

RELAP-7 User's Guide

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RELAP-7 Overview

The RELAP-7 code is the next generation nuclear reactor system safety analysis code being developed at the Idaho National Laboratory (INL). The code is based on the INL's modern scientific software development framework, MOOSE (Multi-Physics Object Oriented Simulation Environment). The overall design goal of RELAP-7 is to take advantage of the previous thirty years of advancements in computer architecture, software design, numerical integration methods, and physical models. The end result will be a reactor systems analysis capability that retains and improves upon RELAP5's capability and extends the analysis capability for all reactor system simulation scenarios.

RELAP-7 will become the main reactor systems simulation toolkit for the LWRS (Light Water Reactor Sustainability) program's RISMIC (Risk Informed Safety Margin Characterization) effort and the next generation tool in the RELAP reactor safety/systems analysis application series. The key to the success of RELAP-7 is the simultaneous advancement of physical models, numerical methods, and software design while maintaining a solid user perspective. Physical models include both PDEs (Partial Differential Equations) and ODEs (Ordinary Differential Equations) and experimental based closure models. RELAP-7 utilizes well-posed governing equations for compressible two-phase flow, which can be strictly verified in a modern verification and validation effort. Closure models used in RELAP5 and newly developed models will be reviewed and selected to reflect the progress made during the past three decades and provide a basis for the closure relations that will be required in RELAP-7. RELAP-7 uses modern numerical methods, which allow implicit time integration, second-order schemes in both time and space, and strongly coupled multi-physics.

RELAP-7 is written with object oriented programming language C++. By using the MOOSE development environment, the RELAP-7 code is developed by following the same modern software design paradigms used for other MOOSE development efforts. The code is easy to read, develop, maintain, and couple with other codes. Most importantly, the modern software design allows the RELAP-7 code to evolve efficiently with time. MOOSE is an HPC development and runtime framework for solving computational engineering problems in a well planned, managed, and coordinated way. By leveraging millions of lines of open source software packages, such as PETSc (a nonlinear solver developed at Argonne National Laboratory) and LibMesh (a Finite Element Analysis package developed at University of Texas), MOOSE reduces the expense and time required to develop new applications. MOOSE provides numerical integration methods and mesh management for parallel computation. Therefore RELAP-7 code developers have been

able to focus more upon the physics and user interface capability. There are currently over 20 different MOOSE based applications ranging from 3-D transient neutron transport, detailed 3-D transient fuel performance analysis, to long-term material aging. Multi-physics and multi-dimensional analysis capabilities, such as radiation transport and fuel performance, can be obtained by coupling RELAP-7 and other MOOSE-based applications through MOOSE and by leveraging with capabilities developed by other DOE programs. This allows restricting the focus of RELAP-7 to systems analysis type simulations and gives priority to retain and significantly extend RELAP5's capabilities.

This document provides a user's guide to help users to learn how to run the RELAP-7 code. A number of example problems and their associated input files are presented in this document to guide users to run the RELAP-7 code starting with simple pipe problems to problems with increasing complexity. Because the code is an ongoing development effort, this RELAP-7 User's Guide will evolve with periodic updates to keep it current with the state of the development, implementation, and model additions/revisions. A complete User's Manual will be developed at a later time when the RELAP-7 code becomes more mature.

1 RELAP-7 Features

An overall description of the RELAP-7 architecture, governing theory, and computational approach is given here as an instructive, and executive overview of the RELAP-7 distinguishing features.

1.1 Software Framework

MOOSE is INL's development and runtime environment for the solution of multi-physics systems that involve multiple physical models or multiple simultaneous physical phenomena. The systems are generally represented (modeled) as a system of fully coupled nonlinear partial differential equation systems (an example of a multi-physics system is the thermal feedback effect upon neutronics cross-sections where the cross-sections are a function of the heat transfer). Inside MOOSE, the Jacobian-Free Newton Krylov (JFNK) method [1, 2] is implemented as a parallel nonlinear solver that naturally supports effective coupling between physics equation systems (or Kernels). The physics Kernels are designed to contribute to the nonlinear residual, which is then minimized inside of MOOSE. MOOSE provides a comprehensive set of finite element support capabilities (LibMesh [3], a Finite Element library developed at University of Texas) and provides for mesh adaptation and parallel execution. The framework heavily leverages software libraries from DOE SC and NNSA, such as the nonlinear solver capabilities in either the the Portable, Extensible Toolkit for Scientific Computation (PETSc [4]) project or the Trilinos project [5] (a collection of numerical methods libraries developed at Sandia National Laboratory). Argonne's PETSc group has recently joined with the MOOSE team in a strong collaboration wherein they are customizing PETSc for our needs. This collaboration is strong enough that Argonne is viewed as a joint developer of MOOSE.

A parallel and tightly coordinated development effort with the RELAP-7 development project is the Reactor Analysis Virtual control ENvironment (RAVEN). This MOOSE-based application is a complex, multi-role software tool that will have several diverse tasks including serving as the RELAP-7 graphical user interface, using RELAP-7 to perform RISMC focused analysis, and controlling the RELAP-7 calculation execution.

Together, MOOSE/RELAP-7/RAVEN comprise the systems analysis capability of LWRS RISMC ToolKit.

1.2 Governing Theory

The primary basis of the RELAP-7 governing theory includes thermal fluids flow, reactor core heat transfer, and reactor kinetics models.

With respect to the thermal fluids flow dynamics models, RELAP-7 incorporates both single- and two-phase flow simulation capabilities encompassing all-speed and all-fluids. The single phase flow models include isothermal flow and nonisothermal flow capabilities. The two-phase flow models include the homogeneous equilibrium flow model (HEM) and the well posed two fluid 7-equation model.

In addition to the fluids flow dynamics model, RELAP-7 necessarily simulates the heat transfer process with reactor kinetics as the heat source. The heat-conduction equation for cylindrical or slab geometries is solved to provide thermal history within metal structures such as fuel and clad. The volumetric power source in the heat conduction equation for the fuel comes from the point kinetics model with thermal hydraulic reactivity feedback considered [6]. The reactor structure is coupled with the thermal fluid through energy exchange (conjugate heat transfer) employing surface convective heat transfer [7] within the fluid. The fluid, heat conduction, conjugate heat transfer and point kinetics equations are solved in a fully coupled fashion in RELAP-7 in contrast to the operator-splitting or loose coupling approach used in the existing system safety analysis codes.

1.3 Computational Approach

Stated previously, the MOOSE framework provides the bulk of the "heavy lifting" available to MOOSE-based applications with a multitude of mathematical and numerical libraries. For RELAP-7, LibMesh [3] provides the second-order accurate spatial discretization by employing linear basis, one-dimensional finite elements. The Message Passing Interface (MPI, from Argonne National Laboratory) provides for distributed parallel processing. Intel Threading Building Blocks (Intel TBB) allows parallel C++ programs to take full advantage of multicore architecture found in most large-scale machines of today. PETSc (from Argonne), Trilinos (from Sandia), and Hypre [8] (from Lawrence Livermore National Laboratory) provide the mathematical libraries and nonlinear solver capabilities for the Jacobian-free Newton-Krylov (JFNK) method. In MOOSE, a stiffly-stable, second-order backward difference (BDF2) formulation is used to provide second-order accurate time integration for strongly coupled physics in JFNK.

The JFNK method easily allows implicit nonlinear coupling of dependent physics under one general computational framework. Besides rapid (second-order) convergence of the iterative procedure, the JFNK method flexibly handles multiphysics problems when time scales of different physics are significantly varied during transients. The key feature of the JFNK method is combining Newton's method to solve implicit nonlinear systems with Krylov subspace iterative methods. The Krylov methods do not require an explicit form of the Jacobian, which eliminates the computationally expensive step of forming Jacobian matrices (which also may be quite difficult to determine analytically), required by Newton's method. The matrix-vector product can be approximated by the numerical differentiation of nonlinear residual functions. Therefore, JFNK readily integrates different physics into one solver framework.

2 Model Description

2.1 Fluids Flow Models

The RELAP-7 code has various flow models implemented. These include:

- 1). a single phase isothermal flow model (`model_type=2`),
- 2). a single phase nonisothermal fluid flow model (`model_type=3`),
- 3). a homogeneous equilibrium (HEM) two phase flow model (`model_type=32`),
- 4). a nonhomogeneous, nonequilibrium seven-equation two phase flow model (`model_type=7`).

2.2 Equation of State

For the **single phase flow models**, various types of equation of state can be used. These include:

- 1). Barotropic equation of state for isothermal flow. This is turned on by setting `type=BarotropicEquationOfState`.
- 2). Linear equation of state for non-isothermal single phase flow. This is turned on by setting `type = NonIsothermalEquationOfState`.
- 3). Stiffened gas equation of state which is turned on by setting `type = StiffenedGasEquationOfStateVapor` for single phase vapor flow and `type = StiffenedGasEquationOfStateLiquid` for single phase water flow.
- 4). Ideal gas equation of state which is turned on by setting `type=IdealGasEquationOfState`. The equation of state for the nitrogen gas has also been implemented into the code. It can be turned on by setting `type=N2Properties`.

For the **homogeneous equilibrium flow model**, only the stiffened gas equation of state is applicable. This is turned on by setting `type = TwoPhaseStiffenedGasEOS`.

For the **seven-equation two phase flow model**, only the stiffened gas equation of state is applicable. This is turned on by setting `type = StiffenedGasEquationOfStateLiquid` for the liquid phase, and `type = StiffenedGasEquationOfStateVapor` for the vapor phase.

2.3 Solution Stabilization Schemes

It is well known that the continuous Galerkin finite element method is unstable when applied directly to hyperbolic systems of equations. Therefore the solution stabilization schemes are required for RELAP-7. Currently available options of solution stabilization for RELAP-7 include:

- 1). Streamline Upwind/Petrov Galerkin method (SUPG). The SUPG scheme works for the single phase flow only. It can be used by setting `stabilization_type = 'SUPG'`.
- 2). Lapidus scheme works for both the single phase flow and the two phase flow cases. It can be used by setting `stabilization_type = 'LAPIDUS'`.
- 3). The entropy viscosity method works for both the single phase flow and the two phase flow. This option can be used by setting `stabilization_type = 'ENTROPY_VISCOSITY'`.

2.4 Time Integration Schemes

There are two types of time integration schemes in RELAP-7 - Implicit Euler and BDF2. Implicit Euler is a first order accurate time integration scheme. This can be turned on by setting: `scheme = 'implicit-euler'` in the Executioner input block (explained later). BDF2 is a second order accurate time integration scheme. This is the default option for RELAP-7.

2.5 Components

A real reactor system is very complex and contains hundreds of different physical components. It is impractical to resolve the real geometry of the entire system. Instead simplified thermal hydraulic models are used to represent (via “nodalization”) the major physical components and describe the major physical processes (such as fluids flow and heat transfer). There are three main types of components developed in RELAP-7: (1) one-dimensional (1-D) components describing the geometry of the reactor system, (2) zero-dimensional (0-D) components for setting boundary conditions, and (3) 0-D components for connecting 1-D components.

2.5.1 Pipe

Pipe is the most basic component in RELAP-7. It is a 1-D component which simulates thermal fluids flow in a pipe. Both a constant cross section area and a variable cross section area options are available for the Pipe component. The wall friction and heat transfer coefficients are either calculated through closure models or provided by user input. The pipe wall temperature can be provided as the wall heat transfer boundary condition. All the thermal fluids dynamic models are available in the Pipe component which includes the isothermal flow model, single-phase non-isothermal flow model, nonequilibrium 7-equation two-phase model, and the much simpler homogeneous equilibrium two-phase flow model.

2.5.2 PipeWithHeatStructure

The PipeWithHeatStructure component simulates fluids flow in a 1-D pipe coupled with 1-D or 2-D heat conduction through the pipe wall. The adiabatic, Dirichlet, or convective boundary conditions at the outer surface of the pipe wall are available. Either a plate type or cylindrical type of heat structure can be selected. Volumetric heat source within the fluids or solid materials can be added.

2.5.3 CoreChannel

The CoreChannel component is a composite component designed to simulate the coolant flow and heat conduction inside a fuel rod as well as the conjugate heat transfer between the coolant and the fuel rod. In this component, the fuel rod is divided into the same number of segments as that of the coolant flow pipe elements. Each fuel rod segment is further simulated as 1-D or 2-D heat conduction model perpendicular to the fluid flow model. Both plate type fuel rod and cylindrical fuel rod type can be simulated. The solid fuel part is able to deal with typical LWR fuel rod with complex clad/gap/fuel pellet geometries. The flow model and conjugate heat transfer model are fully coupled in contrast to loosely coupled in RELAP5.

2.5.4 Subchannel

A fully coupled subchannel channel model for the single-phase has been implemented into RELAP-7. The single-phase subchannel model includes four balance equations: mass, energy, axial momentum, and lateral momentum.

2.5.5 HeatExchanger

A Heat Exchanger component is a combination of two pipes with a solid wall in between. Similar to the CoreChannel model, the fluids flow model and conjugate heat transfer model are fully coupled. More complicated and realistic steam generator component will be developed in the future.

2.5.6 Branch

The branch model is a 0-D component representing a junction model with no volume (inertia) effects considered, and with single/multiple inlets and single/multiple outlets, of which cross section areas can be different. This model conserves the mass and energy among all connecting components.

2.5.7 VolumeBranch

The volume branch model is a 0-D component representing a joint/junction model with volume (inertia) effects considered. This model conserves the mass and energy among all connecting components.

2.5.8 SubchannelBranch

This is a 0D branch model to connect the subchannel component.

2.5.9 Pump

A simple pump model to provide a head and a reverse flow form loss coefficient (K) for either isothermal flow and non-isothermal flow. It can be driven by an user input head or through a driving component which provides shaft work.

2.5.10 Turbine

The turbine model in RELAP-7 is a simplified dynamical turbine model to simulate a reactor core isolation cooling (RCIC) turbine, which drives the RCIC pump through a common shaft.

2.5.11 SeparatorDryer

The separator dryer model in RELAP-7 separates steam and water with mechanical methods. Only an ideal separator dryer model is available in RELAP-7.

2.5.12 DownComer

The down comer component simulates a large volume to mix different streams of water and steam and to track the water level.

2.5.13 Valve

The valve component simulates the open and close behaviors of valves for incompressible flow with user given trigger and response time. The abrupt area change model is used to calculate the form loss.

2.5.14 CompressibleValve

The compressible valve component simulates the open and close behavior of valves for compressible fluid flow, including ckoking. It can be used to simulate the safety relief

valves (SRV) of BWRs.

2.5.15 Check Valves

The check valve component simulates the dynamic behavior of check valves, with the form loss calculated by the abrupt area change model.

2.5.16 WetWell

The wet well component simulates the dynamic response of a BWR suppression pool and its gas space.

2.5.17 TimeDependentVolume

The time dependent volume component provides pressure, temperature, and void fraction boundary conditions as constants or time dependent functions for 1-D components. It is a pure boundary condition type of component and it does not add any entries to the global unknown vector. When acquired by its connected 1-D component, it provides a pressure, a temperature, and a void fraction boundary condition.

2.5.18 TimeDependentJunction

The time dependent junction component provides velocity and temperature boundary conditions as constants or time functions for 1-D components.

2.5.19 Tdm

Tdm sets time dependent mass flow boundary conditions for 1-D components.

2.5.20 Reactor

A virtual component that allows users to input the power for the core channel component.

2.5.21 PointKinetics

The point kinetics model is a lumped parameter neutron kinetics model to calculate the reactor power. User input reactivity or fully coupled feedback reactivity models are available.

3 Running RELAP-7

3.1 Complete Step 1 of MOOSE Environment Setup

The system environment setup for MOOSE can be found with the link:
<http://www.mooseframework.org/getting-started>

3.2 Setup Your SSH Key

SSH key allows you to establish a secure connection between your computer and GitLab. Before generating an SSH key, check to see if your system already has one by running `cat ~/.ssh/id_rsa.pub`. If you see a long string starting with `ssh-rsa` or `ssh-dsa`, you can skip the `ssh-keygen` step below.

To generate a new SSH key, just open your terminal and use the code below. The `ssh-keygen` command prompts you for a location and filename to store the key pair and for a password. When prompted for the location and filename you can press `enter` to use the default. It is a best practice to use a password for an SSH key but it is not required and you can skip creating a password by pressing `enter`. Note that the password you choose here can not be altered or retrieved.

```
ssh-keygen -t rsa -C "$your_email"
```

Use the code below to show your public key.

```
cat ~/.ssh/id_rsa.pub
```

Copy and paste the key to the `My SSH Key` section under the `SSH` tab in your profile. Please copy the complete key starting with `ssh-` and ending with your username and host.

To test your SSH key.

```
$ssh git@hpcgitlab.inl.gov
Welcome to GitLab, <your name here>!
Connection to hpcgitlab.inl.gov closed.
```

3.3 Checking Out the Code

```
$ cd ~/projects/  
$ git clone git@hpcgitlab.inl.gov:idaholab/relap-7.git  
$ cd relap-7  
$ git submodule update --init
```

It is necessary to build libmesh before building any application.

```
$ cd ~/projects/relap-7/moose  
$ ./update_and_rebuild_libmesh.sh
```

Once libmesh has been successfully compiled, you may now compile RELAP-7.

```
cd ~/projects/trunk/relap-7  
make (add -jn to run on multiple "n" processors)
```

Once RELAP-7 has been compiled successfully, it is recommended to run the tests to make sure the version of the code you have is running correctly.

```
cd ~/projects/trunk/relap-7  
./run_tests (add -jn to run "n" jobs at one time)
```

3.4 Executing RELAP-7

When first starting out with running RELAP-7, it is recommended to start from an example problem. Multiple example problems with input files are presented in this document. More examples can be found under the `/relap-7/tests` subdirectory. To demonstrate how to run RELAP-7, consider the `PWR_core_channel.i` test problem.

```
cd ~/projects/trunk/relap-7/examples/components/core_channel/  
# To run with one processor  
~/projects/trunk/relap-7/relap-7-opt -i PWR_core_channel.i  
# To run in parallel (4 processors)  
mpirun -n 4 ../../../../relap-7-opt -i PWR_core_channel.i
```

3.5 Post Processing

RELAP-7 typically writes solution data to an ExodusII file. The solution data may also be written in other formats, one being a comma separated values (CSV) file, which allows the solution data to be saved in a table structured format. The other being a tecplot file in either binary or ASCII format.

Several options exist for viewing ExodusII output files. One good choice is to use open-source software, Paraview (www.paraview.org).

3.6 Graphical User Interface

Another MOOSE based application named RAVEN provides a graphical user interface (GUI) for RELAP-7. RAVEN can be used to generate a text input file. It is also capable of submitting the analysis and provides post processing capabilities.

4 Input Files

RELAP-7 uses a block-structured input file. Each block is identified with square brackets. The opening brackets contain the type of the input block and the empty brackets mark the end of the block. Each block may contain subblocks.

```
[BlockName]
  <block line commands>
  [./subblock_name]
    <subblock line commands>
  [../]
[]
```

Each subblock must have a unique name when compared with all other subblocks in the current block.

Line commands are given as parameter and value pairs with an equal sign between them. They specify parameters to be used by the object being described. The parameter is a string, and the value may be a string, an integer, a real number, or a list of strings, integers, or real numbers. Lists are given in single quotes and are separated by whitespace.

Subblocks normally contain a type line command. This line command specifies the particular type of object being described.

RELAP-7 uses SI units. This standardizes the model input by eliminating the possibility of errors caused by using one set of units for one model and another set of units for a different model.

The following subsections have brief descriptions of each block. More detailed descriptions can be found in the examples section.

4.1 Global Parameters

The GlobalParams block specifies the global parameters used by the code such as the initial pressure (`global_init_P`), velocity (`global_init_V`) and temperature (`global_init_T`) of the system model, the fluid flow model type (`model_type`), the stabilization scheme type (`stabilization_type`), and the scaling factors (`scaling_factor_var`) for the primary variable, etc. The values of global parameters are available to any other block or sub-

block in the input file. If a command line is missing in a block or a subblock but defined in GlobalParams, the block or subblock will use the parameter defined in GlobalParams. However, if the block or subblock has a command line , that will be used regardless of what is in GlobalParams.

The following is an example of the GlobalParams block:

```
[GlobalParams]
  global_init_P = 155.e5
  global_init_V = 0.
  global_init_T = 559.15
  # model_type = 2
  # model_type = 32
  # model_type = 7
  model_type = 3
  stabilization_type = 'SUPG'
  scaling_factor_var = '1e4 1e1 1e-2'
  temperature_sf = '1e-2'
[]
```

symbol indicates comments in the input file and can be located anywhere in the input file.

4.2 Equation of State

The EoS block specifies the equation of state to be used by the code. The following is an example of the EoS block:

```
[EoS]
  [./eos]
  type = NonIsothermalEquationOfState
  p_0 = 155.e5
  rho_0 = 686.
  a2 = 1.e7
  beta = .46e-3
  cv = 5.5e3
  e_0 = 3075325
  T_0 = 559.15
  [../]
```

```
[ ]
```

4.3 Materials

The Materials block specifies the properties of the solid materials for the code. The following is an example of the Materials block:

```
[Materials]
  ./fuel-mat]
    type = SolidMaterialProperties
    k = 2.5
    Cp = 300.
    rho = 1.032e4
  [../]
  ./gap-mat]
    type = SolidMaterialProperties
    k = 0.6
    Cp = 1.
    rho = 1.
  [../]
  ./clad-mat]
    type = SolidMaterialProperties
    k = 21.5
    Cp = 350.
    rho = 6.55e3
  [../]
[ ]
```

4.4 Functions

The Functions block provides the functions to be used by the code during the simulations such as reactor power as a function of time or a boundary condition pressure as a function of time, etc. The following is an example of defining pressure distribution as a function of x .

```
[Functions]
```

```
[./p_func]
axis = 0
type = PiecewiseLinear
x = '0      3'
y = '1.05e5  1e5'
[./]
[]
```

4.5 Components

The Components block specifies the components to be used in the simulations.

```
[Components]
[./reactor]
type = Reactor
initial_power = 77337.69407
[./]

[./CCH1]
type = CoreChannel
eos = eos
position = '0 0 0'
orientation = '0 0 1'
A = 8.7878e-5
Dh = 0.01179
length = 4
n_elems = 20
f = 0.01
Hw = 5.33e4
Phf = 0.029832559676
Ts_init = 559.15
dim_hs = 1
fuel_type = cylinder
name_of_hs = 'fuel gap clad'
n_heatstruct = 3
width_of_hs = '0.004096 0.0001 0.000552'
elem_number_of_hs = '10 1 2'
material_hs = 'fuel-mat gap-mat clad-mat'
power_fraction = '1.0 0.0 0.0'
```

```

[../]

[./inlet]
  type = TimeDependentVolume
  input = 'CCH1(in)'
  p_bc = 155.483e5
  T_bc = 559.15
  eos = eos
[../]

[./outlet]
  type = TimeDependentVolume
  input = 'CCH1(out)'
  p_bc = '155.e5'
  T_bc = 559.15
  eos = eos
[../]
[]

```

4.6 Preconditioner

The Preconditioning block specifies the preconditioner to be used by the preconditioned JFNK solver for the RELAP-7 code. The solution algorithm for RELAP-7 is the Jacobian-free Newton-Krylov (JFNK) method. However the Krylov methods need preconditioning to be efficient. Hence, the solvers available in RELAP-7 are preconditioned JFNK (PJFNK). Two options are available in RELAP-7 to build the preconditioning matrix, the single matrix preconditioner (SMP) and the finite difference preconditioner (FDP). The SMP option uses all the Jacobian terms derived analytically to build one preconditioning matrix. The FDP option uses numerical Jacobian by doing direct finite differences of the residual terms. The SMP option is the more efficient and is the recommended option, while the FDP option is normally slow and inefficient and is recommended to be used for small problems or for debugging purposes. The following is an example of the Preconditioning block:

```

[Preconditioning]
  # Uncomment one of the lines below to activate one of the blocks...
  active = 'SMP_PJFNK'
  #active = 'FDP_PJFNK'

```

```

# The definitions of the above-named blocks follow.
[./SMP_PJFNK]
  type = SMP
  full = true # off diagonal blocks are used
  solve_type = 'PJFNK' # Preconditioned JFNK solver
[../]

[./FDP_PJFNK]
  type = FDP
  full = true
  solve_type = 'PJFNK'
[../]
[]

```

4.7 Executioner

The Executioner block specifies the executioner that will be used in the simulations. The options include Transient, RavenExecutioner, etc.

```

[Executioner]
  type = Transient
  dt = 0.5
  dtmin = 1.e-5
  petsc_options_iname = '-mat_fd_type  -mat_mffd_type  -pc_type'
  petsc_options_value = '  ds                ds                lu'
  nl_rel_tol = 1e-8
  nl_abs_tol = 1e-8
  nl_max_its = 10
  l_tol = 1e-4
  l_max_its = 60
  start_time = 0.0
  num_steps = 30
  [./Quadrature]
    type = TRAP
    order = FIRST
  [../]
[]

```

4.8 Outputs

The `Outputs` block controls the various screen and file output in the simulations.

```
[Outputs]
  [./out]
    type = Exodus
    use_displaced = true
    output_initial = true
    sequence = false
    append_displaced = true
  [../]

  [./console]
    type = Console
    perf_log = true
  [../]
[]
```

5 Examples

5.1 Example 1: A Simple Pipe Flow Problem

5.1.1 Problem Description



Figure 1. A simple pipe flow problem diagram

Example 1 simulates water flowing through a pipe under isothermal conditions with the following parameters:

Hydraulic Diameter = .01 m
Cross section flow area = $7.85e-5 m^2$
Length = 1 m
Wall friction coefficient = .01

The boundary conditions are the following:

Inlet: Pressure = $1.05e5$ Pa
Outlet: Pressure = $1e5$ Pa

A 1D model can be viewed in Paraview to visualize the process better. Once Paraview is opened, on the left, select all of the variables and click "Apply." In the filters tab, under "Data Analysis", select "Plot Over Line" and apply. When this is done, each parameter in the problem can be viewed plotted over the length of the pipe. The figure below shows pressure vs length.

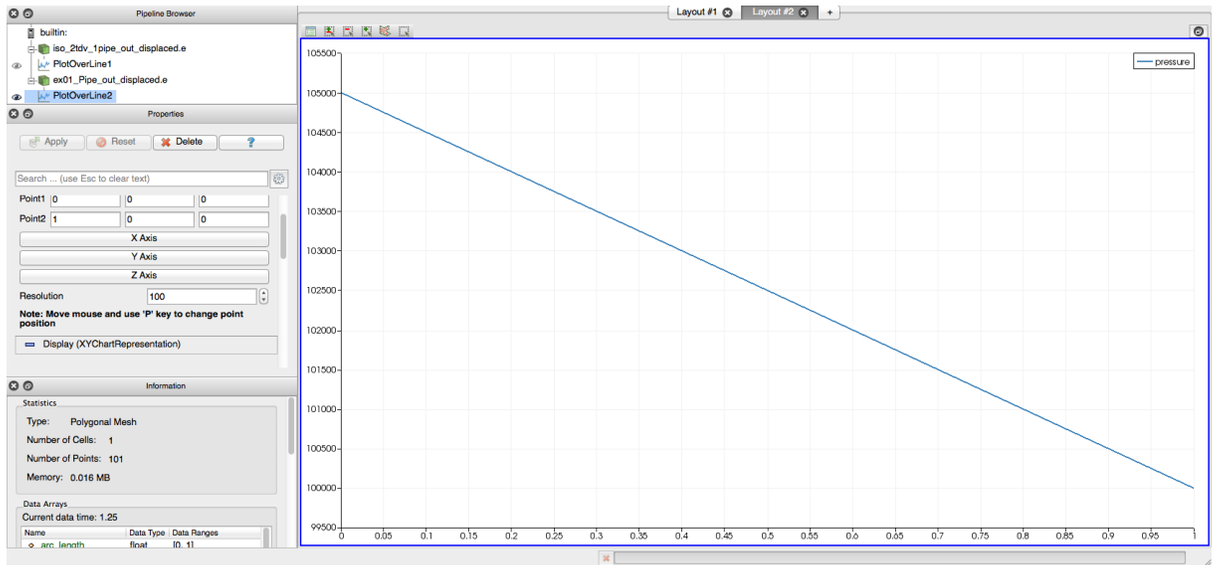


Figure 2. Pressure vs. length for the simple pipe flow problem

The pressure difference can be hand calculated to check the code calculated results using:

$$\Delta P = \frac{fL\rho u^2}{2D_h} \quad (1)$$

Where ρ is the density and u is the velocity. Both values can be found in Paraview to perform the calculation.

5.1.2 Input File

The following shows the input file to run this example problem:

```

#
# Isothermal flow through one simple pipe with B.C. set by TDVs
#

[GlobalParams]
  model_type = 2
  scaling_factor_var = '1e+0 1e-4'
  stabilization_type = 'SUPG'
[]

[EoS]
  [./eos]
    type = BarotropicEquationOfState
    p_0 = 1.e5
    rho_0 = 1.e3
    a2 = 1.e7
  [../]
[]

[Components]
  [./pipe1]
    type = Pipe
    eos = eos
    position = '0 0 0'
    orientation = '1 0 0'
    Dh = 0.01
    length = 1.0
    n_elems = 10
    A = 7.854e-5
    f = 0.01
  [../]

  [./inlet]
    type = TimeDependentVolume
    input = 'pipe1(in)'
    p_bc = 105e3
    eos = eos
  [../]

```

```

[./outlet]
  type = TimeDependentVolume
  input = 'pipe1(out)'
  p_bc = 1e5
  eos = eos
[../]
[]

[Preconditioning]
[./SMP_PJFNK]
  type = SMP
  full = true
  solve_type = 'PJFNK'
[../]
[]

[Executioner]
  type = Transient
  scheme = 'implicit-euler'
  dt = 1.25e-2
  dtmin = 1.e-5

  nl_rel_tol = 1e-8
  nl_abs_tol = 1e-6
  nl_max_its = 15

  l_tol = 1e-6
  l_max_its = 30

  start_time = 0.0
  end_time = 250.0
  num_steps = 500

[./Quadrature]
  type = TRAP
  order = FIRST
[../]

```

```

[]

[Outputs]
  [./out]
    type = Exodus
    use_displaced = true
    output_initial = true
    sequence = false
    append_displaced = true
  [../]

  [./console]
    type = Console
    perf_log = true
  [../]
[]

```

5.1.3 Description of the Input File

The following are the detailed descriptions of the input file block by block.

The Global Parameters block:

```

[GlobalParams]
  model_type = 2
  scaling_factor_var = '1e+0 1e-4'
  stabilization_type = 'SUPG'
[]

```

model_type	Fluid flow model. 2 means single phase isothermal fluid flow.
scaling_factor_var	Scaling factor for the primary variables ρA and $\rho u A$ respectively.
stabilization_type	Stabilization scheme type. Available options are: SUPG for single phase flow, LAPIDUS for both single- and two-phase flow, ENTROPY_VISCOSITY for both single- and two-phase flow.

The Equation of State block:

```
[EoS]
  [./eos]
    type = BarotropicEquationOfState
    p_0 = 1.e5
    rho_0 = 1.e3
    a2 = 1.e7
  [../]
[]
```

[./eos] Subblock for eos

type The type of equation of state to be used. The barotropic equation of state is suitable for isothermal fluid flow model. It describes isentropic (reversible) processes, and implies a constant sound speed. It is given by $p = p_0 + a^2(\rho - \rho_0)$.

p_0 Initial pressure value (*Pa*) in the equation of state calculations.

rho_0 Initial value of density (kg/m^3) in the equation of state calculation.

a2 a is a constant in the barotropic equation of state. The value of a can be taken roughly as the speed of sound. For this example problem with water as the fluids, using $a^2 = 1.e7$ has the same order of magnitude as that of the square of sound speed. This is the recommended value to use.

The Components Block:

```
[Components]
  [./pipe1]
    type = Pipe
    eos = eos
    # geometry
    position = '0 0 0'
    orientation = '1 0 0'
    Dh = 0.01
    length = 1.0
    n_elems = 10
    A = 7.854e-5
```

```

    f = 0.01
[../]

[./inlet]
    type = TimeDependentVolume
    input = 'pipe1(in)'
    p_bc = 105e3
    eos = eos
[../]

[./outlet]
    type = TimeDependentVolume
    input = 'pipe1(out)'
    p_bc = 1e5
    eos = eos
[../]
[]

```

[./pipe1]	Subblock for pipe1.
type	Component type is pipe.
eos	Use the eos equation of state defined in the EoS block.
position	The starting x, y and z coordinate of pipe1.
orientation	The orientation vector of pipe1. Horizontal pipe for this case.
Dh	Hydraulic diameter of of the pipe.
length	Pipe length.
n_elems	Number of elements in the solution of the pipe.
A	The cross section flow area of the pipe.
f	The wall friction coefficient. If this is not provided in the input file, the code built-in correlations will be used to calculate the friction coefficients (for single phase flow only).

<code>[./inlet]</code>	Subblock for the component to set the inlet boundary conditions.
<code>type</code>	Component type is time dependent volume.
<code>input</code>	The pipe inlet end which will be connected with this component.
<code>p_bc</code>	The pressure boundary condition value at the inlet of the pipe.
<code>eos</code>	The equation of state to be used.
<code>[./outlet]</code>	Subblock for the component to set the outlet boundary conditions.
<code>type</code>	Component type is time dependent volume.
<code>input</code>	The pipe outlet end which will be connected with this component.
<code>p_bc</code>	The pressure boundary boundary condition value at the outlet of the pipe.
<code>eos</code>	The equation of state to be used.

The Preconditioning Block:

```
[Preconditioning]
  [./SMP_PJFNK]
    type = SMP
    full = true
    solve_type = 'PJFNK'
  [../]
[]
```

<code>[./SMP_PJFNK]</code>	Subblock for SMP preconditioner and PJFNK solver. This is the recommended user option. For advanced users, please refer to MOOSE and PETSc manuals.
<code>type</code>	Indicating the preconditioning matrix type is SMP for this case.
<code>full</code>	true means the entire preconditioning matrix will be assembled in the calculations.

solve_type PJFNK solver will be used for this problem.

The Executioner Block:

```
[Executioner]
  type = Transient
  scheme = 'implicit-euler'
  dt = 1.25e-2
  dtmin = 1.e-5

  nl_rel_tol = 1e-8
  nl_abs_tol = 1e-6
  nl_max_its = 15

  l_tol = 1e-6
  l_max_its = 30

  start_time = 0.0
  end_time = 250.0
  num_steps = 500

  [./Quadrature]
    type = TRAP
    order = FIRST
  [../]
[]
```

type	Transient executioner type will be used.
scheme	Implicit euler time integration will be used. If this is absent, the default option of BDF2 will be used.
dt	Time step size to be used .
dtmin	Minimum time step size to be used. If the time step size is below this value, the case will stop executing.
nl_rel_tol	Relative tolerance for the nonlinear solve. The recommended value is 1.0E-8.

<code>nl_abs_tol</code>	Absolute tolerance for the nonlinear solve. The recommended value is 1.0E-6.
<code>nl_max_its</code>	Maximum number of nonlinear solve iterations. The recommended value is 15.
<code>l_tol</code>	Relative tolerance for the linear Krylov solve. The recommended value is 1.0E-6.
<code>l_max_its</code>	Maximum number of liner iterations. The recommended value is 30.
<code>start_time</code>	The start time of the simulation.
<code>end_time</code>	The end time of the simulation.
<code>num_steps</code>	The maximum number of time integration steps for the simulation. The simulation will stop either the maximum number of time steps is reached or the <code>end_time</code> is reach.
<code>[./Quadrature]</code>	The quadrature subblock. This is the recommended option.
<code>type</code>	Type of the quadrature rule. Advanced users can refer to the MOOSE manual for other options.
<code>order</code>	Order of the quadrature.

The Outputs Block:

```
[Outputs]
  [./out]
    type = Exodus
    use_displaced = true
    output_initial = true
    sequence = false
    append_displaced = true
  [../]

  [./console]
    type = Console
```

```
perf_log = true
[../]
[]
```

[./out] **Output subblock. This is the recommended option.**

type **Exodus output file type.**

use_displaced **Use the displaced mesh to show the results.**

output_initial **The initial condition is output to the solution file.**

sequence **Enable or disable sequential file output.**

append_displaced **Append '_displaced' to the output file name.**

[./console] **Console subblock.**

type **Console type.**

perf_log **Print all performance logs.**

5.2 Example 2: Use Functions to Set Initial Conditions for a Simple Pipe Flow Problem

5.2.1 Problem Description

This example illustrates how to use functions to set the initial conditions. The example used here is to set the initial pressure distribution in a pipe for an isothermal flow through one simple pipe with boundary conditions set by time dependent volumes (TDV).

Functions:

Initial pressure distribution in the pipe

Axis = 0 The axis used (0, 1, or 2 for x, y, or z) if this is to be a function of position. Note that for RELAP-7 1-D component such as a pipe, we only use x coordinate in the calculations therefore we only need to set Axis = 0.

Type = PiecewiseLinear

x = '0 3' This is the pipe axial direction spatial coordinate relative to the starting point

y = '1.05e5 1e5'

Initial velocity distribution in the pipe

Axis = 0

Type = PiecewiseLinear

x = '0 3'

y = '3 3'

Fig. 3 shows the plot of pressure versus pipe length for this case.

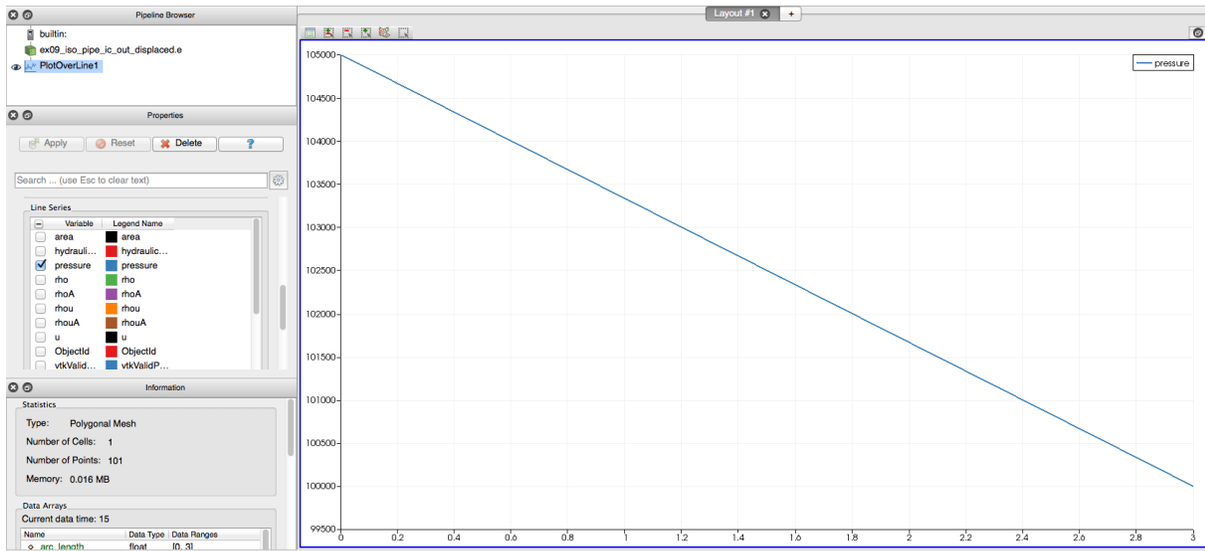


Figure 3. Pressure vs length for an isothermal pipe flow problem with initial pressure set by functions

5.2.2 Input File

The input file for this example is shown as follows:

```
#
# Isothermal flow through one simple pipe with B.C. set by TDVs and I.C. set by functions
#

[GlobalParams]
  model_type = 2
  scaling_factor_var = '1e4 1.0'
  stabilization_type = 'SUPG'
[]

[EoS]
  [./eos]
    type = BarotropicEquationOfState
    p_0 = 1.e5
```

```

    rho_0 = 1.e3
    a2 = 1.e7
[../]
[]

[Functions]
# initial pressure distribution in the pipe
[./p_func]
    axis = 0
    type = PiecewiseLinear
    x = '0      3'
    y = '1.05e5  1e5'
[../]

# initial velocity distribution in the pipe
[./v_func]
    axis = 0
    type = PiecewiseLinear
    x = '0      3'
    y = '3      3'
[../]
[]

[Components]
[./pipe1]
    type = Pipe
    eos = eos
    position = '0 2 0'
    orientation = '1 1 1'
    Dh = 0.01
    length = 3.0
    n_elems = 10
    A = 7.854e-5
    f = 0.01
    P_func = p_func
    V_func = v_func
[../]

[./inlet]
    type = TimeDependentVolume
    input = 'pipe1(in)'
```

```

    eos = eos
    p_bc = 1.05e5
[../]

[./outlet]
    type = TimeDependentVolume
    input = 'pipe1(out)\'
    eos = eos
    p_bc = 1.0e5
[../]
[]

[Preconditioning]
[./SMP_PJFNK]
    type = SMP
    full = true
    solve_type = 'PJFNK\'
[../]
[]

[Executioner]
    type = Transient
    dt = 0.5
    dtmin = 1.e-5

[Executioner]
    type = Transient
    dt = 0.5
    dtmin = 1.e-5

petsc_options_iname = '-mat_fd_type -mat_mffd_type -pc_type -ksp_gmres_restart\'
petsc_options_value = ' ds ds lu 30\'

nl_rel_tol = 1e-8
nl_abs_tol = 1e-8
nl_max_its = 10

l_tol = 1e-4
l_max_its = 60

start_time = 0.0

```

```

num_steps = 30

[./Quadrature]
  type = TRAP
  order = FIRST
[../]
[]

[Outputs]
[./out]
  type = Exodus
  use_displaced = true
  output_initial = true
  sequence = false
  append_displaced = true
[../]

[./console]
  type = Console
  perf_log = true
[../]
[]

```

5.2.3 Description of the Input File

The following will provide explanations to the input parameters that did not exist in Example 1.

The Functions Block:

```

[Functions]
# initial pressure distribution in the pipe
[./p_func]
  axis = 0
  type = PiecewiseLinear
  x = '0          3'
  y = '1.05e5    1e5'
[../]

# initial velocity distribution in the pipe
[./v_func]

```

```

axis = 0
type = PiecewiseLinear
x = '0 3'
y = '3 3'
[../]
[]

```

[./p_func] Subblock for the pressure distribution function.

axis For RELAP-7 1-D components, only the x coordinate is used in the calculations and hence axis is set to be 0 which allows a function to be defined as a function x. If axis is not given in the input file, the data pair of (x, y) will be taken as a time function.

type Type of functions used to interpolate data between data points.

x The coordinate used for the x-axis data.

y The initial pressure distribution function data point values.

The Components Block:

```

[Components]
[./pipe1]
type = Pipe
eos = eos
position = '0 2 0'
orientation = '1 1 1'
Dh = 0.01
length = 3.0
n_elems = 10
A = 7.854e-5
f = 0.01
P_func = p_func
V_func = v_func
[../]

[./inlet]
type = TimeDependentVolume
input = 'pipe1(in)'
eos = eos

```

```

    p_bc = 1.05e5
[../]

[./outlet]
    type = TimeDependentVolume
    input = 'pipe1(out)\'
    eos = eos
    p_bc = 1.0e5
[../]
[]

```

P_func Use the pressure function defined in the Functions block to set the initial pressure value.

V_func Use the velocity function defined in the Functions block to set the initial velocity value.

The Executioner Block:

```

[Executioner]
    type = Transient
    dt = 0.5
    dtmin = 1.e-5

[Executioner]
    type = Transient
    dt = 0.5
    dtmin = 1.e-5

    petsc_options_iname = '-mat_fd_type -mat_mffd_type -pc_type\'
    petsc_options_value = ' ds ds lu\'

    nl_rel_tol = 1e-8
    nl_abs_tol = 1e-8
    nl_max_its = 10

    l_tol = 1e-4
    l_max_its = 60

    start_time = 0.0
    num_steps = 30

```

```

[./Quadrature]
  type = TRAP
  order = FIRST
[../]
[]

nl_rel_tol = 1e-8
nl_abs_tol = 1e-8
nl_max_its = 10

l_tol = 1e-4
l_max_its = 60

start_time = 0.0
num_steps = 30

[./Quadrature]
  type = TRAP
  order = FIRST
[../]
[]

```

`petsc_options_iname` Names of PETSc name and value pairs. `-mat_fd_type = ds` is an option to control how the Jacobian matrix is approximated by using the finite differencing method. `-mat_mffd_type = ds` is an option to control how matrix-free operation is approximated by using the finite difference method. Users are recommended to refer to the PETSc manual for other options.

`petsc_options_value` Values of PETSc name/value pairs.

5.3 Example 3: Two Phase Flow Through a Pipe - HEM Model

5.3.1 Problem Description

Two phase flow with HEM model through one simple pipe with boundary conditions set by time dependent volumes and initial conditions set by functions. The initial conditions are close to steady state solutions; therefore the simulation quickly converges to the steady state.

The following is the description of the functions defined in the `Functions` block:

Initial pressure distribution in the pipe

Axis = 0

Type = PiecewiseLinear

x = '0 1'

y = '7.01e6 7e6'

Initial velocity distribution in the pipe

Axis = 0

Type = PiecewiseLinear

x = '0 1'

y = '1.35 2.1'

Initial temperature distribution in the pipe

Axis = 0

Type = PiecewiseLinear

x = '0 0.04 1'

y = '517 517.343 517.253'

Initial void fraction distribution in the pipe

axis = 0

type = PiecewiseLinear

x = '0 0.04 1'

y = '0 0. 0.37'

The parameters and functions used for the pipe component are the following:

Position = '0 0 0'

Orientation = '1 0 0'

Area = $7.854e - 5m^2$

Diameter = 0.01 m

Length = 1. m

Number of elements = 100

Friction coefficient = 0.1

Convective heat transfer coefficient = 1e4

Heat flux perimeter = 0.031416

Wall temperature = 528 K

The name of EOS to use = two_phase_eos

Initial condition determined by functions which are close to SS solutions

Function which defines pressure as a function of x = p_func

Function which defines velocity as a function of x = v_func

Function which defines temperature as a function of x = temp_func

Function which defines vapor volume fraction as a function of x = void_func

The inlet boundary conditions:

Vapor volume fraction = 0.

The name of EOS to use = two_phase_eos

The outlet boundary conditions:

Vapor volume fraction = 0.5

The name of EOS to use = two_phase_eos

5.3.2 Input File

```
#
# Two phase flow with HEM model through one simple pipe with B.C. set by TDVs and
# I.C. set by functions. The I.C.s are close to steady state solutions; therefore
# the simulation quickly converges to the steady state.
#

[GlobalParams]
  model_type = 32
  stabilization_type = 'NONE'
  scaling_factor_var = '1e+4 1 1e-2'
  gravity = '0.0, 0.0, -9.8'
[]

[EoS]
  [./two_phase_eos]
    type = TwoPhaseStiffenedGasEOS
  [../]
[]

[Functions]

# initial pressure distribution in the pipe
[./p_func]
  axis = 0
  type = PiecewiseLinear
  x = '0      1'
  y = '7.01e6  7e6'
[../]

# initial velocity distribution in the pipe
[./v_func]
  axis = 0
  type = PiecewiseLinear
  x = '0      1'
  y = '1.35  2.1'
[../]

# initial temperature distribution in the pipe
```

```

[./temp_func]
  axis = 0
  type           = PiecewiseLinear
  x              = '0    0.04    1'
  y              = '517    517.343    517.253'
[../]

# initial void fraction distribution in the pipe
[./void_func]
  axis = 0
  type           = PiecewiseLinear
  x              = '0    0.04    1'
  y              = '0    0.    0.37'
[../]
[]

[Components]
[./pipe1]
  type = Pipe
  position = '0 0 0'
  orientation = '1 0 0'
  A = 7.854e-5
  Dh = 0.01
  length = 1.
  n_elems = 100
  f = 0.1
  Hw = 1e4
  Phf = 0.031416000000
  Tw = 528
  eos = two_phase_eos

# initial condition determined by functions which are close to steady state solutions
P_func = p_func
V_func = v_func
T_func = temp_func
volume_fraction_vapor_func = void_func
[../]

[./inlet]
  type = TimeDependentVolume
  input = 'pipe1(in)'

```

```

    p_bc = 7.01e6
    T_bc = 517.
    volume_fraction_vapor_bc = 0.
    eos = two_phase_eos
[../]

[./outlet]
    type = TimeDependentVolume
    input = 'pipe1(out)'
    p_bc = 7.0e6
    T_bc = 517
    volume_fraction_vapor_bc = 0.5
    eos = two_phase_eos
[../]
[]

[Preconditioning]
[./SMP_PJFNK]
    type = SMP
    full = true
    solve_type = 'PJFNK'
    petsc_options_iname = '-mat_fd_type -mat_mffd_type -pc-type'
    petsc_options_value = 'ds ds lu'
[../]
[]

[Executioner]
    type = Transient
    scheme = 'implicit-euler'

    dt = 1e-1
    dtmin = 1.e-3

    nl_rel_tol = 1e-9
    nl_abs_tol = 1e-9
    nl_max_its = 50

    l_tol = 1e-3
    l_max_its = 60

    start_time = 0.0

```

```

num_steps = 50

[./Quadrature]
  type = TRAP
  order = FIRST
[../]
[]

[Outputs]
[./out]
  type = Exodus
  use_displaced = true
  output_initial = true
  sequence = false
  append_displaced = true
[../]

[./console]
  type = Console
  perf_log = true
[../]
[]

```

5.3.3 Description of the Input File

The following are the detailed explanations of the input parameters that did not exist in the previous examples.

The Global Parameters Block:

```

[GlobalParams]
  model_type = 32
  stabilization_type = 'NONE'
  scaling_factor_var = '1e+4 1 1e-2'
  gravity = '0.0, 0.0, -9.8'
[]

```

`model_type` `model_type=32` indicates that the HEM two-phase flow model will be used for this problem.

gravity The gravity acceleration vector.

The Equation of State Block:

```
[EoS]
  [./two_phase_eos]
    type = TwoPhaseStiffenedGasEOS
  [../]
[]
```

type Two phase stiffened gas equation of state will be used for this problem.

The Components Block:

```
[Components]
  [./pipe1]
    type = Pipe
    position = '0 0 0'
    orientation = '1 0 0'
    A = 7.854e-5
    Dh = 0.01
    length = 1.
    n_elems = 100
    f = 0.1
    Hw = 1e4
    Phf = 0.031416000000
    Tw = 528
    eos = two_phase_eos
    P_func = p_func
    V_func = v_func
    T_func = temp_func
    volume_fraction_vapor_func = void_func
  [../]

  [./inlet]
    type = TimeDependentVolume
    input = 'pipe1(in)\'
    p_bc = 7.01e6
    T_bc = 517.
    volume_fraction_vapor_bc = 0.
```

```
    eos = two_phase_eos
[../]

[./outlet]
    type = TimeDependentVolume
    input = 'pipe1(out)'  
    p_bc = 7.0e6  
    T_bc = 517  
    volume_fraction_vapor_bc = 0.5  
    eos = two_phase_eos
[../]
[]
```

Hw	The wall convective heat transfer coefficient.
Phf	Heat flux perimeter.
Tw	Initial pipe wall temperature.
T_bc	Temperature values at the pipe boundaries.
volume_fraction_vapor_bc	Vapor phase volume fraction at the pipe boundaries.

5.4 Example 4: Two Phase Flow Through a Pipe - 7-Equation Model

5.4.1 Problem Description

This example shows a two phase flow with the 7-equation model through one simple pipe with boundary conditions set by Inlet and Outlet components at the pipe inlet and outlet ends respectively. The initial conditions are set by functions. The initial conditions are set to be close to steady state solutions; therefore the simulation quickly converges to steady state solutions.

5.4.2 Input File

The input file for this pipe flow case with the 7-equation two-phase flow model is shown below:

```
[GlobalParams]
model_type = 7
global_init_T = 517.252
global_init_P = 7.0e6
global_init_V = 0
global_init_volume_fraction_vapor = 0.95
scaling_factor_var_2phase = '1
                                1e1 1e1 1e-3
                                1e1 1e1 1e-3'

phase_interaction = true
pressure_relaxation = true
velocity_relaxation = true
interface_transfer = true
wall_mass_transfer = true
stabilization_type = LAPIDUS
specific_interfacial_area_max_value = 1700
[]

[Functions]

# initial pressure distribution in the pipe
[./p_func]
axis = 0
type = PiecewiseLinear
x = '0          3.66'
```

```

    y = '7.005e6 7e6'
[../]

# initial velocity distribution in the pipe
[./v_func]
    axis = 0
    type = PiecewiseLinear
    x = '0      3.66'
    y = '6      6.6'
[../]

# initial temperature distribution in the pipe
[./temp_func]
    axis = 0
    type = PiecewiseLinear
    x = '0      3.66'
    y = '517.252 517.817'
[../]

# initial void fraction distribution in the pipe
[./void_func]
    axis = 0
    type = PiecewiseLinear
    x = '0      3.66'
    y = '0.95  0.956'
[../]

[]

[EoS]
[./eos_liquid]
    type = StiffenedGasEquationOfStateLiquid
    gamma = 2.35
    q = -1167e3
    q_prime = 0
    p_inf = 1.e9
    cv = 1816
[../]

[./eos_vapor]
    type = StiffenedGasEquationOfStateVapor

```

```

gamma = 1.43
q = 2030e3
q_prime = -23e3
p_inf = 0
cv = 1040
[../]
[]

[Components]
[./pipe]
type = Pipe
eos_liquid = eos_liquid
eos_vapor = eos_vapor
position = '0 0 0'
orientation = '0 0 1'
A = 1.907720E-04
Dh = 1.698566E-02
length = 3.66
f = 1.698566E-02
Hw_liquid = 1000.0
Hw_vapor = 1000.0
Phf = 0.0489623
Tw = 550
n_elems = 50

P_liquid_func = p_func
P_vapor_func = p_func
V_liquid_func = v_func
V_vapor_func = v_func
T_liquid_func = temp_func
T_vapor_func = temp_func
volume_fraction_vapor_func = void_func
[../]

[./inlet]
type = Inlet
input = 'pipe(in)'
p_liquid = 7.005e6
p_vapor = 7.005e6
T_liquid = 517.252
T_vapor = 517.252

```

```

    volume_fraction_vapor = 0.95
    eos_liquid = eos_liquid
    eos_vapor = eos_vapor
[../]

[./outlet]
    type = Outlet
    input = 'pipe(out)'
    p_vapor = 7.0e6
    p_liquid = 7.0e6
    eos_liquid = eos_liquid
    eos_vapor = eos_vapor
[../]
[]

[Preconditioning]
[./SMP_PJFNK]
    type = SMP
    full = true
    solve_type = 'PJFNK'
[../]
[]

[Executioner]
    type = Transient
    dt = 1e-1
    dtmin = 1.e-5
    nl_rel_tol = 1e-10
    nl_abs_tol = 1e-8
    nl_max_its = 30
    l_tol = 1e-3
    l_max_its = 30

    PetscOptionsIname = '-mat_mffd_type -ksp_gmres_restart'
    PetscOptionsValue = 'ds 300'

    start_time = 0.0
    num_steps = 100

[./Quadrature]
    type = TRAP

```

```

    order = FIRST
  [../]
[]

[Outputs]
  [./out]
    type = Exodus
    use_displaced = true
    output_initial = true
    sequence = false
    append_displaced = true
  [../]

  [./console]
    type = Console
    perf_log = true
  [../]
[]

```

5.4.3 Description of the Input File

The following provides the explanations of the input parameters that did not exist in the previous examples:

The Global Parameters Block:

```

[GlobalParams]
  model_type = 7
  global_init_T = 517.252
  global_init_P = 7.0e6
  global_init_V = 0
  global_init_volume_fraction_vapor = 0.95
  scaling_factor_var_2phase = '1
                                1e1 1e1 1e-3
                                1e1 1e1 1e-3'

  phase_interaction = true
  pressure_relaxation = true
  velocity_relaxation = true
  interface_transfer = true

```

```

wall_mass_transfer = true
stabilization_type = LAPIDUS
specific_interfacial_area_max_value = 1700
[]

```

model_type = 7	Turn on the 7-equation two phase flow model.
scaling_factor_var_2phase	Scaling factors for each variable of the 7-equation two phase flow model.
phase_interaction	Option to turn on/off the interaction between the two phases.
pressure_relaxation	Option to turn on/off the pressure relaxation terms.
velocity_relaxation	Option to turn on/off the velocity relaxation terms.
interface_transfer	Option to turn on/off the interface mass and heat transfer.
wall_mass_transfer	Option to turn on/off wall boiling model to calculate the interfacial mass transfer near the wall.
specific_interfacial_area_max_value	The maximum value of the specific interfacial area.

The Equation of State Block:

```

[EoS]
[./eos_liquid]
type = StiffenedGasEquationOfStateLiquid
gamma = 2.35
q = -1167e3
q_prime = 0
p_inf = 1.e9
cv = 1816
[../]

[./eos_vapor]
type = StiffenedGasEquationOfStateVapor
gamma = 1.43

```

```

q = 2030e3
q_prime = -23e3
p_inf = 0
cv = 1040
[../]
[]

```

[./eos_liquid] Subblock for the liquid phase equation of state.

type Stiffened gas equation of state (SGEOS) for liquid.

gamma The γ value in SGEOS calculations for liquid.

q The q value in SGEOS calculations for liquid.

q_prime The q' value in SGEOS calculations for liquid.

p_inf The p_∞ value in SGEOS calculations for liquid.

cv The specific heat value in SGEOS calculations for liquid.

The Components Block

```

[Components]
[./pipe]
type = Pipe
eos_liquid = eos_liquid
eos_vapor = eos_vapor
position = '0 0 0'
orientation = '1 0 0'
A = 1.907720E-04
Dh = 1.698566E-02
length = 3.66
f = 1.698566E-02
Hw_liquid = 1000.0
Hw_vapor = 1000.0
Phf = 0.0489623
Tw = 550
n_elems = 50

```

```

P_liquid_func = p_func
P_vapor_func = p_func
V_liquid_func = v_func
V_vapor_func = v_func
T_liquid_func = temp_func
T_vapor_func = temp_func
volume_fraction_vapor_func = void_func
[../]

[./inlet]
type = Inlet
input = 'pipe(in)'
p_liquid = 7.005e6
p_vapor = 7.005e6
T_liquid = 517.252
T_vapor = 517.252
volume_fraction_vapor = 0.95
eos_liquid = eos_liquid
eos_vapor = eos_vapor
[../]

[./outlet]
type = Outlet
input = 'pipe(out)'
p_vapor = 7.0e6
p_liquid = 7.0e6
eos_liquid = eos_liquid
eos_vapor = eos_vapor
[../]
[]

```

eos_liquid	Defines the name of the equation of state to use for the liquid phase.
eos_vapor	Defines the name of the equation of state to use for the vapor phase.
Hw_liquid	Convective heat transfer coefficient for the liquid phase.
Hw_vapor	Convective heat transfer coefficient for the vapor phase.

<code>P_liquid_func</code>	Function which defines liquid phase pressure as a function of x .
<code>P_vapor_func</code>	Function which defines vapor phase pressure as a function of x .
<code>T_liquid_func</code>	Function which defines liquid phase temperature as a function of x .
<code>T_vapor_func</code>	Function which defines vapor phase temperature as a function of x .
<code>volume_fraction_vapor_func</code>	Function which defines vapor phase volume fraction as a function of x .
<code>p_liquid</code>	Prescribed pressure value at the pipe boundary for the liquid phase.
<code>p_vapor</code>	Prescribed pressure value at the pipe boundary for the vapor phase.
<code>T_liquid</code>	Prescribed temperature value at the pipe boundary for the liquid phase.
<code>T_vapor</code>	Prescribed temperature value at the pipe boundary for the vapor phase.
<code>volume_fraction_vapor</code>	Prescribed vapor phase volume fraction.

5.5 Example 5: A Core Channel Problem

5.5.1 Problem Description

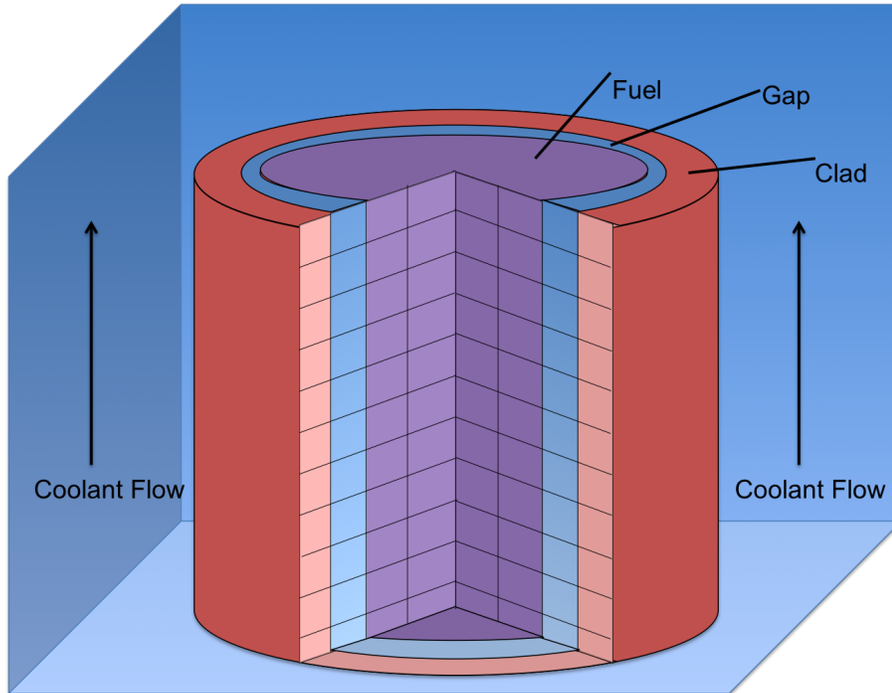


Figure 4. Diagram of a core channel problem

This example simulates single phase fluids flow and heat transfer in a core channel, as schematically shown in Fig. 4. This problem simulates fluids flow in one subchannel with a single fuel rod as the heat source. The boundary conditions are applied to the ends of the core channel. The reactor power = 77337.69407 W is used for the core channel. Other parameters used are the following:

Fluids Flow:

Cross section flow area = $8.7878e-5 \text{ m}^2$

Hydraulic Diameter = 0.01179 m

Length = 4 m

Friction coefficient = 0.01

Convective heat transfer coefficient = $5.33e4 \text{ W/m}^2 * K$

Heat flux perimeter = 0.029832559676 m
Number of elements = 20

Heat Structure:

Dimension of mesh used for heat conduction = 1
Geometry type of fuel = cylinder
Heat structure names = 'fuel gap clad'
Number of heat structures = 3
Width of each heat structure = '0.004096 0.0001 0.000552' m
Number of elements of each heat structure = '10 1 2'
Name of materials used in the heat structure = 'fuel-mat gap-mat clad-mat'
Fraction of reactor power goes into heat structure = '1.0 0.0 0.0'
Initial Solid Temperature = 559.15 K

The boundary conditions are the following:

Inlet:

Pressure = 155.483e5 Pa
Temperature = 559.15 K

Outlet:

Pressure = 155.e5Pa
Temperature = 559.15 K

The solid material properties (k = Thermal conductivity, Cp = heat capacity, rho = Density) are:

Fuel:

k = 2.5
Cp = 300
rho = 1.032e4

Gap:

k = 0.6
Cp = 1.
rho = 1.

Clad:

k = 21.5
Cp = 350.

$\rho = 6.55e3$

5.5.2 Input File

The following shows the input file for the Core Channel example. Please note that the mesh used for this problem does not correspond to what is shown in Fig. 4.

```
[GlobalParams]
  global_init_P = 155.e5
  global_init_V = 0.
  global_init_T = 559.15
  model_type = 3
  stabilization_type = 'SUPG'
  scaling_factor_var = '1e4 1e1 1e-2'
  temperature_sf = '1e-2'
[]

[EoS]
  [./eos]
    type = NonIsothermalEquationOfState
    p_0 = 155.e5
    rho_0 = 686.0
    a2 = 1.e7
    beta = .46e-3
    cv = 5.5e3
    e_0 = 3075325
    T_0 = 559.15
  [../]
[]

[Materials]
  [./fuel-mat]
    type = SolidMaterialProperties
    k = 2.5
    Cp = 300.
    rho = 1.032e4
  [../]
  [./gap-mat]
```

```

    type = SolidMaterialProperties
    k = 0.6
    Cp = 1.
    rho = 1.
[../]
[./clad-mat]
    type = SolidMaterialProperties
    k = 21.5
    Cp = 350.
    rho = 6.55e3
[../]
[]

[Components]
[./reactor]
    type = Reactor
    initial_power = 77337.69407
[../]

[./CCH1]
    type = CoreChannel
    eos = eos
    position = '0 0 0'
    orientation = '0 0 1'
    A = 8.7878e-5
    Dh = 0.01179
    length = 4
    n_elems = 20
    f = 0.01
    Hw = 5.33e4
    Phf = 0.029832559676
    Ts_init = 559.15
    dim_hs = 1
    fuel_type = cylinder
    name_of_hs = 'fuel gap clad'
    n_heatstruct = 3
    width_of_hs = '0.004096 0.0001 0.000552'
    elem_number_of_hs = '10 1 2'
    material_hs = 'fuel-mat gap-mat clad-mat'
    power_fraction = '1.0 0.0 0.0'
[../]

```

```

[./inlet]
  type = TimeDependentVolume
  input = 'CCH1(in)'
  p_bc = 155.483e5
  T_bc = 559.15
  eos = eos
[../]

[./outlet]
  type = TimeDependentVolume
  input = 'CCH1(out)'
  p_bc = '155.e5'
  T_bc = 559.15
  eos = eos
[../]
[]

[Preconditioning]
[./SMP_PJFNK]
  type = SMP
  full = true
  solve_type = 'PJFNK'
[../]
[]

[Executioner]
  type = Transient
  scheme = 'implicit-euler'
  dt = 1.e-2 #1.e-5
  dtmin = 1.e-5

  nl_rel_tol = 1e-9
  nl_abs_tol = 1e-8
  nl_max_its = 20

  l_tol = 1e-3
  l_max_its = 30

  start_time = 0.0
  num_steps = 50

```

```

[./Quadrature]
  type = TRAP
  order = FIRST
[../]
[]

[Outputs]
[./out]
  type = Exodus
  use_displaced = true
  output_initial = true
  sequence = false
  append_displaced = true
[../]

[./console]
  type = Console
  perf_log = true
[../]
[]

```

5.5.3 Description of the Input File

The Global Parameter Block:

```

[GlobalParams]
  global_init_P = 155.e5
  global_init_V = 0.
  global_init_T = 559.15
  model_type = 3
  stabilization_type = 'SUPG'
  scaling_factor_var = '1e4 1e1 1e-2'
  temperature_sf = '1e-2'
[]

```

temperature_sf Scaling factor for the temperature variable in solid materials.

The Materials Block:

```

[Materials]
  ./fuel-mat
    type = SolidMaterialProperties
    k = 2.5
    Cp = 300.
    rho = 1.032e4
  ../
  ./gap-mat
    type = SolidMaterialProperties
    k = 0.6
    Cp = 1.
    rho = 1.
  ../
  ./clad-mat
    type = SolidMaterialProperties
    k = 21.5
    Cp = 350.
    rho = 6.55e3
  ../
[]

```

k **Thermal conductivity.**

Cp **Specific heat.**

rho **Density of solid materials.**

```

[Components]
  ./reactor
    type = Reactor
    initial_power = 77337.69407
  ../

  ./CCH1
    type = CoreChannel
    eos = eos
    position = '0 0 0'
    orientation = '0 0 1'
    A = 8.7878e-5

```

```

Dh = 0.01179
length = 4
n_elems = 20
f = 0.01
Hw = 5.33e4
Phf = 0.029832559676
Ts_init = 559.15
dim_hs = 1
fuel_type = cylinder
name_of_hs = 'fuel gap clad'
n_heatstruct = 3
width_of_hs = '0.004096 0.0001 0.000552'
elem_number_of_hs = '10 1 2'
material_hs = 'fuel-mat gap-mat clad-mat'
power_fraction = '1.0 0.0 0.0'
[../]

[./inlet]
type = TimeDependentVolume
input = 'CCH1(in)'
p_bc = 155.483e5
T_bc = 559.15
eos = eos
[../]

[./outlet]
type = TimeDependentVolume
input = 'CCH1(out)'
p_bc = '155.e5'
T_bc = 559.15
eos = eos
[../]
[]

```

[./reactor] **Subblock for the reactor component.**

initial_power **Prescribed initial reactor power.**

[./CCH1] **Subblock for the core channel component.**

Hw	Wall heat transfer coefficient.
Ts_init	Prescribed initial temperature of the solid materials.
dim_hs	The dimension of the mesh used for the heat conduction calculations in the heat structure. The options are <code>dim_hs=1</code> for 1D heat conduction calculations or <code>dim_hs=2</code> for 2D heat conduction calculations.
fuel_type	Geometry type of the fuel. The available options are <code>fuel_type = cylinder</code> or <code>fuel_type = plate</code> .
name_of_hs	Prescribed heat structure names.
n_heatstruct	Prescribed number of heat structures.
width_of_hs	Width of each heat structure.
elem_number_of_hs	Number of elements of each heat structure.
material_hs	Name of the materials (defined in the Materials block) used in the heat structure.
power_fraction	The fraction of reactor power that goes into each heat structure.

5.6 Example 6: A Two Pipes Flow Problem

5.6.1 Problem Description

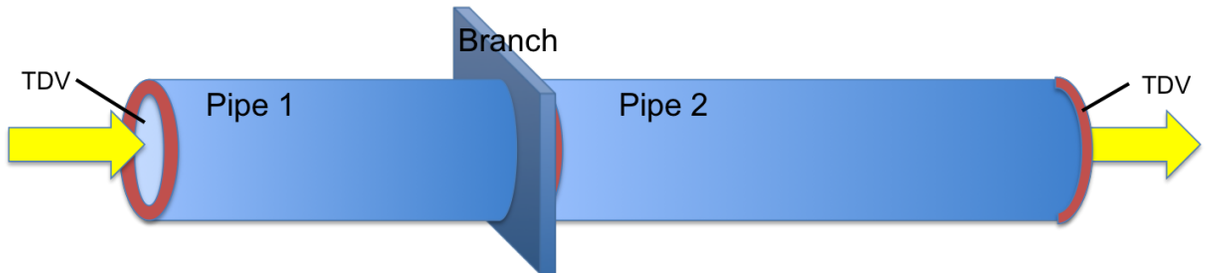


Figure 5. Diagram of a two pipes flow problem

This example demonstrates flow through two pipes with different parameters connected by a branch. There is no heat exchange through the pipe walls for this case. The following are the input parameters for each component:

Pipe 1:
Hydraulic Diameter = .01 m
Area = $1.0e-4 \text{ m}^2$
Length = 1 m
Friction coefficient = .01
Convective heat transfer coefficient = 0

Pipe 2:
Hydraulic Diameter = .02 m
Area = $4e-4 \text{ m}^2$
Length = 1.5 m
Friction = .02
Convective Heat Transfer Coefficient = 0

Branch:
Form loss coefficients = '5 5'
Area = $1.5e-4 \text{ m}^2$

Initial pressure = 1e5 Pa

Inlet:

Pressure = 1.01e5 Pa

Temperature = 300 K

Outlet:

Pressure = 1e5 Pa

The following figure shows the results of the pressure versus pipe length for this case.

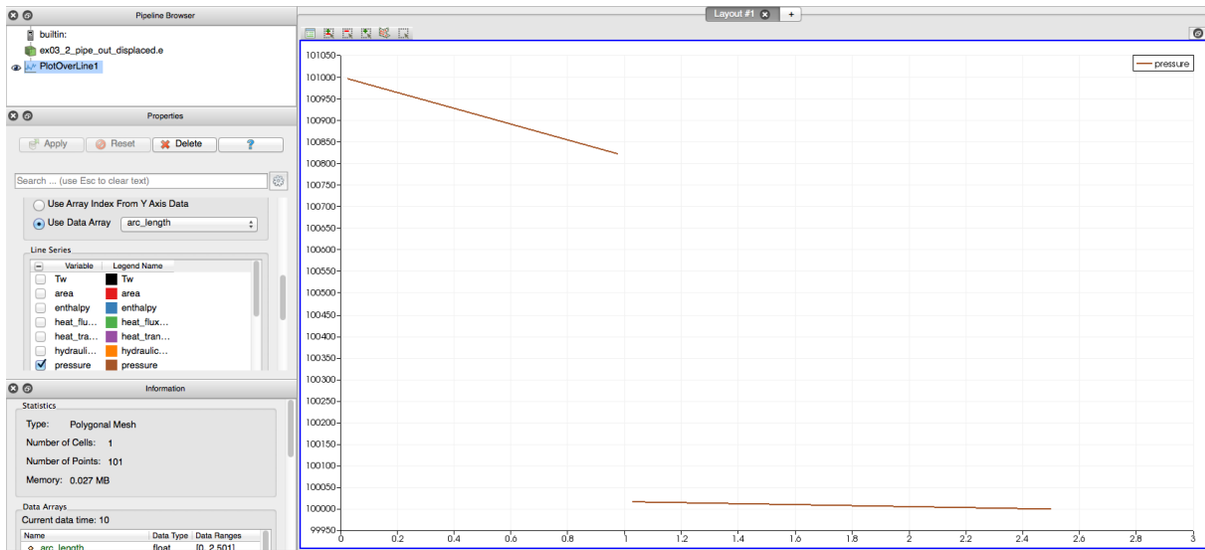


Figure 6. Pressure vs. length for the two pipes flow problem

The results of the pressure difference can be checked by doing hand calculations by summing ΔP using:

$$\Delta P = \frac{fL\rho u^2}{2D_h} \quad (2)$$

and:

$$\Delta P = \frac{k\rho u^2}{2} \quad (3)$$

5.6.2 Input File

The input file for this example problem is listed as follows:

```
[GlobalParams]
  model_type = 3
  gravity = '0 0 0'
  stabilization_type = 'SUPG'
  scaling_factor_var = '1.e3 1.e-1 1.e-3'
  global_init_T = 300
  global_init_P = 1.0e5
  global_init_V = 1
[]

[EoS]
  [./eos]
    type = NonIsothermalEquationOfState
    p_0 = 1.e5
    rho_0 = 1.e3
    a2 = 1.e7
    beta = .46e-3
    cv = 4.18e3
    e_0 = 1.254e6
    T_0 = 300
  [../]
[]

[Components]
  [./pipe1]
    type = Pipe
    eos = eos
    position = '0 0 0'
    orientation = '1 0 0'
    length = 1
    n_elems = 25
    A = 1.0e-4
    Dh = 0.01
    f = .01
    Hw = 0.
  [../]
```

```
[./Branch1]
  type = Branch
  inputs = 'pipe1(out) '
  outputs = 'pipe2(in) '
  K = '5 5'
  Area = 1.5e-4
  Initial_pressure = 1e5
  eos = eos
[../]

[./pipe2]
  type = Pipe
  eos = eos
  position = '1.001 0 0'
  orientation = '1 0 0'
  length = 1.5
  n_elems = 25
  A = 4e-4
  Dh = 0.02
  f = .02
  Hw = 0.
[../]

[./inlet]
  type = TimeDependentVolume
  input = 'pipe1(in) '
  p_bc = 1.01e5
  T_bc = 300
  eos = eos
[../]

[./outlet]
  type = TimeDependentVolume
  input = 'pipe2(out) '
  p_bc = 1e5
  eos = eos
[../]
[]

[Preconditioning]
```

```

[./SMP_PJFNK]
  type = SMP
  full = true
  solve_type = 'PJFNK'
[../]
[]

[Executioner]
  type = Transient
  scheme = 'implicit-euler'

  dt = 1e-2
  dtmin = 1e-5

  nl_rel_tol = 1e-8
  nl_abs_tol = 1e-6
  nl_max_its = 30

  l_tol = 1e-4
  l_max_its = 30

  PetscOptionsIname = '-mat_fd_coloring_err -mat_fd_type -mat_mffd_type -pc_type'
  PetscOptionsValue = '1.e-10                ds                ds                lu'

  start_time = 0.0
  end_time = 10.0
  num_steps = 5000

[./Quadrature]
  type = TRAP
  order = FIRST
[../]
[]

[Outputs]
[./out]
  type = Exodus
  use_displaced = true
  output_initial = true
  sequence = false
  append_displaced = true

```

```

[../]

[./console]
  type = Console
  perf_log = true
[../]
[]

```

5.6.3 Description of the Input File

The only input parameters that need explanations are those in the Branch subblock.

```

[./Branch1]
  type = Branch
  inputs = 'pipe1(out)'
  outputs = 'pipe2(in) '
  K = '5 5'
  Area = 1.5e-4
  Initial_pressure = 1e5
  eos = eos
[../]

```

[./Branch]	Subblock for the Branch component.
type	The component type is Branch.
inputs	Inputs of this branch connects to the outlet end of pipe1.
outputs	Outputs of this branch connects to the inlet end of pipe2.
K	Form loss coefficients at the branch inputs and outputs.
Area	The reference area of this branch.
Initial_pressure	Prescribed initial pressure for the branch.
eos	The name of the equation of state to be used for this branch.

5.7 Example 7: A Volume Branch Case - Three Pipes In and Two Pipes Out

5.7.1 Problem Description

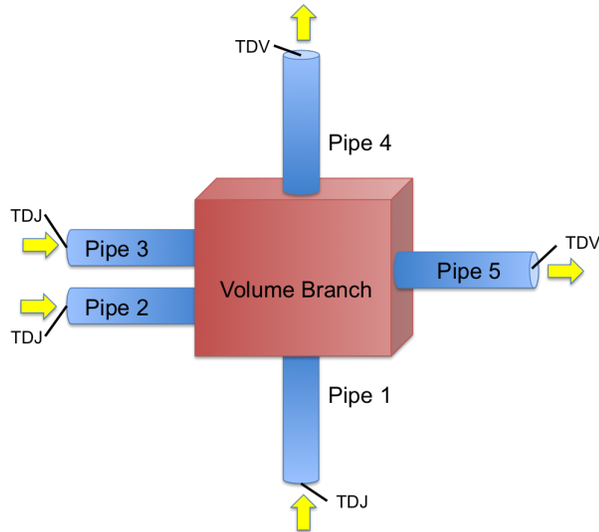


Figure 7. Diagram of a volume branch case with three pipes flowing in and two pipes flowing out

This example shows a volume branch case which connects three pipes with water flowing into them and two pipes with water flowing out of them. The pipe sizes are all the same, but the boundary conditions vary. The volume branch model includes the volume effects of a junction. The input parameters used in this example are listed in the following:

Pipes:

Area = $3.14 \times 10^{-4} \text{ m}^2$

Diameter = 0.02 m

Length = 1 m

Number of elements in pipe = 10

Friction coefficient = 0.01

Branch1:

Type = VolumeBranch

inputs = 'pipe1(out) pipe2(out) pipe3(out)'

outputs = 'pipe4(in) pipe5(in)'

Form loss coefficients = '0.01 0.01 0.01 0.01 100'

Area = $3.14e - 2m^2$

volume = $3.14e - 2m^3$

Initial Temperature = 628.15 K

Inlet 1:

type = TimeDependentJunction

Velocity = 1.0 m/s

Temperature = 628.15 K

Inlet 2:

type = TimeDependentJunction

Velocity = 1.0 m/s

Temperature = 628.15 K

Inlet 3:

type = TimeDependentJunction

Velocity = 10.0 m/s

Temperature = 528.15 K

Outlet 1:

type = TimeDependentVolume

Pressure = $1.0e5$ Pa

Temperature = 628.15 K

Outlet 2:

type = TimeDependentVolume

Pressure = $1.5e5$ Pa

Temperature = 628.15 K

5.7.2 Input File

```
[GlobalParams]
  global_init_P = 1.5e5
  global_init_V = 1
  global_init_T = 628.15
  model_type = 3
  stabilization_type = 'SUPG'
  scaling_factor_var = '1e4 1 1e-3'
[]

[EoS]
  active = 'eos'
  [./eos]
    type = NonIsothermalEquationOfState
    p_0 = 1e5
    rho_0 = 865.51
    a2 = 5.7837e6
    beta = 2.7524e-4
    cv = 1272.0
    e_0 = 7.9898e5
    T_0 = 628.15
  [../]
[]

[Components]

  [./pipe1]
    type = Pipe
    eos = eos
    position = '0 0 0'
    orientation = '0 0 1'
    A = 3.14e-4
    Dh = 0.02
    length = 1
    n_elems = 10
    f = 0.01
    Hw = 0
  [../]
```

```
[./pipe2]
type = Pipe
eos = eos
position = '-1.5 0 1.25'
orientation = '1 0 0'
A = 3.14e-4
Dh = 0.02
length = 1
n_elems = 10
f = 0.01
Hw = 0
[../]
```

```
[./pipe3]
type = Pipe
eos = eos
position = '-1.5 0 1.75'
orientation = '1 0 0'
A = 3.14e-4
Dh = 0.02
length = 1
n_elems = 10
f = 0.01
Hw = 0
[../]
```

```
[./pipe4]
type = Pipe
eos = eos
position = '0 0 2'
orientation = '0 0 1'
A = 3.14e-4
Dh = 0.02
length = 1
n_elems = 10
f = 0.01
Hw = 0
[../]
```

```
[./pipe5]
```

```

type = Pipe
eos = eos
# geometry
position = '0.5 0 1.5'
orientation = '1 0 0'
A = 3.14e-4
Dh = 0.02
length = 1
n_elems = 10
f = 0.01
Hw = 0
[../]

[./branch1]
type = VolumeBranch
eos = eos
center = '0 0 1.5'
inputs = 'pipe1(out) pipe2(out) pipe3(out)'
outputs = 'pipe4(in) pipe5(in)'
K = '0.01 0.01 0.01 0.01 100'
Area = 3.14e-2
volume = 3.14e-2
initial_T = 628.15
[../]

[./inlet1]
type = TimeDependentJunction
input = 'pipe1(in)'
eos = eos
v_bc = 1.0
T_bc = 628.15
[../]

[./inlet2]
type = TimeDependentJunction
input = 'pipe2(in)'
eos = eos
v_bc = 1.0
T_bc = 628.15
[../]

```

```

[./inlet3]
  type = TimeDependentJunction
  input = 'pipe3(in)'
  eos = eos
  v_bc = 10.0
  T_bc = 528.15
[../]

[./outlet1]
  type = TimeDependentVolume
  input = 'pipe4(out)'
  eos = eos
  p_bc = '1.0e5'
  T_bc = 628.15
[../]

[./outlet2]
  type = TimeDependentVolume
  input = 'pipe5(out)'
  eos = eos
  p_bc = '1.5e5'
  T_bc = 628.15
[../]
[]

[Preconditioning]

[./SMP_PJFNK]
  type = SMP
  full = true
  solve_type = 'PJFNK'
[../]

[]

[Executioner]
  type = Transient
  scheme = 'implicit-euler'

  dt = 0.2

```

```

dtmin = 1e-10

# setting time step range
[./TimeStepper]
  type = FunctionDT
  time_t = '0    0.05  0.051    0.3   0.31   2    2.1   1e5'
  time_dt = '1.e-2  1.e-2  5e-2    5e-2  2e-1  0.2  0.5   0.5 '
[../]

nl_rel_tol = 1e-6
nl_abs_tol = 1e-7
nl_max_its = 30

l_tol = 1e-6
l_max_its = 100

petsc_options_iname = '-pc_type -ksp_gmres_restart'
petsc_options_value = 'lu      300'

start_time = 0.0
num_steps = 500

[./Quadrature]
  type = TRAP
  order = FIRST
[../]
[]

[Outputs]
[./out]
  type = Exodus
  use_displaced = true
  output_initial = true
  sequence = false
  append_displaced = true
[../]

[./console]
  type = Console
  perf_log = true
[../]

```

[]

5.7.3 Description of the Input File

```
[./branch1]
  type = VolumeBranch
  eos = eos
  center = '0 0 1.5'
  inputs = 'pipe1(out) pipe2(out) pipe3(out) '
  outputs = 'pipe4(in) pipe5(in) '
  K = '0.01 0.01 0.01 0.01 100'
  Area = 3.14e-2
  volume = 3.14e-2
  initial_T = 628.15
[../]
```

center **The (x, y, z) coordinate of the center of the volume branch.**

volume **The volume of VolumeBranch component.**

```
[./TimeStepper]
  type = FunctionDT
  time_t = '0      0.05  0.051      0.3    0.31    2    2.1    1e5'
  time_dt = '1.e-2  1.e-2 5e-2      5e-2  2e-1  0.2  0.5    0.5 '
[../]
```

[./TimeStepper] **Subblock to define time step size as a function of simulation time.**

type **FunctionDT type to be used.**

time_t **The value of time t.**

time_dt **The value of time step.**

5.8 Example 8: A Simple Pipe Loop with Pump

5.8.1 Problem Description

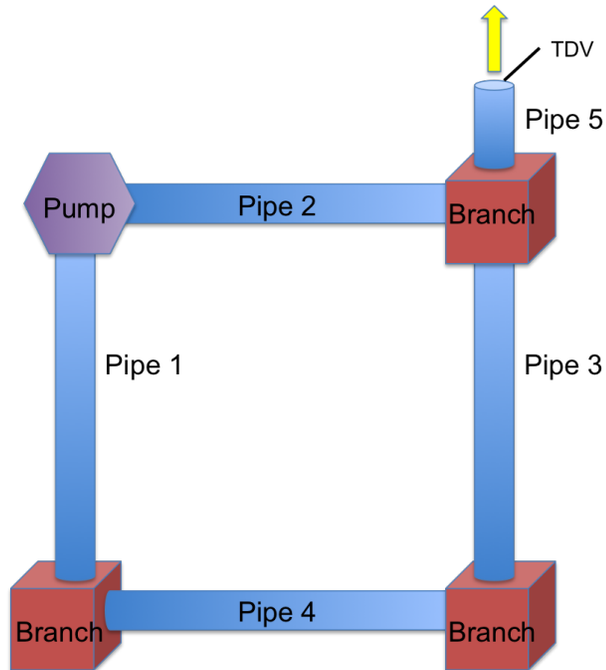


Figure 8. Diagram of a simple loop of pipes connected by branches and a pump

This example shows a loop made up of 5 pipes, 3 branches and a pump. The pump acts as a junction, connecting pipes 1 and 2, and pipe 5 is used as an outlet to help control the pressure. The input parameters used for the Pump model are the following:

Pump head = 1.0

Form loss coefficient (in the reverse direction) = '10. 10.'

Area = $0.785398163e - 3m^2$

Initial pressure = 1.e5 Pa

5.8.2 Input File

```
[GlobalParams]
  model_type = 3
  stabilization_type = 'SUPG'
  scaling_factor_var = '1e4 1e1 1e-2'
[]

[EoS]
  [./eos]
    type = NonIsothermalEquationOfState
    p_0 = 1.e5
    rho_0 = 1.e3
    a2 = 1.e7
    beta = -.46e-3
    cv = 4.18e3
    e_0 = 1.254e6
    T_0 = 300
  [../]
[]

[Components]

  [./pipe1]
    type = Pipe
    eos = eos
    position = '0 0 0'
    orientation = '1 0 0'
    A = 0.785398163e-4
    Dh = 0.01
    length = 1
    n_elems = 10
    f = 0.01
    Hw = 0.0
  [../]

  [./pipe2]
    type = Pipe
    eos = eos
    position = '1 0 0'
```

```
orientation = '0 1 0'  
A = 0.785398163e-4  
Dh = 0.01  
length = 1  
n_elems = 10  
f = 0.01  
Hw = 0.0  
[../]  
  
[./pipe3]  
type = Pipe  
eos = eos  
position = '1 1 0'  
orientation = '-1 0 0'  
A = 0.785398163e-4  
Dh = 0.01  
length = 1  
n_elems = 10  
f = 0.01  
Hw = 0.0  
[../]  
  
[./pipe4]  
type = Pipe  
eos = eos  
# geometry  
position = '0 1 0'  
orientation = '0 -1 0'  
A = 0.785398163e-4  
Dh = 0.01  
length = 1  
n_elems = 10  
f = 0.01  
Hw = 0.0  
[../]  
  
[./pipe5]  
type = Pipe  
eos = eos  
position = '1 1 0'  
orientation = '0 1 0'
```

```

A = 0.785398163e-4
Dh = 0.01
length = .25
n_elems = 10
f = 0.01
Hw = 0.0
[../]

[./pump]
type = Pump
eos = eos
inputs = 'pipe1(out)'
outputs = 'pipe2(in)'
Head = 1.0
K_reverse = '10. 10.'
Area = 0.785398163e-3
Initial_pressure = 1.e5
[../]

[./branch2]
type = Branch
eos = eos
inputs = 'pipe2(out)'
outputs = 'pipe3(in) pipe5(in)'
K = '3. 3. 3.'
Area = 0.785398163e-3
Initial_pressure = 1.e5
[../]

[./branch3]
type = Branch
eos = eos
inputs = 'pipe3(out)'
outputs = 'pipe4(in)'
K = '3. 3.'
Area = 0.785398163e-3
Initial_pressure = 1.e5
[../]

[./branch4]
type = Branch

```

```

eos = eos
inputs = 'pipe4(out)'
outputs = 'pipe1(in)'
K = '3. 3.'
Area = 0.785398163e-3
Initial_pressure = 1.e5
[../]

[./TDV]
type = TimeDependentVolume
input = 'pipe5(out)'
p_bc = '1.e5'
T_bc = 300.0
eos = eos
[../]
[]

[Preconditioning]
[./SMP_PJFNK]
type = SMP
full = true
solve_type = 'PJFNK'
[../]
[]

[Executioner]
type = Transient
scheme = 'implicit-euler'
dt = 1.e-4
dtmin = 1e-7

nl_rel_tol = 1e-6
nl_abs_tol = 1e-8
nl_max_its = 10

l_tol = 1e-8
l_max_its = 60

petsc_options_iname = '-ksp_gmres_restart'
petsc_options_value = '30'

```

```

start_time = 0.0
num_steps = 50

[./Quadrature]
  type = TRAP
  order = FIRST
[../]
[]

[Outputs]
[./out]
  type = Exodus
  use_displaced = true
  output_initial = true
  sequence = false
  append_displaced = true
[../]

[./console]
  type = Console
  perf_log = true
[../]
[]

```

5.8.3 Description of the Input File

```

[./pump]
  type = Pump
  eos = eos
  inputs = 'pipe1(out)'
  outputs = 'pipe2(in)'
  Head = 1.0
  K_reverse = '10. 10.'
  Area = 0.785398163e-3
  Initial_pressure = 1.e5
[../]

```

[./pump] Subblock for the pump component.

type	Component type is a pump.
eos	Name of equation of state to be used by this pump component.
inputs	The inlet of the pump connects to outlet of pipe1.
outputs	The outlet of the pump connects to the inlet of pipe2.
Head	Prescribed pump head.
K_reverse	Form loss coefficient in reverse flow direction.
Area	Reference area for the pump.
Initial_pressure	Prescribed initial pressure for this pump component.

5.9 Example 9: A Heat Exchanger Problem

5.9.1 Problem Description

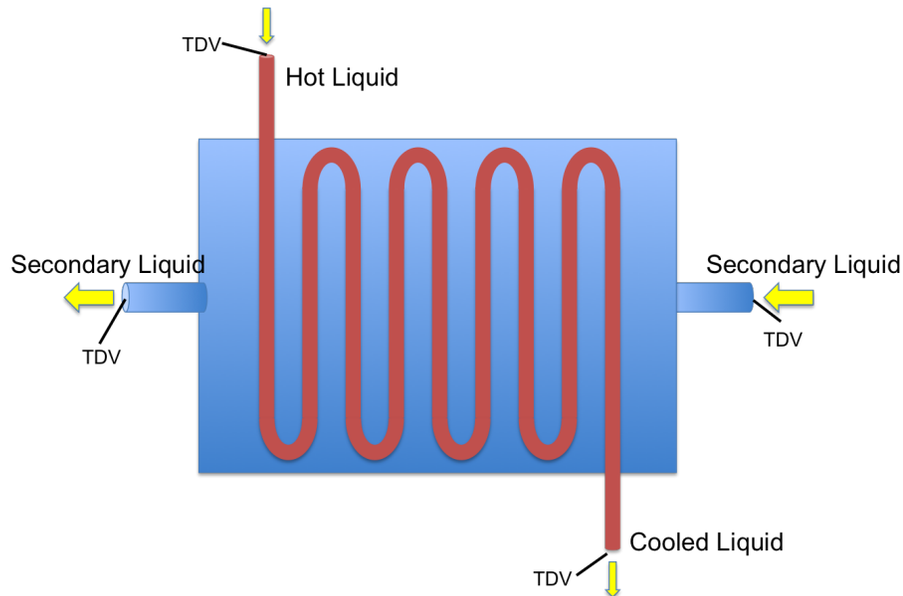


Figure 9. Diagram of a heat exchanger problem

This example illustrates how to run a case with the heat exchanger component. The heat exchanger has two inlets and two outlets, primary and secondary respectively. The primary side inlet lets in hot water, while the secondary side inlet lets in liquid at a cooler temperature. This helps cool the primary side liquid, so the primary side outlet temperature is somewhat lower while the secondary side outlet temperature is somewhat higher. This example problem is for subcooled water only. The input parameters used for this case are listed in the following:

Materials:

Thermal Conductivity = 100.0

Density = 100.0

Specific Heat = 100.0

Heat Exchanger:

Area of primary pipe = $0.785398163e - 4m^2$
Area of secondary pipe = $0.785398163e - 4m^2$
Diameter of primary pipe = 0.01 m
Diameter of secondary pipe = 0.01 m
Length = 1 m
Primary convective heat transfer coefficient = 1.e4
Secondary convective heat transfer coefficient = 1.e4
Primary heat flux perimeter= 0.031415926520
Secondary heat flux perimeter = 0.031415926520
Friction coefficient = 0.01
Initial wall temperature = 300.0 K
Dimension of mesh used for wall = 1
Thickness of the wall between primary and secondary loop = 0.001 m

Primary inlet TDV:
Pressure = 1.05e5 Pa
Temperature = 400.0 K

Primary outlet TDV:
Pressure = 1.e5 Pa
Temperature = 300.0 K

Secondary inlet TDV:
Pressure = 1.05e5 Pa
Temperature = 300.0 K

Secondary outlet TDV:
Pressure = 1.e5 Pa
Temperature = 300.0 K

5.9.2 Input File

```
[GlobalParams]  
  global_init_P = 1.e5
```

```

global_init_V = 0.
global_init_T = 300.
model_type = 3
stabilization_type = 'SUPG'
scaling_factor_var = '1e4 1e1 1e-2'
temperature_sf = '1e-2'
[]

[EoS]
[./eos]
type = NonIsothermalEquationOfState
p_0 = 1.e5
rho_0 = 1.e3
a2 = 1.e7
beta = .46e-3
cv = 4.18e3
e_0 = 1.254e6
T_0 = 300
[../]
[]

[Materials]
[./wall-mat]
type = SolidMaterialProperties
k = 100.0
rho = 100.0
Cp = 100.0
[../]
[]

[Components]
[./HX]
type = HeatExchanger
eos = eos
eos_secondary = eos
position = '0 0 0'
orientation = '1 0 0'
A = 0.785398163e-4
A_secondary = 0.785398163e-4
Dh = 0.01

```

```

Dh_secondary = 0.01
length = 1
n_elems = 10
Hw = 1.e4
Hw_secondary = 1.e4
Phf = 0.031415926520
Phf_secondary = 0.031415926520
f = 0.01
f_secondary = 0.01
Twall_init = 300.0
dim_wall = 1
wall_thickness = 0.001
material_wall = wall-mat
n_wall_elems = 2
[../]

[./primary_inlet_TDV]
type = TimeDependentVolume
input = 'HX(primary_in)\'
p_bc = 1.05e5
T_bc = 400.0
eos = eos
[../]

[./primary_outlet_TDV]
type = TimeDependentVolume
input = 'HX(primary_out)\'
p_bc = '1.e5\'
T_bc = 300.0
eos = eos
[../]

[./secondary_inlet_TDV]
type = TimeDependentVolume
input = 'HX(secondary_in)\'
p_bc = 1.05e5
T_bc = 300.0
eos = eos
[../]

[./secondary_outlet_TDV]

```

```

    type = TimeDependentVolume
    input = 'HX(secondary_out)'
    p_bc = '1.e5'
    T_bc = 300.0
    eos = eos
  [../]
[]

[Preconditioning]
  [./SMP_PJFNK]
    type = SMP
    full = true
    solve_type = 'PJFNK'
    line_search = basic
  [../]
[]

[Executioner]
  type = Transient
  scheme = 'implicit-euler'
  dt = 1.e-3
  dtmin = 1e-7

  nl_rel_tol = 1e-6
  nl_abs_tol = 1e-8
  nl_max_its = 20

  l_tol = 1e-6
  l_max_its = 300

  petsc_options_iname = '-ksp_gmres_restart -pc_type'
  petsc_options_value = '300 lu'

  start_time = 0.0
  num_steps = 1000

  [./Quadrature]
    type = TRAP
    order = FIRST
  [../]
[]

```

```

[Outputs]
  [./out]
    type = Exodus
    use_displaced = true
    output_initial = true
    sequence = false
    append_displaced = true
  [../]

  [./console]
    type = Console
    perf_log = true
  [../]
[]

```

5.9.3 Description of the Input File

The input parameters for the heat exchanger component needs explanantions.

```

[./HX]
  type = HeatExchanger
  eos = eos
  eos_secondary = eos
  position = '0 0 0'
  orientation = '1 0 0'
  A = 0.785398163e-4
  A_secondary = 0.785398163e-4
  Dh = 0.01
  Dh_secondary = 0.01
  length = 1
  n_elems = 10
  Hw = 1.e4
  Hw_secondary = 1.e4
  Phf = 0.031415926520
  Phf_secondary = 0.031415926520
  f = 0.01
  f_secondary = 0.01
  Twall_init = 300.0
  dim_wall = 1

```

```

wall_thickness = 0.001
material_wall = wall-mat
n_wall_elems = 2
[../]

```

[./HX]	Subblock for the heat exchanger component.
eos	Name of the equation of state to be used by the fluids in the primary side of the heat exchanger.
eos_secondary	Name of the equation of state to be used by the fluids in the secondary side of the heat exchanger.
position	The x, y, z coordinate of the start of heat exchanger.
orientation	The orientation vector of heat exchanger.
A	The flow area of the primary side of the heat exchanger.
A_secondary	The flow area of the secondary side of the heat exchanger.
Dh	Hydraulic diameter of the primary side of the heat exchanger.
Dh_secondary	Hydraulic diameter of the secondary side of the heat exchanger.
length	Length of heat exchanger.
n_elems	The number of elements in the heat exchanger.
Hw	The convective heat transfer coefficient on the primary side of the heat exchanger.
hw_secondary	The convective heat transfer coefficient on the secondary side of the heat exchanger.
Phf	Heat flux perimeter of the primary side of the heat exchanger.
Phf_secondary	Heat flux perimeter of the secondary side of the heat exchanger.
f	Wall friction coefficient on the primary side of the heat exchanger.

f_secondary	Wall friction coefficient on the secondary side of the heat exchanger.
Twall_init	Prescribed initial temperature of the solid materials between the fluids.
dim_wall	The dimension of the mesh used in the heat conduction calculations of the wall in the heat exchanger.
wall_thickness	The wall thickness.
material_wall	Name of the material used in the wall.
n_wall_elems	Number of elements in the wall.

5.10 Example 10: A Loop With Core Channel and Heat Exchanger

5.10.1 Problem Description

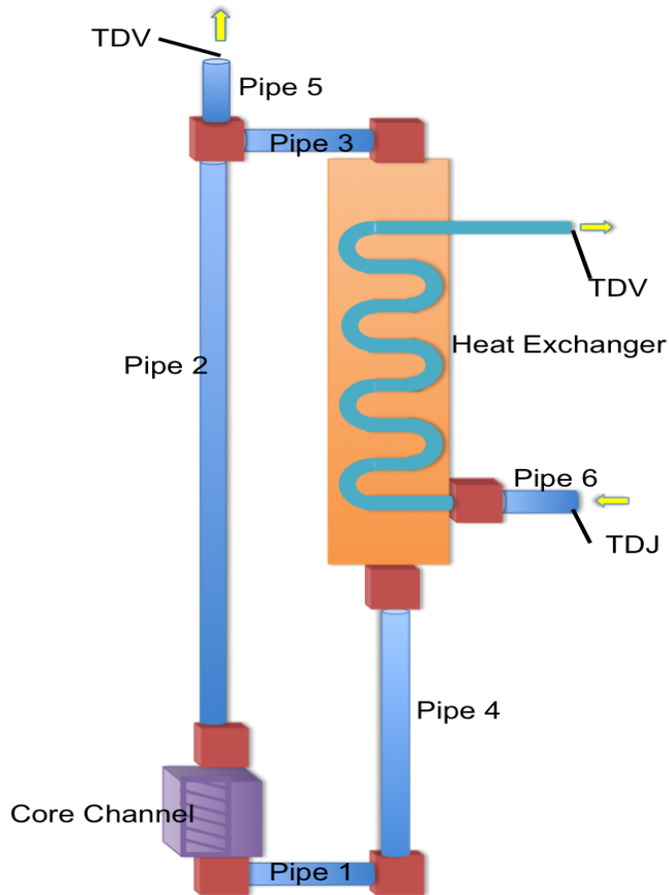


Figure 10. Diagram of a loop with core channel and heat exchanger

This example is a simple loop which consists of 6 pipes, a core channel, and a heat exchanger. Each component differs in length, but has the same area and diameter. The core channel is used to heat up the liquid, while the heat exchanger is used to cool it down. Pipe 5 is used to control the system pressure. The input parameters are listed in the following:

Materials:

Fuel

Thermal conductivity = 29.3 W/m K

Specific heat = 191.67 J/kg K

Density = $1.4583e4 \text{ kg/m}^3$

Clad

Thermal conductivity = 26.3 W/m K

Specific heat = 638 J/kg K

Density = $7.646e3 \text{ kg/m}^3$

Wall

Thermal conductivity = 26.3 W/m K

Specific heat = 638 J/kg K

Density = $7.646e3 \text{ kg/m}^3$

Reactor's initial power = $5.129e7$

Pipe 1:

Area = 0.44934 m^2

Diameter = $2.972e-3 \text{ m}$

Length = 1 m

Friction Coefficient = 0.001

Hw = 0.0

Pipe 2 length = 5.18 m

Pipe 3 length = 1 m

Pipe 4 length = 2.27 m

Pipe 5 length = 0.02 m

Pipe 6 length = 0.3 m

Core Channel:

Length = 0.8 m

Friction coefficient = 0.022

Convective heat transfer coefficient = $1.6129e5$

Heat flux perimeter = 497.778

Initial solid temperature = 628.15 K
Width of each heat structure = '0.00348 0.00052'

Heat Exchanger:

Area of primary pipe = $0.44934m^2$
Area of secondary pipe = $0.44934m^2$
Diameter of primary pipe = 0.0186 m
Diameter of secondary pipe = 0.014 m
Length = 3.71 m
Primary convective heat transfer coefficient = $1.6129e5$
Secondary convective heat transfer coefficient = $1.6129e5$
Primary heat flux perimeter = 327.568
Secondary heat flux perimeter = 327.568
Friction coefficient = 0.022
Initial wall temperature = 628.15 K
Dimension of mesh used for wall = 1
Thickness of the wall between primary and secondary loop = 0.0044 m

Branches:

Form loss coefficients = '0.5 0.5'
Area = $0.44934m^2$
Initial Pressure = $1e5$ Pa (Branch 1 = $2e5$ Pa)

Inlet:

Type = TimeDependentJunction
Input = 'pipe6(in)'
Velocity = 0.5 m/s
Temperature = 628.15 K

Outlet 1:

Type = TimeDependentVolume
Input = 'IHX(secondary_out)'
Pressure = $1.0e5$ Pa
Temperature = 761.15 K

Outlet 2:

Type = TimeDependentVolume
Input = 'pipe5(out)' Pressure = $1e5$ Pa

Temperature = 783.15 K

5.10.2 Input File

```
[GlobalParams]
  model_type = 3
  global_init_P = 1.0e5
  global_init_V = 0.5
  global_init_T = 628.15
  stabilization_type = 'NONE'
[]

[EoS]
  [./eos]
    type = NonIsothermalEquationOfState
    p_0 = 1e5
    rho_0 = 865.51
    a2 = 5.7837e6
    beta = 2.7524e-4
    cv = 1272.0
    e_0 = 7.9898e5
    T_0 = 628.15
  [../]
[]

[Materials]
  [./fuel-mat]
    type = SolidMaterialProperties
    k = 29.3
    Cp = 191.67
    rho = 1.4583e4
  [../]
  [./clad-mat]
    type = SolidMaterialProperties
    k = 26.3
    Cp = 638
    rho = 7.646e3
  [../]
```

```
[./wall-mat]
  type = SolidMaterialProperties
  k = 26.3
  rho = 7.646e3
  Cp = 638
[../]
[]
```

```
[Components]
```

```
[./reactor]
  type = Reactor
  initial_power = 5.1296e7
[../]
```

```
[./pipe1]
  type = Pipe
  eos = eos
  position = '0 1 0'
  orientation = '0 -1 0'
  A = 0.44934
  Dh = 2.972e-3
  length = 1
  n_elems = 5
  f = 0.001
  Hw = 0.0
[../]
```

```
[./CH1]
  type = CoreChannel
  eos = eos
  position = '0 0 0'
  orientation = '0 0 1'
  A = 0.44934
  Dh = 2.972e-3
  length = 0.8
  n_elems = 10
  f = 0.022
  Hw = 1.6129e5
  Phf = 497.778852000000
```

```

dim_hs = 1
name_of_hs = 'fuel clad'
Ts_init = 628.15
n_heatstruct = 2
fuel_type = cylinder
width_of_hs = '0.00348 0.00052'
elem_number_of_hs = '4 1'
material_hs = 'fuel-mat clad-mat'
#peak_power = '1.3366e8 0.'
power_fraction = '1.0 0.0'
[../]

[./pipe2]
type = Pipe
eos = eos
position = '0 0 0.8'
orientation = '0 0 1'
A = 0.44934
Dh = 2.972e-3
length = 5.18
n_elems = 5
f = 0.001
Hw = 0.0
[../]

[./pipe3]
type = Pipe
eos = eos
position = '0 0 5.98'
orientation = '0 1 0'
A = 0.44934
Dh = 2.972e-3
length = 1
n_elems = 5
f = 0.001
Hw = 0.0
[../]

[./IHX]
type = HeatExchanger
eos = eos

```

```
eos_secondary = eos
position = '0 1.0 5.98'
orientation = '0 0 -1'
A = 0.44934
A_secondary = 0.44934
Dh = 0.0186
Dh_secondary = 0.014
length = 3.71
n_elems = 10
Hw = 1.6129e5
Hw_secondary = 1.6129e5
Phf = 327.568860000000
Phf_secondary = 327.568860000000
f = 0.022
f_secondary = 0.022
Twall_init = 628.15
wall_thickness = 0.0044
dim_wall = 1
material_wall = wall-mat
n_wall_elems = 2
[../]
```

```
[./pipe4]
type = Pipe
eos = eos
position = '0 1.0 2.27'
orientation = '0 0 -1'
A = 0.44934
Dh = 2.972e-3
length = 2.27
n_elems = 5
f = 0.001
Hw = 0.0
[../]
```

```
[./pipe5]
type = Pipe
eos = eos
position = '0 0 5.98'
orientation = '0 0 1'
A = 0.44934
```

```

Dh = 2.972e-3
length = 0.02
n_elems = 2
f = 10
Hw = 0.0
[../]

[./Branch1]
type = Branch
inputs = 'pipe1(out)'
outputs = 'CH1(in) '
K = '0.5 0.5'
Area = 0.44934
Initial_pressure = 2e5
eos = eos
[../]

[./Branch2]
type = Branch
inputs = 'CH1(out) '
outputs = 'pipe2(in)'
K = '0.5 0.5'
Area = 0.44934
Initial_pressure = 1.0e5
eos = eos
[../]

[./Branch3]
type = Branch
inputs = 'pipe2(out)'
outputs = 'pipe3(in) pipe5(in)'
K = '0.0 0.0 0.0'
Area = 0.44934
Initial_pressure = 1e5
eos = eos
[../]

[./Branch4]
type = Branch
inputs = 'pipe3(out)'
outputs = 'IHX(primary_in)'
```

```

K = '0.1 0.1'
Area = 0.44934
Initial_pressure = 1e5
eos = eos
[../]

[./Branch5]
type = Branch
inputs = 'IHX(primary_out)'
outputs = 'pipe4(in)'
K = '0.0 0.0'
Area = 0.44934
Initial_pressure = 1e5
eos = eos
[../]

[./Branch6]
type = Branch
inputs = 'pipe4(out)'
outputs = 'pipe1(in)'
K = '0.0 0.0'
Area = 0.44934
Initial_pressure = 1e5
eos = eos
[../]

[./pipe6]
type = Pipe
eos = eos
position = '0 1.5 2.27'
orientation = '0 -1 0'
A = 0.44934
Dh = 2.972e-3
length = 0.3
n_elems = 5
f = 0.001
Hw = 0.0
[../]

[./Branch7]
type = Branch

```

```

    inputs = 'pipe6(out)'
    outputs = 'IHX(secondary_in)'
    K = '0.0 0.0'
    Area = 0.44934
    Initial_pressure = 1e5
    eos = eos
[../]

[./inlet_TDJ]
type = TimeDependentJunction
input = 'pipe6(in)'
v_bc = 0.5
T_bc = 628.15
eos = eos
[../]

[./outlet_TDV]
type = TimeDependentVolume
input = 'IHX(secondary_out)'
p_bc = '1.0e5'
T_bc = 761.15
eos = eos
[../]

[./TDV_p]
type = TimeDependentVolume
input = 'pipe5(out)'
p_bc = '1e5'
T_bc = 783.15
eos = eos
[../]
[]

[Preconditioning]
[./SMP_PJFNK]
type = SMP
full = true
solve_type = 'PJFNK'
[../]
[]

```

```

[Executioner]
  type = Transient
  scheme = 'implicit-euler'
  dt = 1e-2
  dtmin = 1e-10
  nl_rel_tol = 1e-8
  nl_abs_tol = 1e-8
  nl_max_its = 30
  l_tol = 1e-6
  l_max_its = 300
  start_time = 0.0
  num_steps = 500
  end_time = 10.
  petsc_options_iname = '-pc_type -mat_fd_type -mat_mffd_type -ksp_gmres_restart'
  petsc_options_value = '  lu      ds      ds      300'

[./Quadrature]
  type = TRAP
  order = FIRST
[../]
[]

[Outputs]
[./out]
  type = Exodus
  use_displaced = true
  output_initial = true
  sequence = false
  append_displaced = true
[../]

[./console]
  type = Console
  perf_log = true
[../]
[]

```

5.11 Example 11: A Model Pressurized Water Reactor Problem

5.11.1 Problem Description

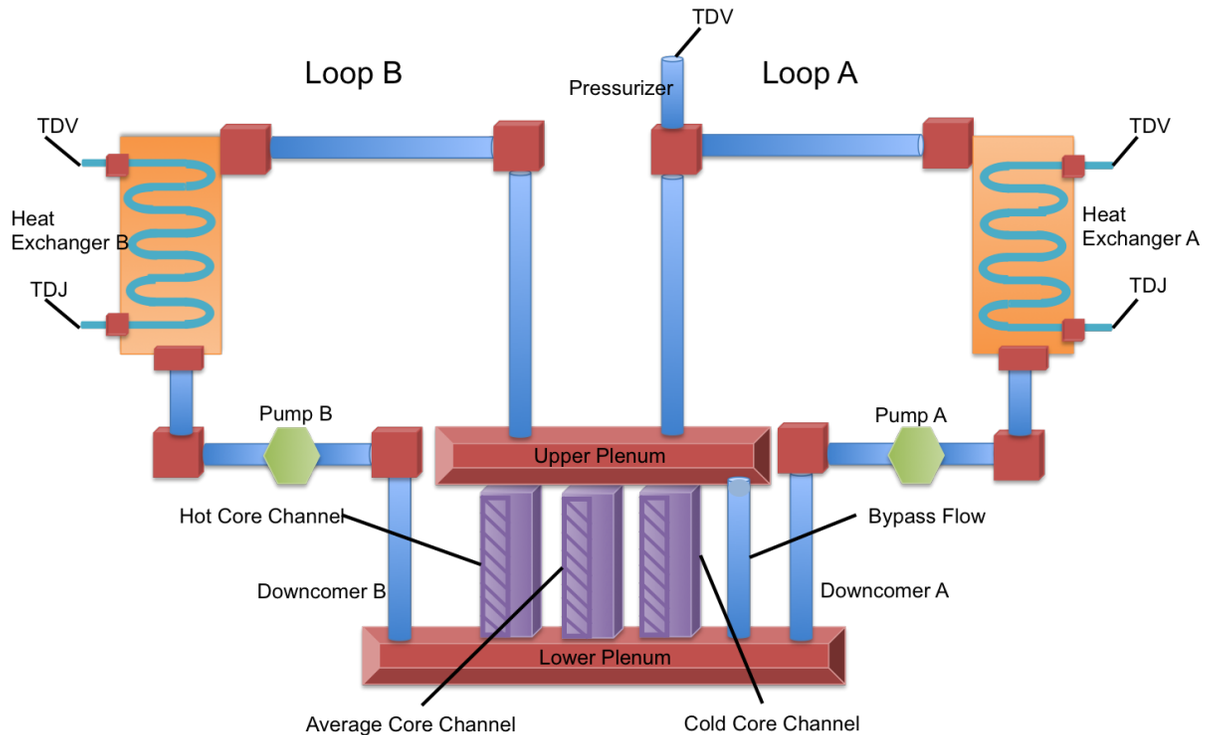


Figure 11. Diagram of a model pressurized water reactor problem

This example shows a simplified PWR plant model. It is made up of three sections - Loop A, Loop B and a reactor vessel model. The reactor vessel region contains three parallel core channels, a bypass flow channel, and an upper and lower plenum. The three core channels represent all the cooling channels and fuel rods in the high power region, average power region and low power region of the reactor core respectively. The upper and lower plenum are modeled with volume branches. The two loops have a Hot Leg, a Heat Exchanger and its secondary side pipes, the Cold Leg and a primary Pump. The heat exchanger secondary side pipes are modeled with subcooled water since the steam generator model is not available yet. Loop A contains a time dependent volume component that works as a pressurizer to help regulate the system pressure.

5.11.2 Input File

```
[GlobalParams]
  model_type = 3
  global_init_P = 15.17e6
  global_init_V = 1.
  global_init_T = 564.15
  scaling_factor_var = '1.e-1 1.e-5 1.e-8'
  temperature_sf = 1e-2
  stabilization_type = 'SUPG'
[]

[EoS]
  [./eos]
    type = NonIsothermalEquationOfState
    p_0 = 15.17e6
    rho_0 = 738.350
    a2 = 1.e7
    beta = .46e-3
    cv = 5.832e3
    e_0 = 3290122.80
    T_0 = 564.15
  [../]
[]

[Materials]
  [./fuel-mat]
    type = SolidMaterialProperties
    k = 3.65
    Cp = 288.734
    rho = 1.0412e2
  [../]

  [./gap-mat]
    type = SolidMaterialProperties
    k = 1.084498
    Cp = 1.0
    rho = 1.0
  [../]
```

```

[./clad-mat]
  type = SolidMaterialProperties
  k = 16.48672
  Cp = 321.384
  rho = 6.6e1
[../]

[./clad3-mat]
  type = SolidMaterialProperties
  k = 16.48672
  Cp = 6.6e3
  rho = 6.6e1
[../]

[./wall-mat]
  type = SolidMaterialProperties
  k = 100.0
  rho = 100.0
  Cp = 100.0
[../]
[]

[Components]
  active = 'reactor
          CH1 CH2 CH3 bypass_pipe LowerPlenum UpperPlenum
          DownComer-A
          pipe1-HL-A pipe2-HL-A
          HX-A
          pipe1-CL-A pipe2-CL-A
          Pump-A
          pipe1-SC-A pipe2-SC-A
          MassFlowRateIn-SC-A PressureOutlet-SC-A
          Branch1-A Branch2-A Branch3-A Branch4-A Branch5-A Branch6-A
          pipe-to-Pressurizer Pressurizer
          DownComer-B
          pipe1-HL-B pipe2-HL-B
          HX-B
          pipe1-CL-B pipe2-CL-B
          Pump-B
          pipe1-SC-B pipe2-SC-B

```

```
MassFlowRateIn-SC-B PressureOutlet-SC-B
Branch1-B Branch2-B Branch3-B Branch4-B Branch5-B Branch6-B
'
```

```
[./reactor]
  type = Reactor
  initial_power = 2.77199979e9
[../]
```

```
#Vessel region components
```

```
[./CH1]
  type = CoreChannel
  eos = eos
  position = '0 -1.2 0'
  orientation = '0 0 1'
  A = 1.161864
  Dh = 0.01332254
  length = 3.6576
  n_elems = 8
  f = 0.01
  Hw = 5.33e4
  Phf = 321.341084980423
  Ts_init = 564.15
  dim_hs = 1
  n_heatstruct = 3
  name_of_hs = 'FUEL GAP CLAD'
  fuel_type = cylinder
  width_of_hs = '0.0046955 0.0000955 0.000673'
  elem_number_of_hs = '3 1 1'
  material_hs = 'fuel-mat gap-mat clad-mat'
  power_fraction = '3.33672612e-1 0 0'
[../]
```

```
[./CH2]
  type = CoreChannel
  eos = eos
  position = '0 0 0'
  orientation = '0 0 1'
  A = 1.549152542
  Dh = 0.01332254
  length = 3.6576
```

```

n_elems = 8 #16
f = 0.01
Hw = 5.33e4
Phf = 428.454929876871
Ts_init = 564.15
dim_hs = 1
n_heatstruct = 3
name_of_hs = 'FUEL GAP CLAD'
fuel_type = cylinder
width_of_hs = '0.0046955 0.0000955 0.000673'
elem_number_of_hs = '3 1 1'
material_hs = 'fuel-mat gap-mat clad-mat'
power_fraction = '3.69921461e-1 0 0'
[../]

```

```

[./CH3]
type = CoreChannel
eos = eos
position = '0 1.2 0'
orientation = '0 0 1'
A = 1.858983051
Dh = 0.01332254
length = 3.6576
n_elems = 8
f = 0.01
Hw = 5.33e4
# aw = 276.5737513
Phf = 514.145916018189
Ts_init = 564.15
dim_hs = 1
n_heatstruct = 3
name_of_hs = 'FUEL GAP CLAD'
fuel_type = cylinder
width_of_hs = '0.0046955 0.0000955 0.000673'
elem_number_of_hs = '3 1 1'
material_hs = 'fuel-mat gap-mat clad3-mat'
#peak_power = '3.401687e8 0. 0.'
power_fraction = '2.96405926e-1 0 0'
[../]

```

```

[./bypass_pipe]

```

```

type = Pipe
eos = eos
position = '0 1.5 0'
orientation = '0 0 1'
A = 1.589571014
Dh = 1.42264
length = 3.6576
n_elems = 5
f = 0.001
Hw = 0.0
[../]

[./LowerPlenum]
type = Branch
eos = eos
inputs = 'DownComer-A(out) DownComer-B(out)'
outputs = 'CH1(in) CH2(in) CH3(in) bypass_pipe(in)'
K = '0.2 0.2 0.2 0.2 0.4 40.0'
Area = 3.618573408
Initial_pressure = 151.7e5
[../]

[./UpperPlenum]
type = Branch
eos = eos
inputs = 'CH1(out) CH2(out) CH3(out) bypass_pipe(out)'
outputs = 'pipe1-HL-A(in) pipe1-HL-B(in)'
K = '0.5 0.5 0.5 80.0 0.5 0.5'
Area = 7.562307456
Initial_pressure = 151.7e5
[../]

#Loop A components
[./DownComer-A]
type = Pipe
eos = eos
position = '0 2.0 4.0'
orientation = '0 0 -1'
A = 3.6185734
Dh = 1.74724302
length = 4

```

```

n_elems = 3
f = 0.001
Hw = 0.
[../]

[./pipe1-HL-A]
type = Pipe
eos = eos
position = '0 0.5 4.0'
orientation = '0 0 1'
A = 7.562307456
Dh = 3.103003207
length = 4.
n_elems = 3
f = 0.001
Hw = 0.0
[../]

[./pipe2-HL-A]
type = Pipe
eos = eos
position = '0 0.5 8.0'
orientation = '0 1 0'
A = 2.624474
Dh = 1.828
length = 3.5
n_elems = 3
f = 0.001
Hw = 0.0
[../]

[./pipe1-CL-A]
type = Pipe
eos = eos
position = '0 3.0 4.0'
orientation = '0 -1 0'
A = 2.624474
Dh = 1.828
length = 1.
n_elems = 3
f = 0.001

```

```
Hw = 0.0
[../]

[./pipe2-CL-A]
type = Pipe
eos = eos
position = '0 4 4.0'
orientation = '0 -1 0'
A = 2.624474
Dh = 1.828
length = 0.8
n_elems = 3
f = 0.001
Hw = 0.0
[../]

[./pipe1-SC-A]
type = Pipe
eos = eos
position = '0 5.2 4.0'
orientation = '0 -1 0'
A = 2.624474
Dh = 1.828
length = 1.
n_elems = 3
f = 0.001
Hw = 0.0
[../]

[./pipe2-SC-A]
type = Pipe
eos = eos
position = '0 4.2 8.0'
orientation = '0 1 0'
A = 2.624474
Dh = 1.828
length = 1.
n_elems = 3
f = 0.001
Hw = 0.0
[../]
```

```

[./Branch1-A]
  type = Branch
  eos = eos
  inputs = 'pipe1-HL-A(out)'
  outputs = 'pipe2-HL-A(in) pipe-to-Pressurizer(in)'
  K = '0.5 0.7 80.'
  Area = 7.562307456
  Initial_pressure = 151.7e5
[../]

[./Branch2-A]
  type = Branch
  eos = eos
  inputs = 'pipe1-CL-A(out)'
  outputs = 'DownComer-A(in)'
  K = '0.5 0.7'
  Area = 3.6185734
  Initial_pressure = 151.7e5
[../]

[./Branch3-A]
  type = Branch
  eos = eos
  inputs = 'pipe2-HL-A(out)'
  outputs = 'HX-A(primary_in)'
  K = '0.5 0.7'
  Area = 2.624474
  Initial_pressure = 151.7e5
[../]

[./Pump-A]
  type = IdealPump
  eos = eos
  inputs = 'pipe2-CL-A(out)'
  outputs = 'pipe1-CL-A(in)'
  Area = 2.624474
  mass_flow_rate = 8801.1
  Initial_pressure = 151.7e5
[../]

```

```

[./HX-A]
type = HeatExchanger
eos = eos
eos_secondary = eos
position = '0 4. 8.'
orientation = '0 0 -1'
A = 5.
A_secondary = 5.
Dh = 0.01
Dh_secondary = 0.01
length = 4.
n_elems = 10
Hw = 1.e4
Hw_secondary = 1.e4
Phf = 2695.100000000000
Phf_secondary = 2695.100000000000
f = 0.01
f_secondary = 0.01
dim_wall = 1
Twall_init = 564.15
wall_thickness = 0.001
n_wall_elems = 2
material_wall = wall-mat
[../]

[./Branch4-A]
type = Branch
eos = eos
inputs = 'pipe1-SC-A(out)'
outputs = 'HX-A(secondary_in)'
K = '0.5 0.7'
Area = 2.624474e2
Initial_pressure = 151.7e5
[../]

[./Branch5-A]
type = Branch
eos = eos
inputs = 'HX-A(secondary_out)'
outputs = 'pipe2-SC-A(in)'
K = '0.5 0.7'

```

```

    Area = 2.624474e2
    Initial_pressure = 151.7e5
[../]

[./Branch6-A]
    type = Branch
    eos = eos
    inputs = 'HX-A(primary_out)\'
    outputs = 'pipe2-CL-A(in)\'
    K = '0.5 0.7\'
    Area = 2.624474e2
    Initial_pressure = 151.7e5
[../]

[./MassFlowRateIn-SC-A]
    type = TimeDependentJunction
    input = 'pipe1-SC-A(in)\'
    v_bc = 4.542
    T_bc = 537.15
    eos = eos
[../]

[./PressureOutlet-SC-A]
    type = TimeDependentVolume
    input = 'pipe2-SC-A(out)\'
    p_bc = '151.7e5\'
    T_bc = 564.15
    eos = eos
[../]

#Loop B components
[./DownComer-B]
    type = Pipe
    eos = eos
    position = '0 -2.0 4.0\'
    orientation = '0 0 -1\'
    A = 3.6185734
    Dh = 1.74724302
    length = 4
    n_elems = 3
    f = 0.001

```

```
Hw = 0.
[../]

[./pipe1-HL-B]
type = Pipe
eos = eos
position = '0 -0.5 4.0'
orientation = '0 0 1'
A = 7.562307456
Dh = 3.103003207
length = 4.
n_elems = 3
f = 0.001
Hw = 0.0
[../]

[./pipe2-HL-B]
type = Pipe
eos = eos
position = '0 -0.5 8.0'
orientation = '0 -1 0'
A = 2.624474
Dh = 1.828
length = 3.5
n_elems = 3
f = 0.001
Hw = 0.0
[../]

[./pipe1-CL-B]
type = Pipe
eos = eos
position = '0 -3.0 4.0'
orientation = '0 1 0'
A = 2.624474
Dh = 1.828
length = 1.
n_elems = 3
f = 0.001
Hw = 0.0
[../]
```

```
[./pipe2-CL-B]
type = Pipe
eos = eos
position = '0 -4.0 4.0'
orientation = '0 1 0'
A = 2.624474
Dh = 1.828
length = 0.8
n_elems = 3
f = 0.001
Hw = 0.0
[../]
```

```
[./pipe1-SC-B]
type = Pipe
eos = eos
position = '0 -5.2 4.0'
orientation = '0 1 0'
A = 2.624474
Dh = 1.828
length = 1.
n_elems = 3
f = 0.001
Hw = 0.0
[../]
```

```
[./pipe2-SC-B]
type = Pipe
eos = eos
position = '0 -4.2 8.0'
orientation = '0 -1 0'
A = 2.624474
Dh = 1.828
length = 1.
n_elems = 3
f = 0.001
Hw = 0.0
[../]
```

```
[./Branch1-B]
```

```

type = Branch
eos = eos
inputs = 'pipe1-HL-B(out)'
outputs = 'pipe2-HL-B(in)'
K = '0.5 0.7'
Area = 7.562307456
Initial_pressure = 151.7e5
[../]

[./Branch2-B]
type = Branch
eos = eos
inputs = 'pipe1-CL-B(out)'
outputs = 'DownComer-B(in)'
K = '0.5 0.7'
Area = 3.6185734
Initial_pressure = 151.7e5
[../]

[./Branch3-B]
type = Branch
eos = eos
inputs = 'pipe2-HL-B(out)'
outputs = 'HX-B(primary_in)'
K = '0.5 0.7'
Area = 2.624474
Initial_pressure = 151.7e5
[../]

[./Pump-B]
type = IdealPump
eos = eos
inputs = 'pipe2-CL-B(out)'
outputs = 'pipe1-CL-B(in)'
Area = 2.624474
mass_flow_rate = 8801.1
Initial_pressure = 151.7e5
[../]

[./HX-B]
type = HeatExchanger

```

```

eos = eos
eos_secondary = eos
position = '0 -4. 8.'
orientation = '0 0 -1'
A = 5.
A_secondary = 5.
Dh = 0.01
Dh_secondary = 0.01
length = 4.
n_elems = 10
Hw = 1.e4
Hw_secondary = 1.e4
Phf = 2695.1000000000000
Phf_secondary = 2695.1000000000000
f = 0.01
f_secondary = 0.01
dim_wall = 1
Twall_init = 564.15
wall_thickness = 0.001
material_wall = wall-mat
n_wall_elems = 2
disp_mode = -1.0
[../]

[./Branch4-B]
type = Branch
eos = eos
inputs = 'pipe1-SC-B(out)'
outputs = 'HX-B(secondary_in)'
K = '0.5 0.7'
Area = 2.624474e2
Initial_pressure = 151.7e5
[../]

[./Branch5-B]
type = Branch
eos = eos
inputs = 'HX-B(secondary_out)'
outputs = 'pipe2-SC-B(in)'
K = '0.5 0.7'
Area = 2.624474e2

```

```

    Initial_pressure = 151.7e5
[../]

[./Branch6-B]
type = Branch
eos = eos
inputs = 'HX-B(primary_out)\'
outputs = 'pipe2-CL-B(in)\'
K = '0.5 0.7\'
Area = 2.624474e2
Initial_pressure = 151.7e5
[../]

[./MassFlowRateIn-SC-B]
#type = TDM
type = TimeDependentJunction
input = 'pipe1-SC-B(in)\'
# massflowrate_bc = 8801.1
v_bc = 4.542
T_bc = 537.15
eos = eos
[../]

[./PressureOutlet-SC-B]
type = TimeDependentVolume
input = 'pipe2-SC-B(out)\'
p_bc = '151.7e5\'
T_bc = 564.15
eos = eos
[../]

# Pressurizer
[./pipe-to-Pressurizer]
type = Pipe
eos = eos
position = '0 0.5 8.0\'
orientation = '0 0 1\'
A = 2.624474
Dh = 1.828
length = 0.5
n_elems = 3

```

```

    f = 10.
    Hw = 0.0
[../]

[./Pressurizer]
    type = TimeDependentVolume
    input = 'pipe-to-Pressurizer(out)'
    p_bc = '151.7e5'
    T_bc = 564.15
    eos = eos
[../]
[]

[Preconditioning]
[./SMP_PJFNK]
    type = SMP
    full = true
    solve_type = 'PJFNK'
[../]
[]

[Executioner]
    type = Transient
    scheme = 'implicit-euler'

#use this parameter to have a better initial guess from previous two time steps, therefore
[./Predictor]
    type = SimplePredictor
    scale = 0.6
[../]

dt = 0.1
dtmin = 1e-7
nl_rel_tol = 1e-9
nl_abs_tol = 1e-8
nl_max_its = 30
l_tol = 1e-4
l_max_its = 300
start_time = 0.0
end_time = 50

```

```

num_steps = 250
petsc_options_iname = '-mat_fd_type -mat_mffd_type -ksp_gmres_restart -pc_type'
petsc_options_value = 'ds ds 300 lu'

[./Quadrature]
  type = TRAP
  order = FIRST
[../]
[]

[Outputs]
[./out]
  type = Exodus
  use_displaced = true
  output_initial = true
  sequence = false
  append_displaced = true
[../]

[./console]
  type = Console
  perf_log = true
[../]
[]

```

5.11.3 Description of the Input File

Most of the input parameters have been explained in the previous examples. Only a few need some explanations here.

```

[./Pump-A]
  type = IdealPump
  eos = eos
  inputs = 'pipe2-CL-A(out)'
  outputs = 'pipe1-CL-A(in)'
  Area = 2.624474
  mass_flow_rate = 8801.1
  Initial_pressure = 151.7e5
[../]

```

type	The ideal pump component is used here. Please note that Ideal Pump component is an interim model. It merely provides a prescribed mass flow rate to the pipes with this component. This component will eventually be replaced with a realistic pump model.
eos	Name of equation of state to be used by IdealPump.
inputs	The inlet of the ideal pump connects to the outlet of pipe2-CL-A.
outputs	The outlet of the ideal pump connects to the inlet of pipe1-CL-A.
Area	The reference flow area of the ideal pump.
mass_flow_rate	The prescribed mass flow rate generated by the ideal pump.
Initial_pressure	The prescribed initial pressure of the ideal pump.

```
[./Predictor]
  type = SimplePredictor
  scale = 0.6
[../]
```

[./Predictor]	Subblock for predictor. The predictor provides a better initial guess from the solution of previous two old time steps. It reduces the number of nonlinear iterations.
type	The SimplePredictor uses an algorithm that will predict the next solution based on previous solutions. The algorithm is: $sol^{n+1} = sol^n + \frac{sol^n - sol^{n-1}}{\delta t^n} * \delta t^{n+1} * scale$.
scale	The scale factor used in the SimplePredictor calculation.

5.12 Example 12: A Simplified Primary System Model of a Boiling Water Reactor

5.12.1 Problem Description

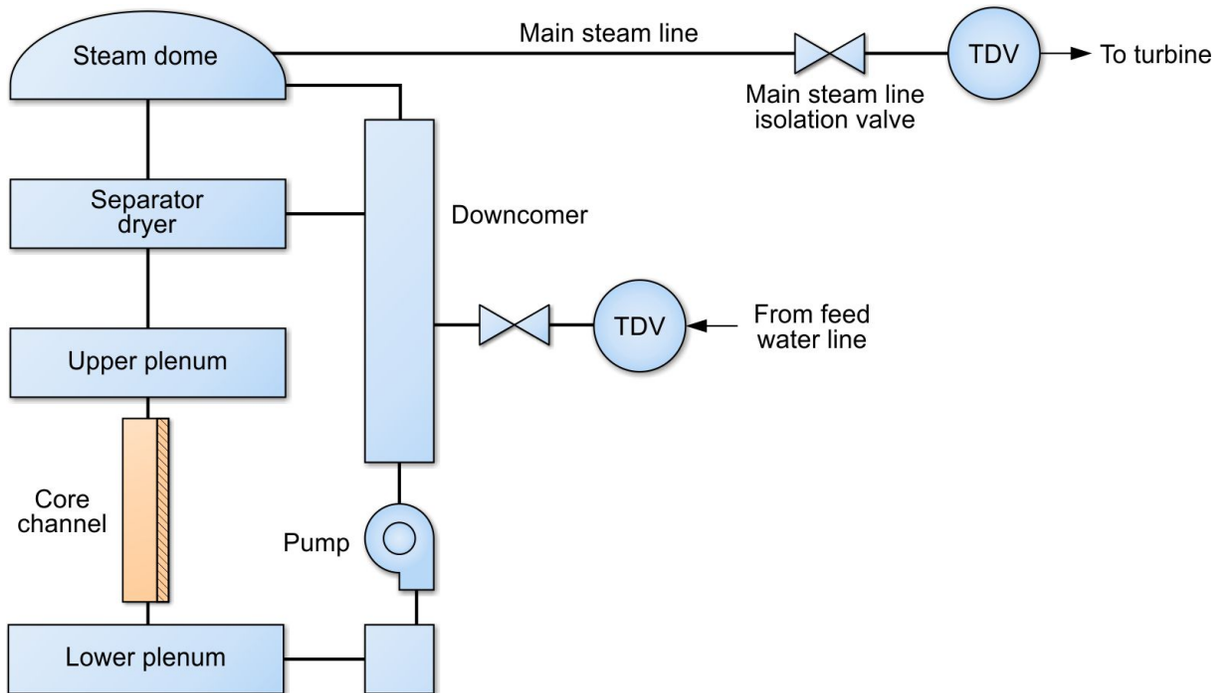


Figure 12. Diagram of a simplified primary system of a BWR model

This example is a simplified primary system of a boiling water reactor. The model consists of downcomer, lower plenum, reactor core, upper plenum, separator dryer, steam dome, main steam line, feedwater line and the primary pump model. One core channel is used to represent the entire reactor core. The lower plenum, upper plenum and steam dome are modeled with volume branch models. External to the reactor vessel, the main steam line is connected to the steam dome. A time dependent volume is attached to the main steam line to provide the necessary boundary conditions for the steam flow. A feedwater line is connected to the downcomer model. A time dependent volume is attached to the feedwater line to provide the necessary boundary conditions for the feedwater. Notably missing from this simplified BWR model are the jet pumps and the recirculation loops that

allow the operator to vary coolant flow rate through the core and change the reactor power. Instead, a pump model is used to represent the functions of the jet pumps and the recirculation loops.

5.12.2 Input File

```
[GlobalParams]
  model_type = 32
  global_init_P = 7.e6
  global_init_V = 3.
  global_init_T = 517.
  scaling_factor_var = '1e-3 1e-4 1e-8'
  temperature_sf = '1e-4'
  gravity = '0 0 -9.8'
  stabilization_type = 'LAPIDUS'
[]

[EoS]

  [./two_phase_eos]
    type = TwoPhaseStiffenedGasEOS
  [../]

  [./vapor_phase_eos]
    type = StiffenedGasEquationOfStateVapor
  [../]

  [./liquid_phase_eos]
    type = StiffenedGasEquationOfStateLiquid
  [../]

[]

[Materials]
  [./fuel-mat]
    type = SolidMaterialProperties
    k = 3.7
    Cp = 3.e2
    rho = 10.42e3
```

```

[../]
[./gap-mat]
  type = SolidMaterialProperties
  k = 0.7
  Cp = 5e3
  rho = 1.0
[../]
[./clad-mat]
  type = SolidMaterialProperties
  k = 16
  Cp = 356.
  rho = 6.551400e3
[../]
[]

[Components]
[./reactor]
  type = Reactor
  initial_power = 3293.0e6
[../]

[./ch1]
  type = CoreChannel
  eos = two_phase_eos
  position = '0 0.0 5.28'
  orientation = '0 0 1'
  A = 7.8
  Dh = 1.3597E-02
  length = 3.66
  n_elems = 100
  f = 0.2 #0.05
  Hw = 5.0e4
  Phf = 1836.8430600
  Ts_init = 517.
  fuel_type = cylinder
  dim_hs = 1
  n_heatstruct = 3
  name_of_hs = 'FUEL GAP CLAD'
  width_of_hs = '6.057900e-3 1.524000e-4 9.398000e-4'
  elem_number_of_hs = '5 1 2'
  material_hs = 'fuel-mat gap-mat clad-mat'

```

```

power_fraction = '1.0 0.0 0.0'
stabilization_type = 'NONE'
model_type = 32
[../]

[./pipe6]
type = Pipe
position = '0.0 0.0 10.82'
orientation = '0 0 1'
A = 3.93
Dh = 1.0
length = 2.72
n_elems = 40
f = 0.1
eos = two_phase_eos
model_type = 32
stabilization_type = 'NONE'
[../]

[./pipe7]
type = Pipe
position = '0.0 0.0 15.42'
orientation = '0 0 1'
A = 3.93
Dh = 1.0
length = 0.1
n_elems = 5
f = 0.1
eos = vapor_phase_eos
model_type = 3
[../]

[./pipe8]
type = Pipe
position = '0.0 2.0 14.48'
orientation = '0 1 0'
A = 3.93
Dh = 1.0
length = 0.5
n_elems = 5
f = 0.1

```

```

eos = liquid_phase_eos
model_type = 3
stabilization_type = 'SUPG'
[../]

[./pipe9]
# main steam line coming out of dome
type = Pipe
position = '0.0 3 18.92'
orientation = '0 1 0'
A = 1.32
Dh = 1.0
length = 1.0
n_elems = 5
f = 0.1
eos = vapor_phase_eos
model_type = 3
[../]

[./pipe10]
type = Pipe
#position = '0.0 5.0 10.51'
position = '0.0 2.75 2.10'
orientation = '0 0 -1'
A = 8.55
Dh = 1.0
#length = 6.42
length = 0.5
n_elems = 5
f = 0.1
eos = liquid_phase_eos
model_type = 3
stabilization_type = 'SUPG'
[../]

[./pipe11]
type = Pipe
position = '0.0 2.75 1.60'
orientation = '0 0 -1'
A = 8.55
Dh = 1.0

```

```

length = 0.5
n_elems = 5
f = 0.1
eos = liquid_phase_eos
model_type = 3
stabilization_type = 'SUPG'
[../]

[./pipe14]
# main steam line to MIV
type = Pipe
position = '0.0 4.0 18.92'
orientation = '0 1 0'
A = 1.32
Dh = 1.0
length = 1.0
n_elems = 5
f = 0.0
eos = vapor_phase_eos
model_type = 3
[../]

[./MainIsolationValve]
type = VolumeBranch
eos = vapor_phase_eos
center = '0.0 5.0 18.92'
inputs = 'pipe14(out)'
outputs = 'pipe_steam_turbine(in)'
K = '0.0 0.0'
volume = 1.32
Area = 1.32
initial_T = 517.0
scale_factors = '1.0E-4 1.0E-8 1.0' # rho, rhoE
[../]

[./pipe_steam_turbine]
# main steam line to TDV
type = Pipe
position = '0.0 5 18.92'
orientation = '0 1 0'
A = 1.32

```

```

Dh = 1.0
length = 1.0
n_elems = 5
f = 0.0
eos = vapor_phase_eos
model_type = 3
[../]

[./Pump]
type = Pump
eos = liquid_phase_eos
inputs = 'pipe10(out)'
outputs = 'pipe11(in)'
Area = 3.0
Initial_pressure = 7.3e6
#mass_flow_rate = 12915.0
Head = 40.0
K_reverse = '10. 10.'
[../]

[./SeparatorDryer]
type = SeparatorDryer
eos = two_phase_eos
center = '0.0 0.0 14.48'
inputs = 'pipe6(out)'
outputs = 'pipe7(in) pipe8(in)'
K = '1.0 1.0 5.0'
volume = 19.30
Area = 10.27
initial_T = 517.0
scale_factors = '1.0E-3 1.0E-8 1.0E-0'
[../]

[./lowerplenum]
type = VolumeBranch
eos = two_phase_eos
center = '0.0 0.0 2.64'
inputs = 'pipe11(out)'
outputs = 'ch1(in)'
K = '1.0 20.0'
volume = 61.48

```

```

Area = 11.64
initial_T = 517.0
scale_factors = '1.0E-3 1.0E-8 1.0E-0'
[../]

[./upperplenum]
type = VolumeBranch
eos = two_phase_eos
center = '0.0 0.0 9.88'
inputs = 'ch1(out)'
outputs = 'pipe6(in)'
K = '3.0 1.0'
volume = 26.99
Area = 14.36
initial_T = 517.0
scale_factors = '1.0E-3 1.0E-8 1.0E-0'
[../]

[./Dome]
type = VolumeBranch
eos = vapor_phase_eos
center = '0.0 0.0 18.92'
inputs = 'pipe7(out)'
outputs = 'pipe9(in)'
K = '1.0 1.0'
volume = 178.19
Area = 26.19
scale_factors = '1.0E-3 1.0E-8 1.0E-0'
[../]

[./DownComer]
type = DownComer
eos = liquid_phase_eos
center = '0.0 2.75 9.81'
inputs = 'pipe8(out) pipe_feedwater3(out)'
outputs = 'pipe10(in)'
K = '1.0 10.0 1.0'
volume = 201.3
Area = 15
initial_level = 13.42
initial_T = 517.0

```

```

dome_component = 'Dome'
dome_eos = vapor_phase_eos
scale_factors = '1.0E-4 1.0E-9 1.0E-0'
[../]

[./SteamLineBranch]
type = VolumeBranch
eos = vapor_phase_eos
center = '0.0 4.0 18.92'
inputs = 'pipe9(out)'
outputs = 'pipe14(in)'
K = '0.0 0.0'
volume = 2.64
Area = 1.32
initial_T = 517.0
scale_factors = '1.0E-4 1.0E-8 1.0'
[../]

[./pipe_feedwater1]
#feedwater line from TDV
type = Pipe
position = '0.0 6.0 12.52' #'0.0 7.0 12.52'
orientation = '0 -1 0'
A = 1.32
Dh = 1.0
length = 1.0
n_elems = 5
f = 0.01 #1
eos = liquid_phase_eos
model_type = 3
stabilization_type = 'SUPG'
[../]

[./FeedWaterValve]
type = Valve
eos = liquid_phase_eos
center = '0.0 5.0 12.52'
inputs = 'pipe_feedwater1(out)'
outputs = 'pipe_feedwater2(in)'
K = '0.0 0.0'
volume = 1.32

```

```

Area = 1.32
initial_T = 517.0
initial_status = open
trigger_time = 1.0E5
response_time = 1.1E5
scale_factors = '1.0E-4 1.0E-8'
[../]

[./pipe_feedwater2]
#feedwater line from feed water valve
type = Pipe
position = '0.0 5.0 12.52' #'0.0 7.0 12.52'
orientation = '0 -1 0'
A = 1.32
Dh = 1.0
length = 1.0
n_elems = 5
f = 0.01
eos = liquid_phase_eos
model_type = 3
stabilization_type = 'SUPG'
[../]

[./branch_feedwater_line]
type = VolumeBranch
eos = liquid_phase_eos
center = '0.0 4.0 12.52'
inputs = 'pipe_feedwater2(out)'
outputs = 'pipe_feedwater3(in)'
K = '0 0'
volume = 1.32
Area = 1.32
initial_T = 517.0
scale_factors = '1.0E-4 1.0E-8 1.0'
[../]

[./pipe_feedwater3]
#feedwater line to downcomer
type = Pipe
position = '0.0 4.0 12.52' #'0.0 6.0 12.52'
orientation = '0 -1 0'

```

```

A = 1.32
Dh = 1.0
length = 1.0
n_elems = 5
f = 0.01
eos = liquid_phase_eos
model_type = 3
stabilization_type = 'SUPG'
[../]

[./inlet]
type = TimeDependentVolume
input = 'pipe_feedwater1(in)'
p_bc = 7.1e6
T_bc = 508.
eos = liquid_phase_eos
[../]

[./outlet1]
type = TimeDependentVolume
input = 'pipe_steam_turbine(out)'
p_bc = 7.0e6
T_bc = 517
eos = vapor_phase_eos
weak_bc = false
[../]
[]

[Preconditioning]
[./SMP_PJFNK]
type = SMP
full = true
solve_type = 'PJFNK'
[../]

[]

[Executioner]
type = Transient
dtmin = 1.e-7
[./TimeStepper]

```

```

    type = SolutionTimeAdaptiveDT
    dt = 0.1
[../]
petsc_options_iname = '-ksp_gmres_restart -pc_type'
petsc_options_value = '30 lu'

nl_rel_tol = 1e-8
nl_abs_tol = 1e-6
nl_max_its = 15

l_tol = 1e-5
l_max_its = 100

start_time = 0.0
end_time = 400
num_steps = 500000

[./Quadrature]
    type = TRAP
    order = FIRST
[../]
[]

[Outputs]
[./out]
    type = Exodus
    use_displaced = true
    output_initial = true
    sequence = false
    append_displaced = true
    interval = 50
[../]

[./console]
    type = Console
    perf_log = true
[../]

[./checkpoint]
    type = Checkpoint
    num_files = 1

```

```
[../]  
[]
```

5.12.3 Description of the Input File

```
[./SeparatorDryer]  
  type = SeparatorDryer  
  eos = two_phase_eos  
  center = '0.0 0.0 14.48'  
  inputs = 'pipe6(out)'  
  outputs = 'pipe7(in) pipe8(in)'  
  K = '1.0 1.0 5.0'  
  volume = 19.30  
  Area = 10.27  
  initial_T = 517.0  
  scale_factors = '1.0E-3 1.0E-8 1.0E-0'  
[../]
```

[./SeparatorDryer]	Subblock for separator dryer component.
type	Component type is SeparatorDryer.
eos	Name of equation of state to be used by the separator dryer component.
center	The (x, y, z) coordinate of the center of the separator dryer component.
inputs	The inlet of the separator dryer component connects with the outlet of pipe6.
outputs	The outlets of the separator dryer component connects with the inlet of pipe7 and pipe8.
K	The form loss coefficients at the junctions with the pipes.
volume	The volume of the separator dryer component.
Area	The flow area of separator dryer component.
initial_T	The initial temperature of the separator dryer component.

scale_factors The scaling factors for the primary variables of the separator dryer component.

```
[./DownComer]
  type = DownComer
  eos = liquid_phase_eos
  center = '0.0 2.75 9.81'
  inputs = 'pipe8(out) pipe_feedwater3(out) '
  outputs = 'pipe10(in) '
  K = '1.0 10.0 1.0'
  volume = 201.3
  Area = 15
  initial_level = 13.42
  initial_T = 517.0
  dome_component = 'Dome'
  dome_eos = vapor_phase_eos
  scale_factors = '1.0E-4 1.0E-9 1.0E-0'
[../]
```

[./DownComer] Subblock for the down comer component.

type The component type is DownComer.

eos Name of the equation of state to be used by the down comer component.

center The (x, y, z) coordinate of the center of the down comer component.

inputs The inlet of down comer connects with the outlet of pipe8 and pipe_feedwater3.

outputs The outlet of down comer connects with the inlet of pipe10.

K The form loss coefficients at the junctions with the pipes.

volume The volume of the down comer component.

Area The reference flow area of the downcomer component.

initial_level The initial liquid level of the down comer component.

initial_T The initial temperature of the down comer component.

dome_component Name of the steam dome component to provide the dome pressure to down comer.

dome_eos Name of the equation of state to be used by the steam dome component.

scale_factors The saling factors for the primary variables of the down comer component.

```
[./FeedWaterValve]
type = Valve
eos = liquid_phase_eos
center = '0.0 5.0 12.52'
inputs = 'pipe_feedwater1(out)'
```

```
outputs = 'pipe_feedwater2(in)'
```

```
K = '0.0 0.0'
```

```
volume = 1.32
```

```
Area = 1.32
```

```
initial_T = 517.0
```

```
initial_status = open
```

```
trigger_time = 1.0E5
```

```
response_time = 1.1E5
```

```
scale_factors = '1.0E-4 1.0E-8'
```

```
[../]
```

[./FeedWater] Subblock for the valve component.

type The component type is Valve.

eos Name od the equation of state to be used by the valve component.

center The (x, y, z) coordinate of the center of the valve component.

inputs The inlet of valve connects with the outlet of pipe_feedwater1.

outputs The outlet of valve connects with the inlet of pipe_feedwater2.

K The form loss coefficients at the junctions with the pipes.

volume	The volume of the valve component.
Area	The reference flow area of the valve component.
initial_T	The initial temperature of the valve component.
initial_status	The initial status of the valve is at the open position.
trigger_time	The time at which the valve open/close action is triggered.
response_time	The time at which the valve open/close action is complete.
scale_factors	The scaling factors for the primary variables of the valve component.

```
[./checkpoint]
  type = Checkpoint
  num_files = 1
[../]
```

[./checkpoint]	Subblock for check point. This is used to create restart files for subsequent runs.
num_files	Number of restart files to save.

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