flowPsi Module Guide

August 31, 2017

Contents

1	Introduction	1		
	1.1 Loci FVM Module Facilities	1		
	1.2 flowPsi facilities and variables	3		
2	Custom Output	6		
3	3 Custom Boundary Condition Input			
4	4 Custom Initial Conditions 1			
5	5 Surface Integration			
6	5 Volume Integration 1			
7	7 Adding a Volume Source 1			
8	Scalar Transport	16		
	8.1 Tracer Variable Example	17		
	8.2 The Menter one equation turbulence model	17		
	8.2.1 Implementation	18		

1 Introduction

This module guide provides a set of tutorials for creating add on modules for the <code>flowPsi</code> solver. It is assumed that the reader has some basic understanding of the Loci programming model. To gain that understanding please work through the tutorial provided with the Loci framework. This document and examples will extend that tutorial to describe how to create common types of extensions to the <code>flowPsi</code> solver. In general modules are loaded into <code>flowPsi</code> by first making sure that the directory that contains the module is either in the <code>LD_LIBRARY_PATH</code> or the <code>LOCI_MODULE_PATH</code> environment variable. Then the modules can be loaded into your <code>flowPsi</code> case by adding a <code>loadModule:</code> followed by the module name at the top of the <code>vars</code> file. The <code>Makefile</code> provided in these example directories are easy to use and can be copied for your module development. The <code>Makefile</code> is designed such that all files that end in with the <code>.loci</code> postfix will be compiled into a module. The file only needs to be edited to provide the module name and the <code>FLOWPSI_BASE</code> directory that points to where the <code>flowPsi</code> solver is installed.

1.1 Loci FVM Module Facilities

Before we begin discussing building modules for flowPsi lets review some of the basic facilities that Loci provides for finite volume solvers that flowPsi utilizes. First, flowPsi uses the face based data-structure that includes the face2node map that provides an ordered listing of the nodes that form each face in the mesh as well as the cl and cr maps that provide the maps from the face to the left and right cells. Faces on the boundary always have normals pointing out of the domain and therefore the cl map points to the cell next to the boundary while the cr cell may point to a ghost cell (in flowPsi, ghost cells are only used to implement periodic boundary conditions). The ci and ref maps are only defined for boundary faces where ci refers to the adjacent cell inside the mesh and ref refers to the reference surface that forms the facets of that boundary. The pmap map associates the periodic boundary face with its periodic pair. The cell to face maps upper, lower, and boundary map is used to form matrices that are based on the face connectivity where upper refers to the faces that form the upper and lower triangular matrix entries and the boundary map identifies boundary faces of the cell.

In addition to these data structure elements, the Loci FVM module provides some generic rules for computing grid metrics such as face areas and cell volumes. Additionally the module provides some basic facilities for computing gradients and interpolating solutions from cells to nodes for plot output. These facilities generally will need the cell center values and the boundary face values. The boundary face values will have the naming convention of having a $_{\perp}$ f postfix annotation denoting that it is a face value. A table that summarizes that facilities provided by the FVM module are provided in table 1.

Table 1: Table of Loci FVM provided utilities

Table 1: Table of Loci FVM provided utilities				
	Supplied by Loci FVM facilities			
Variable	Type	Description		
Mesh Data Structur	Mesh Data Structures			
ci	map	cell next to boundary face		
cl	map	cell on left side of face		
cr	map	cell on right side of face		
ref	map	map from boundary faces to surface		
face2node	multiMap	map from face to nodes		
pmap	map	periodic matching face		
upper	multiMap	map from cell to upper faces		
lower	multiMap	map from cell to lower faces		
boundary_map	multiMap	map from cell to boundary faces		
Mesh Metrics				
pos	vector	node positions		
cellcenter	vector	cell centroid location		
facecenter	vector	face centroid location		
area	composite	face area and normal		
vol	scalar	cell volume		
gridvol	parameter	total grid volume		
Parametric Rules				
grads(X)	vector	gradient of scalar variable X		
gradv3d(X)	tensor	gradient of vector variable X		
gradv(X)	generic vector	gradient of mixture vector X		
<pre>grads_f(X)</pre>	vector	face gradient of scalar X		
gradv3d_f(X)	tensor	face gradient of vector X		
gradv_f(X)	generic vector	gradient of mixture vector X		
cell2node(X)	scalar	cell to node interpolation for scalar X		
cell2node_v3d(X)	vector	cell to node interpolation for vector \mathbf{X}		
cell2nodeMax(X)	scalar	nodal maximum of X for connected cells		
cell2nodeMin(X)	scalar	nodal minimum of X for connected cells		
L1Norm(X)	parameter	volume integrated L_1 norm of cell variable X		
L2Norm(X)	parameter	volume integrated L_2 norm of cell variable X		
LinfNorm(X)	parameter	volume integrated L_{∞} norm of cell variable X		
fileNumber(X)	integer	file numbering for variable X		
		•		

Table 2: Table of flowPsi Fluid Properties

flowPsi fluid properties variables			
Variable Type		Description	
rho	scalar	fluid density	
rho_f	scalar	fluid density at face	
temperature	scalar	fluid temperature	
temperature_f	scalar	fluid temperature at face	
gagePressure	scalar	fluid pressure reference to Pambient	
gagePressure_f	scalar	fluid gage pressure at face	
Pambient	parameter	reference pressure	
pressure	scalar	fluid pressure	
u	vector	fluid velocity in cell	
u_f	vector	fluid velocity at face	
divu	scalar	divergence of u	
vort	vector	fluid vorticity	
vortMag	scalar	fluid vorticity magnitude	
strainRate	scalar	strain rate magnitude, S	
tau	symmetric tensor	fluid stress	
mu	scalar	fluid laminar viscosity	
mu_f	scalar	fluid laminar viscosity at face	
kconduct	scalar	fluid laminar conductivity	
$kconduct_f$	scalar	fluid laminar conductivity at face	
us_n	scalar	face grid velocity dot normal	
us	vector	face grid velocity	
Ср	parameter	fluid C_p	
gamma	parameter	fluid is entropic index, γ	
Rtilde	parameter	fluid gas constant	
tmu	scalar	turbulent viscosity	
soundSpeed	scalar	local speed of sound	

1.2 flowPsi facilities and variables

When flowPsi runs the only the conservative variables (and gage pressure) are carried from time-step to time-step. All other variables that are derived from these conservative variables are only computed during the Newton iteration. Thus most of the computations in flowPsi happen as part of the Newton iteration step. This section outlines some of the facilities that are available during that step. Table 2 gives that generic fluid properties that are available. Solver specific variables that can be replaced through priority rules or used in control are shown in table 3 while the parametric utilities are described in table 4. These tables are provided to give a brief survey of the kinds of data that is available for interfacing to flowPsi modules.

Table 3: Table of flowPsi solver specific variables

flowPsi solver specific variables			
Variable Type		Description	
pORef	scalar	total pressure BC input	
TORef	scalar	total temperature BC input	
massFluxRef	scalar	mass flux BC input	
temperatureRef	scalar	temperature BC input	
gagePressureRef	scalar	gagePressure BC input	
uRef	vector	fluid velocity BC input	
Twall	scalar	viscous wall BC temperature	
qwall	scalar	viscous wall BC heat flux	
wallVelocity	vector	viscous wall tangential velocity	
src	general vector	source array for mass, momentum, and energy	
iflux	Array(5)	inviscid flux	
vflux	Array(4)	viscous flux	
scalar_mdot	scalar	mass flux through face	
srcJ	matrix block	diagonal jacobian term of src	
fjp	matrix block	jacobian of flux from left	
fjm	matrix block	jacobian of flux from right	
min_cell2noslip	map	Map from cell to closest viscous wall	
dist_noslip	scalar	distance from cell to nonslip wall	
dt	parameter	timestep value in seconds	
stime	parameter	simulation time in seconds	
ncycle	parameter	simulation step	
newtonFinished	conditional	Newton iteration complete	
do_restart	conditional	output restart files	
do_output	conditional	text output	
do_plot	conditional	plot file output	
do_boundary_plot	conditional	boundary plot file output	

Table 4: Table of $\mathit{flowPsi}$ parametric variables

flowPsi parametric variables			
Variable	Type	Description	
scalarTransport(X,C)	generic	build transport of scalar X, enabled by C	
<pre>scalarTransportP(X,C)</pre>	generic	like above for positive scalar X	
scalarMean(X)	scalar	time averaged cell value X	
$scalarMean_f(X)$	scalar	time averaged boundary face value X_f	
scalarVariance(X)	scalar	time variance of variable X	
vect3dMean(X)	vector	time averaged vector mean	
<pre>vect3dMean_f(X)</pre>	vector	time averaged boundary vector mean	
<pre>vect3dVariance(X)</pre>	vector	time variance of vector X	
<pre>vect3dVariance_f(X)</pre>	vector	time variance of boundary vector	
<pre>vect3dCoVariance(X)</pre>	vector	co-variance of vector X	
vect3dCoVariance_f(X)	vector	co-variance of boundary vector x	
<pre>scalarFavreMean(X)</pre>	scalar	Favre average of variable X	
scalarFavreMean_f(X)	scalar	Favre average of boundary variable	
scalarFavreVar(X)	scalar	Favre weighted variance	
<pre>vect3dFavreMean(X)</pre>	vector	vector Favre weighted mean	
<pre>vect3dFavreMean_f(X)</pre>	vector	vector Favre weighted boundary mean	
<pre>vect3dFavreVar(X)</pre>	vector	vector Favre weighted variance	
<pre>vect3dFavreVar_f(X)</pre>	vector	vector Favre weighted variance boundary	
<pre>vect3dFavreCoVar(X)</pre>	vector	vector Favre weighted co-variance	
vect3dFavreCoVar_f(X)	vector	vector Favre weighted co-variance boundary	

2 Custom Output

One of the more common needs for add-on modules is the development of derived variable computations. While it is possible to compute derived variables directly in post-processing software, these computations can sometimes be misleading due to the fact that they are performed on interpolated data which may introduce errors in the derived computations. So it is sometimes useful to perform computations directly on the cell values before interpolation in order to get more satisfactory results. For this example we will consider the output of a variable that can be used for numerical schlieren imaging. This is done by computing the logarithm of the density gradient. So in this example we will show how one goes about computing and outputting such a variable in a module.

The first step to this is to copy the Makefile provided in the examples which will automatically compile all .loci files in the directory. The only change that is needed is to set the MODULE_NAME variable in the Makefile to the name that you wish to use in the loadModule line of the vars file. In this case we will call the module schlieren, or

```
# What is the name of the module given to loadModule
MODULE_NAME = schlieren
```

At this point we can begin editing our Loci program which we will conveniently name schlieren.loci. At the beginning of the file we need to include some files that will be needed to access the facilities we will be using. This is accomplished with the following includes:

```
// Include Loci system
#include <Loci.h>
// defines types used by flowPsi
#include "flowTypes.h"
// imports flowPsi IO functionality
#include "flowPsiIO.h"
// defines Loci types for flowPsi variables
$include "flowPsi.lh"
```

It is convenient but not necessary to add the code in the C++ namespace called flowPsi. Alternatively, we can add the code in our own namespace but add code to tell C++ to use variables from the flowPsi namespace such as:

```
namespace schlieren {
  using namespace flowPsi ;
```

We now need to compute the variable that we are going to plot. Since flowPsi computes unknowns at cell centers this will be where we perform the computations. However, to improve nodal interpolation accuracy we also need to compute the target variables for all boundary faces as well. Since these variables are not part of the flowPsi infrastructure we will need to define their types first:

```
// define types for cell and boundary face values
$type loggradrho store<double> ;
$type loggradrho_f store<double> ;
```

Then write rules to define how to compute these variables as well. These computations will use the scalar gradient parametric rules provided by Loci's FVM module. The rules will be defined as:

```
// To interpolate to the nodal values we need both cell values and boundary
// face values. Thus we have two rules to compute the value of interest
$rule pointwise(loggradrho<-grads(rho)) {
    $loggradrho = log10(max(norm($grads(rho)),1e-4)) ;
}
$rule pointwise(loggradrho_f<-grads_f(rho)) {
    $loggradrho_f = log10(max(norm($grads_f(rho)),1e-4)) ;
}</pre>
```

At this point we have defined the variable that we wish to compute but as they are defined now they will not be computed because flowPsi will never request these variables. Instead we will want to tell flowPsi that these variables are ones that we need plot files. For this, flowPsi provides a macro called OUTPUT_SCALAR that interfaces with the flowPsi plotting facilities. It will allow this variable to be written out when requested by placing the given variable name in the plot_output variable in the vars file. This is accomplished with the line:

```
OUTPUT_SCALAR("cell2node(loggradrho)", schlieren) ;
```

This line tells the code to write out the variable cell2node(loggradrho) as a variable named schlieren. Note that the variable that we wrote out is not the variable that we computed. This is because volume data is plotted from nodal values and since flowPsi computes its degrees of freedom at the cells, an interpolation function is needed to compute the nodal values. The cell2node parametric rules provided by the Loci FVM module performs this function. This parametric rule will access the variables that we defined earlier. Also this macro will enable requesting this variable on cutting plane and isosurface output from flowPsi as well.

In addition to volume data, we can also output boundary facet data. This is provided with the macro OUTPUT_BNDRY_SCALAR. So if we wish to output the schlieren variable for the boundary facets we can add the following line to or loci file:

```
OUTPUT_BNDRY_SCALAR("loggradrho_f",schlierenb,"ci") ;
```

The first argument of this macro is the variable that we wish to plot, while the second is the name of the variable (hear with a b appended to differentiate boundary facet data from the nodal data). Finally, the last argument defines the set of boundaries that we wish to write this data. Since all boundary faces have the attribute ci this will write out for all boundary faces. Also not, for outputting 3D vector data replace the SCALAR with VECTOR in the above macros.

3 Custom Boundary Condition Input

Most of the boundary conditions provided by flowPsi can be customized. Typically the inputs for each boundary face are assigned to a Ref variables that can be overridden by modules to change the basic behavior. For this example we will demonstrate this in a simple boundary condition where the pressure is perturbed by a sinusoidal signal. In this particular case we will override the definition of the gagePressureRef variable so that instead of being the constant defined in the boundary condition, it will instead be a time varying value. Note, it is possible to also change the boundary condition to be reactive to the solution itself. However, in these cases the stability of the resulting boundary condition cannot be guaranteed.

To add this capability to an inflow boundary condition we want to be able to add a specification to each boundary where we want to add the perturbation that will give the amplitude and frequency of the disturbance. By default the flowPsi boundary condition checking code will not allow these parameters to be added without generating error messages. Therefore it is necessary to register these new boundary condition arguments with flowPsi. This checker for the boundary conditions are defined in the read_grid.h header file which will need to be included. The checker for these two new variables is added through the following code:

```
// Here we create a checking class that inherits from BC_Check that is defined
// in readGrid.h
class pwave_check : public BC_Check {
  string error_message ;
public:
  // List which boundary conditions this checker is for
  std::string boundaryConditions() { return "inflow,supersonicInflow,farfield" ; }
  // List which variables this checker will verify
  std::string variablesChecked() { return "amplitude, frequency" ; }
  // Check for the existence of amplitude and frequency
  // in our implementation these are optional, but if amplitude is
  // given then frequency must also be given
  bool checkOptions(const options_list& bc_options) {
    error_message = "" ;
    bool check = true ;
    if(bc_options.optionExists("amplitude")) {
      if(!check_scalar_units(bc_options, "amplitude", "Pa")) {
        error_message += "Pressure units needed for 'amplitude' " ;
        check = false ;
      if(!bc_options.optionExists("frequency")) {
        error_message += "'amplitude' must also specify 'frequency' " ;
        check = false ;
      }
    return check ;
  // Print out error messsage. Used if checker fails to generate error
  // message to the user. Usually the error message is remembered from
  // when the checker was run.
```

```
std::ostream &ErrorMessage(std::ostream &s) {
   }
};

// The register_BC template registers this object in the boundary checker
// database. The names are not important only that we create the object
// so that it is registered.
register_BCpwave_check;
```

Next we will need to extract the values for amplitude and pressure from the BC_options variable that is defined for each boundary surface. For this we will use the constraint that is created by the FVM grid reader that indicates what parameter are defined in the BC_options variable. This will ensure that the variables will only be extracted for the boundary conditions where these arguments are specified. The rules that extract these variables are as follows:

```
$type amplitude_BC store<real> ;
$rule pointwise(amplitude_BC<-BC_options),constraint(amplitude_BCoption) {
   $BC_options.getOptionUnits("amplitude","Pa",$amplitude_BC) ;
}
$type frequency_BC store<real> ;
$rule pointwise(frequency_BC<-BC_options),constraint(frequency_BCoption) {
   $BC_options.getOption("frequency",$frequency_BC) ;
}</pre>
```

We can now compute the time varying prescribed pressure and override the gagePressureRef variable that will be used by the inflow boundary condition.

```
$type pgDelta_BC store<real> ;

$rule pointwise(pgDelta_BC<-stime,amplitude_BC,frequency_BC) {
    $pgDelta_BC = $amplitude_BC*sin($stime*$frequency_BC*2.*M_PI) ;
}

// Override the reference condtions for the faces that adds in the
// perturbation. This will be used instead of the default settings
$rule pointwise(distrbance::gagePressureRef<-
ref->(gagePressureRef_BC,pgDelta_BC)) {
    $gagePressureRef = $ref->$gagePressureRef_BC + $ref->$pgDelta_BC ;
}
```

Other variables can also be overridden using a similar process. For a complete list of the Ref variables that can be overridden see table 3.

4 Custom Initial Conditions

```
// For the initial conditons we define temperature_ic, gagePressure_ic, and
// initial velocity (u_ic)
$rule pointwise(temperature_ic,gagePressure_ic,u_ic<-</pre>
                Rtilde,cellcenter,Pambient),constraint(geom_cells) {
  const double x = $cellcenter.x ;
  const double y = $cellcenter.y ;
  double coef = 3.5;
  double u = -1.*coef*cos(x)*sin(y);
  double v = coef*sin(x)*cos(y);
  double Pref = $Pambient ;
  if(Pref < 10)
   Pref = 1e5;
  double rho = 1.0;
  double p0 = Pref-$Pambient ;
  double pg = p0-rho*0.25*coef*coef*(cos(2.*x)+cos(2.*y));
  double P = pg+$Pambient ;
  double T = P/($Rtilde*rho) ;
  $gagePressure_ic = pg ;
  $temperature_ic = T ;
  u_i = vect3d(u,v,0.0);;
}
```

5 Surface Integration

In this example we consider how to add a module that can integrate a function over the boundary surfaces of the mesh. In this example we will compute the area averaged temperature over all of the boundary surfaces of the mesh. In general, Loci is quite flexible and will allow a wide range of possible strategies for this. In effect, an integration is simply a reduction with the summation operator over some set of entities that defines the surface of interest. How does one identify the set of interest? Actually one could identify the set of interest in many different ways. First, one could use the assigned boundary condition type. For example, one could integrate over the set defined by the constraint viscousWall_BC which would include all boundaries that were assigned the viscousWall boundary condition. Alternatively, the user could define a new boundary condition argument that would allow a user to mark that boundary with a special integrate flag (say integrate) and then use a constraint of ref->integrate_BCoption to determine the set. In this example we use the parametric facility to integrate over each boundary surface separately. To do this we use the fact that for each boundary a parametric variable will be created in by the grid reader of the form boundaryName(X) where X is replaced by each boundary surface name. This is a string parameter which is defined over the set of faces that form that boundary surface. For our mean temperature we will use an area weighted average, so at first we will compute the total area for each boundary with the rule:

```
// This sets up the initial value of the boundary area before summing over
// faces. This needs to be set to zero as this is the indentity of the
// summation operator
$rule unit(boundaryArea_X<-boundaryName(X)),parametric(boundaryName(X)) {
   $boundaryArea_X = 0;
}

// Add the areas from each face. In some cases, for example when
// running in axisymmetric mode the area might be zero, so the max
// function is there to keep that case from generating a divide by zero
// condition.
$rule apply(boundaryArea_X<-area,boundaryName(X))[Loci::Summation],
   parametric(boundaryName(X)) {
    // note, the area is the sada component of the area input, the n component
    // is the normal vector
   join($boundaryArea_X,max($area.sada,1e-13));
}</pre>
```

In this set of parametric rules the boundaryArea_X variable will be created for each boundary condition with the X replaced by each boundary name. Now we can compute the weighted temperature with the following rules:

```
// Now we will compute the area weighted average temperature
$type boundaryMeanTemp_X param<real> ;

// set the initial value for the mean temp
$rule unit(boundaryMeanTemp_X<-boundaryName(X)),parametric(boundaryName(X)) {
   $boundaryMeanTemp_X = 0 ;
}</pre>
```

```
// Now compute the weighted sum of the of the temperature. The temperature
// at the boundary face is given by temperature_f
$rule apply(boundaryMeanTemp_X<-area,temperature_f,boundaryArea_X,boundaryName(X))[Loci::Summation],
    parametric(boundaryName(X)) {
    double weight = max($area.sada,1e-13)/$boundaryArea_X;
    join($boundaryMeanTemp_X,weight*$temperature_f);
}</pre>
```

Now we can output the integrated value. Note, that although this appears to be one rule, one will be instantiated for each boundary surface. In the end this will generate output for each boundary.

```
// Now output to stdout the average temperature. Do this output only
// when the code is outputting boundary plot files.
// Note, since this is only output once, we use the prelude and
// because we are in the prelude we must use the dereference operatpr
// '*' to access the computed parameters. The $[Once] tells Loci that
// you only want to output this once even when running in parallel where
// there would be multiple instances of execution
$rule pointwise(OUTPUT<-boundaryMeanTemp_X,boundaryName(X)),</pre>
  conditional(do_boundary_plot),
  parametric(boundaryName(X)), prelude {
  $[Once] {
    cout << "Boundary " << *\$boundaryName(X) << " has mean temperature of "
         << *$boundaryMeanTemp_X << " Kelvin" << endl ;</pre>
  }
} ;
//^ This semicolon is needed because we are not following the prelude
// section with a compute section.
```

6 Volume Integration

In this example we will show how to perform a volume integration over each named component of the mesh. This will work similarly to the surface integration except that we will use the volumeTag(X) parametric variable instead. For this example we will compute the integrated fluid kinetic energy by simply adding up the kinetic energy integrated over each cell. The basic approach is as follows:

```
// KEComponent_X is the integrated kinetic energy for component X. This
// rule is parametric so one of these variables will be created for each
// component.
$type KEComponent_X param<double>;
$rule unit(KEComponent_X),constraint(volumeTag(X)),
    parametric(volumeTag(X)) {
    $KEComponent_X = 0;
}

// Here we integrate the kinetic energy over each cell which by use
// of the midpoint rule is simply vol*rho*0.5*dot(u,u), then the sum
// is the total kinetic energy
$rule apply(KEComponent_X<-vol,rho,u)[Loci::Summation],
    constraint(volumeTag(X)),parametric(volumeTag(X)) {
    double kecell = 0.5*$vol*$rho*dot($u,$u);
    join($KEComponent_X,kecell);
}</pre>
```

Then KEComponent_X can be written out to the screen in a similar fashion as the previous example.

7 Adding a Volume Source

Another common type of addition to the solver is to add a new source term to the equations. In this example we add a body force to return momentum that is lost to viscous walls for the purpose of computing channel flow type of simulations. When using the implicit solver a jacobian may need to also be computed depending on the stiffness of the source term. In *flowPsi* there are two different possible jacobian types that will need to be supported to support all implicit formulations. By default the code uses jacobians with respect to the primitive variables density and gage pressure, but in modes using the local preconditioner it will use temperature and gage pressure instead. If you want to support both modes of operation you will need to define both types of jacobians.

For the channel flow case we first need to find out how much momentum was lost through the fluid interaction with the viscous walls. For this we utilize a convenience variable (AllViscousBCs) that contains all viscous boundary conditions including viscousWall and wallLaw. For the integration of the momentum flux we consider both the inviscid flux, iflux, and the viscous flux vflux. Since these variables contain mass, momentum, and energy fluxes, we need to extract just the part that we are going to use. For the inviscid flux we skip the first item as this is the mass flux. For the viscous flux the first item is the momentum flux as there is no mass flux in the variable vflux

```
$type viscousWallMomentumFlux param<vect3d> ;
$rule unit(viscousWallMomentumFlux),constraint(UNIVERSE) {
  $viscousWallMomentumFlux = vect3d(0,0,0);
// Note viscous and inviscid flux have already been integrated over the
// boundary facet area. We just need to sum up their contributions
$rule apply(viscousWallMomentumFlux<-iflux)[Loci::Summation],</pre>
  constraint(AllViscousBCs) {
  const int mi = 1
  // extract inviscid momentum flux
 vect3d mflux($iflux[mi+0],$iflux[mi+1],$iflux[mi+2]);
  join($viscousWallMomentumFlux,mflux);
$rule apply(viscousWallMomentumFlux<-vflux,qvi)[Loci::Summation],</pre>
  constraint(AllViscousBCs) {
  const int mi = 0 ;
  // extract viscous momentum flux
 vect3d mflux($vflux[mi+0],$vflux[mi+1],$vflux[mi+2]);
  join($viscousWallMomentumFlux,mflux);
}
```

Now we can compute the total force that we need to add back into the system to restore the lost momentum. To do this we also integrate the total mass in the volume using a similar unit/apply rule combination to compute the variable massSum. The body force is then computed for each cell based on the local density. We also add a term to the energy equation to account for the energy added to the system due to this addition (since we do not wish this energy to be deducted from the internal energy of the fluid). Adding the resulting body force to the solver then is implemented as:

```
//add friction loss to the source term
$rule apply(src<-dragAccel,rho,vol,cellcenter,u)[Loci::Summation] {
  const int mi = 1 ;
  const int mj = mi + 1 ;
  const int mk = mi + 2 ;
  const int ei = 4 ;
  $src[mi] += $dragAccel.x*$rho*$vol ;
  $src[mj] += $dragAccel.y*$rho*$vol ;
  $src[mk] += $dragAccel.z*$rho*$vol ;
  $src[ei] += $dragAccel.x*$u.x*$rho*$vol ;
}</pre>
```

The jacobians in *flowPsi* are computed with respect to the primitive variables gage pressure, temperature, and velocity. The jacobians for the diagonal blocks are stored in the variable srcJ. For this simple source term the jacobians are straightforward to implement which is shown below:

```
$rule apply(srcJ<-rho,gagePressure,temperature,Pambient,u,dragAccel,vol)[Loci::Summation],</pre>
 constraint(vol) {
 const int mi = 1 ;
 const int mj = mi + 1 ;
 const int mk = mi + 2;
 const int ei = 4 ;
 const real drdt = -$rho/$temperature ;
 const real drdp = $rho/($gagePressure+$Pambient) ;
 real coefP = drdp*$vol ;
 real coefT = drdt*$vol ;
 real rho_vol = $rho*$vol;
 $srcJ[mi][4] += $dragAccel.x*coefP ;
 $srcJ[mj][4] += $dragAccel.y*coefP ;
 $srcJ[mk][4] += $dragAccel.z*coefP ;
 $srcJ[ei][4] += dot($dragAccel,$u)*coefP ;
 $srcJ[mi][0] += $dragAccel.x*coefT ;
 $srcJ[mj][0] += $dragAccel.y*coefT ;
 $srcJ[mk][0] += $dragAccel.z*coefT ;
 $srcJ[ei][0] += dot($dragAccel,$u)*coefT ;
 $srcJ[ei][mi] += $dragAccel.x*rho_vol ;
 $srcJ[ei][mj] += $dragAccel.y*rho_vol ;
 $srcJ[ei][mk] += $dragAccel.z*rho_vol ;
}
```

8 Scalar Transport

The flowPsi code provides an infrastructure that makes it simple to add new variables that will be transported with the fluid flow. These variables makes it simple to add new models to accompany the basic fluid flow. In this section we will discuss two examples. First the development of a simple tracer variable that can inject spatially distributed sine waves that can be convected with the flow and a second which goes through the implementation of a simple one equation turbulence model. One advantage of using the scalar transport facilities is that these facilities include interfaces with the restart, overset, and time integration facilities of the flowPsi solver. Thus the scalar transport facilities gives a comprehensive way of adding new variables to the flow solver.

Before we begin, we will have a quick review of the implementation of the scalar transport equations. The scalar transport equation in conservative form is written for a convected scalar, ϕ as

$$\frac{\partial \rho \phi}{\partial t} + \nabla \cdot (\rho u \phi) = \nabla \cdot (\lambda \nabla \phi) + S_{\phi}, \tag{1}$$

where ρ is the fluid density, u is the fluid velocity vector, λ is a diffusion coefficient, and S_{ϕ} is a prescribed source term. We note that by substituting $\phi = 1$ into the above equation one recovers the global continuity equation. We discretize the above equation in time and space to arrive at the expression

$$\frac{\mathcal{V}}{\Delta t} \left[(1+\psi)(\phi^{n+1}\rho^{n+1} - \phi^n \rho^n) + \psi(\phi^n \rho^n - \phi^{n-1}\rho^{n-1}) \right] = -\sum \phi_u^{n+1} \dot{m}_f + \sum D_{\phi_f} + S_{\phi}, \tag{2}$$

where ϕ_u is an upwinded extrapolation of ϕ based on the sign of the face mass flux \dot{m} and D_{ϕ_f} is the numerical diffusion flux for scalar ϕ . Note that the discrete continuity equation is given by

$$\frac{\mathcal{V}}{\Delta t} \left[(1 + \psi)(\rho^{n+1} - \rho^n) + \psi(\rho^n - \rho^{n-1}) \right] = -\sum \dot{m}_f. \tag{3}$$

To improve the diagonal dominance of the scheme we employ a trick where we recognize that Eq. (3) is zero when the system of equation is solved, thus we multiply Eq. (3) by ϕ^{n+1} and subtract it from Eq. (3) to arrive at the following expression:

$$\frac{\mathcal{V}}{\Delta t} \left[(1 + \psi) \left(\phi^{n+1} - \phi^n \right) \rho^n + \left(\phi^n \rho^n - \phi^{n-1} \rho^{n-1} \right) \psi - \left(\rho^n - \rho^{n-1} \right) \phi^{n+1} \psi \right] - \left[\phi^{n+1} \sum \dot{m}_f - \sum \phi_u^{n+1} \dot{m}_f + \sum D_{\phi_f} + S_{\phi} \right] = L_{\phi}(\phi^{n+1}) = 0.$$
(4)

The above operator, L_{ϕ} , is solved using a Newton method concurrently with the fluid equations.

These facilities can be enable to track the convection and diffusion of any arbitrary scalar. The facility is provided by the parametric variables <code>scalarTransport(X,C)</code> or <code>scalarTransportP(X,C)</code> where the first version is for variables that can be of any sign while the second is for variables that are always positive. The first <code>X</code> argument to the parameter is the name of the scalar to be transported, while the second <code>C</code> argument is a constraint that can be used to activate/deactivate the scalar integration. Accessing this variable causes an entire iterative infrastructure to be constructed that solves the scalar transport equations which can then be augmented with additional source terms to create a wide variety of models. An arbitrary number of scalar transport equations can be constructed.

8.1 Tracer Variable Example

The first example that we have is simply the convection of a tracer variable that will simply be convected with the flow. This will allow us to show how to set up the boundary and initial conditions are setup. For this we will create the scalar transport for the variable tracer. To setup flowPsi to transport the variable we first need to instantiate the scalar transport equations. This is accomplished with the line:

```
$rule pointwise(OUTPUT<-scalarTransport(tracer,UNIVERSE)) { }</pre>
```

Note that this rule will never actually execute. Instead, the existence of this rule will cause the parametric scalarTransport variable to be instantiated for the variable tracer. Setting the second argument to UNIVERSE ensures that this scalar transport model will be always active.

To complete the scalar transport equations we need to provide the state of the tracer variable at the boundaries. For this we define the boundary variable tracer_bc. In general we will define the variable for whatever our scalar name is followed by the _bc postfix. In this case we will use a spatially sinusoidal function for boundaries that we mark as having the tracer and set the function to zero otherwise. By default the boundary condition will use simple upwinding to determine the value of tracer_f. To define this we simply use the rules:

```
$type tracer_bc store<real> ;

$rule pointwise(tracer_bc),constraint(ref->BC_options) {
    $tracer_bc = 0 ;
}

$rule pointwise(marker::tracer_bc<-facecenter,tracerNormal,tracerPt,tracerInterval),constracerPt real r = 2.*M_PI*dot($facecenter-$tracerPt,$tracerNormal)/$tracerInterval ;
    $tracer_bc = cos(r) ;
}</pre>
```

Finally we need to output the tracer so that we can see where the tracer advected in the simulation. This is accomplished with the line:

```
OUTPUT_SCALAR("cell2node(tracer)",tracer) ;
```

That is all that is needed to add the most basic type of scalar transport equation. Next we will show how to develop a model with more comprehensive set of parts: a turbulence model.

8.2 The Menter one equation turbulence model

The Menter one equation turbulence model[1] is formally derived from the $k - \epsilon$ model after eliminating k and ϵ from the equations by using the definition of the eddy viscosity and by assuming that the turbulent shear stress is proportional to the turbulent kinetic energy. The model solves the following single equation for the undamped eddy viscosity $\tilde{\nu}_t$:

$$\frac{\partial \rho \tilde{\nu}_t}{\partial t} + \nabla \cdot (\rho \vec{u} \tilde{\nu}_t) = \nabla \cdot \left[\rho \left(\nu + \frac{\tilde{\nu}_t}{\sigma} \right) \nabla \tilde{\nu}_t \right] + \rho c_1 D_1 \tilde{\nu}_t S - \rho c_2 E_{1e}$$
 (5)

where S is the strain rate defined by:

$$S = \sqrt{(\nabla \vec{u}) \cdot (\nabla \vec{u} + \nabla \vec{u}^T)}$$
 (6)

The modified destruction term E_{1e} is defined as:

$$E_{1e} = c_3 E_{BB} \tanh\left(\frac{E_{k-\epsilon}}{c_3 E_{BB}}\right), \tag{7}$$

where

$$E_{k-\epsilon} = \tilde{\nu}_t^2 \left(\frac{1}{L_{VK}}\right)^2 = \tilde{\nu}_t^2 \left(\frac{\nabla S \cdot \nabla S}{S^2}\right) \tag{8}$$

involves the inverse of the von Karman length scale, and E_{BB} is the Baldwin-Barth destruction term:

$$E_{BB} = \nabla \tilde{\nu}_t \cdot \nabla \tilde{\nu}_t. \tag{9}$$

The damping terms D_1 and D_2 are defined as:

$$D_1 = \frac{\nu_t + \nu}{\tilde{\nu}_t + \nu} \tag{10}$$

$$D_2 = 1 - e^{-(\tilde{\nu}_t/A^+ \kappa \nu)^2} , \qquad (11)$$

and the model constants are:

$$c_1 = 0.144, c_2 = 1.86, c_3 = 7, \kappa = 0.41, \sigma = 1, A^+ = 13$$
. (12)

The damped eddy viscosity is obtained from:

$$\nu_t = D_2 \tilde{\nu}_t \ . \tag{13}$$

The boundary conditions for $\tilde{\nu}_t$ at a solid wall are $\tilde{\nu}_t = 0$. The initial and free-stream conditions are $\tilde{\nu}_{t\infty} \leq \nu_{\infty}$. The default initial and free-stream value for $\tilde{\nu}_t$ is 5 x $10^{-9} \, m^2/s$. Note that the boundary conditions are given in terms of the undamped eddy viscosity $(\tilde{\nu}_t)$ rather than ν_t .

8.2.1 Implementation

Examining equation (5) we note that the scalar transport will implement the first three terms. We will need to define the diffusion coefficient and the last two terms to implement the turbulence model. First we invoke the scalar transport of the variable nuTM which represents the $\tilde{\nu}_t$ in the equations. This is accomplished with the following code:

```
$rule pointwise(OUTPUT<-scalarTransportP(nuTM,menter)) { }</pre>
```

Here we constraint the model by the variable menter which will be defined as part of the model as the set of constants that are used in the model. For the initial conditions we will extract nu_t from the initial conditions. This is extracted with the following rules:

```
$type nuTM_ic store<real> ;
$type icRegionInfo blackbox<ICparsedInitRegion> ;

$rule pointwise(nuTM_ic<-icRegionInfo,cellcenter),
    constraint(geom_cells,menter) {
    double nu_tm ;

$icRegionInfo.defaultState.get_nu_t(nu_tm) ;
    $nuTM_ic = nu_tm ;
}</pre>
```

To complete the basic transport of **nuTM** we need to setup the boundary conditions for the model For this we define:

```
$type nuTM_bcVal store<real> ;
// extract the boundary condition from BC_options
$rule pointwise(nuTM_bcVal<-BC_options) {
  real nuTM = 5e-9 ;
  if($BC_options.optionExists("nu_t"))
    $BC_options.getOption("nu_t",nuTM) ;
  $nuTM_bcVal = nuTM ;
}
// set boundary condition
$rule pointwise(nuTM_bc<- ref->nuTM_bcVal) {
  $nuTM_bc = $ref->$nuTM_bcVal ;
}
```

This does not complete the boundary condition however, as the default is to use upwinding to define the boundary face values. However for a boundary such as a viscous wall the boundary value for nuTM_f must be set explicitly. This is accomplished with the priority override rule that will replace the upwind definition with this Dirichlet one:

```
$rule pointwise(noslip::nuTM_f),constraint(viscousWall_BC) {
   $nuTM_f = 0.0 ;
}
```

Now we need to define the scalar diffusion term which is the third term in equation (5). This will be defined by a face value for the diffusion coefficient which will have a name derived from the scalar variable name by appending _nu_f. The diffusion term is thus implemented with the following rules:

```
$type nuTM_nu_f store<real> ;
```

```
$rule pointwise(nuTM_nu_f<-rho_f,nuTM_f,mu_f,menter) {
  const real sigma = $menter.sigma;
  $nuTM_nu_f = ($mu_f+$rho_f*$nuTM_f/sigma);
}

// The diffusion coefficient needs nuTM_f at all faces, but the
// infrastructure only defines them at the boundary.
// For interior faces we set the face value to be simply the
// reciprocal volume weighted average of the cell values
$rule pointwise(nuTM_f<-(cr,cl)->(vol,nuTM)) {
  real vrvl = 1.0/($cr->$vol+$cl->$vol);
  $nuTM_f = ($cr->$vol*$cl->$nuTM+$cl->$vol*$cr->$nuTM)*vrvl;
}
```

Now we have completely defined the first three terms of equation (5). Now what remains is to define the destruction and production terms. To add these terms we will use a unit rule to combine them with the previous terms in the variable nuTM_src. First lets consider the production term which which can be added using the rules:

where strainRate is provide by the solver and D1 is the production damping function which is defined by the rule:

```
$type D1 store<real> ;
$rule pointwise(D1<-nuTM,tmu,mu,rho) {
  const real nut = $tmu/$rho ;
  const real nu = $mu/$rho ;
  $D1 = (nut+nu)/($nuTM+nu) ;
}</pre>
```

Note, the turbulent viscosity (tmu) has not yet been defined but will be shortly. The order that rules appear in the file do not matter. Note, we have not considered the jacobian of the production term. In general the sign of the production jacobian is destabilizing and as a general rule it is desirable to leave this term out of an implicit implementation.

Next we will consider the last destruction term. This can be implemented using the following rules:

One point to note on this is that we need the gradient of strainRate to implement this term. For this implementation we are simply using the least squares gradient provided by the FVM module to compute this. However, we will run into a problem here because the gradient computation will require a value for strainRate at the face which is not provided by the solver. For expediency we will assume that strainRate is relatively constant at the boundaries and just copy from the cell to the face. This is accomplished with the rule

```
$rule pointwise(strainRate_f<-ci->strainRate) {
   $strainRate_f = $ci->$strainRate ;
}
```

This implementation is OK in a pinch, but more sophisticated methods for computing this would be suggested for a production level implementation. In particular the recursive application of least squares gradients can be sensitive to numerical noise and the treatment of the boundary values may lead to spurious results. For our limited tests this implementation seems to produce satisfactory results.

For the destruction term it can be helpful to include a jacobian term. However, due to the gradients of nuTM used in the destruction term, the exact jacobian can be expensive and tedious to derive. For this reason we use a diagonalized approximation that can stabilize the solver with a reasonable cost. The diagonal term of the jacobian is in the variable nuTM_srcJ. Our approximation is implemented using the following rule

Now all that remains is defining the turbulent viscosity which is computed from the scalar variable we are solving combined with a the damping term D_2 . This is defined using the rules:

And with this we complete the implementation of the turbulence model.

References

[1] F R. Menter. Eddy viscosity transport and equations and their relation to the k- ϵ model. J. of Fluids Engineering, 119:876–884, 1997.