geosx_mapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >	MeshBody/Region combinations that the solver will be applied to.
Linear-SolverParameters	node	<i>Datastructure: Linear-SolverParameters</i>
Nonlinear-SolverParameters	node	<i>Datastructure: Nonlinear-SolverParameters</i>
Solver-Statistics	node	<i>Datastructure: Solver-Statistics</i>

## 10.257 Datastructure: PhaseFieldFracture

Name	Type	Description
maxStableDt	real64	Value of the Maximum Stable Timestep for this solver.
meshTargets	geosx_mapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >	MeshBody/Region combinations that the solver will be applied to.
Linear-SolverParameters	node	<i>Datastructure: Linear-SolverParameters</i>
Nonlinear-SolverParameters	node	<i>Datastructure: Nonlinear-SolverParameters</i>
Solver-Statistics	node	<i>Datastructure: Solver-Statistics</i>

## 10.258 Datastructure: PorousDelftEgg

Name	Type	Description

## 10.259 Datastructure: PorousDruckerPrager

Name	Type	Description

## 10.260 Datastructure: PorousElasticIsotropic

Name	Type	Description

## 10.261 Datastructure: PorousElasticOrthotropic

Name	Type	Description

## 10.262 Datastructure: PorousElasticTransverselsotropic

Name	Type	Description

## 10.263 Datastructure: PorousExtendedDruckerPrager

Name	Type	Description

## 10.264 Datastructure: PorousModifiedCamClay

Name	Type	Description

## 10.265 Datastructure: PressurePorosity

Name	Type	Description
dPorosity_dPressure	real64_array2d	Derivative of rock porosity with respect to pressure
initialPorosity	real64_array2d	Initial porosity
porosity	real64_array2d	Rock porosity
porosity_n	real64_array2d	Rock porosity at the previous converged time step
referencePorosity	real64_array	Reference porosity

## 10.266 Datastructure: Problem

Name	Type	Description
Benchmarks	node	<i>Datastructure: Benchmarks</i>
Constitutive	node	<i>Datastructure: Constitutive</i>
ElementRegions	node	<i>Datastructure: ElementRegions</i>
Events	node	<i>Datastructure: Events</i>
FieldSpecifications	node	<i>Datastructure: FieldSpecifications</i>
Functions	node	<i>Datastructure: Functions</i>
Geometry	node	<i>Datastructure: Geometry</i>
Included	node	<i>Datastructure: Included</i>
Mesh	node	<i>Datastructure: Mesh</i>
NumericalMethods	node	<i>Datastructure: NumericalMethods</i>
Outputs	node	<i>Datastructure: Outputs</i>
Parameters	node	<i>Datastructure: Parameters</i>
Solvers	node	<i>Datastructure: Solvers</i>
Tasks	node	<i>Datastructure: Tasks</i>
commandLine	node	<i>Datastructure: commandLine</i>
domain	node	<i>Datastructure: domain</i>

## 10.267 Datastructure: ProppantPermeability

Name	Type	Description
dPerm_dDispJump	real64_array4d	Derivative of rock permeability with respect to displacement jump
dPerm_dPressure	real64_array3d	Derivative of rock permeability with respect to pressure
permeability	real64_array3d	Rock permeability
permeabilityMultiplier	real64_array3d	Rock permeability multiplier
proppantPackPermeability	real64	(no description available)

## 10.268 Datastructure: ProppantPorosity

Name	Type	Description
dPorosity_dPressure	real64_array2d	Derivative of rock porosity with respect to pressure
initialPorosity	real64_array2d	Initial porosity
porosity	real64_array2d	Rock porosity
porosity_n	real64_array2d	Rock porosity at the previous converged time step
referencePorosity	real64_array	Reference porosity

## 10.269 Datastructure: ProppantSlurryFluid

Name	Type	Description
FluidDensity	real64_array2d	Fluid density
FluidViscosity	real64_array2d	Fluid viscosity
componentDensity	real64_array3d	Component density
dCompDens_dCompConc	real64_array4d	Derivative of component density with respect to component concentration
dCompDens_dPres	real64_array3d	Derivative of component density with respect to pressure
dDens_dCompConc	real64_array3d	Derivative of density with respect to component concentration
dDens_dProppantConc	real64_array2d	Derivative of density with respect to proppant concentration
dDensity_dPressure	real64_array2d	Derivative of density with respect to pressure
dDensity_dTemperature	real64_array2d	Derivative of density with respect to temperature
dEnthalpy_dPressure	real64_array2d	Derivative of enthalpy with respect to pressure
dEnthalpy_dTemperature	real64_array2d	Derivative of enthalpy with respect to temperature
dFluidDens_dCompConc	real64_array3d	Derivative of fluid density with respect to component concentration
dFluidDens_dPres	real64_array2d	Derivative of fluid density with respect to pressure
dFluidVisc_dCompConc	real64_array3d	Derivative of fluid viscosity with respect to component concentration
dFluidVisc_dPres	real64_array2d	Derivative of fluid viscosity with respect to pressure
dInternalEnergy_dPressure	real64_array2d	Derivative of internal energy with respect to pressure
dInternalEnergy_dTemperature	real64_array2d	Derivative of internal energy with respect to temperature
dVisc_dCompConc	real64_array3d	Derivative of viscosity with respect to component concentration
dVisc_dProppantConc	real64_array2d	Derivative of viscosity with respect to proppant concentration
dViscosity_dPressure	real64_array2d	Derivative of viscosity with respect to pressure
dViscosity_dTemperature	real64_array2d	Derivative of viscosity with respect to temperature
density	real64_array2d	Density
density_n	real64_array2d	Density at the previous converged time step
enthalpy	real64_array2d	Enthalpy
initialDensity	real64_array2d	Initial density
internalEnergy	real64_array2d	Internal energy
internalEnergy_n	real64_array2d	Fluid internal energy at the previous converged step
viscosity	real64_array2d	Viscosity

## 10.270 Datastructure: ProppantSolidProppantPermeability

Name	Type	Description

## 10.271 Datastructure: ProppantTransport

Name	Type	Description
maxStableDt	real64	Value of the Maximum Stable Timestep for this solver.
meshTargets	geosx_mapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >	MeshBody/Region combinations that the solver will be applied to.
Linear-SolverParameters	node	<i>Datastructure: Linear-SolverParameters</i>
Nonlinear-SolverParameters	node	<i>Datastructure: Nonlinear-SolverParameters</i>
Solver-Statistics	node	<i>Datastructure: Solver-Statistics</i>

## 10.272 Datastructure: Python

Name	Type	Description

## 10.273 Datastructure: ReactiveCompositionalMultiphaseOBL

Name	Type	Description
maxStableDt	real64	Value of the Maximum Stable Timestep for this solver.
meshTargets	geosx_mapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >	MeshBody/Region combinations that the solver will be applied to.
Linear-SolverParameters	node	<i>Datastructure: Linear-SolverParameters</i>
Nonlinear-SolverParameters	node	<i>Datastructure: Nonlinear-SolverParameters</i>
Solver-Statistics	node	<i>Datastructure: Solver-Statistics</i>

## 10.274 Datastructure: Restart

Name	Type	Description

## 10.275 Datastructure: Run

Name	Type	Description

## 10.276 Datastructure: Silo

Name	Type	Description

## 10.277 Datastructure: SinglePhaseConstantThermalConductivity

Name	Type	Description
effectiveConductivity	real64_array3d	Effective conductivity

## 10.278 Datastructure: SinglePhaseFVM

Name	Type	Description
maxStableDt	real64	Value of the Maximum Stable Timestep for this solver.
meshTargets	geosx_mapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >	MeshBody/Region combinations that the solver will be applied to.
Linear-SolverParameters	node	<i>Datastructure: Linear-SolverParameters</i>
Nonlinear-SolverParameters	node	<i>Datastructure: Nonlinear-SolverParameters</i>
Solver-Statistics	node	<i>Datastructure: Solver-Statistics</i>

## 10.279 Datastructure: SinglePhaseHybridFVM

Name	Type	Registered On	Description
maxStableDt	real64		Value of the Maximum Stable Timestep for this solver.
meshTargets	geosx_mapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >		MeshBody/Region combinations that the solver will be applied to.
facePressure_n	real64_array	<i>Datastructure: faceManager</i>	Face pressure at the previous converged time step
Linear-SolverParameters	node		<i>Datastructure: Linear-SolverParameters</i>
Non-linear-SolverParameters	node		<i>Datastructure: NonlinearSolverParameters</i>
Solver-Statistics	node		<i>Datastructure: Solver-Statistics</i>



## 10.280 Datastructure: SinglePhasePoromechanics

Name	Type	Description
discretization	string	Name of discretization object (defined in the <i>Numerical Methods</i> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <i>Finite Element Discretization</i> should be specified. If this is a Finite Volume Method, the name of a <i>Finite Volume Discretization</i> discretization should be specified.
maxStableTime	double	Value of the Maximum Stable Timestep for this solver.
meshTargets	std::map< std::pair< string, string >, LvArray< string, 1, camp_int_seq< long, 0l >, int, LvArray< char, 1, camp_int_seq< long, 0l >, std::integral_constant< bool, true > >>	MeshBody/Region combinations that the solver will be applied to.
LinearSolverParameters	node	<i>Datastructure: LinearSolverParameters</i>
NonlinearSolverParameters	node	<i>Datastructure: NonlinearSolverParameters</i>
SolverStatistics	node	<i>Datastructure: SolverStatistics</i>

## 10.281 Datastructure: SinglePhasePoromechanicsEmbeddedFractures

Name	Type	Description
discretization	string	Name of discretization object (defined in the <i>Numerical Methods</i> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <i>Finite Element Discretization</i> should be specified. If this is a Finite Volume Method, the name of a <i>Finite Volume Discretization</i> discretization should be specified.
maxStableTime	double	Value of the Maximum Stable Timestep for this solver.
meshTargets	geosx_mapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >	MeshBody/Region combinations that the solver will be applied to.
LinearSolverParameters	node	<i>Datastructure: LinearSolverParameters</i>
NonlinearSolverParameters	node	<i>Datastructure: NonlinearSolverParameters</i>
SolverStatistics	node	<i>Datastructure: SolverStatistics</i>

## 10.282 Datastructure: SinglePhasePoromechanicsReservoir

Name	Type	Description
discretization	string	Name of discretization object (defined in the <i>Numerical Methods</i> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <i>Finite Element Discretization</i> should be specified. If this is a Finite Volume Method, the name of a <i>Finite Volume Discretization</i> discretization should be specified.
maxStableDt	real64	Value of the Maximum Stable Timestep for this solver.
meshTargets	geosx_mapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >	MeshBody/Region combinations that the solver will be applied to.
Linear-Solver-Parameters	node	<i>Datastructure: LinearSolverParameters</i>
Non-linear-Solver-Parameters	node	<i>Datastructure: NonlinearSolverParameters</i>
Solver-Statistics	node	<i>Datastructure: SolverStatistics</i>

## 10.283 Datastructure: SinglePhaseProppantFVM

Name	Type	Description
maxStableDt	real64	Value of the Maximum Stable Timestep for this solver.
meshTargets	geosx_mapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >	MeshBody/Region combinations that the solver will be applied to.
Linear-SolverParameters	node	<i>Datastructure: Linear-SolverParameters</i>
Nonlinear-SolverParameters	node	<i>Datastructure: Nonlinear-SolverParameters</i>
Solver-Statistics	node	<i>Datastructure: Solver-Statistics</i>

## 10.284 Datastructure: SinglePhaseReservoir

Name	Type	Description
discretization	string	Name of discretization object (defined in the <i>Numerical Methods</i> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <i>Finite Element Discretization</i> should be specified. If this is a Finite Volume Method, the name of a <i>Finite Volume Discretization</i> discretization should be specified.
maxStableDt	double	Value of the Maximum Stable Timestep for this solver.
meshTargets	TargetMapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >	MeshBody/Region combinations that the solver will be applied to.
LinearSolverParameters	node	<i>Datastructure: LinearSolverParameters</i>
NonlinearSolverParameters	node	<i>Datastructure: NonlinearSolverParameters</i>
SolverStatistics	node	<i>Datastructure: SolverStatistics</i>

## 10.285 Datastructure: SinglePhaseStatistics

Name	Type	Description

## 10.286 Datastructure: SinglePhaseWell

Name	Type	Description
discretization	string	Name of discretization object (defined in the <i>Numerical Methods</i> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <i>Finite Element Discretization</i> should be specified. If this is a Finite Volume Method, the name of a <i>Finite Volume Discretization</i> discretization should be specified.
maxStableTime	real64	Value of the Maximum Stable Timestep for this solver.
meshTargets	geosx_mapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >	MeshBody/Region combinations that the solver will be applied to.
LinearSolverParameters	node	<i>Datastructure: LinearSolverParameters</i>
NonlinearSolverParameters	node	<i>Datastructure: NonlinearSolverParameters</i>
SolverStatistics	node	<i>Datastructure: SolverStatistics</i>
WellControls	node	<i>Datastructure: WellControls</i>

## 10.287 Datastructure: SlipDependentPermeability

Name	Type	Description
dPerm_dDispJump	real64_array4d	Derivative of the permeability w.r.t. the displacement jump.
dPerm_dPressure	real64_array3d	Derivative of rock permeability with respect to pressure
permeability	real64_array3d	Rock permeability

## 10.288 Datastructure: SolidInternalEnergy

Name	Type	Description
dInternalEnergy_dTemperature	real64_array2d	Derivative of the solid internal energy w.r.t. temperature [J/(m <sup>3</sup> .K)]
internalEnergy	real64_array2d	Internal energy of the solid per unit volume [J/m <sup>3</sup> ]
internalEnergy_n	real64_array2d	Internal energy of the solid per unit volume at the previous time-step [J/m <sup>3</sup> ]

## 10.289 Datastructure: SolidMechanicsEmbeddedFractures

Name	Type	Description
discretization	string	Name of discretization object (defined in the <i>Numerical Methods</i> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <i>Finite Element Discretization</i> should be specified. If this is a Finite Volume Method, the name of a <i>Finite Volume Discretization</i> discretization should be specified.
maxStableTime	real64	Value of the Maximum Stable Timestep for this solver.
meshTargets	geosx_mapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >	MeshBody/Region combinations that the solver will be applied to.
LinearSolverParameters	node	<i>Datastructure: LinearSolverParameters</i>
NonlinearSolverParameters	node	<i>Datastructure: NonlinearSolverParameters</i>
SolverStatistics	node	<i>Datastructure: SolverStatistics</i>

## 10.290 Datastructure: SolidMechanicsLagrangianSSLE

Name	Type	Description
maxForce	real64	The maximum force contribution in the problem domain.
maxStableDt	real64	Value of the Maximum Stable Timestep for this solver.
meshTargets	geosx_mapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >	MeshBody/Region combinations that the solver will be applied to.
Linear-SolverParameters	node	<i>Datastructure: Linear-SolverParameters</i>
Nonlinear-SolverParameters	node	<i>Datastructure: Nonlinear-SolverParameters</i>
Solver-Statistics	node	<i>Datastructure: Solver-Statistics</i>

## 10.291 Datastructure: SolidMechanicsStateReset

Name	Type	Description

## 10.292 Datastructure: SolidMechanicsStatistics

Name	Type	Description

## 10.293 Datastructure: SolidMechanics\_LagrangianFEM

Name	Type	Description
maxForce	real64	The maximum force contribution in the problem domain.
maxStableDt	real64	Value of the Maximum Stable Timestep for this solver.
meshTargets	geosx_mapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >	MeshBody/Region combinations that the solver will be applied to.
Linear-SolverParameters	node	<i>Datastructure: Linear-SolverParameters</i>
Nonlinear-SolverParameters	node	<i>Datastructure: Nonlinear-SolverParameters</i>
Solver-Statistics	node	<i>Datastructure: Solver-Statistics</i>

## 10.294 Datastructure: SoloEvent

Name	Type	Description
currentSubEvent	integer	Index of the current subevent
eventForecast	integer	Indicates when the event is expected to execute
isTargetExecuting	integer	Index of the current subevent
lastCycle	integer	Last event occurrence (cycle)
lastTime	real64	Last event occurrence (time)
HaltEvent	node	<i>Datastructure: HaltEvent</i>
PeriodicEvent	node	<i>Datastructure: PeriodicEvent</i>
SoloEvent	node	<i>Datastructure: SoloEvent</i>

## 10.295 Datastructure: SolverStatistics

Name	Type	Description
numDiscardedLinearIterations	integer	Cumulative number of discarded linear iterations
numDiscardedNonlinearIterations	integer	Cumulative number of discarded nonlinear iterations
numDiscardedOuterLoopIterations	integer	Cumulative number of discarded outer loop iterations
numSuccessfulLinearIterations	integer	Cumulative number of successful linear iterations
numSuccessfulNonlinearIterations	integer	Cumulative number of successful nonlinear iterations
numSuccessfulOuterLoopIterations	integer	Cumulative number of successful outer loop iterations
numTimeStepCuts	integer	Number of time step cuts
numTimeSteps	integer	Number of time steps



## 10.296 Datastructure: Solvers

Name	Type	Description
AcousticSEM	node	<i>Datastructure: AcousticSEM</i>
CompositionalMultiphaseFVM	node	<i>Datastructure: CompositionalMultiphaseFVM</i>
CompositionalMultiphaseHybridFVM	node	<i>Datastructure: CompositionalMultiphaseHybridFVM</i>
CompositionalMultiphaseReservoir	node	<i>Datastructure: CompositionalMultiphaseReservoir</i>
CompositionalMultiphaseWell	node	<i>Datastructure: CompositionalMultiphaseWell</i>
ElasticSEM	node	<i>Datastructure: ElasticSEM</i>
EmbeddedSurfaceGenerator	node	<i>Datastructure: EmbeddedSurfaceGenerator</i>
FlowProppantTransport	node	<i>Datastructure: FlowProppantTransport</i>
Hydrofracture	node	<i>Datastructure: Hydrofracture</i>
LagrangianContact	node	<i>Datastructure: LagrangianContact</i>
LaplaceFEM	node	<i>Datastructure: LaplaceFEM</i>
MultiphasePoromechanics	node	<i>Datastructure: MultiphasePoromechanics</i>
MultiphasePoromechanicsReservoir	node	<i>Datastructure: MultiphasePoromechanicsReservoir</i>
PhaseFieldDamageFEM	node	<i>Datastructure: PhaseFieldDamageFEM</i>
PhaseFieldFracture	node	<i>Datastructure: PhaseFieldFracture</i>
ProppantTransport	node	<i>Datastructure: ProppantTransport</i>
ReactiveCompositionalMultiphaseOBL	node	<i>Datastructure: ReactiveCompositionalMultiphaseOBL</i>
SinglePhaseFVM	node	<i>Datastructure: SinglePhaseFVM</i>
SinglePhaseHybridFVM	node	<i>Datastructure: SinglePhaseHybridFVM</i>
SinglePhasePoromechanics	node	<i>Datastructure: SinglePhasePoromechanics</i>
SinglePhasePoromechanicsEmbeddedFractures	node	<i>Datastructure: SinglePhasePoromechanicsEmbeddedFractures</i>
SinglePhasePoromechanicsReservoir	node	<i>Datastructure: SinglePhasePoromechanicsReservoir</i>
SinglePhaseProppantFVM	node	<i>Datastructure: SinglePhaseProppantFVM</i>
SinglePhaseReservoir	node	<i>Datastructure: SinglePhaseReservoir</i>
SinglePhaseWell	node	<i>Datastructure: SinglePhaseWell</i>
SolidMechanicsEmbeddedFractures	node	<i>Datastructure: SolidMechanicsEmbeddedFractures</i>
SolidMechanicsLagrangianSSLE	node	<i>Datastructure: SolidMechanicsLagrangianSSLE</i>
SolidMechanics_LagrangianFEM	node	<i>Datastructure: SolidMechanics_LagrangianFEM</i>
SurfaceGenerator	node	<i>Datastructure: SurfaceGenerator</i>

## 10.297 Datastructure: SourceFlux

Name	Type	Description

## 10.298 Datastructure: SurfaceElementRegion

Name	Type	Description
domainBoundaryIndicator	integer_array	(no description available)
ghostRank	integer_array	(no description available)
globalToLocalMap	geosx_mapBase< long long, int, std_integral_constant< bool, false > >	(no description available)
isExternal	integer_array	(no description available)
localToGlobalMap	globalIndex_array	Array that contains a map from localIndex to globalIndex.
elementSubRegions	node	<i>Datastructure: elementSubRegions</i>
neighborData	node	<i>Datastructure: neighborData</i>
sets	node	<i>Datastructure: sets</i>

## 10.299 Datastructure: SurfaceGenerator

Name	Type	Description
discretization	string	Name of discretization object (defined in the <i>Numerical Methods</i> ) to use for this solver. For instance, if this is a Finite Element Solver, the name of a <i>Finite Element Discretization</i> should be specified. If this is a Finite Volume Method, the name of a <i>Finite Volume Discretization</i> discretization should be specified.
fail-Criterion	integer	(no description available)
maxStableTime	double	Value of the Maximum Stable Timestep for this solver.
meshTargets	geosx_mapBase< std_pair< string, string >, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >	MeshBody/Region combinations that the solver will be applied to.
tipEdges	LvArray_SortedArray< int, int, LvArray_ChaiBuffer >	Set containing all the tip edges
tipFaces	LvArray_SortedArray< int, int, LvArray_ChaiBuffer >	Set containing all the tip faces
tipNodes	LvArray_SortedArray< int, int, LvArray_ChaiBuffer >	Set containing all the nodes at the fracture tip
trailingFaces	LvArray_SortedArray< int, int, LvArray_ChaiBuffer >	Set containing all the trailing faces
LinearSolverParameters	node	<i>Datastructure: LinearSolverParameters</i>
NonlinearSolverParameters	node	<i>Datastructure: NonlinearSolverParameters</i>
SolverStatistics	node	<i>Datastructure: SolverStatistics</i>

## 10.300 Datastructure: SymbolicFunction

Name	Type	Description

## 10.301 Datastructure: TableCapillaryPressure

Name	Type	Description
capPresWrappers	LvArray_Array< geosx_TableFunction_KernelWrapper, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >	(no description available)
dPhaseCapPressure_dPhaseVolFraction	real64_array4d	Derivative of phase capillary pressure with respect to phase volume fraction
phaseCapPressure	real64_array3d	Phase capillary pressure
phaseOrder	integer_array	(no description available)
phaseTypes	integer_array	(no description available)

## 10.302 Datastructure: TableFunction

Name	Type	Description

## 10.303 Datastructure: TableRelativePermeability

Name	Type	Description
dPhaseRelPerm_dPhaseVolFraction	real64_array4d	Derivative of phase relative permeability with respect to phase volume fraction
phaseMinVolumeFraction	real64_array	(no description available)
phaseOrder	integer_array	(no description available)
phaseRelPerm	real64_array3d	Phase relative permeability
phaseTrapped-VolFraction	real64_array3d	Phase trapped volume fraction
phaseTypes	integer_array	(no description available)
relPermWrappers	LvArray_Array< geosx_TableFunction_KernelWrapper, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >	(no description available)

## 10.304 Datastructure: TableRelativePermeabilityHysteresis

Name	Type	Description
dPhaseRelPerm_dPhaseVolFraction	real64_array	Derivative of phase relative permeability with respect to phase volume fraction
drainagePhaseMaxVolumeFraction	real64_array	(no description available)
drainagePhaseMinVolumeFraction	real64_array	(no description available)
drainagePhaseRelPermEndPoint	real64_array	(no description available)
drainageRelPermWrappers	LvArray_Array< geosx_TableFunction_KernelWrapper, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >	(no description available)
imbibitionPhaseMaxVolumeFraction	real64_array	(no description available)
imbibitionPhaseMinVolumeFraction	real64_array	(no description available)
imbibitionPhaseRelPermEndPoint	real64_array	(no description available)
imbibitionRelPermWrappers	LvArray_Array< geosx_TableFunction_KernelWrapper, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >	(no description available)
landParameter	real64_array	(no description available)
phaseHasHysteresis	integer_array	(no description available)
phaseMaxHistoricalVolFraction	real64_array2d	Phase max historical phase volume fraction
phaseMinHistoricalVolFraction	real64_array2d	Phase min historical phase volume fraction
phaseOrder	integer_array	(no description available)
phaseRelPerm	real64_array3d	Phase relative permeability
phaseTrappedVolFraction	real64_array3d	Phase trapped volume fraction
phaseTypes	integer_array	(no description available)

## 10.305 Datastructure: Tasks

Name	Type	Description
CompositionalMultiphaseStatistics	node	<i>Datastructure: CompositionalMultiphaseStatistics</i>
PVTDriver	node	<i>Datastructure: PVTDriver</i>
PackCollection	node	<i>Datastructure: PackCollection</i>
SinglePhaseStatistics	node	<i>Datastructure: SinglePhaseStatistics</i>
SolidMechanicsStateReset	node	<i>Datastructure: SolidMechanicsStateReset</i>
SolidMechanicsStatistics	node	<i>Datastructure: SolidMechanicsStatistics</i>
TriaxialDriver	node	<i>Datastructure: TriaxialDriver</i>

## 10.306 Datastructure: ThermalCompressibleSinglePhaseFluid

Name	Type	Description
dDensity_dPressure	real64_array2d	Derivative of density with respect to pressure
dDensity_dTemperature	real64_array2d	Derivative of density with respect to temperature
dEnthalpy_dPressure	real64_array2d	Derivative of enthalpy with respect to pressure
dEnthalpy_dTemperature	real64_array2d	Derivative of enthalpy with respect to temperature
dInternalEnergy_dPressure	real64_array2d	Derivative of internal energy with respect to pressure
dInternalEnergy_dTemperature	real64_array2d	Derivative of internal energy with respect to temperature
dViscosity_dPressure	real64_array2d	Derivative of viscosity with respect to pressure
dViscosity_dTemperature	real64_array2d	Derivative of viscosity with respect to temperature
density	real64_array2d	Density
density_n	real64_array2d	Density at the previous converged time step
enthalpy	real64_array2d	Enthalpy
initialDensity	real64_array2d	Initial density
internalEnergy	real64_array2d	Internal energy
internalEnergy_n	real64_array2d	Fluid internal energy at the previous converged step
viscosity	real64_array2d	Viscosity

## 10.307 Datastructure: ThickPlane

Name	Type	Description

## 10.308 Datastructure: TimeHistory

Name	Type	Description
restart	integer	The current history record to be written, on restart from an earlier time allows use to remove invalid future history.

## 10.309 Datastructure: Traction

Name	Type	Description
component	integer	Component of field (if tensor) to apply boundary condition to.
fieldName	string	Name of field that boundary condition is applied to.

## 10.310 Datastructure: TriaxialDriver

Name	Type	Description

## 10.311 Datastructure: TwoPointFluxApproximation

Name	Type	Description
cellStencil	geosx_CellElementStencilTPFA	(no description available)
coefficient-Name	string	Name of coefficient field
edfmStencil	geosx_EmbeddedSurfaceToCellStencil	(no description available)
faceElement-ToCellStencil	geosx_FaceElementToCellStencil	(no description available)
fieldName	string	Name of primary solution field
fractureStencil	geosx_SurfaceElementStencil	(no description available)
targetRegions	geosx_mapBase< string, LvArray_Array< string, 1, camp_int_seq< long, 0l >, int, LvArray_ChaiBuffer >, std_integral_constant< bool, true > >	List of regions to build the stencil for

## 10.312 Datastructure: VTK

Name	Type	Description

## 10.313 Datastructure: VTKMesh

Name	Type	Description
meshLevels	node	<i>Datastructure: meshLevels</i>

### 10.314 Datastructure: VanGenuchtenBakerRelativePermeability

Name	Type	Description
dPhaseRelPerm_dPhaseVolFraction	real64_array	Derivative of phase relative permeability with respect to phase volume fraction
phaseOrder	integer_array	(no description available)
phaseRelPerm	real64_array	Phase relative permeability
phaseTrappedVol-Fraction	real64_array	Phase trapped volume fraction
phaseTypes	integer_array	(no description available)
volFracScale	real64	Factor used to scale the phase capillary pressure, defined as: one minus the sum of the phase minimum volume fractions.

### 10.315 Datastructure: VanGenuchtenCapillaryPressure

Name	Type	Description
dPhaseCapPressure_dPhaseVolFraction	real64_array	Derivative of phase capillary pressure with respect to phase volume fraction
phaseCapPressure	real64_array	Phase capillary pressure
phaseOrder	integer_array	(no description available)
phaseTypes	integer_array	(no description available)
volFracScale	real64	Factor used to scale the phase capillary pressure, defined as: one minus the sum of the phase minimum volume fractions.

### 10.316 Datastructure: WellControls

Name	Type	Description
currentControl	geosx_WellControls_Control	Current well control



## 10.317 Datastructure: WellElementRegion

Name	Type	Description
domainBoundaryIndicator	integer_array	(no description available)
ghostRank	integer_array	(no description available)
globalToLocalMap	geosx_mapBase< long long, int, std_integral_constant< bool, false > >	(no description available)
isExternal	integer_array	(no description available)
localToGlobalMap	globalIndex_array	Array that contains a map from localIndex to globalIndex.
wellControlName	string	(no description available)
wellGeneratorName	string	(no description available)
elementSubRegions	node	<i>Datastructure: elementSubRegions</i>
neighborData	node	<i>Datastructure: neighborData</i>
sets	node	<i>Datastructure: sets</i>

## 10.318 Datastructure: WellElementRegionUniqueSubRegion

Name	Type	Description
domainBoundaryIndicator	integer_array	(no description available)
elementCenter	real64_array2d	(no description available)
elementVolume	real64_array	(no description available)
ghostRank	integer_array	(no description available)
globalToLocalMap	geosx_mapBase< long long, int, std_integral_constant< bool, false > >	(no description available)
isExternal	integer_array	(no description available)
localToGlobalMap	globalIndex_array	Array that contains a map from localIndex to globalIndex.
nextWellElementIndex	integer_array	(no description available)
nextWellElementIndex-Global	integer_array	(no description available)
nodeList	geosx_InterObjectRelation< LvArray_Array< int, 2, camp_int_seq< long, 0l, 1l >, int, LvArray_ChaiBuffer > >	(no description available)
numEdges-PerElement	integer	(no description available)
numFaces-PerElement	integer	(no description available)
numNodes-PerElement	integer	(no description available)
radius	real64_array	(no description available)
topRank	integer	(no description available)
topWellElementIndex	integer	(no description available)
wellControlsName	string	(no description available)
Constitutive-Models	node	<i>Datastructure: Constitutive-Models</i>
neighborData	node	<i>Datastructure: neighborData</i>
sets	node	<i>Datastructure: sets</i>
wellElementSubRegion	node	<i>Datastructure: wellElementSubRegion</i>

## 10.319 Datastructure: commandLine

Name	Type	Description
beginFrom-Restart	integer	Flag to indicate restart run.
inputFileName	string	Name of the input xml file.
outputDirectory	string	Directory in which to put the output files, if not specified defaults to the current directory.
overridePartitionNumbers	integer	Flag to indicate partition number override
problemName	string	Used in writing the output files, if not specified defaults to the name of the input file.
restartFileName	string	Name of the restart file.
schemaFileName	string	Name of the output schema
suppressPinned	integer	Whether to disallow using pinned memory allocations for MPI communication buffers.
useNonblockingMPI	integer	Whether to prefer using non-blocking MPI communication where implemented (results in non-deterministic DOF numbering).
xPartitionsOverride	integer	Number of partitions in the x-direction
yPartitionsOverride	integer	Number of partitions in the y-direction
zPartitionsOverride	integer	Number of partitions in the z-direction

## 10.320 Datastructure: domain

Name	Type	Description
Neighbors	std_vector< geosx_NeighborCommunicator, std_allocator< geosx_NeighborCommunicator > >	(no description available)
partition-Manager	geosx_PartitionBase	(no description available)
Constitutive	node	<i>Datastructure: Constitutive</i>
MeshBodies	node	<i>Datastructure: MeshBodies</i>

## 10.321 Datastructure: edgeManager

Name	Type	Description
domain-Bound-aryIndicator	integer_array	(no description available)
faceList	geosx_InterObjectRelation< LvArray_ArrayOfSets< int, int, LvArray_ChaiBuffer > >	(no description available)
ghostRank	integer_array	(no description available)
globalToLocalMap	geosx_mapBase< long long, int, std_integral_constant< bool, false > >	(no description available)
isExternal	integer_array	(no description available)
localToGlobalMap	globalIndex_array	Array that contains a map from localIndex to globalIndex.
nodeList	geosx_InterObjectRelation< LvArray_Array< int, 2, camp_int_seq< long, 0l, 1l >, int, LvArray_ChaiBuffer > >	(no description available)
neighborData	node	<i>Datastructure: neighborData</i>
sets	node	<i>Datastructure: sets</i>

## 10.322 Datastructure: elementRegionsGroup

Name	Type	Description

## 10.323 Datastructure: elementSubRegions

Name	Type	Description
WellElementRegionUniqueSubRegion	node	<i>Datastructure: WellElementRegionUniqueSubRegion</i>

## 10.324 Datastructure: embeddedSurfacesEdgeManager

Name	Type	Description
domain-Bound-aryIndicator	integer_array	(no description available)
faceList	geosx_InterObjectRelation< LvArray_ArrayOfSets< int, int, LvArray_ChaiBuffer > >	(no description available)
ghostRank	integer_array	(no description available)
globalToLocalMap	geosx_mapBase< long long, int, std_integral_constant< bool, false > >	(no description available)
isExternal	integer_array	(no description available)
localToGlobalMap	globalIndex_array	Array that contains a map from localIndex to globalIndex.
nodeList	geosx_InterObjectRelation< LvArray_Array< int, 2, camp_int_seq< long, 0l, 1l >, int, LvArray_ChaiBuffer > >	(no description available)
neighborData	node	<i>Datastructure: neighborData</i>
sets	node	<i>Datastructure: sets</i>

## 10.325 Datastructure: embeddedSurfacesNodeManager

Name	Type	Registered By	Description
domain-Bound-aryIndicator	integer_array		(no description available)
edgeList	geosx_InterObjectRelation< LvArray_ArrayOfSets< int, int, LvArray_ChaiBuffer > >		(no description available)
elemList	LvArray_ArrayOfArrays< int, int, LvArray_ChaiBuffer >		(no description available)
elemRegionList	LvArray_ArrayOfArrays< int, int, LvArray_ChaiBuffer >		(no description available)
elemSubRegionList	LvArray_ArrayOfArrays< int, int, LvArray_ChaiBuffer >		(no description available)
ghostRank	integer_array		(no description available)
globalToLocalMap	geosx_mapBase< long long, int, std_integral_constant< bool, false > >		(no description available)
isExternal	integer_array		(no description available)
local-ToGlobalMap	globalIndex_array		Array that contains a map from localIndex to globalIndex.
parentEdge-GlobalIndex	globalIndex_array		(no description available)
reference-Position	real64_array2d		(no description available)
parent-EdgeIndex	integer_array	<i>Datastructure: EmbeddedSurface-Generator</i>	Index of parent edge within the mesh object it is registered on.
neighbor-Data	node		<i>Datastructure: neighborData</i>
sets	node		<i>Datastructure: sets</i>

## 10.326 Datastructure: faceManager

Name	Type	Registered By	Description
domain-Bound-aryIndica-tor	integer_array		(no description avail-able)
edgeList	geosx_InterObjectRelation<LvArray_ArrayOfArrays< int, int, LvArray_ChaiBuffer > >		(no description avail-able)
elemList	integer_array2d		(no description avail-able)
elemRe-gionList	integer_array2d		(no description avail-able)
elemSub-Region-List	integer_array2d		(no description avail-able)
faceArea	real64_array		(no description avail-able)
faceCen-ter	real64_array2d		(no description avail-able)
faceNor-mal	real64_array2d		(no description avail-able)
ghos-tRank	integer_array		(no description avail-able)
global-ToLo-calMap	geosx_mapBase< long long, int, std_integral_constant< bool, false > >		(no description avail-able)
isExternal	integer_array		(no description avail-able)
local-ToGlob-alMap	globalIndex_array		Array that contains a map from localIndex to globalIndex.
nodeList	geosx_InterObjectRelation<LvArray_ArrayOfArrays< int, int, LvArray_ChaiBuffer > >		(no description avail-able)
facePres-sure_n	real64_array	<i>Datastructure: CompositionalMulti-phaseHybridFVM, Datastructure: SinglePhaseHybridFVM</i>	Face pressure at the previous converged time step
mim-Gravity-Coeffi-cient	real64_array	<i>Datastructure: CompositionalMulti-phaseHybridFVM</i>	Mimetic gravity coef-ficient
neighbor-Data	node		<i>Datastructure: neigh-borData</i>
sets	node		<i>Datastructure: sets</i>

### 10.327 Datastructure: lassen

Name	Type	Description
Run	node	<i>Datastructure: Run</i>

### 10.328 Datastructure: meshLevels

Name	Type	Description
Level0	node	<i>Datastructure: Level0</i>

### 10.329 Datastructure: neighborData

Name	Type	Description

### 10.330 Datastructure: nodeManager

Name	Type	Description
ReferencePosition	real64_array2d	(no description available)
domainBoundaryIndicator	integer_array	(no description available)
edgeList	geosx_InterObjectRelation< LvArray_ArrayOfSets< int, int, LvArray_ChaiBuffer > >	(no description available)
elemList	LvArray_ArrayOfArrays< int, int, LvArray_ChaiBuffer >	(no description available)
elemRegionList	LvArray_ArrayOfArrays< int, int, LvArray_ChaiBuffer >	(no description available)
elemSubRegionList	LvArray_ArrayOfArrays< int, int, LvArray_ChaiBuffer >	(no description available)
faceList	geosx_InterObjectRelation< LvArray_ArrayOfSets< int, int, LvArray_ChaiBuffer > >	(no description available)
ghostRank	integer_array	(no description available)
globalToLocalMap	geosx_mapBase< long long, int, std_integral_constant< bool, false > >	(no description available)
isExternal	integer_array	(no description available)
localToGlobalMap	globalIndex_array	Array that contains a map from localIndex to globalIndex.
primaryField	real64_array	Primary field variable
neighborData	node	<i>Datastructure: neighborData</i>
sets	node	<i>Datastructure: sets</i>



### 10.331 Datastructure: quartz

Name	Type	Description
Run	node	<i>Datastructure: Run</i>

### 10.332 Datastructure: sets

Name	Type	Description
externalSet	LvArray_SortedArray< int, int, LvArray_ChaiBuffer >	(no description available)

### 10.333 Datastructure: wellElementSubRegion

Name	Type	Description
domainBound-aryIndicator	integer_array	(no description available)
ghostRank	integer_array	(no description available)
globalToLo-calMap	geosx_mapBase< long long, int, std_integral_constant< bool, false > >	(no description available)
isExternal	integer_array	(no description available)
localToGlob-alMap	globalIndex_array	Array that contains a map from localIndex to globalIndex.
location	real64_array2d	For each perforation, physical location (x,y,z coordinates)
numPerfora-tionsGlobal	globalIndex	(no description available)
reservoirEle-mentIndex	integer_array	For each perforation, element index of the per-forated element
reservoirElemen-tRegion	integer_array	For each perforation, elementRegion index of the perforated element
reservoirEle-mentSubregion	integer_array	For each perforation, elementSubRegion in-dex of the perforated element
wellElementIn-dex	integer_array	For each perforation, index of the well ele-ment
wellTransmissi-bility	real64_array	For each perforation, well transmissibility
neighborData	node	<i>Datastructure: neighborData</i>
sets	node	<i>Datastructure: sets</i>



# CHAPTER 11

## Contributors

An up-to-date list of all GEOSX contributors can be found on our Github page:

[GEOSX Contributors](#)

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**A Multi-Resolution Approach to Hydraulic Fracture Simulation**

A Costa, M Cusini, T Jin, R Settgast, J Dolbow

arXiv preprint

[arXiv:2207.00661](https://arxiv.org/abs/2207.00661)



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### **Smooth implicit hybrid upwinding for compositional multiphase flow in porous media**

SBM Bosma, FP Hamon, BT Mallison, HA Tchelepi

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doi:10.1016/j.cma.2021.114288

### **Phase-field modeling of rock fractures with roughness**

F Fei, J Choo, C Liu, JA White

International Journal for Numerical and Analytical Methods in Geomechanics

doi:10.1002/nag.3317

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A Franceschini, N Castelletto, JA White, HA Tchelepi

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doi:10.1016/j.jcp.2022.111150

### **A scalable preconditioning framework for stabilized contact mechanics with hydraulically active fractures**

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Journal of Computational Physics

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J Huang, Y Hao, RR Settgaest, JA White, K Mateen, H Gross

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SBM Bosma, S Klevtsov, O Møyner, N Castelletto  
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[doi:10.1016/j.jcp.2020.109934](https://doi.org/10.1016/j.jcp.2020.109934)

### **Multigrid reduction preconditioning framework for coupled processes in porous and fractured media**

QM Bui, FP Hamon, N Castelletto, D Osei-Kuffuor, RR Settgaß, JA White  
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[doi:10.1016/j.cma.2021.114111](https://doi.org/10.1016/j.cma.2021.114111)

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[doi:10.1016/j.cma.2020.113432](https://doi.org/10.1016/j.cma.2020.113432)**An inelastic homogenization framework for layered materials with planes of weakness**

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## CHAPTER 13

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## CHAPTER 14

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### Indices and tables

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- `genindex`
- `modindex`
- `search`





### g

`geosx_mesh_tools.abaqus_converter`, [532](#)  
`geosx_xml_tools.attribute_coverage`, [539](#)  
`geosx_xml_tools.main`, [535](#)  
`geosx_xml_tools.regex_tools`, [538](#)  
`geosx_xml_tools.table_generator`, [541](#)  
`geosx_xml_tools.unit_manager`, [538](#)  
`geosx_xml_tools.xml_formatter`, [537](#)  
`geosx_xml_tools.xml_processor`, [535](#)  
`geosx_xml_tools.xml_redundancy_check`,  
[538](#)

### h

`hdf5_wrapper.wrapper`, [530](#)

### p

`pygeosx_tools.file_io`, [545](#)  
`pygeosx_tools.mesh_interpolation`, [545](#)  
`pygeosx_tools.well_log`, [546](#)  
`pygeosx_tools.wrapper`, [542](#)

### t

`timehistory.plot_time_history`, [547](#)



## A

`allgather_wrapper()` (in module `py-geosx_tools.wrapper`), 542  
`apply_regex_to_node()` (in module `geosx_xml_tools.xml_processor`), 535  
`apply_to_bins()` (in module `py-geosx_tools.mesh_interpolation`), 545

## B

`buildUnits()` (`geosx_xml_tools.unit_manager.UnitManager` method), 538

## C

`check_mpi_rank()` (in module `geosx_xml_tools.main`), 535  
`check_redundancy_level()` (in module `geosx_xml_tools.xml_redundancy_check`), 538  
`check_xml_redundancy()` (in module `geosx_xml_tools.xml_redundancy_check`), 539  
`close()` (`hdf5_wrapper.wrapper.hdf5_wrapper` method), 530  
`collect_xml_attributes()` (in module `geosx_xml_tools.attribute_coverage`), 539  
`collect_xml_attributes_level()` (in module `geosx_xml_tools.attribute_coverage`), 539  
`convert_abaqus_to_gmsh()` (in module `geosx_mesh_tools.abaqus_converter`), 532  
`convert_abaqus_to_vtu()` (in module `geosx_mesh_tools.abaqus_converter`), 532  
`convert_E_nu_to_K_G()` (in module `py-geosx_tools.well_log`), 546  
`copy()` (`hdf5_wrapper.wrapper.hdf5_wrapper` method), 530

## D

`DictRegexHandler` (class in `geosx_xml_tools.regex_tools`), 538

## E

`estimate_shmin()` (in module `py-geosx_tools.well_log`), 546  
`extrapolate_nan_values()` (in module `py-geosx_tools.mesh_interpolation`), 546

## F

`format_attribute()` (in module `geosx_xml_tools.xml_formatter`), 537  
`format_file()` (in module `geosx_xml_tools.xml_formatter`), 537  
`format_geosx_arguments()` (in module `geosx_xml_tools.main`), 535  
`format_xml_level()` (in module `geosx_xml_tools.xml_formatter`), 537

## G

`gather_wrapper()` (in module `py-geosx_tools.wrapper`), 542  
`generate_random_name()` (in module `geosx_xml_tools.xml_processor`), 535  
`geosx_mesh_tools.abaqus_converter` (module), 532  
`geosx_xml_tools.attribute_coverage` (module), 539  
`geosx_xml_tools.main` (module), 535  
`geosx_xml_tools.regex_tools` (module), 538  
`geosx_xml_tools.table_generator` (module), 541  
`geosx_xml_tools.unit_manager` (module), 538  
`geosx_xml_tools.xml_formatter` (module), 537  
`geosx_xml_tools.xml_processor` (module), 535  
`geosx_xml_tools.xml_redundancy_check` (module), 538  
`get_copy()` (`hdf5_wrapper.wrapper.hdf5_wrapper` method), 531

get\_global\_value\_range() (in module *pygeosx\_tools.wrapper*), 542  
 get\_group() (*pygeosx.Group* method), 421  
 get\_matching\_wrapper\_path() (in module *pygeosx\_tools.wrapper*), 543  
 get\_random\_realization() (in module *pygeosx\_tools.mesh\_interpolation*), 546  
 get\_realizations() (in module *pygeosx\_tools.mesh\_interpolation*), 546  
 get\_wrapper() (in module *pygeosx\_tools.wrapper*), 543  
 get\_wrapper() (*pygeosx.Group* method), 421  
 get\_wrapper\_par() (in module *pygeosx\_tools.wrapper*), 543  
 getHistorySeries() (in module *timehistory.plot\_time\_history*), 547  
 groups() (*pygeosx.Group* method), 421

## H

hdf5\_wrapper (class in *hdf5\_wrapper.wrapper*), 530  
 hdf5\_wrapper.wrapper (module), 530

## I

insert() (*hdf5\_wrapper.wrapper.hdf5\_wrapper* method), 531

## K

keys() (*hdf5\_wrapper.wrapper.hdf5\_wrapper* method), 531

## L

link() (*hdf5\_wrapper.wrapper.hdf5\_wrapper* method), 531  
 load\_tables() (in module *pygeosx\_tools.file\_io*), 545

## M

main() (in module *geosx\_xml\_tools.attribute\_coverage*), 540  
 main() (in module *geosx\_xml\_tools.xml\_formatter*), 538  
 main() (in module *geosx\_xml\_tools.xml\_redundancy\_check*), 539

merge\_included\_xml\_files() (in module *geosx\_xml\_tools.xml\_processor*), 536  
 merge\_xml\_nodes() (in module *geosx\_xml\_tools.xml\_processor*), 536

## P

parse\_las() (in module *pygeosx\_tools.well\_log*), 547  
 parse\_schema() (in module *geosx\_xml\_tools.attribute\_coverage*), 540  
 parse\_schema\_element() (in module *geosx\_xml\_tools.attribute\_coverage*), 540

plot\_history() (in module *pygeosx\_tools.wrapper*), 543

preprocess\_parallel() (in module *geosx\_xml\_tools.main*), 535  
 preprocess\_serial() (in module *geosx\_xml\_tools.main*), 535  
 print\_global\_value\_range() (in module *pygeosx\_tools.wrapper*), 544  
 process() (in module *geosx\_xml\_tools.xml\_processor*), 536  
 process\_xml\_files() (in module *geosx\_xml\_tools.attribute\_coverage*), 540  
 process\_xml\_files() (in module *geosx\_xml\_tools.xml\_redundancy\_check*), 539

*pygeosx.apply\_initial\_conditions*() (built-in function), 420  
*pygeosx.COMPLETED* (built-in variable), 421  
*pygeosx.finalize*() (built-in function), 420  
*pygeosx.Group* (built-in class), 421  
*pygeosx.initialize*() (built-in function), 420  
*pygeosx.INITIALIZED* (built-in variable), 421  
*pygeosx.READY\_TO\_RUN* (built-in variable), 421  
*pygeosx.reinit*() (built-in function), 420  
*pygeosx.run*() (built-in function), 420  
*pygeosx.UNINITIALIZED* (built-in variable), 421  
*pygeosx.Wrapper* (built-in class), 421  
*pygeosx\_tools.file\_io* (module), 545  
*pygeosx\_tools.mesh\_interpolation* (module), 545  
*pygeosx\_tools.well\_log* (module), 546  
*pygeosx\_tools.wrapper* (module), 542

## R

read\_GEOS\_table() (in module *geosx\_xml\_tools.table\_generator*), 541  
 regexHandler() (*geosx\_xml\_tools.unit\_manager.UnitManager* method), 538  
 register() (*pygeosx.Group* method), 421  
 run\_queries() (in module *pygeosx\_tools.wrapper*), 544

## S

save\_tables() (in module *pygeosx\_tools.file\_io*), 545  
 search\_datastructure\_wrappers\_recursive() (in module *pygeosx\_tools.wrapper*), 544  
 set\_wrapper\_to\_value() (in module *pygeosx\_tools.wrapper*), 544  
 set\_wrapper\_with\_function() (in module *pygeosx\_tools.wrapper*), 544  
 SymbolicMathRegexHandler() (in module *geosx\_xml\_tools.regex\_tools*), 538

## T

`timehistory.plot_time_history` (*module*),  
547

## U

`UnitManager` (*class* *in*  
*geosx\_xml\_tools.unit\_manager*), 538

## V

`validate_xml()` (*in* *module*  
*geosx\_xml\_tools.xml\_processor*), 537

`value()` (*pygeosx.Wrapper method*), 421

`values()` (*hdf5\_wrapper.wrapper.hdf5\_wrapper*  
*method*), 531

## W

`wait_for_file_write_rank_0()` (*in* *module*  
*geosx\_xml\_tools.main*), 535

`wrappers()` (*pygeosx.Group method*), 421

`write_attribute_usage_xml()` (*in* *module*  
*geosx\_xml\_tools.attribute\_coverage*), 541

`write_attribute_usage_xml_level()` (*in*  
*module* *geosx\_xml\_tools.attribute\_coverage*),  
541

`write_GEOS_table()` (*in* *module*  
*geosx\_xml\_tools.table\_generator*), 541

`write_read_GEOS_table_example()` (*in* *mod-*  
*ule* *geosx\_xml\_tools.table\_generator*), 542