## SPHERLS

1.0

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# Chapter 1

# Using and Modifying SPHERLS

This manual is divided into two main parts the current chapter, and the rest of the chapters. All chapters other than the current, contain specific reference material for the SPHERLS code while the current chapter contains a more descriptive how-to approach explaining the usage and modification of SPHERLS. The chapters following chapter 1 will serve as a usefull reference when specific details need to be found, for example a discription of a particular variable in the code. The current chapter on the other hand is the best place to go to get a quick understanding of SPHERLS that will enable you to use it.

#### 1.1 Overview

SPHERLS stands for Stellar Pulsation with a Horizontal Eulearian Radial Lagrangian Scheme. There are three components to SPHERLS: SPHERLS itself which does the hydodynamics calculations, SPHERLSgen which creates starting models, and SPHERLSanal which is able to manipulate the output files. Both SPHERLSgen and SPHERLSanal have there own manuals which can be consulted for their specific uses and installations.

#### 1.1.1 The Basics

SPHERLS calculates the radial pulsation motions together with the horizontal convective flow. The radial pulsation can be described by a radial grid velocity Grid::nU0, moving the grid inward and outward with the pulsation. The movement of the grid is defined by the motion required to maintaining the mass in a spherical shell through out the calculation. This motion is determined so that it will change the volume of the shell so the newly calculated density when multiplied with the new volume will produce the same shell mass. The total motion of the stellar material is simply the combination of the three velocity components, radial Grid::nU, theta Grid::nV, and phi velocities Grid::nW. The convective motion is the radial velocity minus the grid velocity, combined with the theta and phi velocities. This is because the grid velocity describes the bulk motion of the pulsation so subtracting it out leaves only the convective motions.

SPHERLS solves the normal hydodynamic equations of, mass, momentum, and energy conservation. The form of the mass equation, momentum conservation, and energy conservation are:

$$\frac{dM}{dt} + \oint_{\mathbb{S}} (\rho \vec{v}) \cdot \hat{n} d\sigma = 0$$

$$\frac{\partial \vec{v}}{\partial t} + (\vec{v} \cdot \nabla)\vec{v} = -\frac{1}{\rho}\nabla P + \nabla \cdot \boldsymbol{\tau} - \nabla \phi$$

$$\frac{\partial E}{\partial t} + (\vec{v} \cdot \nabla)E + P\frac{d\nabla}{dt} = \epsilon + \frac{1}{\rho} \left[ -\nabla \cdot F + \nabla \cdot (\boldsymbol{\tau} \cdot \vec{v}) - (\nabla \cdot \boldsymbol{\tau}) \cdot \vec{v} \right]$$

where  $\tau$  is the stress tensor for zero bulk viscosity, E is the specific internal energy,  $\mathbb{V}$  is the specific volume, and F is the radiative flux. In addition to these conservation equations an equation of state is needed, in this case the OPAL equation of state and opacities, and the Alaxander opacities at low temperatures are used. The equation of state tables are functions of density and temperature, and produce the energy, pressure, opacity, and adiabatic index of the gas for a given temperature and density. In adiabatic calculations, it is also possible to use a  $\gamma$ -law gas equation of state but in that case an energy profile must also be included.

The simulation grid is broken up into two main sections, the 1D region towards the center of the star, the multi-dimensional region towards the surface. The inner part of the multi-dimensional region solves all the conservation equations explicitly, in that the new values for the conserved quantities are directly calculated from the information in the previous time step. In the outter parts of the multi-dimensional region the energy conservation equation is calculated semi-implicitly, which means that the new values are dependent on the new values averaged with the old values to correctly time cetner the equation. This semi-implicit energy conservervation equation can be preturbed and linearized producing a set of linear equations the size of the region being solved implicitly. The solution of these linear equations provide corrections for the temperature which can be applied and then resolved in an iterative approach until the value of the new temperature converges. The equation of state is a function of temperature and not energy which is why the temperature is pretubed and not the energy. This set of equations for the temperature corrections are solved using the PETSC library.

• Different ways in which SPHERLS can be used, 1D,2D,3D, Adiabatic,Non-adiabatic, implicit, debugging options/test

## 1.2 The Equations

I will want to give a detailed description of the equations used (probably copied from my notes wiki) so that the reader can easily see a 1-1 correspondence between the equation and the terms in SPHERLS.

## 1.3 Program Flow

- Describe the grids
- The order of calculation
- When parts of the grid are updated

#### 1.4 Installing SPHERLS

#### Todo

This should be updated to reflect the use of the GNU build system.

Once the correct libraries are installed, and their paths added to your LD\_LIBRARY\_PATH environment varible, it should just require typing make in the correct directories. SPHERLS is broken up into 3 main codes. SPHERLS it self, which is the main hydrodynamics code which integrates the initial static model, SPHERLSgen which creates the static model, and SPHERLSanal which is used for processing the output of SPHERLS and SPHERLSgen.

To Add

- example .bashrc entries, showing LD\_LIBRARY\_PATH additions, and other SPHERLS related configuration options
- also the make files will need to know where the paths for the libraries are, either describe how the user can do this, or automate it some how.

A few words on installation before we get into the details of the specific packages. In order for the SPHERLS configuration script (needed for installing SPHERLS) to find the required libraries and include files they have to be installed in at least one of the directories that it looks for them. The configuration script looks for the libraries and include files it requires in the following "standard" locations: /lib, /include, /usr/lib /usr/include, /usr/local/lib, /usr/local/include, /home/\$USER/lib, and /home/\$USER/include . If you install the required libraries in places other than these "standard" locations you will have to manualy tell the SPHERLS configuration script where to find them. Running configure -h will list the avaible options to tell the script where to find these include files and libraries.

I am going to assume that the user is installing on a linux system, more over I will be assuming that the linux distribution follows a debian like directory structure (many distributions are based on debian). The install instructions below assume you do not have root access and must do an install to your home directory (a per-user install). The standard install location for per-user level binaries, libraries, include files and documentation (at least far as SPHERLS is concerned) is in  $\sim$ /bin,  $\sim$ /lib,  $\sim$ /include, and  $\sim$ /share directories respectively. Be aware that if you have these directories in your home directory already and are not using them as a standard place to install per-user packages you will likely want to rename your pre-existing directories or a bunch of additional files will be added to them from the installations of various packages below. Alternatively, you can install the libraries and binaries to any directory on your machine just by changing

```
--prefix=/home/$USER/
```

, mentioned below, to point to where you want to install it.

If you have root access and want to install for all users of the current machine you will likely want to install the libraries into /usr/local which can be achieved by setting

```
--prefix=/usr/local
```

instead of

```
--prefix=/home/$USER/
```

used in the below installations and SPHERLS will automatically check this location for the install libraries.

#### 1.4.1 Requirements

- gcc/g++
- openMPI
- PETSc library, used as the core matrix solver

#### 1.4.2 Optional Requirements

- python for analysis scripts numpy used by matplotlib matplotlib for creating plots scipy for interpolating in equation of state files
- fftw3 library for frequency analysis
- hdf4 library for converting to hdf4 file format (not yet implemented)
- Doxygen used to create documentation from source code via "make docs"

#### 1.4.3 Installing PETSC Library

Version petsc-lite-3.1-p8, has been tested to work with SPHERLS. petsc-lite-3.2-p7 is known to be incompatible, which as of this writting is the current version of the petsc library. At some point in the future support for the newer version of the library maybe added. The below commands will install PETSc into your home directory. ASIDE: I have also had difficulties installing PETSc on Fundy, and Placentia ACENet machines.

- Download PETSc library, from the PETSc website.
- Then untar and unzip it with tar -xzf petsc-lite-3.1-p8.tar
- To install the library change into the directory made when you extracted the archive and type the following commands:
  - 1. ./configure PETSC\_DIR=\$PWD --prefix=/home/\$USER/ --with-c++-support --with-c-support --with-shared --download-f-blas-lapack=1 --with-x11=no --with-x11=no \$USER is the environment varible coresponding to your username.
  - 2. make all
    - \ Often at the end of the configuration stage the configuration script will give the command to make the library. One should use this over the above if given.
  - 3. make install

\ as with the

make all

the makeFile will also likely tell you the command needed for the installation, which should be used over the one provided here.

- 4. make PETSC\_DIR=/home/\$USER/lib test
- You will then need to add the following line to you .bashrc file to assure that you will pick up the library

export PETSC\_DIR=/home/\$USER/lib

#### 1.4.4 Installing FFTW Library

- Download the FFTW Library from the FFTS website. Version fftw-3.2.2 has been tested to work with SPHERLS.
- The downloaded FFTW file (e.g. fftw-3.2.2.tar.gz) will need to be unziped to do so type gunzip fftw-3.2.2.tar.gz
- $\bullet$  Then untar it with tar -xf fftw-3.2.2.tar.gz
- To install the library change into the directory made when you extracted the archive and type the following commands:
  - $1. \ \ \, \hbox{\tt ./configure --prefix=<path-to-final-location-of-library>}$
  - 2. make
  - 3. make install

#### 1.4.5 Installing HDF4 Library

#### 1.4.6 Installing Doxygen

The latest version seems to hang while creating documents. doxygen-1.5.6 is known to work.

#### 1.4.7 Installing Python

#### 1.4.8 Installing SPHERLS

## 1.5 Using SPHERLS

- Generating a starting model (see SPHERLSgen documentation for details)
- The XML configuration file
- Starting a calculation and the "makeFile"
- getting data
  - watchzones
  - model dumps
- post calculation analysis (see SPHERLSanal documention for details)
- Adiabatic Calculations

- 1D, 2D, and 3D
- \$gamma\$-law gas
- Sedov Blast wave test
- Non-Adiabatic Calculations
  - 1D, 2D, and 3D
  - Tabulate EOS
  - Different versions of the energy equation
  - LES models

### 1.6 Modifing or Developing SPHERLS

- Basic layout/design of the code
  - model output
  - data monitoring
    - \* watch zones
    - \* peak KE tracking
  - internal/versus external variables
  - message passing
  - grid layout
  - ranges of grids
  - boundary regions
  - grid updating
- How to document SPHERLS
- Premade test for SPHERLS after modification
  - reference calculations
  - restart test
  - calculation test (if not modifying calculation part of SPHERLS)
- How to modify SPHERLS
  - Common changes
    - \* How to add a new internal variable
      - 1. Add to the internal variable count: Decide in what cases the variable will be needed, 1D calculations, 2D calculations, when there is a gamma law gas or a tabulated equation of state, adiabatic or non-adiabatic etc. Then once decided it can be added to the total number of internal variables Grid::nNumIntVars by increasing the value by one in the function modelRead in the section below the comment "set number of internal variables ..." under the appropriate if block. If the specific if block for the situation you need isn't there, you can create your own, and add it there.

- 2. Create a new variable ID: In the grid::h file under the Grid class are variable ID's. These ID's simply indicate the location of the variable in the array. One must add a new ID for the new variable as an integer. The value of the ID is set in the function modelRead in the same section as the number of internal variables. The value used should be the last integer after the last pre-existing variable ID. This should also be Grid::nNumVars + Grid::nNumIntVars -1. The ID should also be initalized to -1, so that the code knows when it isn't being used. This is done in the grid class constructor, Grid::Grid. Simply add a line in the constructor setting your new ID = -1.
- Set variable infos: Decide what the dimensions of the new variable will be. It can be cell centered or interface centered. It can also be only 1D, 2D, or 3D. Of course it will be only 1D if the entire calculation is 1D, or 2D if the calculation is 2D, but if the calculation is 3D it could also only be 2D, or 1D, and if 2D it could be only 1D. Also decide if the variable will change with time, dependent variables are only initialized and not updated during the calculations. This information is given to SPHERLS in the setInternal VarInf function in the physEquations.cpp file. The variable that is set is Grid::nVariables. It is a 2D array, the first index corresponds to the particular variable in question, the ID you made in the previous step can be used as the first index of this array. The second index referes to one of the three directions (0-2) or the time dimension (3). If the variable is cenertered in the grid in direction 0 (r-direction) then this array element should have a value of 0. If the variable is interface centered in the grid in direction 0, then this array element should have a value of 1. If it isn't defined in direction 0 (for example the theta independent variable isn't defined in the 0 direction) then it should be -1. This is the same for the other 2directions. The last element (3) should be either 0 not updated every time step, or 1 if updated every timestep. There are various sections here which allows one to set variable information based on which conditions are the varible is defined in. Put these varible infos into the most general case in which the varible is defined. At the end of this function variables are atomatically adjusted depending on what the number of dimensions the model uses, so this does not need to be considered unless the variable is not used at all for a specific case of dimensions. For example a varible which is defined at cell center for all three cases for the number of dimensions (1D, 2D, 3D) will be atomatically adjusted to be not defined in the 3rd direction when only doing 2D calculations, and similarly for 1D only defined in 1st direction and not defined in the 2nd or 3rd directions.
- 4. Add functions: Finally to do anything usefull with your new internal variable functions must be added to initialize the values of the variables, and to update them with time if needed. Initiliazation functions are called within the initInternalVars function in the physEquations.cpp file. The details of these functions will depend on what the individual variables are intended for. Functions to be called every timestep must be called from the main program loop in the file main.cpp in the appropriate order.
- \* How to add a new external variable
- \* How to add a new physics functions
  - · Function naming conventions
  - · Grid variables
  - indecies and their ranges
- SPHERLS debugging tips

# 1.7 Message Passing

 $\bullet~$  Explain message passing in SPHERLS

Chapter 2

Todo List

10 Todo List

page Using and Modifying SPHERLS This should be updated to reflect the use of the GNU build system.

Member Time::nTimeStepIndex should probably make this an unsigned variable, and perhaps also use the keyword long to help ensure there are enough values. Often need 7 decimal places.

**Member initImplicitCalculation** isFrom, isTo, matCoeff,vecTCorrections, vecTCorrections,vecRHS,vecTCorrectionsLocal ,kspContext,vecscatTCorrections all need to be destroyed before program finishes.

Member modelRead At some point should get it working with only 1 processor

Member updateLocalBoundaries Shouldn't need MPI::COMM\_WORLD.Barrier() may want to test out removing this at some point as it might produce a bit of a speed up.

Member updateLocalBoundariesNewGrid May want to do some waiting on this message at some point before the end of the timestep, but it doesn't need to be done in this function. It might also be that this is built into the code by waiting at some other point. This is something that should be checked out at somepoint, perhaps once the preformance starts to be analyzed. I would think that if the send buffer was being modified before the send was completed, that there would be some errors poping up that would likely kill the program.

Member calNewU0\_R At some point I will likely want to make this function compatiable with a 3D domain decomposition instead of a purely radial domain decomposition.

Member calNewU0\_RT At some point I will likely want to make this function compatiable with a 3D domain decomposition instead of a purely radial domain decomposition.

Member calNewU0\_RTP At some point I will likely want to make this function compatiable with a 3D domain decomposition instead of a purely radial domain decomposition.

Member dImplicitEnergyFunction\_R\_LES this function should probably be turffed, the LES terms aren't needed in 1D. keeping it for now though.

Chapter 3

**Boundary Conditions** 

Member calNewD R doesn't allow mass flux through outter interface

Member calNewD RT doesn't allow mass flux through outter interface

Member calNewD RTP doesn't allow mass flux through outter interface

Member calNewE\_R\_NA Missing grid.dLocalGridOld[grid.nE][i+1][j][k] in calculation of  $E_{i+1/2,j,k}$  setting it equal to value at i.

 $\label{lem:member calNewE} \begin{tabular}{ll} Member calNewE \_R \_NA & $\operatorname{grid.dLocalGridOld[grid.nDM][i+1][0][0]} & \text{and} \\ & \operatorname{grid.dLocalGridOld[grid.nE][i+1][j][k]} & \text{missing in the calculation of upwind gradient in} \\ & \operatorname{dA1. Using the centered gradient instead.} \\ \end{tabular}$ 

Member calNewE R NA Missing grid.dLocalGridOld[grid.nT][i+1][0][0]

Member calNewE\_R\_NA\_LES Missing grid.dLocalGridOld[grid.nE][i+1][j][k] in calculation of  $E_{i+1/2,j,k}$  setting it equal to value at i.

 $\label{lember calNewE} \begin{tabular}{ll} \bf Member\ calNewE\ R\ \_NA\ \_LES \ grid.dLocalGridOld[grid.nDM][i+1][0][0] \ and \\ grid.dLocalGridOld[grid.nE][i+1][j][k] \ missing \ in \ the \ calculation \ of \ upwind \ gradient \ in \ dA1. \ Using the \ centered \ gradient \ instead. \end{tabular}$ 

Member calNewE R NA LES Missing grid.dLocalGridOld[grid.nT][i+1][0][0]

Member calNewE\_R\_NA\_LES missing energy outside the model, assuming it is the same as that in the last zone. That causes this term to be zero.

 $\begin{tabular}{ll} \bf Member\ calNewE & \bf RT & \bf AD & grid.dLocalGridOld[grid.nE][i+1][j][k] \ is \ missing \end{tabular}$ 

 $\begin{array}{ll} \mathbf{Member\ calNewE\_RT\_AD} & \mathrm{grid.dLocalGridOld[grid.nDM][i+1][0][0]} \\ & \mathrm{grid.dLocalGridOld[grid.nE][i+1][j][k]} & \mathrm{missing\ using\ inner\ gradient\ for\ both} \\ \end{array}$ 

and

Member calNewE\_RT\_NA Missing grid.dLocalGridOld[grid.nE][i+1][j][k] in calculation of  $E_{i+1/2,j,k}$  setting it equal to value at i.

Member calNewE\_RT\_NA grid.dLocalGridOld[grid.nDM][i+1][0][0] and grid.dLocalGridOld[grid.nE][i+1][j][k] missing in the calculation of upwind gradient in dA1. Using the centered gradient instead.

Member calNewE\_RT\_NA Missing grid.dLocalGridOld[grid.nT][i+1][0][0] using flux equals  $2\sigma T^4$  at surface.

Member calNewE\_RT\_NA\_LES Missing  $\Delta M_r$  outside model using Parameters::dAlpha times  $\Delta M_r$  in the last zone instead.

Member calNewE RT NA LES Setting energy at surface equal to energy in last zone.

Member calNewE\_RT\_NA\_LES missing eddy viscosity outside the model setting it to zero

Member calNewE\_RT\_NA\_LES missing energy outside the model, assuming it is the same as that in the last zone. That causes this term to be zero.

Member calNewE\_RTP\_AD Missing grid.dLocalGridOld[grid.nE][i+1][j][k] in calculation of  $E_{i+1/2,j,k}$  setting it equal to zero.

 $\begin{tabular}{ll} \bf Member\ calNewE\_RTP\_AD & grid.dLocalGridOld[grid.nDM][i+1][0][0] & and grid.dLocalGridOld[grid.nE][i+1][j][k] & missing in the calculation of upwind gradient in dA1.Using the centered gradient. \\ \end{tabular}$ 

**Member calNewE\_RTP\_NA** Missing grid.dLocalGridOld[grid.nE][i+1][j][k] in calculation of  $E_{i+1/2,j,k}$  setting it equal to the value at i.

 $\label{lem:member calNewE} \begin{array}{ll} \textbf{Member calNewE} \_ \textbf{RTP} \_ \textbf{NA} & grid.dLocalGridOld[grid.nDM][i+1][0][0] & and \\ grid.dLocalGridOld[grid.nE][i+1][j][k] & missing in the calculation of upwind gradient in dA1. Using the centered gradient instead. \\ \end{array}$ 

Member calNewE RTP NA Missing grid.dLocalGridOld[grid.nT][i+1][0][0]

Member calNewE\_RTP\_NA\_LES Missing  $\Delta M_r$  outside model using Parameters::dAlpha times  $\Delta M_r$  in the last zone instead.

Member calNewE RTP NA LES Missing W at i+1, assuming the same as at i

Member calNewE\_RTP\_NA\_LES Missing grid.dLocalGridOld[grid.nE][i+1][j][k] in calculation of  $E_{i+1/2,j,k}$  setting it equal to the value at i.

Member calNewE RTP NA LES missing density outside model, setting it to zero

 $\label{lem:member calNewE} \begin{array}{lll} \textbf{Member calNewE} \_ \textbf{RTP} \_ \textbf{NA} \_ \textbf{LES} & grid.dLocalGridOld[grid.nDM][i+1][0][0] & and \\ grid.dLocalGridOld[grid.nE][i+1][j][k] & missing in the calculation of upwind gradient in dA1. Using the centered gradient instead. \\ \end{array}$ 

Member calNewE RTP NA LES Missing grid.dLocalGridOld[grid.nT][i+1][0][0]

Member calNewEddyVisc RTP SM assume phi velocity is constant across surface

Member calNewU0\_R assuming density outside the star is 0 for the purposes of calculating the grid velocity. This boundary condition works, but is probably not strictly correct. The proper condition can be calculated by knowing that the mass conservation is independent of u0\_ip1half at the surface. This allows one to simply use the new densities and calculate the grid velocity from them.

Member calNewU0 RT grid.dLocalGridOld[grid.nD][i+1][j][k] is missing

Member calNewU\_R Missing grid.dLocalGridOld[grid.nD][nICen+1][j][k] in calculation of  $\rho_{i+1/2}$ , setting it to 0.0

**Member calNewU\_R** Missing grid.dLocalGridOld[grid.nP][nICen+1][j][k] in calculation of  $S_1$ , setting it to -1.0\*grid.dLocalGridOld[grid.nP][nICen][j][k].

Member calNewU\_R Missing grid.dLocalGridOld[grid.nDM][nICen+1][0][0] in calculation of centered  $A_1$  gradient, setting it to zero.

 $\begin{array}{llll} \textbf{Member calNewU} \underline{\textbf{R}} & \text{Missing} & \text{grid.dLocalGridOld[grid.nU][i+1][j][k]} & \text{and} \\ & \text{grid.dLocalGridOld[grid.nDM][nICen+1][0][0]} & \text{in calculation of upwind gradient, when} \\ & \text{moving inward. Using centered gradient instead.} \end{array}$ 

Member calNewU\_R\_LES Missing grid.dLocalGridOld[grid.nD][nICen+1][j][k] in calculation of  $\rho_{i+1/2}$ , setting it to 0.0

Member calNewU R LES Assuming eddy viscosity outside model is zero.

- Member calNewU\_R\_LES Missing grid.dLocalGridOld[grid.nP][nICen+1][j][k] in calculation of  $S_1$ , setting it to -1.0\*grid.dLocalGridOld[grid.nP][nICen][j][k].
- Member calNewU\_R\_LES Missing grid.dLocalGridOld[grid.nDM][nICen+1][0][0] in calculation of centered  $A_1$  gradient, setting it to zero.
- $\label{lem:lember_calNewU} \begin{tabular}{ll} \bf R\_LES & Missing & grid.dLocalGridOld[grid.nU][i+1][j][k] & and & grid.dLocalGridOld[grid.nDM][nICen+1][0][0] & in calculation of upwind gradient, when moving inward. Using centered gradient instead. \\ \end{tabular}$
- Member calNewU\_RT Missing grid.dLocalGridOld[grid.nD][nICen+1][j][k] in calculation of  $\rho_{i+1/2,j,k}$ , setting it to zero.
- Member calNewU RT assuming theta velocity is constant across surface
- Member calNewU\_RT Missing grid.dLocalGridOld[grid.nDenAve][nICen+1][0][0] in calculation of  $\langle \rho \rangle_{i+1/2}$ , setting it to zero.
- Member calNewU\_RT Missing grid.dLocalGridOld[grid.nP][nICen+1][j][k] in calculation of  $S_1$ , setting it to -1.0\*dP ijk n.
- Member calNewU\_RT Missing grid.dLocalGridOld[grid.nDM][nICen+1][0][0] in calculation of centered  $A_1$  gradient, setting it to zero.
- Member calNewU\_RT Missing grid.dLocalGridOld[grid.nU][i+1][j][k] and grid.dLocalGridOld[grid.nDM][nICen+1][0][0] in calculation of upwind gradient, when moving inward. Using centered gradient instead.
- $\begin{tabular}{ll} \bf Member\ calNewU\_RT & Missing\ grid.dLocalGridOld[grid.nDM][i+1][0][0] \ in\ calculation\ of\ S_1\\ using\ Parameters::dAlpha\ *grid.dLocalGridOld[grid.nDM][nICen][0][0] \ instead. \\ \end{tabular}$
- $\begin{tabular}{ll} \bf Member\ calNewU\_RT\_LES & Missing\ grid.dLocalGridOld[grid.nDM][i+1][0][0] & in\ calculation\ of\ S_1\ using\ Parameters::dAlpha\ *grid.dLocalGridOld[grid.nDM][nICen][0][0] & instead. \\ \end{tabular}$

Member calNewU RT LES Missing density outside of surface, setting it to zero.

Member calNewU RT LES Missing density outside model, setting it to zero.

Member calNewU RT LES assuming theta and phi velocity same outside star as inside.

Member calNewU RT LES Assuming theta velocities are constant across surface.

Member calNewU RT LES assuming that \$V\$ at \$i+1\$ is equal to \$v\$ at \$i\$.

Member calNewU\_RT\_LES Missing pressure outside surface setting it equal to negative pressure in the center of the first cell so that it will be zero at surface.

Member calNewU RT LES assume viscosity is zero outside the star.

Member calNewU RT LES Missing mass outside model, setting it to zero.

 $\label{lem:lember_calNewU_RT_LES} \begin{tabular}{ll} Member calNewU_RT_LES & Missing & grid.dLocalGridOld[grid.nU][i+1][j][k] & and & grid.dLocalGridOld[grid.nDM][nICen+1][0][0] & in calculation of upwind gradient, when moving inward. Using centered gradient instead. \\ \end{tabular}$ 

Member calNewU\_RTP missing grid.dLocalGridOld[grid.nU][i+1][j][k] in calculation of  $u_{i+1,j,k}$  setting  $u_{i+1,j,k}=u_{i+1/2,j,k}$ .

**Member calNewU\_RTP** Missing grid.dLocalGridOld[grid.nD][i+1][j][k] in calculation of  $\rho_{i+1/2,j,k}$ , setting it to zero.

Member calNewU RTP assuming theta velocity is constant across the surface.

Member calNewU RTP assuming phi velocity is constant across the surface.

Member calNewU\_RTP Missing grid.dLocalGridOld[grid.nDenAve][nICen+1][0][0] in calculation of  $\langle \rho \rangle_{i+1/2}$  setting it to zero.

**Member calNewU\_RTP** Missing grid.dLocalGridOld[grid.nDM][nICen+1][0][0] in calculation of centered  $A_1$  gradient, setting it equal to Parameters::dAlpha grid.dLocalGridOld[grid.nDM][nICen][0][0].

 $\label{lem:member calNewU} \begin{array}{ll} \textbf{Member calNewU} \_ \textbf{RTP} & \text{Missing} & \text{grid.dLocalGridOld[grid.nU][i+1][j][k]} & \text{and} \\ & \text{grid.dLocalGridOld[grid.nDM][nICen+1][0][0]} & \text{in calculation of upwind gradient, when} \\ & \text{moving inward. Using centered gradient instead.} \end{array}$ 

Member calNewU\_RTP Missing grid.dLocalGridOld[grid.nDM][i+1][0][0] in calculation of  $S_1$  using Parameters::dAlpha \*grid.dLocalGridOld[grid.nDM][nICen][0][0] instead.

Member calNewU RTP LES Missing density outside of surface, setting it to zero.

Member calNewU RTP LES Missing density outside model, setting it to zero.

Member calNewU RTP LES assuming theta and phi velocity same outside star as inside.

Member calNewU RTP LES Assuming theta velocities are constant across surface.

Member calNewU\_RTP\_LES Missing pressure outside surface setting it equal to negative pressure in the center of the first cell so that it will be zero at surface.

Member calNewU RTP LES assume viscosity is zero outside the star.

Member calNewU RTP LES Missing mass outside model, setting it to zero.

 $\label{lem:member calNewU} \begin{tabular}{ll} \bf RTP\_LES & Missing & grid.dLocalGridOld[grid.nU][i+1][j][k] & and & grid.dLocalGridOld[grid.nDM][nICen+1][0][0] & in calculation of upwind gradient, when moving inward. Using centered gradient instead. \\ \end{tabular}$ 

Member calNewV RT grid.dLocalGridOld[grid.nV][i+1][j+1][k] is missing

Member calNewV RT missing upwind gradient, using centred gradient instead

Member calNewV RT LES Assuming density outside star is zero

Member calNewV RT LES Assuming theta velocity is constant across surface.

Member calNewV RT LES Assuming eddy viscosity is zero at surface.

Member calNewV\_RTP Assuming theta and phi velocities are the same at the surface of the star as just inside the star.

Member calNewV RTP ussing cetnered gradient for upwind gradient outside star at surface.

Member calNewV RTP LES Assuming density outside star is zero

Member calNewV RTP LES Assuming theta velocity is constant across surface.

Member calNewV RTP LES Assuming eddy viscosity is zero at surface.

Member calNewW\_RTP missing grid.dLocalGridOld[grid.nW][i+1][j][k] assuming that the phi velocity at the outter most interface is the same as the phi velocity in the center of the zone.

Member calNewW\_RTP missing grid.dLocalGridOld[grid.nW][i+1][j][k] in outter most zone. This is needed to calculate the upwind gradient for donnor cell. The centered gradient is used instead when moving in the negative direction.

Member calNewW RTP LES assume theta and phi velocities are constant across surface

Member calNewW RTP LES assume eddy viscosity is zero at surface

Member calOldEddyVisc RTP SM assuming that theta velocity is constant across surface

Member calOldEddyVisc RTP SM assume phi velocity is constant across surface

Member dImplicitEnergyFunction\_R\_LES\_SB Missing grid.dLocalGridOld[grid.nE][i+1][j][k] in calculation of  $E_{i+1/2,j,k}$  setting it equal to value at i.

Member dImplicitEnergyFunction\_R\_LES\_SB grid.dLocalGridOld[grid.nDM][i+1][0][0] and grid.dLocalGridOld[grid.nE][i+1][j][k] missing in the calculation of upwind gradient in dA1. Using the centered gradient instead.

```
 \mathbf{Member\ dImplicitEnergyFunction\ R\ LES\ SB\ } \operatorname{Missing\ grid.dLocalGridOld[grid.nT][i+1][0][0] }
```

Member dImplicitEnergyFunction\_R\_SB Missing grid.dLocalGridOld[grid.nE][i+1][j][k] in calculation of  $E_{i+1/2,j,k}$  setting it equal to value at i.

Member dImplicitEnergyFunction\_R\_SB A1 upwind set to zero as no material is flowing into the star

Member dImplicitEnergyFunction R SB Missing grid.dLocalGridOld[grid.nT][i+1][0][0]

Member dImplicitEnergyFunction\_RT\_LES\_SB Missing  $\Delta M_r$  outside model using Parameters::dAlpha times  $\Delta M_r$  in the last zone instead.

Member dImplicitEnergyFunction\_RT\_LES\_SB missing density outside model assuming it is zero

Member dImplicitEnergyFunction\_RT\_LES\_SB missing desnity outside model assuming it is zero

Member dImplicitEnergyFunction\_RT\_LES\_SB assuming V at ip1half is the same as V at i

Member dImplicitEnergyFunction\_RT\_LES\_SB Assuming energy outside model is the same as the energy in the last zone inside the model.

Member dImplicitEnergyFunction\_RT\_LES\_SB Using centered gradient for upwind gradient when motion is into the star at the surface

Member dImplicitEnergyFunction\_RT\_LES\_SB Missing grid.dLocalGridOld[grid.nT][i+1][0][0] using flux equals  $2\sigma T^4$  at surface.

- Member dImplicitEnergyFunction\_RT\_SB Using centered gradient for upwind gradient when motion is into the star at the surface
- Member dImplicitEnergyFunction\_RT\_SB Missing grid.dLocalGridOld[grid.nT][i+1][0][0] using flux equals  $2\sigma T^4$  at surface.
- Member dImplicitEnergyFunction\_RTP\_LES\_SB Missing  $\Delta M_r$  outside model using Parameters::dAlpha times  $\Delta M_r$  in the last zone instead.
- Member dImplicitEnergyFunction\_RTP\_LES\_SB missing density outside model assuming it is zero
- Member dImplicitEnergyFunction\_RTP\_LES\_SB missing desnity outside model assuming it is zero
- Member dImplicitEnergyFunction\_RTP\_LES\_SB assuming V at ip1half is the same as V at i
- Member dImplicitEnergyFunction\_RTP\_LES\_SB assuming W at ip1half is the same as W at i
- Member dImplicitEnergyFunction\_RTP\_LES\_SB Assuming energy outside model is the same as the energy in the last zone inside the model.
- Member dImplicitEnergyFunction\_RTP\_LES\_SB A1 upwind set to zero as no material is flowing into the star
- Member dImplicitEnergyFunction\_RTP\_LES\_SB Missing grid.dLocalGridOld[grid.nT][i+1][0][0] using flux equals  $2\sigma T^4$  at surface.
- $\begin{tabular}{ll} \bf Member \ dImplicitEnergyFunction\_RTP\_SB \ Using \ \$E_{i,j,k}^{n+1/2}\$ \ for \ \$E_{-i+1/2,j,k}^{n+1/2}\$ \\ \hline \end{tabular}$

Member dImplicitEnergyFunction\_RTP\_SB Using centered gradient for upwind gradient when motion is into the star at the surface

 $\begin{array}{ll} \textbf{Member dImplicitEnergyFunction\_RTP\_SB} & \textbf{Missing grid.dLocalGridOld[grid.nT][i+1][0][0]} \\ & \textbf{using flux equals } 2\sigma T^4 \text{ at surface.} \end{array}$ 

# Chapter 4

# Class Index

## 4.1 Class List

Here are the classes, structs, unions and interfaces with brief descriptions:

eos												 										29
Functions																						
Global												 										43
Grid .												 										45
Implicit												 										67
MessPass																						
Output												 										74
Paramete	rs											 										77
Performa	nce	е										 										83
ProcTop												 										85
Time .												 										88
WatchZo:	ne																					92

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# Chapter 5

# File Index

## 5.1 File List

Here is a list of all documented files with brief descriptions:

/home/cgeroux/WORK/SPHERLS/src/eos.cpp	93
/home/cgeroux/WORK/SPHERLS/src/eos.h	94
/home/cgeroux/WORK/SPHERLS/src/exception2.h	??
/home/cgeroux/WORK/SPHERLS/src/xmlFunctions.h	??
lataManipulation.cpp	95
lataManipulation.h	05
${ m lataMonitoring.cpp}$	14
lataMonitoring.h	19
${ m ileExists.h}$	??
${f global.cpp}$	<b>23</b>
${ m global.h}$	24
nain.cpp	$^{27}$
nain.h	<b>3</b> 0
m physEquations.cpp	32
${ m physEquations.h}$	89
${ m procTop.cpp}$	46
${ m procTop.h}$	47
${ m profile Data.h}$	48
ime.cpp	49
ime.h	50
m serguide.h	51
${f vatch zone.cpp}$	52
vatchzone.h	53

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## Chapter 6

## Class Documentation

## 6.1 eos Class Reference

#include <eos.h>

### **Public Member Functions**

- eos ()
- eos (int nNumT, int nNumRho)
- eos (const eos &ref)
- ~eos ()
- eos & operator= (const eos &eosRightSide)
- void readAscii (std::string sFileName)
- void readBobsAscii (std::string sFileName)
- void writeAscii (std::string sFileName)
- void readBin (std::string sFileName) throw (exception2)
- void writeBin (std::string sFileName)
- double dGetPressure (double dT, double dRho)
- double dGetEnergy (double dT, double dRho)
- double dGet Opacity (double dT, double dRho)
- double dDRhoDP (double dT, double dRho)
- double dSoundSpeed (double dT, double dRho)
- void getEKappa (double dT, double dRho, double &dE, double &dKappa)
- void getPEKappa (double dT, double dRho, double &dP, double &dE, double &dKappa)
- void getPEKappaGamma (double dT, double dRho, double &dP, double &dE, double &dKappa, double &dGamma)
- void getPEKappaGammaCp (double dT, double dRho, double &dP, double &dE, double &dKappa, double &dGamma, double &dCp)
- void getPKappaGamma (double dT, double dRho, double &dP, double &dKappa, double &dGamma)
- void gamma1DelAdC\_v (double dT, double dRho, double &dGamma1, double &dDelAd, double &dC\_v)
- void getPAndDRhoDP (double dT, double dRho, double &dP, double &dDRhoDP)

- void getEAndDTDE (double dT, double dRho, double &dE, double &dDTDE)
- void getDlnPDlnTDlnPDlnPDEDT (double dT, double dRho, double &dDlnPDlnT, double &dDlnPDlnRho, double &dDEDT)

## Public Attributes

- int nNumRho
- int nNumT
- double dXMassFrac
- double dYMassFrac
- double dLogRhoMin
- double dLogRhoDelta
- double dLogTMin
- double dLogTDelta
- double \*\* dLogP
- double \*\* dLogE
- double \*\* dLogKappa

## 6.1.1 Detailed Description

This class holds an equation of state as well as many functions useful for manipulating it

## 6.1.2 Constructor & Destructor Documentation

#### 6.1.2.1 eos::eos ()

Constructor, doesn't really do anything

References dLogE, dLogKappa, dLogP, nNumRho, and nNumT.

## 6.1.2.2 eos::eos (int nNumT, int nNumRho)

Constructor, allocates memory for the 2D arrays

#### Parameters:

- $\leftarrow nNumT$  number of temperatures in the equaiton of state table
- $\leftarrow nNumRho$  number of densities in the equaiton of state table

## 6.1.2.3 eos::eos (const eos & ref)

Copy constructor, simply constructs a new eos object from another eos object

References dLogE, dLogKappa, dLogP, dLogRhoDelta, dLogRhoMin, dLogTDelta, dLogTMin, dXMassFrac, dYMassFrac, nNumRho, and nNumT.

## 6.1.2.4 eos:: $\sim$ eos ()

Destructor, delets dynamic arrays

References dLogE, dLogKappa, dLogP, and nNumRho.

### 6.1.3 Member Function Documentation

## 6.1.3.1 eos & eos::operator= (const eos & eosRightSide)

Assignment operator, assigns one eos object to another.

References dLogE, dLogKappa, dLogP, dLogRhoDelta, dLogRhoMin, dLogTDelta, dLogTMin, nNumRho, and nNumT.

## 6.1.3.2 void eos::readAscii (std::string sFileName)

This fuction reads in an ascii file and stores it in the current object.

#### Parameters:

 $\leftarrow$  **sFileName** name of the equation of state file to read from.

References dLogE, dLogKappa, dLogP, dLogRhoDelta, dLogRhoMin, dLogTDelta, dLogTMin, dXMassFrac, dYMassFrac, nNumRho, and nNumT.

### 6.1.3.3 void eos::readBobsAscii (std::string sFileName)

This fuction reads in an ascii file and stores it in the current object. The ascii file is in Bob's format.

#### Parameters:

 $\leftarrow sFileName$  name of the equation of state file to read from.

References dLogE, dLogKappa, dLogP, dLogRhoDelta, dLogRhoMin, dLogTDelta, dLogTMin, dXMassFrac, dYMassFrac, nNumRho, and nNumT.

### 6.1.3.4 void eos::writeAscii (std::string sFileName)

This fuction writes the equation of state stored in the current object to an ascii file.

### Parameters:

 $\leftarrow$  sFileName name of the file to write the equation of state to.

References dLogE, dLogKappa, dLogP, dLogRhoDelta, dLogRhoMin, dLogTDelta, dLogTMin, dXMassFrac, dYMassFrac, nNumRho, and nNumT.

## 6.1.3.5 void eos::readBin (std::string sFileName) throw (exception2)

This fuction reads in a binary file and stores it in the current object.

#### Parameters:

 $\leftarrow$  **sFileName** name of the equation of state file to read from.

References dLogE, dLogKappa, dLogP, dLogRhoDelta, dLogRhoMin, dLogTDelta, dLogTMin, dXMassFrac, dYMassFrac, nNumRho, and nNumT.

Referenced by init().

## 6.1.3.6 void eos::writeBin (std::string sFileName)

This fuction writes the equation of state stored in the current object to a binary file.

### Parameters:

 $\leftarrow$  sFileName name of the file to write the equaiton of state to.

References dLogE, dLogKappa, dLogP, dLogRhoDelta, dLogRhoMin, dLogTDelta, dLogTMin, dXMassFrac, dYMassFrac, nNumRho, and nNumT.

## 6.1.3.7 double eos::dGetPressure (double dT, double dRho)

This function linearly interpolates the pressure to a given temperature and density. Note that both dT and dRho are not in log space.

#### Parameters:

- $\leftarrow dT$  temperature to interpolate to.
- $\leftarrow$  dRho density to interpolate to.

### Returns:

the interpolated pressure.

 $\label{eq:control_relation} References \ dLogP, \ dLogRhoDelta, \ dLogRhoMin, \ dLogTDelta, \ dLogTMin, \ nNumRho, \ and \ nNumT.$ 

```
\label{eq:referenced} Referenced & by & dImplicitEnergyFunction\_R(), & dImplicitEnergyFunction\_R\_LES(), & dImplicitEnergyFunction\_R\_LES\_SB(), & dImplicitEnergyFunction\_RT\_LES_SB(), & dImplicitEnergyFunction\_RT\_LES(), & dImplicitEnergyFunction\_RT\_LES\_SB(), & dImplicitEnergyFunction\_RT\_SB(), & dImplicitEnergyFunction\_RTP_LES(), & dImplicitEnergyFunction\_RTP\_LES(), & dImplicitEne
```

## 6.1.3.8 double eos::dGetEnergy (double dT, double dRho)

This function linearly interpolates the energy to a given temperature and and density. Note that both dT and dRho are not in log space.

#### Parameters:

 $\leftarrow dT$  temperature to interpolate to.

 $\leftarrow dRho$  density to interpolate to.

#### Returns:

the interpolated energy.

References dLogE, dLogRhoDelta, dLogRhoMin, dLogTDelta, dLogTMin, nNumRho, and nNumT.

```
Referenced
                       dImplicitEnergyFunction_R(),
               by
                                                          dImplicitEnergyFunction R -
LES(),
               dImplicitEnergyFunction R LES SB(),
                                                             dImplicitEnergyFunction -
R SB(),
                  dImplicitEnergyFunction RT(),
                                                         dImplicitEnergyFunction RT -
LES(),
            dImplicitEnergyFunction RT LES SB(),
                                                         dImplicitEnergyFunction RT -
            dImplicitEnergyFunction RTP(),
                                                  dImplicitEnergyFunction RTP LES(),
SB(),
dImplicitEnergyFunction RTP LES SB(), and dImplicitEnergyFunction RTP SB().
```

## 6.1.3.9 double eos::dGetOpacity (double dT, double dRho)

This function linearly interpolates the opacity to a given temperature and and density. Note that both dT and dRho are not in log space.

#### Parameters:

- $\leftarrow dT$  temperature to interpolate to.
- $\leftarrow dRho$  density to interpolate to.

#### Returns:

the interpolated opacity.

 $References\ dLogKappa,\ dLogRhoDelta,\ dLogRhoMin,\ dLogTDelta,\ dLogTMin,\ nNumRho,\ and\ nNumT.$ 

```
Referenced
                        dImplicitEnergyFunction R(),
                                                          dImplicitEnergyFunction R -
LES(),
               dImplicitEnergyFunction R LES SB(),
                                                             dImplicitEnergyFunction -
R SB(),
                  dImplicitEnergyFunction RT(),
                                                         dImplicitEnergyFunction RT
LES(),
                                                         dImplicitEnergyFunction RT -
            dImplicitEnergyFunction RT LES SB(),
            dImplicitEnergyFunction RTP(),
                                                  dImplicitEnergyFunction RTP LES(),
SB(),
dImplicitEnergyFunction RTP LES SB(), and dImplicitEnergyFunction RTP SB().
```

### 6.1.3.10 double eos::dDRhoDP (double dT, double dRho)

This function calculates the partial derivative of density w.r.t. pressure

### Parameters:

- $\leftarrow$  dT temperature at which the derivative is to be computed
- $\leftarrow dRho$  density at which the derivative is to be computed

## Returns:

the partial derivative of density w.r.t. pressure.

References dLogP, dLogRhoDelta, dLogRhoMin, dLogTDelta, dLogTMin, nNumRho, and nNumT.

## 6.1.3.11 double eos::dSoundSpeed (double dT, double dRho)

This function calculates the adiabatic sound speed

#### Parameters:

- $\leftarrow dT$  temperature at which the derivative is to be computed
- $\leftarrow$  dRho density at which the derivative is to be computed

### Returns:

the sound speed.

References dLogE, dLogP, dLogRhoDelta, dLogRhoMin, dLogTDelta, dLogTMin, nNumRho, and nNumT.

## 6.1.3.12 void eos::getEKappa (double dT, double dRho, double & dE, double & dKappa)

This function linearly interpolates the three dependent quantities (Pressure, Energy, Opacity) to a given temperature and density. Note that both dT and dRho are not in log space.

#### Parameters:

- $\leftarrow dT$  temperature to interpolate to.
- $\leftarrow dRho$  density to interpolate to.
- $\rightarrow$  **dE** energy at dT and dRho.
- $\rightarrow dKappa$  opacity at dT and dRho.

References dLogE, dLogKappa, dLogRhoDelta, dLogRhoMin, dLogTDelta, dLogTMin, nNumRho, and nNumT.

## 6.1.3.13 void eos::getPEKappa (double dT, double dRho, double & dP, double & dKappa)

This function linearly interpolates the three dependent quantities (Pressure, Energy, Opacity) to a given temperature and density. Note that both dT and dRho are not in log space.

#### Parameters:

- $\leftarrow dT$  temperature to interpolate to.
- $\leftarrow dRho$  density to interpolate to.
- $\rightarrow$  **dP** pressure at dT and dRho.
- $\rightarrow dE$  energy at dT and dRho.
- $\rightarrow dKappa$  opacity at dT and dRho.

References dLog<br/>E, dLog Kappa, dLog P, dLog Rho<br/>Delta, dLog RhoMin, dLog TDelta, dLog TMin, nNum<br/>Rho, and nNum<br/>T.

## 6.1.3.14 void eos::getPEKappaGamma (double dT, double dRho, double & dP, double & dE, double & dKappa, double & dGamma)

This function linearly interpolates the energy and opacity to a given temperature and density. Note that both dT and dRho are not in log space.

#### Parameters:

- $\leftarrow dT$  temperature to interpolate to.
- $\leftarrow dRho$  density to interpolate to.
- $\rightarrow$  **dP** pressure at dT and dRho.
- $\rightarrow dE$  energy at dT and dRho.
- $\rightarrow dKappa$  opacity at dT and dRho.
- $\rightarrow dGamma$  adiabatic index at dT and dRho.

References dLogE, dLogKappa, dLogP, dLogRhoDelta, dLogRhoMin, dLogTDelta, dLogTMin, nNumRho, and nNumT.

Referenced by calNewPEKappaGamma\_TEOS(), and calOldPEKappaGamma\_TEOS().

## 6.1.3.15 void eos::getPEKappaGammaCp (double dT, double dRho, double & dP, double & dE, double & dKappa, double & dGamma, double & dCp)

This function linearly interpolates the energy and opacity to a given temperature and density. Note that both dT and dRho are not in log space.

#### Parameters:

- $\leftarrow dT$  temperature to interpolate to.
- $\leftarrow dRho$  density to interpolate to.
- $\rightarrow$  **dP** pressure at dT and dRho.
- $\rightarrow dE$  energy at dT and dRho.
- $\rightarrow dKappa$  opacity at dT and dRho.
- $\rightarrow dGamma$  adiabatic index at dT and dRho.
- $\rightarrow dCp$  specific heat at constant pressure at dT and dRho.

References dLogE, dLogKappa, dLogP, dLogRhoDelta, dLogRhoMin, dLogTDelta, dLogTMin, nNumRho, and nNumT.

## 6.1.3.16 void eos::getPKappaGamma (double dT, double dRho, double & dP, double & dKappa, double & dGamma)

This function linearly interpolates the energy and opacity to a given temperature and density. Note that both dT and dRho are not in log space.

#### Parameters:

- $\leftarrow dT$  temperature to interpolate to.
- $\leftarrow$  dRho density to interpolate to.
- $\rightarrow$  dP pressure at dT and dRho.

- $\rightarrow dKappa$  opacity at dT and dRho.
- $\rightarrow dGamma$  adiabatic index at dT and dRho.

References dLogE, dLogKappa, dLogP, dLogRhoDelta, dLogRhoMin, dLogTDelta, dLogTMin, nNumRho, and nNumT.

Referenced by calNewTPKappaGamma\_TEOS().

## 6.1.3.17 void eos::gamma1DelAdC\_v (double dT, double dRho, double & dGamma1, double & dDelAd, double & dC v)

This function calculates gammal and the adiabatic gradient

#### Parameters:

- $\leftarrow$  dT temperature at which the derivative is to be computed
- $\leftarrow dRho$  density at which the derivative is to be computed
- $\rightarrow$  dGamma1 gamma1
- ightarrow dDelAd adiabatic gradient
- ightarrow dC v specific heat at constant volume

References dLogE, dLogRhoDelta, dLogRhoMin, dLogTDelta, dLogTMin, nNumRho, and nNumT.

## 6.1.3.18 void eos::getPAndDRhoDP (double dT, double dRho, double & dP, double & dDRhoDP)

This function calculates the partial derivative of density w.r.t. pressure and the pressure

### Parameters:

- $\leftarrow dT$  temperature at which the derivative is to be computed
- $\leftarrow dRho$  density at which the derivative is to be computed
- ightarrow dP pressure at dT and dRho
- ightarrow dDRhoDP derivative of density w.r.t. pressure at conatant temperature

References dLogP, dLogRhoDelta, dLogRhoMin, dLogTDelta, dLogTMin, nNumRho, and nNumT.

## 6.1.3.19 void eos::getEAndDTDE (double dT, double dRho, double & dE, double & dDTDE)

This function calculates the partial derivative of temperature w.r.t. energy and the energy

## Parameters:

- $\leftarrow$  dT temperature at which the derivative is to be computed
- $\leftarrow$  dRho density at which the derivative is to be computed
- $\rightarrow dE$  energy at dT and dRho
- $\rightarrow dDTDE$  derivative of temperature w.r.t. energy at constant density

References dLogE, dLogRhoDelta, dLogRhoMin, dLogTDelta, dLogTMin, nNumRho, and nNumT.

Referenced by calNewTPKappaGamma TEOS().

## 6.1.3.20 void eos::getDlnPDlnTDlnPDlnPDEDT (double dT, double dRho, double & dDlnPDlnT, double & dDlnPDlnRho, double & dDEDT)

This function calculates various partial derivatives

#### Parameters:

- $\leftarrow dT$  temperature at which the derivative is to be computed
- $\leftarrow$  dRho density at which the derivative is to be computed
- $\rightarrow dD \ln PD \ln T$  derivative of ln(P) w.r.t. ln(T)
- $\rightarrow dDlnPDlnRho$  derivative of ln(P) w.r.t. ln(Rho)
- $\rightarrow dDEDT$  derivative of temperature w.r.t. energy at constant density

References dLogE, dLogRappa, dLogP, dLogRhoDelta, dLogRhoMin, dLogTDelta, dLogTMin, nNumRho, and nNumT.

#### 6.1.4 Member Data Documentation

## 6.1.4.1 int eos::nNumRho

Number of densities in the equation of state table

#### 6.1.4.2 int eos::nNumT

Number of temperatures in the equation of state table

Referenced by dDRhoDP(), dGetEnergy(), dGetOpacity(), dGetPressure(), dSoundSpeed(), eos(), gamma1DelAdC\_v(), getDlnPDlnTDlnPDlnPDEDT(), getEAndDTDE(), getEKappa(), getPAndDRhoDP(), getPEKappa(), getPEKappaGamma(), getPEKappaGammaCp(), getPKappaGamma(), operator=(), readAscii(), readBin(), readBobsAscii(), writeAscii(), and writeBin().

### 6.1.4.3 double eos::dXMassFrac

Hydrogen mass fraction of the composition used to generate the equation of state table. Referenced by eos(), readAscii(), readBin(), readBobsAscii(), writeAscii(), and writeBin().

#### 6.1.4.4 double eos::dYMassFrac

Helium mass fraction of the composition used to generate the equation of state table.

Referenced by eos(), readAscii(), readBin(), readBobsAscii(), writeAscii(), and writeBin().

## 6.1.4.5 double eos::dLogRhoMin

Minimum density of the table in log10.

Referenced by dDRhoDP(), dGetEnergy(), dGetOpacity(), dGetPressure(), dSoundSpeed(), eos(), gamma1DelAdC\_v(), getDlnPDlnTDlnPDlnPDEDT(), getEAndDTDE(), getEKappa(), getPAndDRhoDP(), getPEKappa(), getPEKappaGamma(), getPEKappaGammaCp(), getPKappaGamma(), operator=(), readAscii(), readBin(), readBobsAscii(), writeAscii(), and writeBin().

### 6.1.4.6 double eos::dLogRhoDelta

Increment of the density between table entries in log10.

Referenced by dDRhoDP(), dGetEnergy(), dGetOpacity(), dGetPressure(), dSoundSpeed(), eos(), gamma1DelAdC\_v(), getDlnPDlnTDlnPDlnPDEDT(), getEAndDTDE(), getEKappa(), getPAndDRhoDP(), getPEKappa(), getPEKappaGamma(), getPEKappaGamma(), operator=(), readAscii(), readBin(), readBobsAscii(), writeAscii(), and writeBin().

## 6.1.4.7 double eos::dLogTMin

Minimum temperature of the table in log10.

Referenced by dDRhoDP(), dGetEnergy(), dGetOpacity(), dGetPressure(), dSoundSpeed(), eos(), gamma1DelAdC\_v(), getDlnPDlnTDlnPDlnPDEDT(), getEAndDTDE(), getEKappa(), getPAndDRhoDP(), getPEKappa(), getPEKappaGamma(), getPEKappaGammaCp(), getPKappaGamma(), operator=(), readAscii(), readBin(), readBobsAscii(), writeAscii(), and writeBin().

### 6.1.4.8 double eos::dLogTDelta

Increment of the temperature between table entries in log10.

 $\label{eq:control_read_equation} Referenced by dDRhoDP(), dGetEnergy(), dGetOpacity(), dGetPressure(), dSoundSpeed(), eos(), gamma1DelAdC_v(), getDlnPDlnTDlnPDlnPDEDT(), getEAndDTDE(), getEKappa(), getPAndDRhoDP(), getPEKappa(), getPEKappaGamma(), getPEKappaGammaCp(), getPKappaGamma(), operator=(), readAscii(), readBin(), readBobsAscii(), writeAscii(), and writeBin().$ 

## 6.1.4.9 double\*\* eos::dLogP

2D array of log10 pressures. dLogP[i][j] gives the log10 pressure at log10 density of eos::dLogRhoDelta\*i+eos::dLogRhoMin, and at log10 temperature of eos::dLogTDelta\*j+eos::dLogTMin.

Referenced by dDRhoDP(), dGetPressure(), dSoundSpeed(), eos(), gamma1DelAdC\_v(), getDlnPDlnTDlnPDEDT(), getPAndDRhoDP(), getPEKappa(), getPEKappaGamma(), getPEKappaGammaCp(), getPKappaGamma(), operator=(), readAscii(), readBin(), readBobsAscii(), writeAscii(), writeBin(), and  $\sim$ eos().

#### 6.1.4.10 double\*\* eos::dLogE

2D array of log10 energies. dLogE[i][j] gives the log10 energy at log10 density of eos::dLogRhoDelta\*i+eos::dLogRhoMin, and at log10 temperature of eos::dLogTDelta\*j+eos::dLogTMin.

Referenced by dGetEnergy(), dSoundSpeed(), eos(), gamma1DelAdC\_v(), getDlnPDlnTDlnPDlnPDEDT(), getEAndDTDE(), getEKappa(), getPEKappa(), getPEKappaGamma(), getPEKappaGammaCp(), getPKappaGamma(), operator=(), readAscii(), readBin(), readBobsAscii(), writeAscii(), writeBin(), and  $\sim$ eos().

### 6.1.4.11 double\*\* eos::dLogKappa

Referenced by dGetOpacity(), eos(), getDlnPDlnTDlnPDlnPDEDT(), getEKappa(), getPEKappaGamma(), getPEKappaGammaCp(), getPKappaGamma(), operator=(), readAscii(), readBin(), readBobsAscii(), writeAscii(), writeBin(), and ~eos().

The documentation for this class was generated from the following files:

- /home/cgeroux/WORK/SPHERLS/src/eos.h
- /home/cgeroux/WORK/SPHERLS/src/eos.cpp

## 6.2 Functions Class Reference

#include <global.h>

#### Public Member Functions

• Functions ()

### Public Attributes

- void(\* fpCalculateNewVelocities )(Grid &, Parameters &, Time &, ProcTop &)
- void(\* fpCalculateNewGridVelocities )(Grid &, Parameters &, Time &, ProcTop &, MessPass &)
- void(\* fpCalculateNewRadii )(Grid &, Time &)
- void(\* fpCalculateNewDensities)(Grid &, Parameters &, Time &, ProcTop &)
- void(\* fpCalculateNewEnergies )(Grid &, Parameters &, Time &, ProcTop &)
- void(\* fpCalculateDeltat )(Grid &, Parameters &, Time &, ProcTop &)
- void(\* fpCalculateAveDensities )(Grid &)
- void(\* fpCalculateNewEOSVars )(Grid &, Parameters &)
- void(\* fpCalculateNewAV )(Grid &, Parameters &)
- void(\* fpModelWrite)(std::string sFileName, ProcTop &, Grid &, Time &, Parameters &)
- void(\* fpWriteWatchZones )(Output &, Grid &, Parameters &, Time &, ProcTop &)
- void(\* fpUpdateLocalBoundaryVelocitiesNewGrid )(ProcTop &, MessPass &, Grid &)
- void(\* fpImplicitSolve )(Grid &, Implicit &, Parameters &, Time &, ProcTop &, MessPass &, Functions &)
- void(\* fpCalculateNewEddyVisc )(Grid &, Parameters &)

### 6.2.1 Detailed Description

This class holds function pointers used to indicate the functions which should be used to calculate the various needed quantities. These functions can be different from processor to processor. For example ProcTop::nRank=0 processor will have only 1D verions of the conservation equations, while the rest of the processors will have 3D versions. These functions will also change depending on what kind of model is being calculated and the number of dimensions the calculation includes.

#### 6.2.2 Constructor & Destructor Documentation

## 6.2.2.1 Functions::Functions ()

Constructor for the class Functions.

References fpCalculateAveDensities, fpCalculateDeltat, fpCalculateNewAV, fpCalculateNewDensities, fpCalculateNewEnergies, fpCalculateNewEOSVars, fpCalculateNewGridVelocities, fpCalculateNewRadii, and fpCalculateNewVelocities.

## 6.2.3 Member Data Documentation

## 6.2.3.1 void(\* Functions::fpCalculateNewVelocities)(Grid &, Parameters &, Time &, ProcTop &)

Function pointer to the function used to calculate new velocities.

Referenced by Functions(), main(), and setMainFunctions().

## 6.2.3.2 void(\* Functions::fpCalculateNewGridVelocities)(Grid &, Parameters &, Time &, ProcTop &, MessPass &)

Function pointer to the function used to calculate new grid velocities.

Referenced by Functions(), main(), and setMainFunctions().

## 6.2.3.3 void(\* Functions::fpCalculateNewRadii)(Grid &, Time &)

Functin pointer to the function used to calculate new radii.

Referenced by Functions(), main(), and setMainFunctions().

## 6.2.3.4 void(\* Functions::fpCalculateNewDensities)(Grid &, Parameters &, Time &, ProcTop &)

Function pointer to the function used to calculate the new densities.

Referenced by Functions(), main(), and setMainFunctions().

## 6.2.3.5 void(\* Functions::fpCalculateNewEnergies)(Grid &, Parameters &, Time &, ProcTop &)

Function pointer to the function used to calculate the new energies.

Referenced by Functions(), main(), and setMainFunctions().

## 6.2.3.6 void(\* Functions::fpCalculateDeltat)(Grid &, Parameters &, Time &, ProcTop &)

Function pointer to the function used to calculate the new time step.

Referenced by Functions(), main(), and setMainFunctions().

### 6.2.3.7 void(\* Functions::fpCalculateAveDensities)(Grid &)

Function pointer to the function used to calculate the new average density.

Referenced by Functions(), main(), and setMainFunctions().

## 6.2.3.8 void(\* Functions::fpCalculateNewEOSVars)(Grid &, Parameters &)

Function pointer to the function used to calculate the new variables depending on the equation of state.

Referenced by Functions(), main(), and setMainFunctions().

## 6.2.3.9 void(\* Functions::fpCalculateNewAV)(Grid &, Parameters &)

Function pointer to the function used to calculate new Artificial viscosity.

Referenced by Functions(), main(), and setMainFunctions().

## 6.2.3.10 void(\* Functions::fpModelWrite)(std::string sFileName, ProcTop &, Grid &, Time &, Parameters &)

Function pointer to the function used to write out models.

Referenced by fin(), main(), and setMainFunctions().

## 6.2.3.11 void(\* Functions::fpWriteWatchZones)(Output &, Grid &, Parameters &, Time &, ProcTop &)

Function pointer to the function that is used to write out watch zone files

Referenced by main(), and setMainFunctions().

## 6.2.3.12 void(\* Functions::fpUpdateLocalBoundaryVelocitiesNewGrid)(ProcTop &, MessPass &, Grid &)

Function pointer to the faction that is used to update velocities across boundaries.

Referenced by main(), and setMainFunctions().

## 6.2.3.13 void(\* Functions::fpImplicitSolve)(Grid &, Implicit &, Parameters &, Time &, ProcTop &, MessPass &, Functions &)

Funciton pointer to the function that is used to implicitly solve for the temperature, then uses the equation of state to solve for energy, opacity, and pressure.

Referenced by main(), and setMainFunctions().

#### 6.2.3.14 void(\* Functions::fpCalculateNewEddyVisc)(Grid &, Parameters &)

Function pointer to the function that is used to calculate the new eddy viscosity.

Referenced by main(), and setMainFunctions().

The documentation for this class was generated from the following files:

- global.h
- global.cpp

## 6.3 Global Class Reference

#include <global.h>

## **Public Member Functions**

• Global ()

## Public Attributes

- ProcTop procTop
- $\bullet \ \operatorname{MessPass} \ \operatorname{messPass}$
- Grid grid
- Time time
- Parameters parameters
- Output output
- Performance performance
- Functions functions
- Implicit implicit

## 6.3.1 Detailed Description

This class is simply a class that holds the other classes.

## 6.3.2 Constructor & Destructor Documentation

## 6.3.2.1 Global::Global ()

Constructor for the class Global.

## 6.3.3 Member Data Documentation

## 6.3.3.1 ProcTop Global::procTop

An instance of the ProcTop class.

Referenced by main().

#### 6.3.3.2 MessPass Global::messPass

An instance of the MessPass class.

Referenced by main().

## 6.3.3.3 Grid Global::grid

An instance of the Grid class.

Referenced by main().

## 6.3.3.4 Time Global::time

An instance of the Time class.

Referenced by main().

## 6.3.3.5 Parameters Global::parameters

An instance of the Parameters class.

Referenced by main().

## 6.3.3.6 Output Global::output

An instance of the Output class.

Referenced by main().

### 6.3.3.7 Performance Global::performance

An instance of the Performance class.

Referenced by main().

## 6.3.3.8 Functions Global::functions

An instance of the Functions class.

Referenced by main().

## 6.3.3.9 Implicit Global::implicit

An instance of the Implicit class.

Referenced by main().

The documentation for this class was generated from the following files:

- global.h
- global.cpp

## 6.4 Grid Class Reference

#include <global.h>

## **Public Member Functions**

• Grid ()

## Public Attributes

- $\bullet$  int nM
- int nTheta
- int nPhi
- int nDM
- int nR
- int nD
- int nU
- int nU0
- int nV
- int nW
- int nT
- int nE
- int nP
- int nKappa
- int nGamma
- int nDenAve
- int nQ0
- int nQ1
- int nQ2
- $\bullet$  int nDTheta
- int nDPhi
- $\bullet$  int nSinThetaIJK
- int nSinThetaIJp1halfK
- int nCotThetaIJp1halfK
- int nCotThetaIJK
- int nDCosThetaIJK
- int nEddyVisc
- int nDonorCellFrac
- int nNumDims
- int nNumVars
- int nNumIntVars
- int nNum1DZones
- int nNumGhostCells
- $\bullet$  int \* nGlobalGridDims
- int \*\* nVariables
- $\bullet$  int \*\*\* nLocalGridDims

- double \*\*\*\* dLocalGridNew
- double \*\*\*\* dLocalGridOld
- int \*\* nStartUpdateExplicit
- int \*\* nEndUpdateExplicit
- $\bullet \ \, int \, ** \, nStartUpdateImplicit \\$
- int \*\* nEndUpdateImplicit
- $\bullet$  int \*\*\* nStartGhostUpdateExplicit
- $\bullet \ \ int \ *** nEndGhostUpdateExplicit$
- $\bullet$  int \*\*\* nStartGhostUpdateImplicit
- $\bullet \ \ int \ *** nEndGhostUpdateImplicit$
- $\bullet$  int \* nCenIntOffset
- int nGlobalGridPositionLocalGrid [3]

## 6.4.1 Detailed Description

This class manages information which pertains to grid data.

External variables used with Gamma Law (GL) gas equaiton of state and their array indexes:

1D (nNumVars=7)			2D (nNumVa	rs=9)	3D (nNumVars=11)					
Variable	Index		Variable	Index	Variable	Index				
nM	0		nM	0	nM	0				
nDM	1		nTheta	1	nTheta	1				
nR	2		nDM	2	nPhi	2				
nD	3		nR	3	nDM	3				
nU	4		nD	4	nR	4				
nU0	5		nU	5	nD	5				
nE	6		nU0	6	nU	6				
	•		nV	7	nU0	7				
			nE	8	nV	8				
			<u>,                                    </u>	•	 nW	9				
					nE	10				
						·				

External variables used with Tabulated Equation Of State (TEOS) and their array indexes:

1D (nNumVa	rs=7)	2D (nNu	mVars=9)	3D (nNumVars=11)					
Variable	Index	Variabl	e Index	Variable	Index				
nM	0	nM	0	nM	0				
nDM	1	nTheta	1	nTheta	1				
nR	2	nDM	2	nPhi	2				
nD	3	nR	3	nDM	3				
nU	4	nD	4	nR	4				
nU0	5	nU	5	nD	5				
nT	6	nU0	6	nU	6				
<u>-</u>	•	nV	7	nU0	7				
		nT	8	nV	8				
			•	nW	9				
				nT	10				
				<u> </u>					

Internal variables with GL gas equation of state:

1D (nNumIntVars=2)		Т	2D (nNumIntVars=8)	
		T		
Variable	Index		Variable	Index
nP	nNumVars+0		nP	nNumVars+0
nQ0	nNumVars+1		nQ0	nNumVars+1
			nDenAve	nNumVars+2
			nDCosThetaIJK	nNumVars+3
			nQ1	$ m nNumVars{+}4$
			nDTheta	nNumVars+5
			nSinThetaIJK	nNumVars+6
				nNumVars+7
			nSinThetaIJp1halfK	
3D (nNumIntVars=12)		$\dagger$		
	<u> </u>	$^{\dagger}$		
Variable	Index			
nP	nNumVars+0			
nQ0	nNumVars+1			
nDenAve	nNumVars+2			
nDPhi	nNumVars+3			
${ m nDCosThetaIJK}$	nNumVars+4			
nQ1	m nNumVars+5			
nDTheta	nNumVars+6			
nSinThetaIJK	nNumVars+7			
	nNumVars+8			
nSinThetaIJp1halfK				
${ m nCotThetaIJK}$	nNumVars+9			
	nNumVars+10			
nCotThetaIJp1halfK				
nQ2	nNumVars+11			

Internal variables with TEOS:

1D (nNumIntVars	s=5)	2D (nNumInt Va	urs=11)
Variable	Index	Variable	Index
nP	nNumVars+0	nP	nNumVars+0
nQ0	$ m nNumVars{+}1$	nQ0	nNumVars+1
nE	$ m nNumVars{+}2$	nDenAve	nNumVars+2
nKappa	nNumVars+3	nDCosThetaIJ	K nNumVars+3
nGamma	$ m nNumVars{+}4$	nE	$ m nNumVars{+}4$
	•	nKappa	nNumVars+5
		nGamma	nNumVars+6
		nQ1	nNumVars+7
		nDTheta	nNumVars+8
		nSinThetaIJK	nNumVars+9
			nNumVars+10
		nSinThetaIJp1	halfK
3D (nNumIntVar	s=15)		
Variable	Index	$\neg$	
nP	nNumVars+0	-	
nQ0	nNumVars+1	$\dashv$	
nDenAve	$\frac{\text{nNumVars}+2}{\text{nNumVars}+2}$	$\dashv$	
nDPhi	nNumVars+3	$\dashv$	
nDCosThetaIJK		<del> </del>	
nE	nNumVars+5	-	
nKappa	nNumVars+6	<b>-</b>	
nGamma	nNumVars+7	<b>- </b>	
nQ1	nNumVars+8	7	
nDTheta	nNumVars+9	71	
nSinThetaIJK	nNumVars+10	71	
nSinThetaIJp1h	nNumVars+11		
nCotThetaIJK	nNumVars+12	-	
noot metaijK	nNumVars+12 nNumVars+13	-	
nCotThetaIJp1l	nalfK		
nQ2	nNumVars+14		

The variable indexes are set in modelRead based on the input model.

## 6.4.2 Constructor & Destructor Documentation

### 6.4.2.1 Grid::Grid ()

sets how many zones out from the 1D-multi-D boundary that theta/phi velocities are not updated and thus kept at zero. Constructor for the class Grid.

References dLocalGridNew, dLocalGridOld, nCenIntOffset, nCotThetaIJK, nCotThetaIJp1halfK, nD, nDCosThetaIJK, nDenAve, nDM, nDonorCellFrac, nDPhi, nDTheta, nE, nEddyVisc, nEndGhostUpdateExplicit, nEndGhostUpdateImplicit, nEndUpdateExplicit, nEndUpdateImplicit, nGamma, nGlobalGridDims, nKappa, nLocalGridDims, nM, nP, nPhi, nQ0, nQ1, nQ2, nR, nSinThetaIJK, nSinThetaIJp1halfK, nStartGhostUpdateExplicit, nStartGhostUpdateImplicit, nStartGhostUpdateImplic

plicit, nStartUpdateExplicit, nStartUpdateImplicit, nT, nTheta, nU, nU0, nV, nVariables, and nW.

### 6.4.3 Member Data Documentation

## 6.4.3.1 int Grid::nM

Index of  $M_r$  independent variable in grid Grid::dLocalGridOld and Grid::dLocalGridNew. This is an external grid variable included in the count Grid::nNumVars. This is an independent grid variable.

 $\label{eq:referenced_by_calNewU_R(), calNewU_R_LES(), calNewU_RT(), calNewU_RT_-LES(), calNewU_RTP(), calNewU_RTP_LES(), dImplicitEnergyFunction_RT_LES(), dImplicitEnergyFunction_RTP_LES(), dImplicitEnergyFunction_RTP_LES_SB(), Grid(), and modelRead().}$ 

### 6.4.3.2 int Grid::nTheta

Index of  $\theta$  independent variable in grid Grid::dLocalGridOld and Grid::dLocalGridNew. This is an external grid variable included in the count Grid::nNumVars. This is an independent grid variable.

Referenced by Grid(), initInternalVars(), and modelRead().

### 6.4.3.3 int Grid::nPhi

Index of  $\phi$  independent variable in grid Grid::dLocalGridOld and Grid::dLocalGridNew. This is an external grid variable included in the count Grid::nNumVars. This is an independent grid variable.

Referenced by Grid(), initInternalVars(), and modelRead().

#### 6.4.3.4 int Grid::nDM

Index of  $\delta M$  in grids, Grid::dLocalGridOld and Grid::dLocalGridNew. This is an external grid variable included in the count Grid::nNumVars

calNewE R NA(), Referenced by calNewE R AD(),calNewE R NA LES(),  ${\tt calNewE\_RT\_NA()}, \quad {\tt calNewE\_RT\_NA()}, \quad {\tt calNewE\_RT\_NA\_LES()}, \quad {\tt calNewE\_RTP\_-level}, \quad {\tt calNewE\_RTP\_-level$ calNewE RTP NA(), calNewE RTP NA LES(), calNewU R(), AD(),calNewU -R LES(),calNewU RT(), calNewU RT LES(), calNewU RTP(), calNewU -RTP LES(), calNewV RT(), calNewV RT LES(), calNewV RTP(), calNewV calNewW RTP(), calNewW RTP LES(),  $RTP_LES()$ , dImplicitEnergyFunction dImplicitEnergyFunction R LES(), dImplicitEnergyFunction R LES SB(), dImplicitEnergyFunction R SB(), dImplicitEnergyFunction RT(), dImplicitEnergyFunction -RT LES(), dImplicitEnergyFunction RT LES SB(), dImplicitEnergyFunction -RT SB(),dImplicitEnergyFunction RTP(), dImplicitEnergyFunction RTP LES(), dImplicitEnergyFunction RTP LES SB(), dImplicitEnergyFunction RTP SB(), Grid(), modelRead(), writeWatchZones R GL(), writeWatchZones R TEOS(), writeWatchZones RT -GL(), writeWatchZones RT TEOS(), writeWatchZones RTP GL(), and writeWatchZones -RTP TEOS().

#### 6.4.3.5 int Grid::nR

Index of r in grids, Grid::dLocalGridOld and Grid::dLocalGridNew. This is an external grid variable included in the count Grid::nNumVars

Referenced by average3DTo1DBoundariesNew(), average3DTo1DBoundariesOld(), calDelt -R GL()calDelt R TEOS(), calDelt RT GL(), calDelt RT TEOS(),calDelt -RTP GL(), calDelt\_RTP\_TEOS(), calNewD\_R(), calNewD RT(), calNewD RTP(), calNewDenave RTP(), calNewE R AD(), calNewE R NA(), calNewDenave RT(), calNewE R NA LES(), calNewE RT AD(), calNewE RT NA(), calNewE RT calNewE RTP AD(), calNewE RTP NA(), calNewE RTP NA LES(), NA LES(), calNewEddyVisc R CN(), calNewEddyVisc R SM(), calNewEddyVisc RT CN(), calNewEddyVisc RT SM(), calNewEddyVisc RTP CN(), calNewEddyVisc RTP SM(), calNewQ0 R GL(),calNewQ0 R TEOS(), calNewQ0Q1 RT GL(), calNewQ0Q1 - $RT \quad TEOS(), \quad calNewQ0Q1Q2 \quad RTP \quad GL(), \quad calNewQ0Q1Q2\_RTP\_TEOS(),$ calNewR(), calNewU0 R(), calNewU0 RT(), calNewU0 RTP(), calNewU R(), calNewU R LES(), calNewU RT(), calNewU RT LES(), calNewU RTP(), calNewU RTP LES(), calNewV -RT(), calNewV RT LES(), calNewV RTP(), calNewV RTP LES(), calNewW RTP(), calNewW RTP LES(), calOldDenave RT(),calOldDenave RTP(), calOldEddyVisc calOldEddyVisc RT CN(), calOldEddyVisc -R CN(),calOldEddyVisc R SM(), RT SM(),calOldEddyVisc RTP CN(), calOldEddyVisc RTP SM(), calOldQ0 calOldQ0 R TEOS(), calOldQ0Q1\_RT\_GL(), R GL()calOldQ0Q1 RT TEOS(), calOldQ0Q1Q2 RTP GL(), calOldQ0Q1Q2 RTP TEOS(), dImplicitEnergyFunction dImplicitEnergyFunction R LES(), dImplicitEnergyFunction R LES SB(), dImplicitEnergyFunction R SB(), dImplicitEnergyFunction RT(), dImplicitEnergyFunction -RT LES(), dImplicitEnergyFunction RT LES SB(), dImplicitEnergyFunction -RT SB(), dImplicitEnergyFunction RTP(), dImplicitEnergyFunction RTP LES(), dImplicitEnergyFunction RTP LES SB(), dImplicitEnergyFunction RTP SB(), modelRead(). writeWatchZones R GL(), writeWatchZones R TEOS(). writeWatchZones RT GL(), writeWatchZones RT TEOS(), writeWatchZones RTP GL(), and writeWatchZones RTP TEOS().

### 6.4.3.6 int Grid::nD

Index of  $\rho$  in grids, Grid::dLocalGridOld and Grid::dLocalGridNew. This is an external grid variable included in the count Grid::nNumVars

calDelt R TEOS(), calDelt RT GL(), Referenced by calDelt R GL(), RT TEOS(), calDelt RTP GL(), calDelt RTP TEOS(), calNewD R(), calNewD -RT(), calNewD RTP(), calNewDenave R(), calNewDenave RTP(), calNewDenave RTP(), calNewE R AD(), calNewE R NA(), calNewE R NA LES(), calNewE RT AD(), calNewE\_RT\_NA(), calNewE\_RT\_NA\_LES(), calNewE\_RTP\_AD(), calNewE\_RTP\_-NA(), calNewE RTP NA LES(), calNewEddyVisc R SM(), calNewEddyVisc RT SM(), calNewEddyVisc RTP SM(), calNewP GL(), calNewPEKappaGamma TEOS(), calNewQ0 calNewQ0 R TEOS(), calNewQ0Q1 RT GL(), calNewQ0Q1 RT TEOS(), calNewQ0Q1Q2 RTP GL(),  $calNewQ0Q1Q2\_RTP\_TEOS(),$ calNewTPKappaGamma -TEOS(), calNewU0 R(), calNewU0 RT(), calNewU0 RTP(), calNewU\_R(), calNewU\_calNewU RT(), calNewU RT LES(), calNewU RTP(), calNewU RTP -R LES(),calNewV RT(), calNewV RT LES(), calNewV RTP(), calNewV RTP LES(), calNewW RTP LES(), calOldDenave R(), calOldDenave -LES(), calNewW RTP(), calOldEddyVisc R SM(), calOldEddyVisc RT SM(), calOldDenave RTP(), calOldEddyVisc RTP SM(), calOldP GL(), calOldPEKappaGamma TEOS(), calOldQ0 calOldQ0 R TEOS(), calOldQ0Q1 RT GL(), calOldQ0Q1 RT TEOS(), calOldQ0Q1Q2 RTP TEOS(), calOldQ0Q1Q2 RTP GL(), dImplicitEnergyFunction - R()dImplicitEnergyFunction R LES(), dImplicitEnergyFunction R LES SB(), dImplicitEnergyFunction R SB(), dImplicitEnergyFunction RT(), dImplicitEnergyFunction -RT LES(), dImplicitEnergyFunction RT LES SB(), dImplicitEnergyFunction -RT SB(), dImplicitEnergyFunction RTP(), dImplicitEnergyFunction RTP LES(), dImplicitEnergyFunction RTP LES SB(), dImplicitEnergyFunction RTP SB(), initDonorFracAndMaxConVel R GL(), initDonorFracAndMaxConVel R TEOS(), initDonorFracAndMaxConVel RT GL(), initDonorFracAndMaxConVel RT TEOS(), initDonorFracAndMaxConVel RTP GL(), initDonorFracAndMaxConVel RTP TEOS(), initWatchZones(), modelRead(), setupLocalGrid(), writeWatchZones R GL(), main(), writeWatchZones R TEOS(), writeWatchZones RT GL(), writeWatchZones RT TEOS(), writeWatchZones RTP GL(), and writeWatchZones RTP TEOS().

### 6.4.3.7 int Grid::nU

Index of u in grids, Grid::dLocalGridOld and Grid::dLocalGridNew. This is an external grid variable included in the count Grid::nNumVars

Referenced by calDelt R GL(), calDelt R TEOS(), calDelt RT GL(), calDelt RT - $TEOS(), \quad calDelt\_RTP\_GL(), \quad calDelt\_RTP\_TEOS(), \quad calNewD\_R(), \quad calNewD\_RT(),$ calNewD RTP(), calNewE R AD(), calNewE R NA(), calNewE R NA LES(), calNewE RT AD(), calNewE RT NA(), calNewE RT NA LES(), calNewE RTP calNewE RTP NA LES(), calNewE RTP NA(), calNewEddyVisc R SM(),  $calNewEddyVisc\_RT\_SM(),\ calNewEddyVisc\_RTP\_SM(),\ calNewQ0\_R\_GL(),\ calNewQ0\_-R_GL(),\ calNewQ0\_-R_GL(),\ calNewRDR_GL(),\ calNewRDR_GL(),\$ R TEOS(), calNewQ0Q1 RT GL(), calNewQ0Q1 RT TEOS(), calNewQ0Q1Q2 RTP -GL(), calNewQ0Q1Q2 RTP TEOS(), calNewU0 R(), calNewU0 RT(), calNewU0 RTP(), calNewU R(), calNewU R LES(), calNewU RT(), calNewU RT LES(), calNewU -RTP(), calNewU RTP LES(), calNewV RT(), calNewV RT LES(), calNewV RTP(), calNewV RTP LES(), calNewW RTP(), calNewW RTP LES(), calOldEddvVisc calOldEddyVisc RT SM(), R SM(),calOldEddyVisc RTP SM(), calOldQ0 R -GL(), calOldQ0 R TEOS(), calOldQ0Q1 RT GL(), calOldQ0Q1 RT TEOS(), calOldQ0Q1Q2 RTP GL(), calOldQ0Q1Q2 RTP TEOS(), dImplicitEnergyFunction dImplicitEnergyFunction R LES(), dImplicitEnergyFunction R LES SB(), dImplicitEnergyFunction R SB(), dImplicitEnergyFunction RT(), dImplicitEnergyFunction -RT LES(),dImplicitEnergyFunction RT LES SB(), dImplicitEnergyFunction -RT SB(), dImplicitEnergyFunction RTP(), dImplicitEnergyFunction RTP LES(), dImplicitEnergyFunction RTP LES SB(), dImplicitEnergyFunction RTP SB(), initDonorFracAndMaxConVel R GL(), init Donor Frac And Max Con Vel R -Grid(), initDonorFracAndMaxConVel RT GL(), init Donor Frac And Max Con Vel RTTEOS(),initDonorFracAndMaxConVel RTP GL(), TEOS(), initDonorFracAndMaxConVel -RTP TEOS(), modelRead(), updateLocalBoundaryVelocitiesNewGrid R(), updateLocalBoundaryVelocitiesNewGrid RT(), updateLocalBoundaryVelocitiesNewGrid -RTP(), writeWatchZones\_R\_GL(), writeWatchZones\_R\_TEOS(), writeWatchZones\_RT\_-GL(), writeWatchZones RT TEOS(), writeWatchZones RTP GL(), and writeWatchZones -RTP TEOS().

### 6.4.3.8 int Grid::nU0

Index of  $u_0$  in grids, Grid::dLocalGridOld and Grid::dLocalGridNew. This is an external grid variable included in the count Grid::nNumVars

LES(), calNewE RT AD(), calNewE RT NA(), calNewE RT NA LES(), calNewE -RTP AD(). calNewE RTP NA(), calNewE RTP NA LES(), calNewEddyVisc -RT SM(), calNewEddyVisc RTP SM(), calNewR(), calNewU0 R(), calNewU0 RT(), calNewU0 RTP(), calNewU R(), calNewU R LES(), calNewU RT(), calNewU RT -LES(), calNewU RTP(), calNewU RTP LES(), calNewV RT(), calNewV RT LES(), calNewV RTP(), calNewV RTP LES(), calNewW RTP(), calNewW RTP LES(), calOldEddyVisc RT SM(), calOldEddyVisc RTP SM(), dImplicitEnergyFunction dImplicitEnergyFunction R LES(), dImplicitEnergyFunction R LES SB(), R(),dImplicitEnergyFunction R SB(), dImplicitEnergyFunction RT(), dImplicitEnergyFunction dImplicitEnergyFunction RT LES SB(), dImplicitEnergyFunction -RT LES(), RT SB(), dImplicitEnergyFunction RTP(), dImplicitEnergyFunction RTP LES(), dImplicitEnergyFunction RTP LES SB(), dImplicitEnergyFunction RTP SB(), initDonorFracAndMaxConVel R GL(), initDonorFracAndMaxConVel R -TEOS(),initDonorFracAndMaxConVel RT GL(), initDonorFracAndMaxConVel RT initDonorFracAndMaxConVel RTP GL(), initDonorFracAndMaxConVel RTP -TEOS(), TEOS(), main(), modelRead(), writeWatchZones R GL(), writeWatchZones R TEOS(), writeWatchZones RT GL(), writeWatchZones RT TEOS(), writeWatchZones RTP GL(), and writeWatchZones RTP TEOS().

### 6.4.3.9 int Grid::nV

Index of v in grids, Grid::dLocalGridOld and Grid::dLocalGridNew. This is an external grid variable included in the count Grid::nNumVars

Referenced by calDelt\_RT\_GL(), calDelt RT TEOS(), calDelt RTP GL(), calDelt -RTP TEOS(), calNewD RT(),calNewD RTP(), calNewE RT AD(), calNewE calNewE RT NA LES(), calNewE RTP AD(), RT NA(),calNewE RTP NA(), calNewEddyVisc RTP calNewE RTP NA LES(), calNewEddyVisc RT SM(), calNewQ0Q1 RT TEOS(), calNewQ0Q1 RT GL(),calNewQ0Q1Q2 RTP SM(), calNewQ0Q1Q2\_RTP\_TEOS(), calNewU0 RT(), calNewU0 RTP(), calNewU GL(), RT(),calNewU RT LES(), calNewU RTP(), calNewU RTP LES(), calNewV calNewV RT LES(), calNewV RTP(), calNewV RTP LES(), calNewW -RT(), calNewW RTP LES(), calOldEddyVisc RT SM(), RTP(),calOldEddyVisc RTP calOldQ0Q1 RT GL(), calOldQ0Q1 RT TEOS(), calOldQ0Q1Q2 RTP GL(), SM(), calOldQ0Q1Q2 RTP TEOS(), dImplicitEnergyFunction RT(), dImplicitEnergyFunction -RT LES(), dImplicitEnergyFunction RT LES SB(), dImplicitEnergyFunction -RT SB(), dImplicitEnergyFunction RTP(), dImplicitEnergyFunction RTP LES(), dImplicitEnergyFunction RTP LES SB(), dImplicitEnergyFunction RTP SB(), Grid(),initDonorFracAndMaxConVel RT GL(), initDonorFracAndMaxConVel RT TEOS(), initDonorFracAndMaxConVel RTP GL(), initDonorFracAndMaxConVel RTP TEOS(), initUpdateLocalBoundaries(), modelRead(), updateLocalBoundaryVelocitiesNewGrid -RT(), updateLocalBoundaryVelocitiesNewGrid RTP(), writeWatchZones RT GL(), writeWatchZones RT TEOS(), writeWatchZones RTP GL(), and writeWatchZones -RTP TEOS().

### 6.4.3.10 int Grid::nW

Index of w in grids, Grid::dLocalGridOld and Grid::dLocalGridNew. This is an external grid variable included in the count Grid::nNumVars

Referenced by calDelt\_RTP\_GL(), calDelt\_RTP\_TEOS(), calNewD\_RTP(), calNewE\_-RTP\_AD(), calNewE\_RTP\_NA(), calNewE\_RTP\_NA\_LES(), calNewEddyVisc\_RTP\_SM(), calNewQ0Q1Q2\_RTP\_GL(), calNewQ0Q1Q2\_RTP\_TEOS(), calNewU0\_RTP(), calNewU\_-

 $\label{eq:reconstruction} RTP(), \quad calNewU\_RTP\_LES(), \quad calNewV\_RTP(), \quad calNewV\_RTP\_LES(), \quad calNewW\_RTP\_LES(), \quad calNewW\_RTP\_LES(), \quad calOldEddyVisc\_RTP\_SM(), \quad calOldQ0Q1Q2\_RTP\_GL(), \\ calOldQ0Q1Q2\_RTP\_TEOS(), \quad dImplicitEnergyFunction\_RTP(), \quad dImplicitEnergyFunction\_RTP\_LES(), \quad dImplicitEnergyFunction\_RTP\_LES\_SB(), \quad dImplicitEnergyFunction\_RTP\_SB(), \\ Grid(), \quad initDonorFracAndMaxConVel\_RTP\_GL(), \quad initDonorFracAndMaxConVel\_RTP\_TEOS(), \\ initUpdateLocalBoundaries(), \quad modelRead(), \quad updateLocalBoundaryVelocitiesNewGrid\_RTP(), \\ writeWatchZones\_RTP\_GL(), \quad and \quad writeWatchZones\_RTP\_TEOS(). \\ \end{cases}$ 

#### 6.4.3.11 int Grid::nT

Index of T in grids, Grid::dLocalGridOld and Grid::dLocalGridNew. This is an external grid variable included in the count Grid::nNumVars. This variable is defined at cell centers.

calDelt R TEOS(), calDelt RT TEOS(), calDelt RTP TEOS(), Referenced by calNewE R NA LES(), calNewE\_RT\_NA(), calNewE R NA(),calNewE RT NA calNewE RTP NA(), calNewE RTP NA LES(), calNewPEKappaGamma -LES(), TEOS(), calNewTPKappaGamma TEOS(), calOldPEKappaGamma TEOS(), dImplicitEnergyFunction R(), dImplicitEnergyFunction R LES(), dImplicitEnergyFunction -R LES SB(), dImplicitEnergyFunction R SB(), dImplicitEnergyFunction -RT(),dImplicitEnergyFunction RT LES(), dImplicitEnergyFunction RT -LES SB(), dImplicitEnergyFunction RT SB(), dImplicitEnergyFunction RTP(), dImplicitEnergyFunction RTP LES(), dImplicitEnergyFunction RTP LES SB(), Grid(), dImplicitEnergyFunction RTP SB(), implicitSolve R(),implicitSolve RT(), implicitSolve RTP(), initImplicitCalculation(), initUpdateLocalBoundaries(), main(), model-Read(), writeWatchZones R TEOS(), writeWatchZones RT TEOS(), and writeWatchZones -RTP TEOS().

### 6.4.3.12 int Grid::nE

Index of E in grids, Grid::dLocalGridOld and Grid::dLocalGridNew. This is an internal grid variable included in the count Grid::nNumIntVars, unless the calculation is adiabatic in which case it is an external grid variable. This variable is defined at cell centers.

Referenced by calDelt R GL(), calDelt RT GL(), calDelt RTP GL(), calNewE R AD(), calNewE R NA(), calNewE R NA LES(), calNewE RT AD(), calNewE RT NA(), calNewE RT NA LES(), calNewE RTP AD(), calNewE RTP NA(), calNewE RTP -NA LES(), calNewP GL(), calNewPEKappaGamma TEOS(), calNewTPKappaGamma -TEOS(), calOldPEKappaGamma TEOS(), calOldP GL(), dImplicitEnergyFunction dImplicitEnergyFunction R LES(), dImplicitEnergyFunction R LES SB(), dImplicitEnergyFunction R SB(), dImplicitEnergyFunction RT(), dImplicitEnergyFunction -RT LES(), dImplicitEnergyFunction RT LES SB(), dImplicitEnergyFunction -RT SB(), dImplicitEnergyFunction RTP(), dImplicitEnergyFunction RTP LES(), dImplicitEnergyFunction RTP LES SB(), dImplicitEnergyFunction RTP SB(), initUpdateLocalBoundaries(), modelRead(), setInternalVarInf(), writeWatchZones R GL(),  $writeWatchZones\_R\_TEOS(), \ writeWatchZones\_RT\_GL(), \ writeWatchZones \ RT \ TEOS(),$ writeWatchZones RTP GL(), and writeWatchZones RTP TEOS().

## 6.4.3.13 int Grid::nP

Index of Pressure in grids, Grid::dLocalGridOld and Grid::dLocalGridNew. This is an internal grid variable and is included in the count of Grid::nNumIntVars. This variable is defined at cell centers.

Referenced by calDelt R GL(), calDelt R TEOS(), calDelt RT GL(), calDelt RT -TEOS(), calDelt RTP GL(), calDelt RTP TEOS(), calNewE R AD(), calNewE R -NA(), calNewE R NA LES(), calNewE RT AD(), calNewE RT NA(), calNewE -RT NA LES(), calNewE RTP AD(), calNewE RTP NA(), calNewE RTP NA LES(), calNewP GL(), calNewPEKappaGamma TEOS(), calNewQ0 R GL(), calNewQ0 R calNewQ0Q1 RT TEOS(), calNewQ0Q1Q2 RTP -TEOS(), calNewQ0Q1 RT GL(), GL(), calNewQ0Q1Q2 RTP TEOS(), calNewTPKappaGamma TEOS(), calNewU R(),  ${\tt calNewU\_R\_LES()}, \quad {\tt calNewU\_RT()}, \quad {\tt calNewU\_RT\_LES()}, \quad {\tt calNewU\_RTP()}, \quad {\tt$ RTP LES(), calNewV RT(), calNewV RT LES(), calNewV RTP(), calNewV RTP -LES(), calNewW RTP(), calNewW RTP LES(), calOldP GL(), calOldPEKappaGamma -TEOS(), calOldQ0\_R\_GL(), calOldQ0\_R\_TEOS(), calOldQ0Q1\_RT\_GL(), calOldQ0Q1\_-RT TEOS(), calOldQ0Q1Q2\_RTP\_GL(), calOldQ0Q1Q2\_RTP\_TEOS(), Grid(). initDonorFracAndMaxConVel R GL(), initDonorFracAndMaxConVel R TEOS(), initDonorFracAndMaxConVel RT GL(), initDonorFracAndMaxConVel RT TEOS(), initDonorFracAndMaxConVel RTP GL(),  $initDonorFracAndMaxConVel \ RTP \ TEOS(),$ initUpdateLocalBoundaries(), main(), modelRead(), setInternalVarInf(), writeWatchZones R -GL(), writeWatchZones R TEOS(), writeWatchZones RT GL(), writeWatchZones RT -TEOS(), writeWatchZones RTP GL(), and writeWatchZones RTP TEOS().

#### 6.4.3.14 int Grid::nKappa

Index of Opacity in grids, Grid::dLocalGridOld and Grid::dLocalGridNew. This is an internal grid variable and is included in the count of Grid::nNumIntVars. This variable is defined at cell centers.

 $\label{eq:referenced} Referenced by calNewE_R_NA(), calNewE_R_NA_LES(), calNewE_RT_NA(), calNewE_RT_NA_LES(), calNewE_RTP_NA(), calNewE_RTP_NA_-LES(), calNewPEKappaGamma_TEOS(), calNewTPKappaGamma_TEOS(), calOldPEKappaGamma_TEOS(), Grid(), initUpdateLocalBoundaries(), modelRead(), and setInternalVarInf().$ 

#### 6.4.3.15 int Grid::nGamma

Index of adiabatic index in grids, Grid::dLocalGridOld and Grid::dLocalGridNew. This is an internal grid variable and is included in the count of Grid::nNumIntVars. This variable is defined at cell centers.

 $\label{local_Referenced_by_calDelt_R_TEOS(), calDelt_RT_TEOS(), calDelt_RTP_TEOS(), calNewPEKappaGamma_TEOS(), calNewQ0_R_TEOS(), calNewQ0Q1_RT_TEOS(), calNewQ0Q1Q2_RTP_TEOS(), calNewTPKappaGamma_TEOS(), calOldPEKappaGamma_TEOS(), calOldQ0_R_TEOS(), calOldQ0Q1_RT_TEOS(), calOldQ0Q1Q2_RTP_TEOS(), Grid(), initDonorFracAndMaxConVel_R_TEOS(), initDonorFracAndMaxConVel_RT_TEOS(), initDonorFracAndMaxConVel_RTP_TEOS(), main(), modelRead(), and setInternalVarInf().$ 

#### 6.4.3.16 int Grid::nDenAve

Index of  $\langle \rho \rangle$  in grids, Grid::dLocalGridOld and Grid::dLocalGridNew. This is an internal grid variable and is included in the count of Grid::nNumIntVars. This variable is defined at cell centers only in the radial direction.

Referenced by calNewDenave\_R(), calNewDenave\_RT(), calNewDenave\_RTP(), calNewE\_-R\_NA(), calNewE\_R\_NA\_LES(), calNewE\_RT\_AD(), calNewE\_RT\_NA(), calNewE\_RT\_NA(), calNewE\_RTP\_NA(), calNewE\_RTP\_NA(

calNewU RTP(), LES(), calNewU RT(), calNewU RT LES(), calNewU RTP calNewV RT(), calNewV RT LES(), calNewV RTP(), calNewV RTP LES(), calNewW RTP(), calNewW RTP LES(), calOldDenave R(), calOldDenave RT(), dImplicitEnergyFunction R(), dImplicitEnergyFunction R LES(), calOldDenave RTP(), dImplicitEnergyFunction R SB(), dImplicitEnergyFunction RT(), dImplicitEnergyFunction dImplicitEnergyFunction RT LES SB(), dImplicitEnergyFunction -RT SB(),dImplicitEnergyFunction RTP(), dImplicitEnergyFunction RTP LES(), dImplicitEnergyFunction\_RTP\_LES\_SB(), dImplicitEnergyFunction RTP SB(), main(), modelRead(), setInternalVarInf(), writeWatchZones RT GL(), writeWatchZones -RT TEOS(), writeWatchZones RTP GL(), and writeWatchZones RTP TEOS().

#### 6.4.3.17 int Grid::nQ0

Index of the radial artificial viscosity in grids, Grid::dLocalGridOld and Grid::dLocalGridNew. This is an internal grid variable and is included in the count of Grid::nNumIntVars. This variable is defined at cell centers.

Referenced by calDelt R GL(), calDelt R TEOS(), calDelt RT GL(), calDelt RT - $TEOS(), \quad calDelt\_RTP\_GL(), \quad calDelt\_RTP\_TEOS(), \quad calNewE\_R\_AD(), \quad calNewE\_R\_-AD(), \quad calNewE\_R\_-AD(),$ calNewE R NA LES(), calNewE RT AD(), calNewE RT NA(), calNewE calNewE RTP\_AD(), RT NA LES(), calNewE RTP NA(), calNewE RTP calNewQ0 R TEOS(), NA LES(), calNewQ0 R GL(),calNewQ0Q1 RT GL(), calNewQ0Q1 RT TEOS(), calNewQ0Q1Q2 RTP GL(), calNewQ0Q1Q2 RTP TEOS(), calNewU R(),calNewU R LES(), calNewU RT(), calNewU RT LES(), calNewU calNewU RTP LES(), calNewV RT(), calNewV RT LES(), RTP(),calNewV calNewV RTP LES(), calNewW RTP(), calNewW RTP LES(), RTP(),calOldQ0 calOldQ0 R TEOS(), calOldQ0Q1 RT GL(), R GL()calOldQ0Q1 RT TEOS(), calOldQ0Q1Q2 RTP TEOS(), calOldQ0Q1Q2 RTP GL(), dImplicitEnergyFunction dImplicitEnergyFunction R LES(), dImplicitEnergyFunction R LES SB(), dImplicitEnergyFunction R SB(), dImplicitEnergyFunction RT(), dImplicitEnergyFunction -RT LES(), dImplicitEnergyFunction RT LES SB(), dImplicitEnergyFunction -RT SB(),dImplicitEnergyFunction RTP(), dImplicitEnergyFunction RTP LES(), dImplicitEnergyFunction RTP LES SB(), dImplicitEnergyFunction RTP SB(), Grid(). initDonorFracAndMaxConVel R GL(), initDonorFracAndMaxConVel R TEOS(), initDonorFracAndMaxConVel RT GL(), initDonorFracAndMaxConVel RT TEOS(), initDonorFracAndMaxConVel RTP GL(), initDonorFracAndMaxConVel RTP TEOS(), modelRead(), setInternalVarInf(), writeWatchZones R GL(), writeWatchZones R TEOS(), writeWatchZones RT GL(), writeWatchZones RT TEOS(), writeWatchZones RTP GL(), and writeWatchZones RTP TEOS().

#### 6.4.3.18 int Grid::nQ1

Index of the theta artificial viscosity in grids, Grid::dLocalGridOld and Grid::dLocalGridNew. This is an internal grid variable and is included in the count of Grid::nNumIntVars. This variable is defined at cell centers.

 

## 6.4.3.19 int Grid::nQ2

Index of the phi artificial viscosity in grids, Grid::dLocalGridOld and Grid::dLocalGridNew. This is an internal grid variable and is included in the count of Grid::nNumIntVars. This variable is defined at cell centers.

 $\label{eq:referenced_solution} Referenced by calDelt_RTP\_GL(), calDelt\_RTP\_TEOS(), calNewE\_RTP\_AD(), calNewE\_RTP\_NA(), calNewE\_RTP\_NA\_LES(), calNewQ0Q1Q2\_RTP\_GL(), calNewQ0Q1Q2\_RTP\_TEOS(), calNewU\_RTP\_LES(), calNewV\_RTP(), calNewV\_RTP\_LES(), calOldQ0Q1Q2\_RTP\_GL(), calOldQ0Q1Q2\_RTP\_TEOS(), dImplicitEnergyFunction\_RTP(), dImplicitEnergyFunction\_RTP\_LES(), dImplicitEnergyFunction\_RTP\_LES_SB(), dImplicitEnergyFunction\_RTP\_SB(), Grid(), initDonorFracAndMaxConVel\_RTP\_TEOS(), modelRead(), setInternalVarInf(), writeWatchZones\_RTP\_GL(), and writeWatchZones\_RTP\_TEOS().$ 

#### 6.4.3.20 int Grid::nDTheta

Index of  $\Delta\theta$  in grids, Grid::dLocalGridOld and Grid::dLocalGridNew. This is an internal grid variable and is included in the count of Grid::nNumIntVars. This variable is defined at cell centers.

Referenced by calDelt RT GL(), calDelt RT TEOS(), calDelt RTP GL(), calDelt -RTP TEOS(), calNewD RTP(), calNewE RT AD(), calNewE RT NA(), calNewE -RT NA LES(), calNewE RTP AD(), calNewE RTP NA(), calNewE RTP NA -LES(), calNewEddyVisc\_RT\_CN(), calNewEddyVisc\_RT\_SM(), calNewEddyVisc\_RTP\_-CN(), calNewEddyVisc RTP SM(), calNewU0 RTP(), calNewU RT(), calNewU RT LES(), calNewU RTP(), calNewU RTP LES(), calNewV RT(), calNewV RT LES(), calNewV RTP(), calNewV RTP LES(), calNewW RTP(), calNewW RTP LES(), calOldEddyVisc RT CN(), calOldEddyVisc RT SM(), calOldEddyVisc RTP CN(), calOldEddyVisc RTP SM(), dImplicitEnergyFunction RT(), dImplicitEnergyFunction -RT LES(), dImplicitEnergyFunction RT LES SB(), dImplicitEnergyFunction -RT SB(),dImplicitEnergyFunction RTP(), dImplicitEnergyFunction RTP LES(), dImplicitEnergyFunction RTP LES SB(), dImplicitEnergyFunction RTP SB(), initInternalVars(), modelRead(), and setInternalVarInf().

## 6.4.3.21 int Grid::nDPhi

Index of  $\Delta \phi$  in grids, Grid::dLocalGridOld and Grid::dLocalGridNew. This is an internal grid variable and is included in the count of Grid::nNumIntVars. This variable is defined at cell centers.

Referenced by average3DTo1DBoundariesNew(), average3DTo1DBoundariesOld(), calDelt\_RTP\_GL(), calDelt\_RTP\_TEOS(), calNewD\_RTP(), calNewDenave\_RTP(), calNewE\_-

 $\label{eq:rtp_ad} RTP\_AD(), \quad calNewE\_RTP\_NA(), \quad calNewE\_RTP\_NA\_LES(), \quad calNewEddyVisc\_RTP\_CN(), \quad calNewEddyVisc\_RTP\_SM(), \quad calNewU0\_RTP(), \quad calNewU\_RTP(), \quad calNewU\_RTP_LES(), \quad calNewV\_RTP(), \quad calNewV\_RTP\_LES(), \quad calNewW\_RTP(), \quad calNewW\_RTP\_LES(), \quad calOldDenave\_RTP(), \quad calOldEddyVisc\_RTP\_CN(), \quad calOldEddyVisc\_RTP\_SM(), \quad dImplicitEnergyFunction\_RTP\_LES(), \quad dImplicitEnergyFunction\_RTP\_LES(), \quad dImplicitEnergyFunction\_RTP\_LES(), \quad dImplicitEnergyFunction\_RTP\_SB(), \quad Grid(), \quad initInternalVars(), \quad modelRead(), \quad and \quad setInternalVarInf().$ 

#### 6.4.3.22 int Grid::nSinThetaIJK

Index of  $\sin \theta$  defined at zone center in grids, Grid::dLocalGridOld and Grid::dLocalGridNew. This is an internal grid variable and is included in the count of Grid::nNumIntVars.

 $calDelt RTP\_TEOS(),$ calDelt RTP GL(), Referenced by calNewE RT AD(), calNewE RT NA(), calNewE RT NA LES(), calNewE RTP AD(), calNewE RTP calNewE RTP NA LES(), calNewEddyVisc RTP CN(), calNewEddyVisc -NA(),calNewQ0Q1 RT GL(), calNewQ0Q1 RT TEOS(), RTP SM(), calNewQ0Q1Q2 -RTP GL(), calNewQ0Q1Q2 RTP TEOS(), calNewU RT LES(), calNewU RTP(), calNewV RTP LES(), calNewU RTP LES(), calNewV RT LES(), calNewW calNewW RTP LES(), calOldEddyVisc RTP CN(), calOldEddyVisc RTP -RTP(),calOldQ0Q1 RT GL(), calOldQ0Q1 RT TEOS(), calOldQ0Q1Q2 RTP GL(), calOldQ0Q1Q2 RTP TEOS(), dImplicitEnergyFunction RT(), dImplicitEnergyFunction -RT LES(), dImplicitEnergyFunction RT LES SB(), dImplicitEnergyFunction -RT SB(),dImplicitEnergyFunction RTP(), dImplicitEnergyFunction\_RTP\_LES(), dImplicitEnergyFunction RTP LES SB(), dImplicitEnergyFunction RTP SB(), initInternalVars(), modelRead(), and setInternalVarInf().

#### 6.4.3.23 int Grid::nSinThetaIJp1halfK

Index of  $\sin \theta$  at  $\theta$  interfaces in grids. This is an internal grid variable and is included in the count of Grid::nNumIntVars.

Referenced by calNewD RT(), calNewD RTP(), calNewE RT AD(), calNewE -RT NA(), calNewE RTP AD(), calNewE RTP calNewE RT NA LES(), NA(), calNewE RTP NA LES(), calNewQ0Q1 RT GL(),calNewQ0Q1 RT -TEOS(), calNewQ0Q1Q2 RTP GL(), calNewQ0Q1Q2 RTP TEOS(), calNewU0 calNewU0 RTP(), calNewU RT LES(), calNewU RTP LES(), RT(),calNewV calNewV RTP LES(), calNewW RTP LES(), RT LES(), calNewV RTP(), calOldQ0Q1 RT GL(), calOldQ0Q1 RT TEOS(), calOldQ0Q1Q2 RTP GL(), calOldQ0Q1Q2 RTP TEOS(), dImplicitEnergyFunction RT(), dImplicitEnergyFunction -RT LES(), dImplicitEnergyFunction RT LES SB(), dImplicitEnergyFunction dImplicitEnergyFunction RTP(), dImplicitEnergyFunction RTP LES(), dImplicitEnergyFunction RTP LES SB(), dImplicitEnergyFunction RTP SB(), initInternalVars(), modelRead(), and setInternalVarInf().

#### 6.4.3.24 int Grid::nCotThetaIJp1halfK

Index of  $\cot \theta$  at  $\theta$  interfaces in grids. This is an internal grid variable and is included in the count of Grid::nNumIntVars.

Referenced by calNewV\_RT\_LES(), calNewV\_RTP(), calNewV\_RTP\_LES(), Grid(), initInternalVars(), modelRead(), and setInternalVarInf().

#### 6.4.3.25 int Grid::nCotThetaIJK

Index of  $\cot \theta$  at cell centeres of grids. This is an internal grid variable and is included in the count of Grid::nNumIntVars.

Referenced by calNewEddyVisc\_RTP\_SM(), calNewU\_RT\_LES(), calNewU\_RTP\_LES(), calNewW\_RTP(), calNewW\_RTP\_LES(), calOldEddyVisc\_RTP\_SM(), Grid(), initInternal-Vars(), modelRead(), and setInternalVarInf().

#### 6.4.3.26 int Grid::nDCosThetaIJK

Index of  $\Delta \cos \theta$  defined at zone center in grids. This is an internal grid variable and is included in the count of Grid::nNumIntVars.

Referenced by average3DTo1DBoundariesNew(), average3DTo1DBoundariesOld(), calNewD\_-RT(), calNewD\_RTP(), calNewDenave\_RT(), calNewDenave\_RTP(), calNewUo\_RTP(), calNewUo\_RTP(), calOldDenave\_RT(), calOldDenave\_RTP(), Grid(), initInternalVars(), modelRead(), and setInternalVarInf().

## 6.4.3.27 int Grid::nEddyVisc

Index of the eddy viscosity in the grid, it is defined at zone centers in the grids. This is an internal grid variable and is included in the count of Grid::nNumIntVars.

calNewE RT NA LES(), Referenced by calNewE R NA LES(), calNewE RTP -NA LES(),calNewEddyVisc R CN(), calNewEddyVisc R SM(), calNewEddyVisc -RT CN(), calNewEddyVisc RT SM(), calNewEddyVisc RTP CN(), calNewEddyVisc -RTP SM(), calNewU R LES(), calNewU RT LES(), calNewU RTP LES(), calNewV calNewW RTP LES(), RT LES(), calNewV RTP LES(), calOldEddyVisc R CN(), calOldEddyVisc R SM(), calOldEddyVisc RT CN(), calOldEddyVisc RT SM(), calOldEddyVisc RTP CN(), calOldEddyVisc RTP SM(), dImplicitEnergyFunction -R LES(), dImplicitEnergyFunction R LES SB(), dImplicitEnergyFunction RT dImplicitEnergyFunction RT LES SB(), dImplicitEnergyFunction RTP LES(), dImplicitEnergyFunction RTP LES SB(), Grid(), main(), modelRead(), setInternalVarInf(), writeWatchZones RT GL(), writeWatchZones RT TEOS(), writeWatchZones RTP GL(), and writeWatchZones RTP TEOS().

#### 6.4.3.28 int Grid::nDonorCellFrac

Index of the amount of donor cell to use at that particular radial zone. It is defined at zone centers, and is an internal grid variable and is included in the count of Grid::nNumIntVars.

Referenced by calDelt R GL(), calDelt R TEOS(), calDelt RT GL(), calDelt RT TEOS(), calDelt RTP GL(), calDelt RTP TEOS(), calNewD R(), calNewD RT(), calNewD RTP(), calNewE R AD(), calNewE R NA(), calNewE R NA LES(), calNewE RT AD(), calNewE\_RT\_NA(), calNewE\_RT\_NA\_LES(), calNewE\_RTP\_AD(), calNewE\_RTP\_-NA(), calNewE RTP NA LES(), calNewU0 R(), calNewU0 RT(), calNewU0 RTP(), calNewU R(), calNewU R LES(), calNewU RT(), calNewU RT LES(), RTP(), calNewU RTP LES(), calNewV RT(), calNewV RT LES(), calNewV RTP(), calNewV RTP LES(), calNewW RTP(), calNewW RTP LES(), dImplicitEnergyFunction dImplicitEnergyFunction R LES SB(), dImplicitEnergyFunction R LES(), dImplicitEnergyFunction R SB(), dImplicitEnergyFunction RT(), dImplicitEnergyFunction dImplicitEnergyFunction RT LES SB(), RT LES(),dImplicitEnergyFunction -RT SB(),dImplicitEnergyFunction RTP(), dImplicitEnergyFunction RTP LES(),

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 \begin{array}{lll} dImplicitEnergyFunction\_RTP\_LES\_SB(), & dImplicitEnergyFunction\_RTP\_SB(), & Grid(), \\ initDonorFracAndMaxConVel\_R\_GL(), & initDonorFracAndMaxConVel\_R\_TEOS(), \\ initDonorFracAndMaxConVel\_RTP\_GL(), & initDonorFracAndMaxConVel\_RT\_TEOS(), \\ initDonorFracAndMaxConVel\_RTP\_TEOS(), & initDonorFracAndMaxConVel\_RTP\_TEOS(), \\ modelRead(), and setInternalVarInf(). & \\ \end{array}
```

#### 6.4.3.29 int Grid::nNumDims

Number of dimensions of the grid. It is used to chose the appropriate conservation equations. The value of this variable is independent of processor ProcTop::nRank.

Referenced by fin(), initImplicitCalculation(), initInternalVars(), initWatchZones(), main(), modelRead(), setInternalVarInf(), setMainFunctions(), setupLocalGrid(), and updateNewGridWith-Old().

#### 6.4.3.30 int Grid::nNumVars

Number of grid variables. This is set when reading in the model input file in the function modelRead. It is the number of variables that are printed and read from a file. The total number of variables also includes Grid::nNumIntVars. The value of this variable is independent of processor ProcTop::nRank.

Referenced by average3DTo1DBoundariesOld(), initUpdateLocalBoundaries(), modelRead(), modelWrite\_GL(), modelWrite\_TEOS(), setInternalVarInf(), setupLocalGrid(), updateNew-GridWithOld(), and updateOldGrid().

#### 6.4.3.31 int Grid::nNumIntVars

Number of internal variables. Internal variables are variables which are not reported in model dumps, and are not required to fully specify a starting model. They are used to save important information required during computation, an example is  $\sin \theta$ . The value of this variable is independent of processor ProcTop::nRank. This variable is set depending on the model read in (adiabatic/non-adiabatic/number of dimensions) in the function modelRead located in the file dataManipulation.cpp.

Referenced by average3DTo1DBoundariesOld(), initUpdateLocalBoundaries(), modelRead(), set-InternalVarInf(), setupLocalGrid(), updateNewGridWithOld(), and updateOldGrid().

#### 6.4.3.32 int Grid::nNum1DZones

Number of zones in 1D region of grid. The number of zones in 3D region is (Grid::nGlobalGridDims[0]-Grid::nNum1DZones). This is set when reading in the model input file in the function modelRead. The value of this variable is independent of processor ProcTop::nRank.

 $\label{lem:cond_equal} Referenced \quad by \quad init(), \quad initUpdateLocalBoundaries(), \quad initWatchZones(), \quad modelRead(), \\ modelWrite\_GL(), \\ modelWrite\_TEOS(), \\ and \\ setupLocalGrid().$ 

#### 6.4.3.33 int Grid::nNumGhostCells

Number of cells which are not included in local grid updating. This number is used in all dimensions to add to each local grid. When variables are not defined in a given direction ghost cells are not included in that direction. This is set when reading in the model input file in the function modelRead. The value of this variable is independent of processor ProcTop::nRank.

Referenced by calNewD\_R(), calNewD\_RTP(), calNewE\_R\_NA(), calNewE\_RTP\_NA\_-LES(), calNewU0\_RT(), calNewU0\_RTP(), calNewV\_RTP\_LES(), calNewW\_RTP\_LES(), calNewW\_RTP\_LES(), calOldEddyVisc\_R\_CN(), calOldEddyVisc\_RT\_SM(), calOldEddyVisc\_RT\_SM(), calOldEddyVisc\_RTP\_CN(), calOldEddyVisc\_RTP\_SM(), calOldEddyVisc\_RTP\_SM(), calOldPEKappaGamma\_TEOS(), dImplicitEnergyFunction\_R(), dImplicitEnergyFunction\_RLES(), dImplicitEnergyFunction\_R\_SB(), dImplicitEnergyFunction\_RTP\_LES(), dImplicitEnergyFunction\_RTP\_LES\_SB(), initImplicitCalculation(), initInternalVars(), initUpdateLocalBoundaries(), initWatchZones(), modelRead(), modelWrite\_GL(), modelWrite\_TEOS(), setupLocalGrid(), and updateNewGridWithOld().

#### 6.4.3.34 int\* Grid::nGlobalGridDims

Size of the entire global grid. It is an array of size 3 to hold size of each dimension of global grid. This size does not include Grid::nNumGhostCells or the extra size required for interface centered quantities. The values of this variable are independent of processor ProcTop::nRank. In the case of 1D or 2D calculations the  $\theta$  and  $\phi$  dimensions are set to 1 or just the  $\phi$  dimensions is set to 1 depending on the number of dimensions. The r,  $\theta$  and  $\phi$  dimensions are in the 0, 1 and 2 indices of the array respectively.

Referenced by Grid(), init(), initImplicitCalculation(), initUpdateLocalBoundaries(), initWatch-Zones(), modelWrite GL(), modelWrite TEOS(), and setupLocalGrid().

#### 6.4.3.35 int\*\* Grid::nVariables

Provides information on grid variables. A 2D array of size Grid::nNumVars+Grid::nNumIntVars by 3+1. nVariables[n][1] has values:

- -1: indicating that variable n is not defined
- 0: indicating that variable n is zone centered quantity
- 1: indicating that variable n is an interface centered quantity

in directions 1=0,1,2 which corresponding to  $\hat{r}$ ,  $\hat{\theta}$ , and  $\hat{\phi}$  respectively. nVariables[n][1] with 1=3 is used to indicate if a variable is dependent on time (1) or not(0). The values of this variable are independent of processor ProcTop::nRank.

Referenced by average3DTo1DBoundariesNew(), average3DTo1DBoundariesOld(), Grid(), initUpdateLocalBoundaries(), modelRead(), modelWrite\_GL(), modelWrite\_TEOS(), setInternalVar-Inf(), setupLocalGrid(), and updateNewGridWithOld().

## 6.4.3.36 int\*\*\* Grid::nLocalGridDims

Local grid dimensions. It is An array of size ProcTop::nNumProcs by Grid::nNumVars+Grid::nNumIntVars by 3. nLocalGridDims[p][n][l] gives the dimension of the local grid on processor p for variable n in direction 1. This variable does not include Grid::nNumGhostCells. The values of this variable are independent of processor ProcTop::nRank.

Referenced by calNewU0\_RT(), calNewU0\_RTP(), Grid(), initImplicitCalculation(), initInternalVars(), initUpdateLocalBoundaries(), initWatchZones(), modelRead(), modelWrite\_GL(), modelWrite\_TEOS(), setupLocalGrid(), and updateNewGridWithOld().

#### 6.4.3.37 double\*\*\*\* Grid::dLocalGridNew

Updated local grid values. An array of size Grid::nNumVars+Grid::nNumIntVars by Grid::nLocalGridDims[0] +2\*Grid::nNumGhostCells by Grid::nLocalGridDims[1]+2\*Grid::nNumGhostCells by Grid::nLocalGridDims[2]+2\*Grid::nNumGhostCells provided that the variable is defined in all 3 directions. Variables that are not defined in all 3 directions will have the additional two ghost cells left out in that direction and will also have a dimension of size 1 in that direction. This array contains the current grid state as it is being updated through calculations. This is a processor dependent variable and contains only the local grid for the current processor plus ghost cells.

average3DTo1DBoundariesNew(), calDelt R GL(), Referenced calDelt R by calDelt RT TEOS(), TEOS(), calDelt RT GL(), calDelt RTP GL(), calDelt calNewD R(), RTP TEOS(), calNewD RT(), calNewD RTP(), calNewDenave R(),calNewDenave RT(), calNewDenave RTP(), calNewE R AD(), calNewE R NA(), calNewE RT calNewE R NA LES(), calNewE RT AD(), calNewE RT NA(), NA LES(), calNewE RTP AD(), calNewE RTP NA(), calNewE RTP NA LES(), calNewEddyVisc RT CN(), calNewEddyVisc R SM(), calNewEddyVisc R CN(), calNewEddyVisc RT SM(), calNewEddyVisc RTP CN(), calNewEddyVisc RTP SM(), calNewP GL(), calNewPEKappaGamma TEOS(), calNewQ0 R GL(),calNewQ0Q1 RT TEOS(), R TEOS(), calNewQ0Q1 RT GL(),calNewQ0Q1Q2 -RTP GL(), calNewQ0Q1Q2 RTP TEOS(), calNewR(), calNewTPKappaGamma -TEOS(), calNewU0 R(), calNewU0 RT(), calNewU0 RTP(), calNewU R(), calNewU calNewU RT(), calNewU RT LES(), calNewU RTP(), R LES(),calNewU RTP LES(), calNewV RT(), calNewV RT LES(), calNewV RTP(), calNewV -RTP LES(), calNewW RTP(), calNewW RTP LES(), calOldEddyVisc R CN(), calOldEddyVisc RT CN(), calOldEddyVisc RTP CN(), dImplicitEnergyFunction dImplicitEnergyFunction R LES(), dImplicitEnergyFunction R LES SB(), R()dImplicitEnergyFunction R SB(), dImplicitEnergyFunction RT(), dImplicitEnergyFunction dImplicitEnergyFunction RT LES SB(), dImplicitEnergyFunction -RT LES(). RT SB(), dImplicitEnergyFunction RTP(), dImplicitEnergyFunction RTP LES(), dImplicitEnergyFunction\_RTP\_LES\_SB(), dImplicitEnergyFunction RTP SB(), implicitSolve R(), implicitSolve RT(), implicitSolve RTP(), initUpdateLocalBoundaries(), setupLocalGrid(), updateLocalBoundaries(), updateLocalBoundariesNewGrid(), updateNew-GridWithOld(), and updateOldGrid().

#### 6.4.3.38 double\*\*\*\* Grid::dLocalGridOld

Grid values from previous time step. An array the same size as Grid::dLocalGridNew but instead of containing the current grid state, it contains the last complete grid state. This is a processor dependent variable and contains only the local grid for the current processor plus ghost cells.

Referenced by average3DTo1DBoundariesNew(), average3DTo1DBoundariesOld(), calDelt -R GL(), calDelt R TEOS(), calDelt RT GL(), calDelt RT TEOS(), calDelt RTP GL(), calDelt RTP TEOS(), calNewD R(), calNewD RT(), calNewD RTP(), calNewDenave -RT(), calNewDenave RTP(), calNewE R AD(), calNewE R NA(), calNewE R NA -LES(), calNewE\_RT\_AD(), calNewE\_RT\_NA(), calNewE\_RT\_NA\_LES(), calNewE\_-RTP AD(), calNewE RTP NA(), calNewE RTP NA LES(), calNewEddyVisc R calNewEddyVisc RT CN(), calNewEddyVisc RT SM(), SM(), calNewEddyVisc calNewEddyVisc RTP SM(), RTP CN(), calNewQ0Q1 RT GL(), calNewQ0Q1 RT TEOS(),calNewQ0Q1Q2 RTP GL(), calNewQ0Q1Q2 RTP TEOS(), calNewR(), calNewTPKappaGamma TEOS(), calNewU0 R(), calNewU0 RT(),calNewU0 RTP(), calNewU R LES(), calNewU RT(), calNewU RT LES(), calNewU -RTP(), calNewU RTP LES(), calNewV RT(), calNewV RT LES(), calNewV RTP(), calNewW RTP LES(), calNewV RTP LES(), calNewW RTP(), calOldDenave R(),

calOldDenave RT(), calOldDenave RTP(), calOldEddyVisc R CN(), calOldEddyVisc R -SM(), calOldEddyVisc RT CN(), calOldEddyVisc RT SM(), calOldEddyVisc RTP CN(),  $calOldEddyVisc\ RTP\ SM(),\ calOldP\_GL(),\ calOldPEKappaGamma\_TEOS(),\ calOldQ0\_-temperature (Application of the control of$ calOldQ0 R TEOS(), calOldQ0Q1 RT GL(), calOldQ0Q1 RT TEOS(), R GL()calOldQ0Q1Q2 RTP TEOS(), calOldQ0Q1Q2 RTP GL(), dImplicitEnergyFunction dImplicitEnergyFunction R LES(), dImplicitEnergyFunction R LES SB(), dImplicitEnergyFunction R SB(), dImplicitEnergyFunction RT(), dImplicitEnergyFunction -RT LES(), dImplicitEnergyFunction\_RT\_LES\_SB(), dImplicitEnergyFunction -RT SB(), dImplicitEnergyFunction RTP LES(), dImplicitEnergyFunction RTP(), dImplicitEnergyFunction RTP LES SB(), dImplicitEnergyFunction RTP SB(), initDonorFracAndMaxConVel R GL(), initDonorFracAndMaxConVel R TEOS(), initDonorFracAndMaxConVel\_RT\_TEOS(), initDonorFracAndMaxConVel RT GL(), initDonorFracAndMaxConVel RTP GL(), initDonorFracAndMaxConVel RTP TEOS(), initInternalVars(), initUpdateLocalBoundaries(), modelRead(), modelWrite GL(), modelWrite -TEOS(), setupLocalGrid(), updateLocalBoundaries(), updateNewGridWithOld(), updateOld-Grid(), writeWatchZones R GL(), writeWatchZones R TEOS(), writeWatchZones RT -GL(), writeWatchZones RT TEOS(), writeWatchZones RTP GL(), and writeWatchZones -RTP TEOS().

#### 6.4.3.39 int\*\* Grid::nStartUpdateExplicit

Positions to begin updating grid with explicit calculations. It is an array of size nNumVars+nNumIntVars by 3. The start positions are defined in initUpdateLocalBoundaries(). These start values are dependent on processor ProcTop::nRank.

Referenced by calDelt R GL(), calDelt R TEOS(), calDelt RT GL(), calDelt RT TEOS(), calDelt RTP GL(), calDelt RTP TEOS(), calNewD R(), calNewD RT(), calNewD -RTP(), calNewDenave R(), calNewDenave RT(), calNewDenave RTP(), calNewE R AD(), calNewE R NA(), calNewE R NA LES(), calNewE RT AD(), calNewE RT NA(), calNewE RT NA LES(), calNewE RTP AD(), calNewE RTP NA(), calNewE RTP calNewEddyVisc R CN(), calNewEddyVisc R SM(), NA LES(),calNewEddyVisc -RT CN(), calNewEddyVisc RT SM(), calNewEddyVisc RTP CN(), calNewEddyVisc -RTP SM(), calNewP GL(), calNewQ0 R GL(), calNewQ0 R TEOS(), calNewQ0Q1 calNewQ0Q1 RT TEOS(), calNewQ0Q1Q2 RTP GL(), calNewQ0Q1Q2 -RTP TEOS(), calNewR(), calNewTPKappaGamma TEOS(), calNewU0 R(), calNewU0 -RT(), calNewU0\_RTP(), calNewU\_R(), calNewU R LES(), calNewU RT(), calNewU calNewU RTP(), calNewU RTP LES(), calNewV RT(), calNewV RT RT LES(), LES(), calNewV RTP(), calNewV RTP LES(), calNewW RTP(), calNewW RTP LES(), calOldDenave R(), calOldDenave RT(), calOldDenave RTP(), calOldEddyVisc calOldEddyVisc R SM(), calOldEddyVisc RT CN(), R CN()calOldEddyVisc -RT SM(), calOldEddyVisc RTP CN(), calOldEddyVisc RTP SM(), calOldP GL(), calOldPEKappaGamma TEOS(), calOldQ0 R GL(), calOldQ0 R TEOS(), calOldQ0Q1 -RT GL(), calOldQ0Q1 RT TEOS(), calOldQ0Q1Q2 RTP GL(), calOldQ0Q1Q2 RTP -TEOS(), Grid(), initDonorFracAndMaxConVel R GL(), initDonorFracAndMaxConVel R -TEOS(), initDonorFracAndMaxConVel RT GL(), initDonorFracAndMaxConVel RT TEOS(), initDonorFracAndMaxConVel RTP GL(), initDonorFracAndMaxConVel RTP TEOS(), initUpdateLocalBoundaries(), and updateOldGrid().

#### 6.4.3.40 int\*\* Grid::nEndUpdateExplicit

Positions to stop updating grid with explicit calculations. It is an array of size nNumVars+nNumIntVars by 3. The end positions are defined in initUpdateLocalBoundaries(). These start values are dependent on processor ProcTop::nRank.

Referenced by calDelt R GL(), calDelt R TEOS(), calDelt RT GL(), calDelt RT TEOS(), calDelt RTP GL(), calDelt RTP TEOS(), calNewD R(), calNewD RT(), calNewD -RTP(), calNewDenave R(), calNewDenave RT(), calNewDenave RTP(), calNewE R AD(), calNewE R NA(), calNewE R NA LES(), calNewE RT AD(), calNewE RT NA(), calNewE RT NA LES(), calNewE RTP AD(), calNewE RTP NA(), calNewE RTP calNewEddyVisc R CN(), calNewEddyVisc R SM(), calNewEddyVisc -RT CN(), calNewEddyVisc RT SM(), calNewEddyVisc RTP CN(), calNewEddyVisc -RTP SM(), calNewP GL(), calNewQ0 R GL(), calNewQ0 R TEOS(), calNewQ0Q1 calNewQ0Q1 RT TEOS(), calNewQ0Q1Q2 RTP GL(), RT GL(),calNewQ0Q1Q2 -RTP TEOS(), calNewR(), calNewTPKappaGamma TEOS(), calNewU0 R(), calNewU0 -RT(), calNewU0 RTP(), calNewU R(), calNewU R LES(), calNewU RT(), calNewU calNewU RTP(), calNewU RTP LES(), calNewV RT(), calNewV RT LES(), calNewV RTP(), calNewV RTP LES(), calNewW RTP(), calNewW RTP -LES(), calOldDenave R(), calOldDenave RT(), calOldDenave RTP(), calOldEddyVisc calOldEddyVisc R SM(), calOldEddyVisc RT CN(), calOldEddyVisc -R CN()RT SM(),calOldEddyVisc RTP CN(), calOldEddyVisc RTP SM(), calOldP GL(), calOldPEKappaGamma TEOS(), calOldQ0 R GL(), calOldQ0 R TEOS(), calOldQ0Q1 -RT GL(), calOldQ0Q1 RT TEOS(), calOldQ0Q1Q2 RTP GL(), calOldQ0Q1Q2 RTP -TEOS(), Grid(), initDonorFracAndMaxConVel R GL(), initDonorFracAndMaxConVel R -TEOS(), initDonorFracAndMaxConVel RT GL(), initDonorFracAndMaxConVel RT TEOS(), initDonorFracAndMaxConVel RTP GL(), initDonorFracAndMaxConVel RTP TEOS(), initUpdateLocalBoundaries(), and updateOldGrid().

#### 6.4.3.41 int\*\* Grid::nStartUpdateImplicit

Positions to begin updating grid with implicit calculations. It is an array of size nNumVars+nNumIntVars by 3. The start positions are defined in initUpdateLocalBoundaries(). These start values are dependent on processor ProcTop::nRank.

 $Referenced \ by \ calNewPEKappaGamma\_TEOS(), \ calOldDenave\_R(), \ calOldDenave\_RT(), \ calOldDenave\_RTP(), \ calOldPEKappaGamma\_TEOS(), \ Grid(), \ initUpdateLocalBoundaries(), \ and \ updateOldGrid().$ 

#### 6.4.3.42 int\*\* Grid::nEndUpdateImplicit

Positions to stop updating grid with implicit calculations. It is an array of size nNumVars+nNumIntVars by 3. The end positions are defined in initUpdateLocalBoundaries(). These start values are dependent on processor ProcTop::nRank.

Referenced by calNewPEKappaGamma\_TEOS(), calOldDenave\_R(), calOldDenave\_RT(), calOldDenave\_RTP(), calOldPEKappaGamma\_TEOS(), Grid(), initUpdateLocalBoundaries(), and updateOldGrid().

#### 6.4.3.43 int\*\*\* Grid::nStartGhostUpdateExplicit

Positions to begin updating ghost cells with explicit calculations. It is an array of size Grid::nNumVars+Grid::nNumIntVars by 2\*3 by 3. The second dimension indicates a particular ghost region. There are 2\*3 since each direction can have two ghost regions. The ghost region 0, is the outter ghost region in direction 0, 1 is the inner ghost region in direction 0, etc.

 $\label{eq:referenced_by_average3DTo1DBoundariesNew()} Referenced_by_average3DTo1DBoundariesOld(), calNewD_-R(), calNewD_RTP(), calNewD_RTP(), calNewDenave_R(), calNewDenave_RT(), calNewDenave_RTP(), calNewE_R_AD(), calNewE_R_NA(), calNe$ 

LES(), calNewE RT AD(), calNewE RT NA(), calNewE RT NA LES(), calNewE -RTP AD(), calNewE RTP NA(), calNewE RTP NA LES(), calNewEddyVisc R -CN(), calNewEddyVisc R SM(), calNewEddyVisc RT CN(), calNewEddyVisc RT calNewEddyVisc RTP\_SM(), calNewP GL(), calNewEddyVisc RTP CN(), SM(),calNewQ0 R TEOS(), calNewQ0 R GL(),calNewQ0Q1 RT GL(), calNewQ0Q1 -RT TEOS(), calNewQ0Q1Q2 RTP GL(), calNewQ0Q1Q2 RTP TEOS(), calNewTPKappaGamma TEOS(), calNewU0 R(), calNewU0 RT(), calNewU0 RTP(),  $calNewU_R()$ ,  $calNewU_R_LES()$ ,  $calNewU_RT()$ ,  $calNewU_RT_LES()$ , RTP(), calNewU RTP LES(), calNewV RT(), calNewV RT LES(), calNewV RTP(), calNewV RTP LES(), calNewW RTP(), calNewW RTP LES(), calOldDenave R(), calOldDenave RT(), calOldDenave RTP(), calOldEddyVisc R CN(), calOldEddyVisc - $R_SM()$ , calOldEddyVisc\_RT\_CN(), calOldEddyVisc\_RT\_SM(), calOldEddyVisc\_RTP calOldEddyVisc RTP\_SM(), calOldP GL(), calOldPEKappaGamma TEOS(), calOldQ0 R GL(), calOldQ0 R TEOS(), calOldQ0Q1 RT GL(), calOldQ0Q1 RT -TEOS(), calOldQ0Q1Q2 RTP GL(), calOldQ0Q1Q2 RTP TEOS(), Grid(), initUpdateLocal-Boundaries(), and updateOldGrid().

#### 6.4.3.44 int\*\*\* Grid::nEndGhostUpdateExplicit

Positions to end updating ghost cells with explicit calculations. Is an array of size Grid::nNumVars+Grid::nNumIntVars by 2\*3 by 3. The second dimension corresponds to which ghost region, since each dimension can have two ghost regions. The ghost region 0, is the outter ghost region in direction 0, 1 is the inner ghost region in direction 0, etc.

Referenced by average3DTo1DBoundariesNew(), average3DTo1DBoundariesOld(), calDelt -R GL(), calDelt R TEOS(), calDelt RT GL(), calDelt RT TEOS(), calDelt RTP GL(), calDelt RTP TEOS(), calNewD R(), calNewD RT(), calNewD RTP(), calNewDenave calNewDenave RT(), calNewDenave RTP(), calNewE R AD(), calNewE R -R()calNewE R NA LES(), calNewE RT AD(), calNewE RT NA(), NA(),calNewE -RT NA LES(), calNewE RTP AD(), calNewE RTP NA(), calNewE RTP NA calNewEddyVisc R CN(), calNewEddyVisc R SM(), calNewEddyVisc RT -LES(), calNewEddyVisc RT SM(), calNewEddyVisc RTP CN(), CN(). calNewEddyVisc -RTP SM(), calNewP GL(), calNewQ0 R GL(), calNewQ0 R TEOS(), calNewQ0Q1 calNewQ0Q1 RT TEOS(), calNewQ0Q1Q2 RTP GL(), calNewQ0Q1Q2 -RTP TEOS(), calNewR(), calNewTPKappaGamma TEOS(), calNewU0 R(), calNewU0 -RT(), calNewU0\_RTP(), calNewU\_R(), calNewU\_R\_LES(), calNewU\_RT(), calNewU calNewU RTP(), calNewU RTP LES(), calNewV RT(), calNewV RT RT LES(), LES(), calNewV RTP(), calNewV RTP LES(), calNewW RTP(), calNewW RTP LES(), calOldDenave\_R(), calOldDenave\_RT(), calOldDenave\_RTP(), calOldEddyVisc calOldEddyVisc R SM(), calOldEddyVisc RT CN(), R CN()calOldEddyVisc -RT SM(), calOldEddyVisc RTP CN(), calOldEddyVisc RTP SM(), calOldP GL(), calOldPEKappaGamma TEOS(), calOldQ0 R GL(), calOldQ0 R TEOS(), calOldQ0Q1 -RT GL(), calOldQ0Q1 RT TEOS(), calOldQ0Q1Q2 RTP GL(), calOldQ0Q1Q2 RTP - $TEOS(), \;\; Grid(), \;\; initDonorFracAndMaxConVel \;\; R \;\; GL(), \;\; initDonorFracAndMaxConVel \;\; R \;\; - 1000 \;\; and \;\;\; and \;\;\; and \;\;\; and \;\;\; and \;\; and \;\; and \;\; and \;\; and \;\;\; and \;\; and \;\; and \;\; and \;\; and \;$ TEOS(), initDonorFracAndMaxConVel RT GL(), initDonorFracAndMaxConVel RT TEOS(), initDonorFracAndMaxConVel RTP GL(), initDonorFracAndMaxConVel RTP TEOS(), initUpdateLocalBoundaries(), and updateOldGrid().

#### 6.4.3.45 int\*\*\* Grid::nStartGhostUpdateImplicit

Positions to begin updating ghost cells with implicit calculations. It is an array of size Grid::nNumVars+Grid::nNumIntVars by 2\*3 by 3. The second dimension indicates a particular ghost region. There are 2\*3 since each direction can have two ghost regions. The ghost region

0, is the outter ghost region in direction 0, 1 is the inner ghost region in direction 0, etc.

Referenced by calNewPEKappaGamma\_TEOS(), calOldDenave\_R(), calOldDenave\_RT(), calOldDenave\_RTP(), calOldPEKappaGamma\_TEOS(), Grid(), initUpdateLocalBoundaries(), and updateOldGrid().

#### 6.4.3.46 int\*\*\* Grid::nEndGhostUpdateImplicit

Positions to end updating ghost cells with implicit calculations. Is an array of size Grid::nNumVars+Grid::nNumIntVars by 2\*3 by 3. The second dimension corresponds to which ghost region, since each dimension can have two ghost regions. The ghost region 0, is the outter ghost region in direction 0, 1 is the inner ghost region in direction 0, etc.

 $Referenced \ by \ calNewPEKappaGamma\_TEOS(), \ calOldDenave\_R(), \ calOldDenave\_RT(), \ calOldDenave\_RTP(), \ calOldPEKappaGamma\_TEOS(), \ Grid(), \ initUpdateLocalBoundaries(), \ and \ updateOldGrid().$ 

#### 6.4.3.47 int \* Grid::nCenIntOffset

Indicates the offset between interface and center quantities. If nCenIntOffset[1]=0 then the outter interface quantities have the same index as zone centered quantities in direction 1. If nCenIntOffset[1]=1 then the outter interface quantities are given by the index for the zone centered quantities +1, in direction 1. The values are dependent on ProcTop::nRank and ProcTop::nPeriodic.

Referenced by average3DTo1DBoundariesNew(), average3DTo1DBoundariesOld(), calDelt -R GL()calDelt R TEOS(), calDelt RT GL(), calDelt RT TEOS(), calDelt calDelt RTP TEOS(), RTP GL(), calNewD R(), calNewD RT(), calNewD RTP(), calNewDenave RTP(), calNewE R AD(), calNewE R NA(). calNewDenave RT(), calNewE RT AD(), calNewE RT NA(),calNewE RT calNewE R NA LES(), calNewE\_RTP\_AD(), calNewE\_RTP\_NA(), calNewE RTP NA LES(), NA LES(), calNewEddyVisc R CN(), calNewEddyVisc R SM(), calNewEddyVisc RT CN(), calNewEddyVisc RT SM(), calNewEddyVisc RTP CN(), calNewEddyVisc RTP SM(), calNewQ0 R TEOS(), calNewQ0 R GL(),calNewQ0Q1 RT GL(),calNewQ0Q1 -RT TEOS(), calNewQ0Q1Q2 RTP GL(), calNewQ0Q1Q2 RTP TEOS(), calNewU0 -R(), calNewU0 RT(), calNewU0 RTP(), calNewU R(), calNewU R LES(), calNewU -RT(), calNewU RT LES(), calNewU RTP(), calNewU RTP LES(), calNewV RT(), calNewV RTP(), calNewV RTP LES(), calNewW RTP(), calNewV RT LES(), calOldDenave RTP(), calNewW RTP LES(), calOldDenave RT(), calOldEddyVisc -R CN(). calOldEddyVisc R SM(), calOldEddyVisc RT CN(), calOldEddyVisc -RT SM(),calOldEddyVisc RTP CN(), calOldEddyVisc RTP SM(), calOldQ0 -R GL() $calOldQ0_R_TEOS()$ , calOldQ0Q1 RT GL(),calOldQ0Q1 RT TEOS(), calOldQ0Q1Q2 RTP TEOS(), calOldQ0Q1Q2 RTP GL(), dImplicitEnergyFunction dImplicitEnergyFunction R LES(), dImplicitEnergyFunction R LES SB(), dImplicitEnergyFunction R SB(), dImplicitEnergyFunction RT(), dImplicitEnergyFunction dImplicitEnergyFunction RT LES SB(), dImplicitEnergyFunction -RT LES(), RT SB(),dImplicitEnergyFunction RTP(), dImplicitEnergyFunction RTP LES(), dImplicitEnergyFunction RTP LES SB(), dImplicitEnergyFunction RTP SB(), initDonorFracAndMaxConVel R TEOS(), initDonorFracAndMaxConVel R GL(),  $initDonorFracAndMaxConVel\_RT\_TEOS(),$ initDonorFracAndMaxConVel RT GL(), initDonorFracAndMaxConVel RTP GL(), initDonorFracAndMaxConVel RTP TEOS(), initInternalVars(), initUpdateLocalBoundaries(), setupLocalGrid(), writeWatchZones R GL(), writeWatchZones R TEOS(), writeWatchZones RT GL(), writeWatchZones RT TEOS(), writeWatchZones RTP GL(), and writeWatchZones RTP TEOS().

## 6.4.3.48 int Grid::nGlobalGridPositionLocalGrid[3]

The location at which the local grid starts in the global grid. This starts at 0, for the inner most cell, including ghost zones.

Referenced by calNewD\_R(), calNewD\_RTP(), calNewE\_R\_NA(), calNewE\_RTP\_NA\_-LES(), calNewU\_R(), calNewU\_RTP\_LES(), calNewV\_RTP\_LES(), calNewW\_RTP\_LES(), dImplicitEnergyFunction\_R(), dImplicitEnergyFunction\_R\_LES(), dImplicitEnergyFunction\_RTP\_LES(), dImplicitEnergyFunction\_RTP\_LES\_SB(), initUpdateLocalBoundaries(), and setupLocalGrid().

The documentation for this class was generated from the following files:

- global.h
- global.cpp

# 6.5 Implicit Class Reference

#include <global.h>

## **Public Member Functions**

• Implicit ()

#### Public Attributes

- int nNumImplicitZones
- Mat matCoeff
- Vec vecTCorrections
- Vec vecRHS
- Vec vecTCorrectionsLocal
- KSP kspContext
- ullet VecScatter vecscatTCorrections
- int nMaxNumIterations
- double dTolerance
- int nNumRowsALocal
- int nNumRowsALocalSB
- int \* nNumDerPerRow
- int \*\* nTypeDer
- int \*\*\* nLocDer
- int \*\* nLocFun
- double dDerivativeStepFraction
- double dCurrentRelTError
- int nCurrentNumIterations
- int nMaxNumSolverIterations
- double dMaxErrorInRHS
- $\bullet \ \ double \ dAverageRHS$

## 6.5.1 Detailed Description

This class holds data required for the implicit calculation.

## 6.5.2 Constructor & Destructor Documentation

## 6.5.2.1 Implicit::Implicit ()

constructor the the class Implicit.

 $References \quad dCurrentRelTError, \quad dDerivativeStepFraction, \quad dMaxErrorInRHS, \quad dTolerance, \\ nCurrentNumIterations, \quad nLocDer, \quad nLocFun, \quad nMaxNumIterations, \quad nMaxNumSolverIterations, \quad nNumDerPerRow, \quad nNumImplicitZones, \quad nNumRowsALocal, \quad nNumRowsALocalSB, \quad and \quad nTypeDer.$ 

#### 6.5.3 Member Data Documentation

#### 6.5.3.1 int Implicit::nNumImplicitZones

The number of zones in the region near the surface which should used the implicit calculation of the energy equation. If zero no zones will use the implicit calculation of energy.

Referenced by fin(), Implicit(), init(), initImplicitCalculation(), initUpdateLocalBoundaries(), main(), and setMainFunctions().

#### 6.5.3.2 Mat Implicit::matCoeff

Parallel coeffecient matrix (spread across all processors)

Referenced by implicitSolve\_R(), implicitSolve\_RT(), implicitSolve\_RTP(), and initImplicitCalculation().

#### 6.5.3.3 Vec Implicit::vecTCorrections

Temperature corrections solution vector (spread across all processors)

Referenced by implicitSolve\_R(), implicitSolve\_RT(), implicitSolve\_RTP(), and initImplicitCalculation().

## 6.5.3.4 Vec Implicit::vecRHS

RHS vector (spread across all processors)

Referenced by implicitSolve\_R(), implicitSolve\_RT(), implicitSolve\_RTP(), and initImplicitCalculation().

## 6.5.3.5 Vec Implicit::vecTCorrectionsLocal

Corrections to local temperatures only (on local processor only).

Referenced by implicitSolve\_R(), implicitSolve\_RT(), implicitSolve\_RTP(), and initImplicitCalculation().

#### 6.5.3.6 KSP Implicit::kspContext

PETSc solver context.

Referenced by implicitSolve\_R(), implicitSolve\_RT(), implicitSolve\_RTP(), and initImplicitCalculation().

#### 6.5.3.7 VecScatter Implicit::vecscatTCorrections

Scatter context, used to hold information about retrieving the distributed temperature corrections from vecTCorrections and placing them into the local vector vecTCorrectionsLocal.

Referenced by implicitSolve\_R(), implicitSolve\_RT(), implicitSolve\_RTP(), and initImplicitCalculation().

#### 6.5.3.8 int Implicit::nMaxNumIterations

The maximum number of iterations to try to get the largest value of vecTCorrections relative to the temperature below dTolerance. Ater which the calculation continues.

Referenced by Implicit(), implicitSolve\_R(), implicitSolve\_RT(), implicitSolve\_RTP(), init(), and initImplicitCalculation().

#### 6.5.3.9 double Implicit::dTolerance

The amount of relative error that is allowed in the calculation of the temperature with the implicit calculation.

Referenced by Implicit(), implicitSolve\_R(), implicitSolve\_RT(), implicitSolve\_RTP(), init(), and initImplicitCalculation().

#### 6.5.3.10 int Implicit::nNumRowsALocal

The number of rows of the coeffecient matrix which is on the local processor.

Referenced by Implicit(),  $implicitSolve_R()$ ,  $implicitSolve_RT()$ ,  $implicitSolve_RTP()$ , and initImplicitCalculation().

#### 6.5.3.11 int Implicit::nNumRowsALocalSB

The number or rows of the coeffecient matrix which is on the local processor, and that are in the surface boundary region.

Referenced by Implicit(),  $implicitSolve_R()$ ,  $implicitSolve_RT()$ ,  $implicitSolve_RTP()$ , and initImplicitCalculation().

#### 6.5.3.12 int\* Implicit::nNumDerPerRow

An array of size nNumRowsALocal which contains the number of non-zero derivatives for a given row of A.

Referenced by Implicit(),  $implicitSolve_R()$ ,  $implicitSolve_RT()$ ,  $implicitSolve_RTP()$ , and initImplicitCalculation().

#### 6.5.3.13 int\*\* Implicit::nTypeDer

An array of size nNumRowsALocal by nNumDerPerRow [q], where q is a row index. Thus each row of the array can have a different length. This gives the type of derivative of row q for each derivative in that row. The value of this variable is set in the function initImplicitCalculation.

Referenced by Implicit(), implicitSolve\_R(), implicitSolve\_RT(), implicitSolve\_RTP(), and initImplicitCalculation().

#### 6.5.3.14 int\*\*\* Implicit::nLocDer

An array of size nNumRowsALocal by 2 by nNumDerPerRow [q], where q is a row index. This array holds the global position of the current row q for the current derivative e.g. the p th derivative in the q th row would be in row and column (nLocDer[q][0][p], nLocDer[q][1][p]). The value of this variable is set in the function initImplicitCalculation.

Referenced by Implicit(), implicitSolve\_R(), implicitSolve\_RT(), implicitSolve\_RTP(), and initImplicitCalculation().

#### 6.5.3.15 int\*\* Implicit::nLocFun

An array of size nNumRowsALocal by 3 [q] , where q is a row index. This array holds the local grid position of the current row q e.g. the (i,j,k) location of the the current row in the local grid. The value of this variable is set in the function initImplicitCalculation.

 $Referenced \ by \ Implicit(), \ implicitSolve\_R(), \ implicitSolve\_RT(), \ implicitSolve\_RT(), \ and \ initImplicitCalculation().$ 

#### 6.5.3.16 double Implicit::dDerivativeStepFraction

Dicates the size of the step that should be used to evaluate the numerical derivitives of the energy equation, for solving for the temperature implicitily. This value multiplies the temperature to produce the step size. A good value is around 5e-7.

Referenced by Implicit(),  $implicitSolve_R()$ ,  $implicitSolve_RT()$ ,  $implicitSolve_RTP()$ , and init().

## 6.5.3.17 double Implicit::dCurrentRelTError

keeps track of the largest relative error in the calculation of the temperature

Referenced by fin(), Implicit(), implicitSolve\_R(), implicitSolve\_RT(), implicitSolve\_RTP(), and main().

## 6.5.3.18 int Implicit::nCurrentNumIterations

keeps track of the number of iterations needed to converge to a solution

Referenced by fin(), Implicit(), implicitSolve\_R(), implicitSolve\_RT(), implicitSolve\_RTP(), and main().

#### 6.5.3.19 int Implicit::nMaxNumSolverIterations

If TRACKMAXSOLVERERROR set to 1, then this will be the current maximum number of iterations required for the linear equaiton solver to solve for the temperature correction over all iterations and time steps since the last model dump.

Referenced by fin(), Implicit(), implicitSolve\_R(), implicitSolve\_RT(), implicitSolve\_RTP(), and main().

#### 6.5.3.20 double Implicit::dMaxErrorInRHS

If TRACKMAXSOLVERERROR set to 1, then this will be the current maximum absolute error between the RHS as calculated from the solution and the coeffecient matrix, and the actual RHS. This value is the maximum from all values at each iteration of the solution, from each time step since the last model dump.

Referenced by fin(), Implicit(), implicitSolve\_R(), implicitSolve\_RT(), implicitSolve\_RTP(), and main().

## 6.5.3.21 double Implicit::dAverageRHS

Holds the average value of the right hand side for the timestep where the error in the RHS is the largest dMaxErrorInRHS. Only set if TRACKMAXSOLVERERROR is set to 1.

Referenced by fin(), implicitSolve R(), implicitSolve RT(), implicitSolve RTP(), and main().

The documentation for this class was generated from the following files:

- global.h
- global.cpp

## 6.6 MessPass Class Reference

```
#include <global.h>
```

## **Public Member Functions**

• MessPass ()

## Public Attributes

- MPI::Datatype \* typeSendNewGrid
- MPI::Datatype \* typeRecvOldGrid
- MPI::Datatype \*\* typeSendNewVar
- MPI::Datatype \*\* typeRecvNewVar
- MPI::Request \* requestSend
- MPI::Request \* requestRecv
- MPI::Status \* statusSend
- MPI::Status \* statusRecv

## 6.6.1 Detailed Description

This class manages information which pertains to message passing between processors.

#### 6.6.2 Constructor & Destructor Documentation

#### 6.6.2.1 MessPass::MessPass ()

Constructor for class MessPass.

 $References\ request Recv,\ request Send,\ status Recv,\ status Send,\ type Recv New Var,\ type Recv Old-Grid,\ type Send New Grid,\ and\ type Send New Var.$ 

#### 6.6.3 Member Data Documentation

## 6.6.3.1 MPI::Datatype\* MessPass::typeSendNewGrid

Send data types for entire grid. It is of size ProcTop::nNumNeighbors.

Referenced by initUpdateLocalBoundaries(), MessPass(), and updateLocalBoundaries().

## 6.6.3.2 MPI::Datatype\* MessPass::typeRecvOldGrid

Recv data types for entire grid. It is of sizee ProcTop::nNumNeighbors.

Referenced by initUpdateLocalBoundaries(), MessPass(), and updateLocalBoundaries().

## $\textbf{6.6.3.3} \quad \textbf{MPI::Datatype} ** \textbf{MessPass::typeSendNewVar}$

Send data types for variables. It is of size ProcTop::nNumNeighbors by Grid::nNumVars.

Referenced by calNewU0\_R(), calNewU0\_RT(), calNewU0\_RTP(), initUpdateLocalBoundaries(), MessPass(), and updateLocalBoundariesNewGrid().

#### 6.6.3.4 MPI::Datatype\*\* MessPass::typeRecvNewVar

Recieve data types for variables. It is of size ProcTop::nNumNeighbors by Grid::nNumVars.

Referenced by calNewU0\_R(), calNewU0\_RT(), calNewU0\_RTP(), initUpdateLocalBoundaries(), MessPass(), and updateLocalBoundariesNewGrid().

## 6.6.3.5 MPI::Request\* MessPass::requestSend

Message handles.

Referenced by initUpdateLocalBoundaries(), MessPass(), and updateLocalBoundaries().

## 6.6.3.6 MPI::Request\* MessPass::requestRecv

Message handles.

Referenced by initUpdateLocalBoundaries(), MessPass(), updateLocalBoundaries(), and updateLocalBoundariesNewGrid().

## 6.6.3.7 MPI::Status\* MessPass::statusSend

Message status.

Referenced by initUpdateLocalBoundaries(), MessPass(), and updateLocalBoundaries().

## 6.6.3.8 MPI::Status\* MessPass::statusRecv

Message status.

Referenced by initUpdateLocalBoundaries(), MessPass(), updateLocalBoundaries(), and updateLocalBoundariesNewGrid().

The documentation for this class was generated from the following files:

- global.h
- global.cpp

# 6.7 Output Class Reference

#include <global.h>

#### **Public Member Functions**

• Output ()

## Public Attributes

- int nDumpFrequencyStep
- double dDumpFrequencyTime
- double dTimeLastDump
- $\bullet \ \ int \ nNumTimeStepsSinceLastPrint$
- bool bDump
- bool bPrint
- int nPrintMode
- std::string sBaseOutputFileName
- $\bullet \ \ {\rm std::ofstream} * \ \ {\rm ofWatchZoneFiles}$
- std::vector< WatchZone > watchzoneList
- int nPrintFrequencyStep
- double dPrintFrequencyTime
- double dTimeLastPrint

## 6.7.1 Detailed Description

This class manages information pertianing to the output of data to files.

## 6.7.2 Constructor & Destructor Documentation

## 6.7.2.1 Output::Output ()

Constructor for this class.

 $References\ bDump,\ nDumpFrequencyStep,\ nNumTimeStepsSinceLastPrint,\ of WatchZoneFiles, and\ sBaseOutputFileName.$ 

## 6.7.3 Member Data Documentation

#### 6.7.3.1 int Output::nDumpFrequencyStep

How ofter a the grid state should be written to a file according to time step index. If it is 1 the will state will be written every time step, if it equals 2 it will be written every other time step etc. If it is 0 no dumps will be made according to the time step index.

Referenced by init(), main(), and Output().

#### 6.7.3.2 double Output::dDumpFrequencyTime

How ofter a the grid state should be written to a file according to simulation time in seconds. If it is 0 no dumps will be made according to simulation time.

Referenced by init(), and main().

#### 6.7.3.3 double Output::dTimeLastDump

The simulation time at which the last dump was made using the Output::dDumpFrequencyTime criterion.

Referenced by init(), and main().

#### 6.7.3.4 int Output::nNumTimeStepsSinceLastPrint

The number of time steps since the last model dump.

Referenced by fin(), main(), and Output().

## 6.7.3.5 bool Output::bDump

The number of time steps since the last print. Should the grid state be written to a file at a frequency of Output::nDumpFrequencyStep timesteps, and/or every Output::dDumpFrequencyTime seconds of simulation time. This is set to true by putting a "<dump>" node into the "SPHERLS.xml" configuration file.

Referenced by init(), main(), and Output().

## 6.7.3.6 bool Output::bPrint

Should status updates be printed to the screen.

Referenced by init(), and main().

#### 6.7.3.7 int Output::nPrintMode

Sets the way in which information should be printed to the standard output during the run. If it is 0, it will print the standard information reporting on the progress of the code. If it is 1 it will print out information to diagnose timestepping problems.

Referenced by fin(), init(), and main().

#### 6.7.3.8 std::string Output::sBaseOutputFileName

Base filename used for output, default is "out". All model dumps, and output information will contain this file name and extend it to indicate their specific information. The value of this variable is independent of processor ProcTop::nRank.

Referenced by fin(), init(), initWatchZones(), main(), and Output().

#### 6.7.3.9 std::ofstream\* Output::ofWatchZoneFiles

An array of output streams of size Output::watchzoneList .size() which are used to write out the information of the watched zones.

 $\label{lem:referenced_by_finWatchZones} Referenced by finWatchZones(), initWatchZones(), Output(), writeWatchZones_R_GL(), writeWatchZones_RT_GL(), writeWatchZones_RT_GL(), writeWatchZones_RT_TEOS(), writeWatchZones_RTP_GL(), and writeWatchZones_RTP_TEOS().$ 

#### 6.7.3.10 std::vector<WatchZone> Output::watchzoneList

A vector used to keep information used to specify the zones to be watched.

 $\label{lem:referenced_solution} Referenced & by & finWatchZones(), & initWatchZones(), & writeWatchZones\_R\_GL(), \\ writeWatchZones\_R\_TEOS(), & writeWatchZones\_RT\_GL(), & writeWatchZones\_RT\_TEOS(), \\ writeWatchZones & RTP\_GL(), & and & writeWatchZones & RTP\_TEOS(). \\ \end{cases}$ 

## 6.7.3.11 int Output::nPrintFrequencyStep

How often the status is printed to the screen in time steps.

Referenced by init(), and main().

#### 6.7.3.12 double Output::dPrintFrequencyTime

How often the status is printed to the screen in simulation time.

Referenced by init(), and main().

#### 6.7.3.13 double Output::dTimeLastPrint

Simulation time when last status was printed.

Referenced by init(), and main().

The documentation for this class was generated from the following files:

- global.h
- global.cpp

## 6.8 Parameters Class Reference

#include <global.h>

#### **Public Member Functions**

• Parameters ()

#### Public Attributes

- bool bEOSGammaLaw
- bool bAdiabatic
- int nTypeTurbulanceMod
- double dPi
- double dSigma
- $\bullet$  double dG
- double dGamma
- std::string sEOSFileName
- eos eosTable
- double dA
- double dAVThreshold
- double dDonorCellMultiplier
- double dDonorCellMin
- double dAlpha
- double dTolerance
- int nMaxIterations
- double dEddyViscosity
- double dMaxConvectiveVelocity
- double dMaxConvectiveVelocity c
- double dPrt
- double dT cut
- double dDEDM cut
- int nDEDM cut zone
- $\bullet$  bool bDEDM cut set
- std::string sDebugProfileOutput

## 6.8.1 Detailed Description

This class holds parameters and constants used for calculation.

## 6.8.2 Constructor & Destructor Documentation

#### 6.8.2.1 Parameters::Parameters ()

Constructor for the class Parameters

References bDEDM\_cut\_set, dA, dAlpha, dAVThreshold, dDonorCellMin, dEddyViscosity, dG, dMaxConvectiveVelocity, dMaxConvectiveVelocity c, dPi, dPrt, dSigma, and dT cut.

#### 6.8.3 Member Data Documentation

#### 6.8.3.1 bool Parameters::bEOSGammaLaw

If true SPHERLS will use a gamma law gas instead of a tabulated equation of state. This is set in the starting model.

Referenced by init(), initInternalVars(), initWatchZones(), modelRead(), setInternalVarInf(), and setMainFunctions().

#### 6.8.3.2 bool Parameters::bAdiabatic

If true SPHERLS will use adiabatic functions to calculate the energy. This can be used for both gamma law gas and tabulated equations of state (see Parameters::bEOSGammaLaw).

Referenced by init(), and setMainFunctions().

#### 6.8.3.3 int Parameters::nTypeTurbulanceMod

This varible indicates the type of turbulance model to be used. If 0, no turbulance model will be used, if 1 it will use a constant times the zoning size, and if 2 it will use the Smagorinksy turbulance model which increases the value of the eddy viscosity parameter when there are large amounts of shear, and decrease it when there isn't.

Referenced by init(), initInternalVars(), modelRead(), setInternalVarInf(), setMainFunctions(), writeWatchZones\_RT\_GL(), writeWatchZones\_RT\_TEOS(), writeWatchZones\_RTP\_GL(), and writeWatchZones\_RTP\_TEOS().

### 6.8.3.4 double Parameters::dPi

The value of  $\pi$ .

calNewE R AD(), calNewE R NA(), calNewE R NA LES(), Referenced by calNewE RT AD(), calNewE RT NA(),calNewE RT NA LES(), calNewE RTP calNewE RTP NA(), calNewE RTP NA LES(), calNewU R(), calNewU -AD()calNewU RT(), calNewU RT LES(), R LES(), calNewU RTP(), calNewU calNewV RT LES(), calNewV RTP(), RTP LES(), calNewV RT(), calNewV -RTP LES(), calNewW RTP(), calNewW RTP LES(), dImplicitEnergyFunction dImplicitEnergyFunction R LES(), dImplicitEnergyFunction R LES SB(), dImplicitEnergyFunction R SB(), dImplicitEnergyFunction RT(), dImplicitEnergyFunction -RT LES(),dImplicitEnergyFunction RT LES SB(), dImplicitEnergyFunction dImplicitEnergyFunction RTP LES(), RT SB(),dImplicitEnergyFunction RTP(), dImplicitEnergyFunction RTP LES SB(), dImplicitEnergyFunction RTP SB(), Parameters(), writeWatchZones R GL(), writeWatchZones R TEOS(), writeWatchZones RT GL(),  $writeWatchZones\_RT\_TEOS(), \ writeWatchZones\_RTP\_GL(), \ and \ writeWatchZones\_RTP\_-TEOS().$ 

#### 6.8.3.5 double Parameters::dSigma

The value of  $\sigma$ , the Stefan-Boltzmann constant.

calNewE R NA(), Referenced calNewE R NA LES(), calNewE RT NA(), calNewE RT NA LES(), calNewE RTP NA(), calNewE RTP NA LES(), dImplicitEnergyFunction R(), dImplicitEnergyFunction R LES(), dImplicitEnergyFunction -R LES SB(), dImplicitEnergyFunction R SB(), dImplicitEnergyFunction -RT(),dImplicitEnergyFunction RT LES(), dImplicitEnergyFunction RT LES SB(), dImplicitEnergyFunction RT SB(), dImplicitEnergyFunction RTP(), dImplicitEnergyFunction RTP LES(), dImplicitEnergyFunction RTP LES SB(), dImplicitEnergyFunction RTP SB(), and Parameters().

#### 6.8.3.6 double Parameters::dG

The Gravitational constant G.

Referenced by calNewU\_R(), calNewU\_R\_LES(), calNewU\_RT(), calNewU\_RT\_LES(), calNewU\_RTP(), calNewU\_RTP\_LES(), and Parameters().

#### 6.8.3.7 double Parameters::dGamma

The adiabatic  $\gamma$ , used in calculating the equation of state. If using a gamma law gas.

 $\label{eq:referenced_problem} Referenced by calDelt\_R\_GL(), calDelt\_RT\_GL(), calDelt\_RTP\_GL(), calNewQ0\_R\_GL(), calNewQ0Q1\_RT\_GL(), calNewQ0Q1Q2\_RTP\_GL(), calOldQ0\_R\_GL(), calOldQ0Q1\_RT\_GL(), calOldQ0Q1Q2\_RTP\_GL(), dEOS\_GL(), initDonorFracAndMaxConVel\_R\_GL(), initDonorFracAndMaxConVel\_RT\_GL(), initDonorFracAndMaxConVel\_RTP\_GL(), initDonorFracAndMaxConVel\_RTP\_GL(), init-WatchZones(), modelRead(), and modelWrite\_GL().$ 

#### 6.8.3.8 std::string Parameters::sEOSFileName

File name of equation of state table. This value is set either by the configuration file, SPHERLS.xml or in the model file read in. If it is specified in SPHERLS.xml it will overide the file name set in the model.

Referenced by init(), modelRead(), and modelWrite TEOS().

#### 6.8.3.9 eos Parameters::eosTable

Holds the equation of state table. If using a tabulated equation of state.

calNewPEKappaGamma TEOS(), calNewTPKappaGamma TEOS(), Referenced calOldPEKappaGamma TEOS(), dImplicitEnergyFunction\_R(), dImplicitEnergyFunction\_-R LES(),dImplicitEnergyFunction R LES SB(), dImplicitEnergyFunction -R SB(),dImplicitEnergyFunction RT(), dImplicitEnergyFunction RT -LES(), dImplicitEnergyFunction RT LES SB(), dImplicitEnergyFunction RT dImplicitEnergyFunction RTP LES(), dImplicitEnergyFunction RTP(), SB(),dImplicitEnergyFunction RTP LES SB(), dImplicitEnergyFunction RTP SB(), and init().

#### 6.8.3.10 double Parameters::dA

Artificial viscosity parameter, reasonable values range from 0 to  $\sim$ 3.

 $\label{eq:referenced} Referenced by calNewQ0_R_GL(), calNewQ0_R_TEOS(), calNewQ0Q1_RT_GL(), calNewQ0Q1_RT_TEOS(), calNewQ0Q1Q2_RTP_GL(), calNewQ0Q1Q2_RTP_TEOS(), calNewQ0Q1Q2_RTP_TEOS(), calOldQ0_R_GL(), calOldQ0_R_TEOS(), calOldQ0Q1_RT_GL(), calOldQ0Q1_RT_TEOS(), calOldQ0Q1Q2_RTP_GL(), calOldQ0Q1Q2_RTP_TEOS(), init(), modelRead(), modelWrite_GL(), modelWrite_TEOS(), and Parameters().$ 

#### 6.8.3.11 double Parameters::dAVThreshold

The amount of compression before AV is turned on. It is in terms of a velocity difference between zone sides and is in fractions of the local sound speed.

 $\label{eq:referenced_by_calNewQ0_R_GL(), calNewQ0_R_TEOS(), calNewQ0Q1_RT_GL(), calNewQ0Q1_RT_TEOS(), calNewQ0Q1Q2_RTP_GL(), calNewQ0Q1Q2_RTP_TEOS(), calNewQ0Q1Q2_RTP_TEOS(), calOldQ0_R_GL(), calOldQ0_R_TEOS(), calOldQ0Q1_RT_GL(), calOldQ0Q1_RT_TEOS(), calOldQ0Q1Q2_RTP_GL(), calOldQ0Q1Q2_RTP_TEOS(), init(), modelRead(), modelWrite_GL(), modelWrite_TEOS(), and Parameters(). }$ 

#### 6.8.3.12 double Parameters::dDonorCellMultiplier

Multiplier used to determine the faction of the sound speed at which donor cell is full. e.g. a value of 1.0 means the donor cell will be full when the convective velocity is equal to the sound speed. A value of 0.5 will mean that it will be full donor cell when the convective velocity is twice the sound speed. A value of 2.0 will mean that it will use full donor cell when the convective velocity is half the sound speed.

 $\label{localDelt_R_GL()} Referenced by calDelt_R_GL(), calDelt_R_TEOS(), calDelt_RT_GL(), calDelt_RT_TEOS(), init(), initDonorFracAndMaxConVel_R_GL(), initDonorFracAndMaxConVel_R_TEOS(), initDonorFracAndMaxConVel_RT_GL(), initDonorFracAndMaxConVel_RT_TEOS(), initDonorFracAndMaxConVel_RT_GL(), and initDonorFracAndMaxConVel_RTP_TEOS(). \\$ 

## 6.8.3.13 double Parameters::dDonorCellMin

The minimum amount of donor cell allowed. Set in constructor, Parameters::Parameters

 $\label{eq:referenced} Referenced by calDelt\_R\_GL(), calDelt\_R\_TEOS(), calDelt\_RT\_GL(), calDelt\_RT\_TEOS(), calDelt\_RTP\_GL(), calDelt\_RTP\_TEOS(), initDonorFracAndMaxConVel\_R\_GL(), initDonorFracAndMaxConVel\_RT\_TEOS(), initDonorFracAndMaxConVel\_RT\_GL(), initDonorFracAndMaxConVel\_RT\_TEOS(), initDonorFracAndMaxConVel\_RTP\_GL(), initDonorFracAndMaxConVel\_RTP\_GL(), initDonorFracAndMaxConVel\_RTP\_GL(), initDonorFracAndMaxConVel\_RTP\_GL(), initDonorFracAndMaxConVel\_RTP\_GL(), initDonorFracAndMaxConVel_RTP\_GL(), initDonorFracAndMaxConVel_RTP$ 

#### 6.8.3.14 double Parameters::dAlpha

This parameter controls the amount of extra mass above the outter interface. it is read in from the starting model, so that it will be consistent with the value used in calculating the starting model.

Referenced by  $calNewE_RT_NA_LES()$ ,  $calNewE_RTP_NA_LES()$ ,  $calNewU_R()$ ,  $calNewU_RLES()$ ,  $calNewU_RT()$ ,  $calNewU_RT_LES()$ ,  $calNewU_RTP()$ ,  $calNewU_RTP()$ ,  $calNewU_RTP()$ ,  $calNewU_RTP()$ 

RTP\_LES(), dImplicitEnergyFunction\_RT\_LES\_SB(), dImplicitEnergyFunction\_RTP\_-LES\_SB(), modelRead(), modelWrite\_GL(), modelWrite\_TEOS(), and Parameters().

#### 6.8.3.15 double Parameters::dTolerance

Amount of error to tolerate when calculating temperature from the equation of state.

Referenced by calNewTPKappaGamma TEOS(), and init().

#### 6.8.3.16 int Parameters::nMaxIterations

The maximum number of iterations to try to get the the relative error in the temperture below parameters::dTolerance.

Referenced by calNewTPKappaGamma TEOS(), and init().

#### 6.8.3.17 double Parameters::dEddyViscosity

Used in calculating the eddy viscosity, larger values will produce a larger value of the eddy viscosity, causing the rethermalization to happen at larger scales. This value should be kept small, a good value is 0.17, which seems to correspond with experiments.

 $\label{lem:control_relation} Referenced by calNewEddyVisc\_R\_CN(), calNewEddyVisc\_R\_SM(), calNewEddyVisc\_RT\_CN(), calNewEddyVisc\_RT\_SM(), calNewEddyVisc\_RTP\_CN(), calNewEddyVisc\_RTP\_CN(), calNewEddyVisc\_RTP\_SM(), calOldEddyVisc\_RTSM(), calOldEddyVisc\_RTSM(), calOldEddyVisc\_RTSM(), calOldEddyVisc\_RTP\_SM(), calOldEddyVisc\_RTP\_SM(), init(), and Parameters().$ 

#### 6.8.3.18 double Parameters::dMaxConvectiveVelocity

Holds the maximum convective velocity, it is set in the functions which calculate the timestep (see calDelt\_R\_GL, calDelt\_R\_TEOS, calDelt\_RT\_GL, calDelt\_RT\_TEOS, calDelt\_RTP\_GL, calDelt\_RTP\_TEOS, calDelt\_CONST).

 $\label{lem:reconstruction} Referenced by calDelt\_R\_GL(), calDelt\_R\_TEOS(), calDelt\_RT\_GL(), calDelt\_RT\_TEOS(), calDelt\_RTP\_GL(), calDelt\_RTP\_TEOS(), calNewEddyVisc\_R\_CN(), calNewEddyVisc\_RT\_CN(), calNewEddyVisc\_RTP\_CN(), calOldEddyVisc\_R\_CN(), calOldEddyVisc\_RT\_CN(), calOldEddyVisc\_RTP\_CN(), initDonorFracAndMaxConVel\_R\_GL(), initDonorFracAndMaxConVel\_RT\_TEOS(), initDonorFracAndMaxConVel\_RT\_GL(), initDonorFracAndMaxConVel\_RTT\_GS(), initDonorFracAndMaxConVel\_RTP\_GL(), initDonorFracAndMaxConVel$ 

#### 6.8.3.19 double Parameters::dMaxConvectiveVelocity c

Holds the maximum of convective velocity divided by the sound speed. It is set in the functions which calculate the timestep (see calDelt\_R\_GL, calDelt\_R\_TEOS, calDelt\_RT\_GL, calDelt\_RT\_TEOS, calDelt\_RT\_GL, calDelt\_RT\_TEOS, calDelt\_RTP\_GL, calDelt\_RTP\_GL

Referenced by Parameters().

#### 6.8.3.20 double Parameters::dPrt

This is the value of the Prandtl number, a value of 0.7 is what is suggested by Lawrence D. Cloutman in "The LUVD11 Large Eddy Simulation Model" April 15, 1991 a Lawrence Livermore National Labratory report.

 $\label{lem:referenced} Referenced & by & calNewE\_RT\_NA\_LES(), & calNewE\_RTP\_NA\_LES(), \\ dImplicitEnergyFunction\_RT\_LES(), & dImplicitEnergyFunction\_RT\_LES\_SB(), \\ dImplicitEnergyFunction\_RTP\_LES(), & dImplicitEnergyFunction\_RTP\_LES\_SB(), & and \\ Parameters(). & \\ Referenced & by & calNewE\_RTP\_NA\_LES(), & dImplicitEnergyFunction\_RTP\_LES\_SB(), & and \\ Referenced & by & calNewE\_RTP\_NA\_LES(), & dImplicitEnergyFunction\_RTP\_LES\_SB(), & and \\ Referenced & by & calNewE\_RTP\_LES\_SB(), & dImplicitEnergyFunction\_RTP\_LES\_SB(), & and \\ Referenced & dImplicitEnergyFunction\_RTP\_$ 

## 6.8.3.21 double Parameters::dT cut

The temperature at which to cut the DEDM gradient back Referenced by Parameters().

#### 6.8.3.22 double Parameters::dDEDM cut

The value to use for DEDM in energy conservation quation when DEDM becomes too large.

### 6.8.3.23 int Parameters::nDEDM cut zone

The zone at which the DEDM cut was made

#### 6.8.3.24 bool Parameters::bDEDM cut set

If Paramters::dDEDM\_cut has been set for this time step it will be true.

Referenced by main(), and Parameters().

## 6.8.3.25 std::string Parameters::sDebugProfileOutput

output file name for debuging profile, only used if DEBUG\_EQUATIONS is set to 1 Referenced by fin(), init(), and main().

The documentation for this class was generated from the following files:

- global.h
- global.cpp

# 6.9 Performance Class Reference

#include <global.h>

## **Public Member Functions**

• Performance ()

## Public Attributes

- double dStartTimer
- double dEndTimer

## 6.9.1 Detailed Description

This class manages information pertianing to performace analysis of the code.

#### 6.9.2 Constructor & Destructor Documentation

## 6.9.2.1 Performance::Performance ()

Constructor for the class Performance.

References dEndTimer, and dStartTimer.

## 6.9.3 Member Data Documentation

#### 6.9.3.1 double Performance::dStartTimer

The time that the code timer was started.

Referenced by fin(), init(), and Performance().

### 6.9.3.2 double Performance::dEndTimer

The time that the code timer was ended. The difference between Performance::dStartTimer and dEndTimer gives the total run time

Referenced by fin(), and Performance().

The documentation for this class was generated from the following files:

• global.h

• global.cpp

# 6.10 ProcTop Class Reference

#include cTop.h>

#### **Public Member Functions**

• ProcTop ()

#### Public Attributes

- int nNumProcs
- int \* nProcDims
- int \* nPeriodic
- int \*\* nCoords
- int nRank
- int nNumNeighbors
- int \* nNeighborRanks
- int nNumRadialNeighbors
- int \* nRadialNeighborRanks
- $\bullet$  int \* nRadialNeighborNeighborIDs

## 6.10.1 Detailed Description

This class manages information which pertains to the processor topology.

## 6.10.2 Constructor & Destructor Documentation

#### 6.10.2.1 ProcTop::ProcTop ()

Constructor for class ProcTop.

References nCoords, nNeighborRanks, nNumNeighbors, nNumRadialNeighbors, nPeriodic, nProcDims, nRadialNeighborNeighborIDs, and nRadialNeighborRanks.

## 6.10.3 Member Data Documentation

## 6.10.3.1 int ProcTop::nNumProcs

Number of processors in global communicator MPI::COMM\_WORLD. The value of this variable is independent of processor ProcTop::nRank.

Referenced by init(), initImplicitCalculation(), initUpdateLocalBoundaries(), initWatchZones(), modelRead(), and setupLocalGrid().

#### 6.10.3.2 int\* ProcTop::nProcDims

Dimensions of the processor topology. It is an array of size 3 to hold the size of the processor grid in each dimension. The value of this variable is set in the configuration file "config.xml" which is parsed by the function init. The values of this variable are independent of processor ProcTop::nRank.

Referenced by init(), initImplicitCalculation(), initUpdateLocalBoundaries(), initWatchZones(), modelRead(), modelWrite GL(), modelWrite TEOS(), ProcTop(), and setupLocalGrid().

#### 6.10.3.3 int\* ProcTop::nPeriodic

Periodic boundary conditions. It is an array of size 3 to tell if a dimension is periodic (wraps) or not. It contains an interger value of 0 or 1. 0, the boundary condition is not periodic, 1 the boundary condition is periodic. The value of this variable is set in the configuration file "config.xml" which is parsed by the function init. The values of this variable are independent of processor ProcTop::nRank.

Referenced by initUpdateLocalBoundaries(), modelRead(), modelWrite\_GL(), modelWrite\_-TEOS(), ProcTop(), and setupLocalGrid().

#### 6.10.3.4 int\*\* ProcTop::nCoords

Coordinates of the processors. It is of size ProcTop::nNumProcs by 3. The values of this variable are independent of processor ProcTop::nRank.

Referenced by calNewU0\_R(), calNewU0\_RT(), calNewU0\_RTP(), initImplicitCalculation(), initUpdateLocalBoundaries(), initWatchZones(), modelRead(), modelWrite\_GL(), modelWrite\_TEOS(), ProcTop(), and setupLocalGrid().

#### 6.10.3.5 int ProcTop::nRank

Is a unique integer which identifies the processor. The values of ProcTop::nRank range from 0 to ProcTop::nNumProcs-1 depending on the processor.

Referenced by calDelt\_CONST(), calDelt\_R\_GL(), calDelt\_R\_TEOS(), calDelt\_RT\_GL(), calDelt\_RT\_TEOS(), calDelt\_RTP\_GL(), calDelt\_RTP\_TEOS(), calNewD\_R(), calNewD\_RT(), calNewD\_RTP(), calNewE\_R\_AD(), calNewE\_R\_NA(), calNewE\_R\_NA\_-LES(), calNewE\_RT\_AD(), calNewE\_RT\_NA(), calNewE\_RT\_NA\_LES(), calNewE\_RTP\_-AD(), calNewE\_RTP\_NA(), calNewE\_RTP\_NA\_LES(), calNewU0\_R(), calNewU0\_RT(), calNewU0\_RTP(), fin(), implicitSolve\_R(), implicitSolve\_RT(), implicitSolve\_RTP(), init(), initImplicitCalculation(), initInternalVars(), initUpdateLocalBoundaries(), initWatchZones(), main(), modelRead(), modelWrite\_GL(), modelWrite\_TEOS(), setMainFunctions(), setupLocalGrid(), updateLocalBoundaries(), updateLocalBoundariesNewGrid(), and updateNewGridWithOld().

#### 6.10.3.6 int ProcTop::nNumNeighbors

The number of neighbors surrounding the current processor. The maximum number of neighbors possible is 27, 3x3x3 don't forget the current processor itself can be its own neighbor because of periodic boundary conditions. The value of this variable is dependent on processor ProcTop::nRank.

Referenced by initUpdateLocalBoundaries(), ProcTop(), updateLocalBoundaries(), and updateLocalBoundariesNewGrid().

### 6.10.3.7 int\* ProcTop::nNeighborRanks

ProcTop::nRank s of the neighboring processors. An array of size nNumNeighbors to hold ranks of neighbouring processors.

Referenced by initUpdateLocalBoundaries(), ProcTop(), updateLocalBoundaries(), and updateLocalBoundariesNewGrid().

## 6.10.3.8 int ProcTop::nNumRadialNeighbors

The number of neighbors in the radial direction. Can range from 1 to 2 depending on weather there is a processor beneath or above the current precessor.

Referenced by calNewU0\_R(), calNewU0\_RT(), calNewU0\_RTP(), initUpdateLocalBoundaries(), and ProcTop().

#### 6.10.3.9 int\* ProcTop::nRadialNeighborRanks

ProcTop::nRank s of the neighboring radial processors. It is an array of size ProcTop::nNumRadialNeighbors.

Referenced by calNewU0\_R(), calNewU0\_RT(), calNewU0\_RTP(), initUpdateLocalBoundaries(), and ProcTop().

#### 6.10.3.10 int\* ProcTop::nRadialNeighborNeighborIDs

 $\label{eq:holds} \begin{tabular}{ll} Holds the ID of a radialial neighbor, to be used to obtain their $\operatorname{ProcTop::nRank}$ from $\operatorname{ProcTop::nNeighborRanks}$ \\ \end{tabular}$ 

Referenced by  $calNewU0_R()$ ,  $calNewU0_RT()$ ,  $calNewU0_RTP()$ , initUpdateLocalBoundaries(), and ProcTop().

The documentation for this class was generated from the following files:

- procTop.h
- procTop.cpp

## 6.11 Time Class Reference

#include <time.h>

#### **Public Member Functions**

• Time ()

## Public Attributes

- double dDeltat np1half
- double dDeltat nm1half
- ullet double dDeltat n
- $\bullet$  double dt
- double dEndTime
- int nEndTimeStep
- double dTimeStepFactor
- int nTimeStepIndex
- bool bVariableTimeStep
- double dConstTimeStep
- double dPerChange
- double dDelRho t Rho max
- $\bullet \ \, double \, dDelT\_t\_T\_max \\$
- double dDelE t E max

## 6.11.1 Detailed Description

This class manages information which pertains to time variables.

#### 6.11.2 Constructor & Destructor Documentation

#### 6.11.2.1 Time::Time ()

Constructor for the class Time.

## 6.11.3 Member Data Documentation

#### 6.11.3.1 double Time::dDeltat np1half

The time step centered at n + 1/2 in seconds. It is used for calculating new variables defined at time step n, e.g. the density Grid::nD.

Referenced by calDelt CONST(), calDelt R GL(), calDelt R TEOS(), calDelt RT GL(), calDelt RT TEOS(), calDelt RTP GL(), calDelt RTP TEOS(), calNewD R(), calNewD - $RT(), \quad calNewD_RTP(), \quad calNewE_R_AD(), \quad calNewE_R_NA(), \quad calNewE_R_NA_LES(), \quad calNewE_R_NA_LES(), \quad calNewE_R_NA_LES(), \quad calNewE_R_NA(), \quad calNewE_R$ calNewE RT AD(), calNewE RT NA(), calNewE RT NA LES(), calNewE RTP AD(), calNewE RTP NA(), calNewE RTP NA LES(), calNewR(), dImplicitEnergyFunction - ${\it dImplicitEnergyFunction} \ \ {\it R} \ \ {\it LES()},$ dImplicitEnergyFunction R LES SB(), dImplicitEnergyFunction R SB(), dImplicitEnergyFunction RT(), dImplicitEnergyFunction -RT LES(), dImplicitEnergyFunction RT LES SB(), dImplicitEnergyFunction -RT SB(), dImplicitEnergyFunction RTP(), dImplicitEnergyFunction RTP LES(), dImplicitEnergyFunction RTP LES SB(), dImplicitEnergyFunction RTP SB(), main(), modelRead(), modelWrite TEOS(), and Time().

#### 6.11.3.2 double Time::dDeltat nm1half

The previously used timestep centered at n-1/2 in seconds. It is used for calculating dDeltat\_n the n centered time step.

 $\label{eq:const_const_const_const_const_const_const_const_const_relation} Referenced by calDelt_CONST(), calDelt_R_GL(), calDelt_R_T_TEOS(), calDelt_RTP_GL(), calDelt_RTP_TEOS(), modelRead(), modelWrite_GL(), and modelWrite_TEOS(). \\$ 

#### 6.11.3.3 double Time::dDeltat n

The time step centered at n in seconds. It is used for calculating new variables defined at time step n + 1/2, e.g. the radial velocity Grid::nU. This value is determined by averaging the current Time::dDeltat np1half, and the last Time::dDeltat np1half.

Referenced by calDelt\_CONST(), calDelt\_R\_GL(), calDelt\_R\_TEOS(), calDelt\_RT\_GL(), calDelt\_RT\_TEOS(), calDelt\_RTP\_GL(), calDelt\_RTP\_TEOS(), calNewU\_R(), calNewU\_-R\_LES(), calNewU\_RT(), calNewU\_RT\_LES(), calNewU\_RTP(), calNewU\_RTP\_LES(), calNewV\_RTP(), calNewV\_RTP\_LES(), calNewV\_RTP\_LES(), calNewV\_RTP(), calNewV\_RTP(),

#### 6.11.3.4 double Time::dt

The current time of the simulation in seconds.

 $\label{eq:referenced} Referenced by calDelt\_CONST(), calDelt\_R\_GL(), calDelt\_RT\_TEOS(), calDelt\_RT\_GL(), calDelt\_RT\_TEOS(), calDelt\_RTP\_GL(), calDelt\_RTP\_TEOS(), fin(), init(), main(), modelRead(), modelWrite\_GL(), modelWrite\_TEOS(), Time(), writeWatchZones\_R\_GL(), writeWatchZones\_R_TEOS(), writeWatchZones\_RT\_GL(), writeWatchZones\_RT\_TEOS(), writeWatchZones\_RTP\_GL(), and writeWatchZones\_RTP\_TEOS().$ 

#### 6.11.3.5 double Time::dEndTime

The end time of the current calculation in seconds.

Referenced by init(), main(), and Time().

#### 6.11.3.6 int Time::nEndTimeStep

The last time step to calculate, will stop if the current time step is larger than this. The default value is the largest integer of the system.

Referenced by init(), main(), and Time().

#### 6.11.3.7 double Time::dTimeStepFactor

Used for determining the time step. It is the factor which the courrant time step is multiplied by in order to determine Time::dDeltat np1half.

Referenced by calDelt\_R\_GL(), calDelt\_R\_TEOS(), calDelt\_RT\_GL(), calDelt\_RT\_TEOS(), calDelt\_RTP\_GL(), calDelt\_RTP\_TEOS(), init(), and Time().

#### 6.11.3.8 int Time::nTimeStepIndex

An index indecating the current time step. An index of zero corresponds to a Time::dt=0.

#### Todo

should probably make this an unsigned variable, and perhaps also use the keyword long to help ensure there are enough values. Often need 7 decimal places.

#### 6.11.3.9 bool Time::bVariableTimeStep

If true a variable time step is used as specified by the Courant condition, times the dTimeStepFactor.

Referenced by init(), and setMainFunctions().

#### 6.11.3.10 double Time::dConstTimeStep

If set to a value other than 0, will use that constant time step in place of the courant time step. Referenced by calDelt\_CONST(), and init().

#### 6.11.3.11 double Time::dPerChange

A percentage amount to allow the maximum horizontal temperture variation and radial, theta and phi convective velocities to change by from one time step to the next. The time step is reduced accordingly to keep this precent change intact.

Referenced by calDelt\_R\_GL(), calDelt\_R\_TEOS(), calDelt\_RT\_GL(), calDelt\_RT\_TEOS(), calDelt\_RTP\_GL(), calDelt\_RTP\_TEOS(), init(), and Time().

#### 6.11.3.12 double Time::dDelRho t Rho max

Keeps track of the maximum relative change in density from one time step to the next.

 $Referenced \ by \ calDelt\_R\_GL(), \ calDelt\_R\_TEOS(), \ calDelt\_RT\_GL(), \ calDelt\_RT\_TEOS(), \ calDelt\_RTP\_GL(), \ calDelt\_RTP\_TEOS(), \ fin(), \ main(), \ and \ Time().$ 

#### $6.11.3.13 \quad double \ Time:: dDelT \quad t \quad T \quad max \\$

Keeps track of the maximum relative change in temperature from one time step to the next. This quantity is only tracked if the calculation is non-adiabatic, else the energy is tracked instead, see Time::dDelE\_t\_E\_max

Referenced by calDelt\_R\_TEOS(), calDelt\_RT\_TEOS(), calDelt\_RTP\_TEOS(), fin(), main(), and Time().

#### 6.11.3.14 double Time::dDelE t E max

Keeps track of the maximum relative change in energy from one time step to the next. This quantity is only tracked if the calculation is adiabatic, else the temperature is tracked instead, see  $Time::dDelT\_t\_T\_max$ 

Referenced by calDelt R GL(), calDelt RT GL(), calDelt RTP GL(), and Time().

The documentation for this class was generated from the following files:

- time.h
- time.cpp

### 6.12 WatchZone Class Reference

#include <watchzone.h>

### 6.12.1 Detailed Description

This class contains information used to monitor a particular zone of the grid.

The documentation for this class was generated from the following files:

- $\bullet$  watchzone.h
- $\bullet \ \ watchzone.cpp$

### Chapter 7

### File Documentation

# $7.1 \quad /home/cgeroux/WORK/SPHERLS/src/eos.cpp \quad \quad File \\ Reference$

```
#include <string>
#include <fstream>
#include <sstream>
#include <iostream>
#include <cmath>
#include "eos.h"
#include "exception2.h"
```

### 7.1.1 Detailed Description

Implements the eos (equation of state) class defined in eos.h

# $7.2 \quad /home/cgeroux/WORK/SPHERLS/src/eos.h \ \, File \ \, Reference$

```
#include <string>
#include "exception2.h"
```

#### Classes

• class eos

### 7.2.1 Detailed Description

 ${\rm Header\ file\ for\ } \frac{}{\cos . cpp}$ 

### 7.3 dataManipulation.cpp File Reference

```
#include <cmath>
#include <sstream>
#include <fstream>
#include <iomanip>
#include <vector>
#include <fenv.h>
#include "dataManipulation.h"
#include "global.h"
#include "xmlFunctions.h"
#include "exception2.h"
#include "dataMonitoring.h"
#include "physEquations.h"
#include <string>
```

#### **Functions**

- void init (ProcTop &procTop, Grid &grid, Output &output, Time &time, Parameters &parameters, MessPass &messPass, Performance &performance, Implicit &implicit, int nNumArgs, char \*cArgs[])
- void setupLocalGrid (ProcTop &procTop, Grid &grid)
- void fin (bool bWriteCurrentStateToFile, Time &time, Output &output, ProcTop &procTop, Grid &grid, Parameters &parameters, Functions &functions, Performance &performance, Implicit &implicit)
- void modelWrite\_GL (std::string sFileName, ProcTop &procTop, Grid &grid, Time &time, Parameters &parameters)
- void modelWrite\_TEOS (std::string sFileName, ProcTop &procTop, Grid &grid, Time &time, Parameters &parameters)
- void modelRead (std::string sFileName, ProcTop &procTop, Grid &grid, Time &time, Parameters &parameters)
- void initUpdateLocalBoundaries (ProcTop &procTop, Grid &grid, MessPass &messPass, Implicit &implicit)
- void updateLocalBoundaries (ProcTop &procTop, MessPass &messPass, Grid &grid)
- void updateLocalBoundariesNewGrid (int nVar, ProcTop &procTop, MessPass &messPass, Grid &grid)
- void updateOldGrid (ProcTop &procTop, Grid &grid)
- void updateNewGridWithOld (Grid &grid, ProcTop &procTop)
- void average3DTo1DBoundariesOld (Grid &grid)
- void average3DTo1DBoundariesNew (Grid &grid, int nVar)
- void updateLocalBoundaryVelocitiesNewGrid\_R (ProcTop &procTop, MessPass &mess-Pass, Grid &grid)

• void updateLocalBoundaryVelocitiesNewGrid\_RT (ProcTop &procTop, MessPass &mess-Pass, Grid &grid)

- void updateLocalBoundaryVelocitiesNewGrid\_RTP (ProcTop &procTop, MessPass &mess-Pass, Grid &grid)
- void initImplicitCalculation (Implicit & Emplicit, Grid & Grid, ProcTop & procTop, int nNumArgs, char \*cArgs[])

#### 7.3.1 Detailed Description

This file holds functions for manipulating data. This includes initializing the program, parsing the configuration file "config.xml", allocating memory for the model to be read in, reading in the input model, etc.

#### 7.3.2 Function Documentation

#### 7.3.2.1 void average3DTo1DBoundariesNew (Grid & grid, int nVar)

This function averages the 3D boundary recieved by the 1D processor (ProcTop::nRank ==0) into 1D. This average is volume weighted. This function only needs to be called by the 1D processor, and if called by other processors may have unexpected results. This function calculates the average from the new grid, and places the average into new old grid. It does so for only the specified variable. This function is used every time the grid boundaries are updated with updateLocalBoundariesNewGrid.

#### Parameters:

- $\leftrightarrow grid$  supplies the information for calculating the averages and recieves the averages.
- $\leftarrow n \, Var$  index of the variable to be averaged with in the grid.

 $\label{lem:condition} References & Grid::dLocalGridNew, & Grid::dLocalGridOld, & Grid::nCenIntOffset, \\ Grid::nDCosThetaIJK, & Grid::nDPhi, & Grid::nEndGhostUpdateExplicit, & Grid::nR, \\ Grid::nStartGhostUpdateExplicit, and & Grid::nVariables. \\ \\$ 

Referenced by updateLocalBoundariesNewGrid().

#### 7.3.2.2 void average3DTo1DBoundariesOld (Grid & grid)

This function averages the 3D boundary recieved by the 1D processor (ProcTop::nRank ==0) into 1D. This average is volume weighted. This function only needs to be called by the 1D processor, and if called by other processors may have unexpected results. This function calculates the average from the old grid, and places the average into the old grid. It does so for all variables external and internal. This function is used every time the grid boundaries are updated with updateLocalBoundaries.

#### Parameters:

 $\leftrightarrow grid$  supplies the information for calculating the averages and recieves the averages.

References Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nDCosThetaIJK, Grid::nDPhi, Grid::nEndGhostUpdateExplicit, Grid::nNumIntVars, Grid::nNumVars, Grid::nNumVars, Grid::nNumVars, Grid::nStartGhostUpdateExplicit, and Grid::nVariables.

Referenced by updateLocalBoundaries().

# 7.3.2.3 void fin (bool bWriteCurrentStateToFile, Time & time, Output & output, ProcTop & procTop, Grid & grid, Parameters & parameters, Functions & functions, Performance & performance, Implicit & implicit)

Finishes program execution by writing out last grid state, closing output files, and writting out run time.

#### Parameters:

- ← **bWriteCurrentStateToFile** is a bool value which indicates wheather or not to write out current model state.
- $\leftarrow time$
- $\leftarrow output$
- $\leftarrow procTop$
- $\leftarrow qrid$
- $\leftarrow$  parameters
- $\leftarrow functions$
- $\leftarrow performance$
- $\leftarrow implicit$

References Implicit::dAverageRHS, Implicit::dCurrentRelTError, Time::dDelRho t Rho -Time::dDelT t T max, Time::dDeltat np1half, Time::dDelUmU0 t UmU0 -Time::dDelW t W max, Performance::dEndTimer, Pa-Time::dDelV t V max, rameters::dMaxConvectiveVelocity, Implicit::dMaxErrorInRHS, Performance::dStartTimer, finWatchZones(), Time::dt, Functions::fpModelWrite, Implicit::nCurrentNumIterations, Implicit::nMaxNumSolverIterations, Grid::nNumDims, Implicit::nNumImplicitZones, Output::nNumTimeStepsSinceLastPrint, Output::nPrintMode, ProcTop::nRank, Time::nTimeStepIndex, Output::sBaseOutputFileName, and Parameters::sDebugProfileOutput.

Referenced by main().

# 7.3.2.4 void init (ProcTop & procTop, Grid & grid, Output & output, Time & time, Parameters & parameters, MessPass & messPass, Performance & performance, Implicit & implicit, int argc, char \* argv[])

Initializes the program. It does this by reading a number of configuration options from the config file "SPHERLS.xml". It also reads in the starting model, as specified in the "SPHERLS.xml" file, using the function modelRead. During the reading of the initial model the modelRead function also calls setupLocalGrid to determine the sizes of the local grids and allocate memory for them.

Other things of note that are done in this function are:

- the calulation timer is started, Performance::dStartTimer
- It also reads in the equation of state table if using a tabulated equation of state (Parameters::bEOSGammaLaw = false) by calling eos::readBin

• Initilizes the watchZones, i.e. figure out which processors have which watch zones, opens the files and prints headers.

#### Parameters:

- $\rightarrow$  procTop all parts of this stucture are set, and do not change thoughout the rest of the calculation.
- $\rightarrow$  grid through the function modelRead the function setupLocalGrid is called to allocate memory for the grid, and set sizes of it.
- $\rightarrow output$
- $\rightarrow time$
- $\rightarrow$  parameters
- ightarrow messPass
- $\rightarrow$  performance
- $\rightarrow implicit$
- $\leftarrow argc$
- $\leftarrow argv$

References Parameters::bAdiabatic, Output::bDump, Parameters::bEOSGammaLaw, put::bPrint, Time::bVariableTimeStep, Parameters::dA, Parameters::dAlphaExtra, Parameters::dAVThreshold, Time::dConstTimeStep, Implicit::dDerivativeStepFraction, eters::dDonorCellMultiplier, Output::dDumpFrequencyTime, Parameters::dEddyViscosity, Time::dEndTime, Time::dPerChange, Output::dPrintFrequencyTime, Performance::dStartTimer, Time∷dt, Output::dTimeLastDump, Output::dTimeLastPrint, Time::dTimeStepFactor, Implicit::dTolerance, Parameters::dTolerance, Parameters::eosTable, initImplicitCalculation(), initInternalVars(), initUpdateLocalBoundaries(), initWatchZones(), Read(),Output::nDumpFrequencyStep, Time::nEndTimeStep, Grid::nGlobalGridDims, Parameters::nMaxIterations, Implicit::nMaxNumIterations, Grid::nNum1DZones, Implicit::nNumImplicitZones, ProcTop::nNumProcs, Output::nPrintFrequencyStep, Output::nPrintMode, ProcTop::nProcDims, ProcTop::nRank, Parameters::nTypeTurbulanceMod, eos::readBin(), Output::sBaseOutputFileName, Parameters::sDebugProfileOutput, and Parameters::sEOSFileName.

Referenced by main().

### 7.3.2.5 void initImplicitCalculation (Implicit & implicit, Grid & grid, ProcTop & procTop, int nNumArgs, char \* cArgs[])

This function initilizes data structures and defines indixes of non-zero elements in the coeffecient matrix. It also sets up pathways for collection of the temperature corrections back to the processors which need them for their local grids.

- $\leftrightarrow implicit$
- $\leftarrow$  *grid* size information of the grid is used
- $\leftarrow procTop$
- $\leftarrow nNumArgs$  number of command line arguments, PETSc wants them
- $\leftarrow cArgs$  a list of command line arguments, PETSc wants them

#### Todo

isFrom, isTo, matCoeff,vecTCorrections, vecTCorrections,vecRHS,vecTCorrectionsLocal,kspContext,vecscatTCorrections all need to be destroyed before program finishes.

References Implicit::dTolerance, Implicit::kspContext, Implicit::matCoeff, Proc-Grid::nLocalGridDims, Top::nCoords, Grid::nGlobalGridDims, Implicit::nLocDer, Implicit::nLocFun, Implicit::nMaxNumIterations, Implicit::nNumDerPerRow, Grid::nNumDims, Grid::nNumGhostCells, Implicit::nNumImplicitZones, ProcTop::nNumProcs, Im-Implicit::nNumRowsALocalSB, ProcTop::nProcDims, plicit::nNumRowsALocal, Proc-Top::nRank, Grid::nT, Implicit::nTypeDer, Implicit::vecRHS, Implicit::vecscatTCorrections, Implicit::vecTCorrections, and Implicit::vecTCorrectionsLocal.

Referenced by init().

## 7.3.2.6 void initUpdateLocalBoundaries (ProcTop & procTop, Grid & grid, MessPass & messPass, Implicit & implicit)

Sets up MPI derived data types used for updating the local grid boundaries between processors. It sets where the local grids should start/stop updating the local grids (Grid::nStartUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nStartUpdateImplicit, Grid::nEndUpdateImplicit, Grid::nEndUpdateExplicit, Grid::nEndGhostUpdateExplicit, Grid::nStartGhostUpdateImplicit, Grid::nEndGhostUpdateImplicit). It sets the radial processor neighbors (ProcTop::nNumRadialNeighbors).

It also allocates memeory for:

- MessPass::requestSend
- MessPass::requestRecv
- statusSend
- statusRecv

#### Parameters:

- $\leftrightarrow procTop$
- $\leftrightarrow grid$
- $\leftrightarrow messPass$
- $\leftrightarrow implicit$

Grid::dLocalGridNew, Grid::nCenIntOffset, References Grid::dLocalGridOld, Proc-Top::nCoords, Grid::nE, Grid::nEndGhostUpdateExplicit, Grid::nEndGhostUpdateImplicit, Grid::nEndUpdateExplicit, Grid::nEndUpdateImplicit, Grid::nGlobalGridDims, Grid::nGlobalGridPositionLocalGrid, Grid::nKappa, Grid::nLocalGridDims, ProcTop::nNeighborRanks, Grid::nNumGhostCells, Grid::nNum1DZones, plicit::nNumImplicitZones, Grid::nNumIntVars, ProcTop::nNumNeighbors, ProcTop::nNumProcs, ProcTop::nNumRadialNeighbors, Grid::nNumVars, Grid::nNumZones1DBoundaryZeroHorizontalVelocity, Grid::nP, ProcTop::nPeriodic, Top::nProcDims, ProcTop::nRadialNeighborNeighborIDs, ProcTop::nRadialNeighborRanks, Grid::nStartGhostUpdateExplicit, Grid::nStartGhostUpdateImplicit, ProcTop::nRank, Grid::nStartUpdateExplicit, Grid::nStartUpdateImplicit, Grid::nT, Grid::nV, Grid::nVariables, Mess-Grid::nW, MessPass::requestRecv, MessPass::requestSend, MessPass::statusRecv, Pass::statusSend. MessPass::tvpeRecvNewVar. MessPass::typeRecvOldGrid, Mess-Pass::typeSendNewGrid, and MessPass::typeSendNewVar.

Referenced by init().

### 7.3.2.7 void modelRead (std::string sFileName, ProcTop & procTop, Grid & grid, Time & time, Parameters & parameters)

Reads in a collected binary file into the local grid and calls setupLocalGrid to allocate memory and set various parameters of the model. Works for both gamma-law gas, and tabulated equation of state models.

#### Parameters:

- $\leftarrow sFileName$  name of the file containing the model to be read in
- $\rightarrow procTop$
- $\rightarrow grid$
- $\rightarrow time$
- $\rightarrow$  parameters

#### Todo

At some point should get it working with only 1 processor

References Parameters::bEOSGammaLaw, Parameters::dA, Parameters::dAlpha, Parameters: ters::dAVThreshold, Time::dDeltat n, Time::dDeltat nm1half, Time::dDeltat np1half, Parameters::dGamma, Grid::dLocalGridOld, Time::dt, DUMP VERSION, ProcTop::nCoords, Grid::nCotThetaIJK, Grid::nCotThetaIJp1halfK, Grid::nD, Grid::nDCosThetaIJK, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nE, Grid::nEddyVisc, Grid::nGamma, Grid::nGlobalGridDims, Grid::nKappa, Grid::nLocalGridDims, Grid::nM, Grid::nNum1DZones, Grid::nNumDims, Grid::nNumGhostCells, Grid::nNumIntVars, ProcTop::nNumProcs, Grid::nNumVars, Grid::nP, ProcTop::nPeriodic, ProcTop::nProcDims, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR,ProcTop::nRank, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nT, Grid::nTheta, Time::nTimeStepIndex, Parameters::nTypeTurbulanceMod, Grid::nU, Grid::nU0, Grid::nV, Grid::nVariables, Grid::nW, Parameters::sEOSFileName, setInternalVarInf(), and setupLocalGrid().

Referenced by init().

## 7.3.2.8 void modelWrite\_GL (std::string sFileName, ProcTop & procTop, Grid & grid, Time & time, Parameters & parameters)

Writes out a model in distributed model format, meaning that each processor writes it's own local grid to a file in binary format. They can be combined, and or converted to ascii format using SPHERLSanal. This is for a gamma-law gas model.

#### Parameters:

- $\leftarrow sFileName$  base name of the output files
- $\leftarrow procTop$
- $\leftarrow grid$
- $\leftarrow time$
- $\leftarrow$  parameters

References Parameters::dA, Parameters::dAlpha, Parameters::dAVThreshold, Time::dDeltat\_-nm1half, Parameters::dGamma, Grid::dLocalGridOld, Time::dt, DUMP\_VERSION, ProcTop::nCoords, Grid::nGlobalGridDims, Grid::nLocalGridDims, Grid::nNum1DZones,

 $Grid::nNumGhostCells, \quad Grid::nNumVars, \quad ProcTop::nPeriodic, \quad ProcTop::nProcDims, \quad ProcTop::nRank, \quad Time::nTimeStepIndex, \quad and \quad Grid::nVariables.$ 

Referenced by setMainFunctions().

### 7.3.2.9 void modelWrite\_TEOS (std::string sFileName, ProcTop & procTop, Grid & grid, Time & time, Parameters & parameters)

Writes out a model in distributed model format, meaning that each processor writes it's own local grid to a file in binary format. They can be combined, and or converted to ascii format using SPHERLSanal. This is for a tabulated equation of state model.

#### Parameters:

- $\leftarrow sFileName$  base name of the output files
- $\leftarrow procTop$
- $\leftarrow grid$
- $\leftarrow time$
- $\leftarrow parameters$

References Parameters::dA, Parameters::dAlpha, Parameters::dAVThreshold, Time::dDeltat\_nm1half, Time::dDeltat\_np1half, Grid::dLocalGridOld, Time::dt, DUMP\_VERSION, ProcTop::nCoords, Grid::nGlobalGridDims, Grid::nLocalGridDims, Grid::nNum1DZones, Grid::nNumGhostCells, Grid::nNumVars, ProcTop::nPeriodic, ProcTop::nProcDims, ProcTop::nRank, Time::nTimeStepIndex, Grid::nVariables, and Parameters::sEOSFileName.

Referenced by setMainFunctions().

#### 7.3.2.10 void setupLocalGrid (ProcTop & procTop, Grid & grid)

Determins size of local grids (Grid::nLocalGridDims) based on processor topology, and allocates memory for the local grids (Grid::dLocalGridNew, Grid::dLocalGridOld). It sets various other quantities aswell such as,

- the coordinates of all processors (ProcTop::nCoords)
- the offset for interface centered quantities (Grid::nCenIntOffset, which depends on zoning and boundary conditions
- the position the local grid is in relative to the global grid (Grid::nGlobalGridPositionLocalGrid).

#### Parameters:

- $\leftrightarrow procTop$  contains information about the processor topology
- $\leftrightarrow$  grid contains information about gird

References Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nCenIntOffset, ProcTop::nCoords, Grid::nD, Grid::nGlobalGridDims, Grid::nGlobalGridPositionLocalGrid, Grid::nLocalGridDims, Grid::nNum1DZones, Grid::nNumDims, Grid::nNumGhostCells, Grid::nNumIntVars, ProcTop::nNumProcs, Grid::nNumVars, ProcTop::nPeriodic, ProcTop::nProcDims, ProcTop::nRank, and Grid::nVariables.

Referenced by modelRead().

### 7.3.2.11 void updateLocalBoundaries (ProcTop & procTop, MessPass & messPass, Grid & grid)

Updates the boundaries of the local grids from the data in the local grids of other processors. It does this for all variables and updates to the old grid. It also has processor ProcTop::nRank=0 call average3DTo1DBoundariesOld which averages the 3D information into the 1D boundaries.

#### Parameters:

 $\leftarrow procTop \\ \leftarrow messPass \\ \leftrightarrow grid$ 

#### Todo

Shouldn't need MPI::COMM\_WORLD.Barrier() may want to test out removing this at some point as it might produce a bit of a speed up.

 $References \quad average 3DTo1DBoundaries Old(), \quad Grid::dLocalGridNew, \quad Grid::dLocalGridOld, \\ ProcTop::nNeighborRanks, \quad ProcTop::nNumNeighbors, \quad ProcTop::nRank, \quad Mess-Pass::requestRecv, \quad Mess-Pass::statusRecv, \quad Mess-Pass::statusSend, \\ Mess-Pass::type-RecvOldGrid, \quad Mess-Pass::type-SendNewGrid, \\ and \quad updateOldGrid().$ 

Referenced by main().

### 7.3.2.12 void updateLocalBoundariesNewGrid (int nVar, ProcTop & procTop, MessPass & messPass, Grid & grid)

Updates the boundaries of the local grids from the data in the local grids of other processors. It does this for a specific variable specified by nVar and updates to the new grid. It also has processor ProcTop::nRank=0 call average3DTo1DBoundariesNew which averages the 3D information into the 1D boundaries for that specific variable.

#### Parameters:

 $\leftarrow procTop \\ \leftarrow messPass \\ \leftrightarrow grid$ 

#### Todo

May want to do some waiting on this message at some point before the end of the timestep, but it doesn't need to be done in this function. It might also be that this is built into the code by waiting at some other point. This is something that should be checked out at somepoint, perhaps once the preformance starts to be analyzed. I would think that if the send buffer was being modified before the send was completed, that there would be some errors poping up that would likely kill the program.

 $References\ average 3DTo 1DBoundaries New (),\ Grid:: dLocal Grid New,\ ProcTop:: nNeighbor Ranks,\ ProcTop:: nNumNeighbors,\ ProcTop:: nRank,\ MessPass:: request Recv,\ MessPass:: status Recv,\ MessPass:: type RecvNew Var,\ and\ MessPass:: type SendNew Var.$ 

 $Referenced \quad by \quad implicitSolve\_R(), \quad implicitSolve\_RT(), \quad implicitSolve\_RTP(), \quad main(), \\ updateLocalBoundaryVelocitiesNewGrid\_R(), \quad updateLocalBoundaryVelocitiesNewGrid\_RT(), \\ and \quad updateLocalBoundaryVelocitiesNewGrid \quad RTP().$ 

### 7.3.2.13 void updateLocalBoundaryVelocitiesNewGrid\_R (ProcTop & procTop, MessPass & messPass, Grid & grid)

Updates velocity boundaries of the new grid in a 1D calculations after the velocities have been newly calculated.

References Grid::nU, and updateLocalBoundariesNewGrid().

Referenced by setMainFunctions().

### 7.3.2.14 void updateLocalBoundaryVelocitiesNewGrid\_RT (ProcTop & procTop, MessPass & messPass, Grid & grid)

Updates velocity boundaries of the new grid in a 2D calculations after the velocities have been newly calculated.

References Grid::nU, Grid::nV, and updateLocalBoundariesNewGrid().

Referenced by setMainFunctions().

## 7.3.2.15 void updateLocalBoundaryVelocitiesNewGrid\_RTP (ProcTop & procTop, MessPass & messPass, Grid & grid)

Updates velocity boundaries of the new grid in a 3D calculations after the velocities have been newly calculated.

References Grid::nU, Grid::nV, Grid::nW, and updateLocalBoundariesNewGrid().

Referenced by setMainFunctions().

#### 7.3.2.16 void updateNewGridWithOld (Grid & grid, ProcTop & procTop)

Copies the contents of the old grid to the new grid including ghost cells.

#### Parameters:

- $\leftrightarrow grid$
- $\leftarrow procTop$

References Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nLocalGridDims, Grid::nNumDims, Grid::nNumGhostCells, Grid::nNumIntVars, Grid::nNumVars, Proc-Top::nRank, and Grid::nVariables.

Referenced by main().

#### 7.3.2.17 void updateOldGrid (ProcTop & procTop, Grid & grid)

Updates the old grid with the new grid, not including boundaries.

- $\leftarrow procTop$
- $\leftrightarrow \textit{grid}$

References Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nEndGhostUpdateExplicit, Grid::nEndGhostUpdateImplicit, Grid::nEndUpdateExplicit, Grid::nEndUpdateImplicit, Grid::nNumIntVars, Grid::nNumVars, Grid::nStartGhostUpdateExplicit, Grid::nStartGhostUpdateImplicit, Grid::nStartUpdateExplicit, and Grid::nStartUpdateImplicit.

Referenced by updateLocalBoundaries().

### 7.4 dataManipulation.h File Reference

```
#include <mpi.h>
#include "global.h"
```

#### **Functions**

- void init (ProcTop &procTop, Grid &grid, Output &output, Time &time, Parameters &parameters, MessPass &messPass, Performance &performance, Implicit &implicit, int argc, char \*argv[])
- void setupLocalGrid (ProcTop &procTop, Grid &grid)
- void fin (bool bWriteCurrentStateToFile, Time &time, Output &output, ProcTop &procTop, Grid &grid, Parameters &parameters, Functions &functions, Performance &performance, Implicit &implicit)
- void modelWrite\_GL (std::string sFileName, ProcTop &procTop, Grid &grid, Time &time, Parameters &parameters)
- void modelWrite\_TEOS (std::string sFileName, ProcTop &procTop, Grid &grid, Time &time, Parameters &parameters)
- void modelRead (std::string sFileName, ProcTop &procTop, Grid &grid, Time &time, Parameters &parameters)
- void initUpdateLocalBoundaries (ProcTop &procTop, Grid &grid, MessPass &messPass, Implicit &implicit)
- void updateLocalBoundaries (ProcTop &procTop, MessPass &messPass, Grid &grid)
- void updateLocalBoundariesNewGrid (int nVar, ProcTop &procTop, MessPass &messPass, Grid &grid)
- void updateOldGrid (ProcTop &procTop, Grid &grid)
- void updateNewGridWithOld (Grid &grid, ProcTop &procTop)
- void average3DTo1DBoundariesOld (Grid &grid)
- void average3DTo1DBoundariesNew (Grid &grid, int nVar)
- void updateLocalBoundaryVelocitiesNewGrid\_R (ProcTop &procTop, MessPass &mess-Pass, Grid &grid)
- void updateLocalBoundaryVelocitiesNewGrid\_RT (ProcTop &procTop, MessPass &mess-Pass, Grid &grid)
- void updateLocalBoundaryVelocitiesNewGrid\_RTP (ProcTop &procTop, MessPass &mess-Pass, Grid &grid)
- void initImplicitCalculation (Implicit & Emplicit, Grid & Grid, ProcTop & procTop, int nNumArgs, char \*cArgs[])

#### 7.4.1 Detailed Description

Header file for dataManipulation.cpp

#### 7.4.2 Function Documentation

#### 7.4.2.1 void average3DTo1DBoundariesNew (Grid & grid, int nVar)

This function averages the 3D boundary recieved by the 1D processor (ProcTop::nRank ==0) into 1D. This average is volume weighted. This function only needs to be called by the 1D processor, and if called by other processors may have unexpected results. This function calculates the average from the new grid, and places the average into new old grid. It does so for only the specified variable. This function is used every time the grid boundaries are updated with updateLocalBoundariesNewGrid.

#### Parameters:

- $\leftrightarrow grid$  supplies the information for calculating the averages and recieves the averages.
- $\leftarrow n Var$  index of the variable to be averaged with in the grid.

 $\label{lem:condition} References & Grid::dLocalGridNew, & Grid::dLocalGridOld, & Grid::nCenIntOffset, \\ Grid::nDCosThetaIJK, & Grid::nDPhi, & Grid::nEndGhostUpdateExplicit, & Grid::nR, \\ Grid::nStartGhostUpdateExplicit, & and Grid::nVariables. \\ \\$ 

Referenced by updateLocalBoundariesNewGrid().

#### 7.4.2.2 void average3DTo1DBoundariesOld (Grid & grid)

This function averages the 3D boundary recieved by the 1D processor (ProcTop::nRank ==0) into 1D. This average is volume weighted. This function only needs to be called by the 1D processor, and if called by other processors may have unexpected results. This function calculates the average from the old grid, and places the average into the old grid. It does so for all variables external and internal. This function is used every time the grid boundaries are updated with updateLocalBoundaries.

#### Parameters:

 $\leftrightarrow$  grid supplies the information for calculating the averages and receives the averages.

References Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nDCosThetaIJK, Grid::nDPhi, Grid::nEndGhostUpdateExplicit, Grid::nNumIntVars, Grid::nNumVars, Grid::nNumVars, Grid::nStartGhostUpdateExplicit, and Grid::nVariables.

Referenced by updateLocalBoundaries().

7.4.2.3 void fin (bool bWriteCurrentStateToFile, Time & time, Output & output, ProcTop & procTop, Grid & grid, Parameters & parameters, Functions & functions, Performance & performance, Implicit & implicit)

Finishes program execution by writing out last grid state, closing output files, and writting out run time.

- $\leftarrow b \, Write Current State To File$  is a bool value which indicates wheather or not to write out current model state.
- $\leftarrow time$
- $\leftarrow output$
- $\leftarrow procTop$

- $\leftarrow grid$
- $\leftarrow parameters$
- $\leftarrow functions$
- $\leftarrow performance$
- $\leftarrow implicit$

References Implicit::dAverageRHS, Implicit::dCurrentRelTError, Time::dDelRho t Rho -Time::dDeltat np1half, Time::dDelUmU0 t UmU0 - $Time::dDelT_t_T_max,$ Time::dDelV t V max, Time::dDelW t W max, Performance::dEndTimer, Pamax, rameters::dMaxConvectiveVelocity, Implicit::dMaxErrorInRHS, Performance::dStartTimer, finWatchZones(), Time::dt, Functions::fpModelWrite, Implicit::nCurrentNumIterations, Implicit::nMaxNumSolverIterations, Grid::nNumDims, Implicit::nNumImplicitZones, Output::nNumTimeStepsSinceLastPrint, Output::nPrintMode, ProcTop::nRank, Time::nTimeStepIndex, Output::sBaseOutputFileName, and Parameters::sDebugProfileOutput. Referenced by main().

# 7.4.2.4 void init (ProcTop & procTop, Grid & grid, Output & output, Time & time, Parameters & parameters, MessPass & messPass, Performance & performance, Implicit & implicit, int argc, char \* argv[])

Initializes the program. It does this by reading a number of configuration options from the config file "SPHERLS.xml". It also reads in the starting model, as specified in the "SPHERLS.xml" file, using the function modelRead. During the reading of the initial model the modelRead function also calls setupLocalGrid to determine the sizes of the local grids and allocate memory for them.

Other things of note that are done in this function are:

- the calulation timer is started, Performance::dStartTimer
- It also reads in the equation of state table if using a tabulated equation of state (Parameters::bEOSGammaLaw = false) by calling eos::readBin
- Initilizes the watchZones, i.e. figure out which processors have which watch zones, opens the files and prints headers.

- $\rightarrow procTop$  all parts of this stucture are set, and do not change thoughout the rest of the calculation.
- $\rightarrow$  grid through the function modelRead the function setupLocalGrid is called to allocate memory for the grid, and set sizes of it.
- $\rightarrow output$
- $\rightarrow time$
- ightarrow parameters
- $\rightarrow messPass$
- $\rightarrow$  performance
- $\rightarrow implicit$
- $\leftarrow argc$
- $\leftarrow argv$

References Parameters::bAdiabatic, Output::bDump, Parameters::bEOSGammaLaw, put::bPrint, Time::bVariableTimeStep, Parameters::dA, Parameters::dAlphaExtra, Param-Time::dConstTimeStep, Implicit::dDerivativeStepFraction, eters::dAVThreshold, eters::dDonorCellMultiplier, Output::dDumpFrequencyTime, Parameters::dEddyViscosity, Time::dEndTime, Time::dPerChange, Output::dPrintFrequencyTime, Performance::dStartTimer, Output::dTimeLastDump, Output::dTimeLastPrint, Time::dTimeStepFactor, Implicit::dTolerance, Parameters::dTolerance, Parameters::eosTable, initImplicitCalculation(), initInternalVars(), initUpdateLocalBoundaries(), initWatchZones(), model-Output::nDumpFrequencyStep, Time::nEndTimeStep, Grid::nGlobalGridDims, Read(),Parameters::nMaxIterations, Implicit::nMaxNumIterations, Grid::nNum1DZones, plicit::nNumImplicitZones, ProcTop::nNumProcs, Output::nPrintFrequencyStep, Output::nPrintMode, ProcTop::nProcDims, ProcTop::nRank, Parameters::nTypeTurbulanceMod, eos::readBin(), Output::sBaseOutputFileName, Parameters::sDebugProfileOutput, and Parameters::sEOSFileName.

Referenced by main().

### 7.4.2.5 void initImplicitCalculation (Implicit & implicit, Grid & grid, ProcTop & procTop, int nNumArgs, char \* cArgs[])

This function initilizes data structures and defines indixes of non-zero elements in the coeffecient matrix. It also sets up pathways for collection of the temperature corrections back to the processors which need them for their local grids.

#### Parameters:

- $\leftrightarrow implicit$
- $\leftarrow$  *grid* size information of the grid is used
- $\leftarrow procTop$
- $\leftarrow nNumArgs$  number of command line arguments, PETSc wants them
- $\leftarrow cArgs$  a list of command line arguments, PETSc wants them

#### Todo

 $is From, \quad is To, \quad mat Coeff, vec T Corrections, \quad vec T Corrections, vec R H S, vec T Corrections Local \\ , ksp Context, vec scat T Corrections all need to be destroyed before program finishes.$ 

Implicit::dTolerance, References Implicit::kspContext, Implicit::matCoeff, Proc-Top::nCoords, Grid::nGlobalGridDims, Grid::nLocalGridDims, Implicit::nLocDer, Implicit::nLocFun, Implicit::nMaxNumIterations, Implicit::nNumDerPerRow, Grid::nNumDims, ProcTop::nNumProcs, Grid::nNumGhostCells, Implicit::nNumImplicitZones, Implicit::nNumRowsALocal, Implicit::nNumRowsALocalSB, ProcTop::nProcDims, Proc-Top::nRank, Grid::nT, Implicit::nTypeDer, Implicit::vecRHS, Implicit::vecscatTCorrections, Implicit::vecTCorrections, and Implicit::vecTCorrectionsLocal.

Referenced by init().

# 7.4.2.6 void initUpdateLocalBoundaries (ProcTop & procTop, Grid & grid, MessPass & messPass, Implicit & implicit)

Sets up MPI derived data types used for updating the local grid boundaries between processors. It sets where the local grids should start/stop updating the local grids (Grid::nStartUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nStartUpdateImplicit,

Grid::nEndUpdateImplicit, Grid::nStartGhostUpdateExplicit, Grid::nEndGhostUpdateExplicit, Grid::nEndGhostUpdateImplicit, Grid::nEndGhostUpdateImplicit). It sets the radial processor neighbors (ProcTop::nNumRadialNeighbors).

It also allocates memeory for:

- MessPass::requestSend
- MessPass::requestRecv
- statusSend
- statusRecv

#### Parameters:

- $\leftrightarrow procTop$
- $\leftrightarrow qrid$
- $\leftrightarrow messPass$
- $\leftrightarrow implicit$

References Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nCenIntOffset, Top::nCoords, Grid::nE, Grid::nEndGhostUpdateExplicit, Grid::nEndGhostUpdateImplicit, Grid::nEndUpdateExplicit, Grid::nEndUpdateImplicit, Grid::nGlobalGridDims, Grid::nGlobalGridPositionLocalGrid, Grid::nKappa, Grid::nLocalGridDims, ProcTop::nNeighborRanks, Grid::nNum1DZones, Grid::nNumGhostCells, plicit::nNumImplicitZones, Grid::nNumIntVars, ProcTop::nNumNeighbors, ProcTop::nNumProcs, ProcTop::nNumRadialNeighbors, Grid::nNumVars, Grid::nNumZones1DBoundaryZeroHorizontalVelocity, Grid::nP, ProcTop::nPeriodic, ProcTop::nRadialNeighborNeighborIDs, Top::nProcDims, ProcTop::nRadialNeighborRanks, Grid:: nStartGhostUpdateExplicit,ProcTop::nRank, Grid::nStartGhostUpdateImplicit, Grid::nStartUpdateExplicit, Grid::nStartUpdateImplicit, Grid::nT, Grid::nV, Grid::nVariables, Grid∷nW, MessPass::requestRecv, MessPass::requestSend, MessPass::statusRecv, Mess-Pass::statusSend, MessPass::typeRecvNewVar, MessPass::typeRecvOldGrid, Mess-Pass::typeSendNewGrid, and MessPass::typeSendNewVar.

Referenced by init().

### 7.4.2.7 void modelRead (std::string sFileName, ProcTop & procTop, Grid & grid, Time & time, Parameters & parameters)

Reads in a collected binary file into the local grid and calls setupLocalGrid to allocate memory and set various parameters of the model. Works for both gamma-law gas, and tabulated equation of state models.

- $\leftarrow$  sFileName name of the file containing the model to be read in
- $\rightarrow procTop$
- $\rightarrow grid$
- $\rightarrow time$
- $\rightarrow$  parameters

#### Todo

At some point should get it working with only 1 processor

References Parameters::bEOSGammaLaw, Parameters::dA, Parameters::dAlpha, ters::dAVThreshold, Time::dDeltat n, Time::dDeltat nm1half, Time::dDeltat np1half, Parameters::dGamma, Grid::dLocalGridOld, Time::dt, DUMP VERSION, ProcTop::nCoords, Grid::nCotThetaIJK, Grid::nCotThetaIJp1halfK, Grid::nD, Grid::nDCosThetaIJK, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nE, Grid::nEddyVisc, Grid::nGamma, Grid::nGlobalGridDims, Grid::nKappa, Grid::nLocalGridDims, Grid::nM, Grid::nNum1DZones, Grid::nNumDims, Grid::nNumGhostCells, Grid::nNumIntVars, ProcTop::nNumProcs, Grid::nNumVars, Grid::nP, ProcTop::nPeriodic, ProcTop::nProcDims, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, ProcTop::nRank, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nT, Grid::nTheta, Time::nTimeStepIndex, Parameters::nTypeTurbulanceMod, Grid::nU, Grid::nU0, Grid::nV, Grid::nVariables, Grid::nW, Parameters::sEOSFileName, setInternalVarInf(), and setupLocalGrid().

Referenced by init().

### 7.4.2.8 void modelWrite\_GL (std::string sFileName, ProcTop & procTop, Grid & grid, Time & time, Parameters & parameters)

Writes out a model in distributed model format, meaning that each processor writes it's own local grid to a file in binary format. They can be combined, and or converted to ascii format using SPHERLSanal. This is for a gamma-law gas model.

#### Parameters:

- $\leftarrow sFileName$  base name of the output files
- $\leftarrow procTop$
- $\leftarrow \textit{grid}$
- $\leftarrow time$
- $\leftarrow$  parameters

References Parameters::dA, Parameters::dAlpha, Parameters::dAVThreshold, Time::dDeltat\_-nm1half, Parameters::dGamma, Grid::dLocalGridOld, Time::dt, DUMP\_VERSION, ProcTop::nCoords, Grid::nGlobalGridDims, Grid::nLocalGridDims, Grid::nNum1DZones, Grid::nNumGhostCells, Grid::nNumVars, ProcTop::nPeriodic, ProcTop::nProcDims, ProcTop::nRank, Time::nTimeStepIndex, and Grid::nVariables.

Referenced by setMainFunctions().

### 7.4.2.9 void modelWrite\_TEOS (std::string sFileName, ProcTop & procTop, Grid & grid, Time & time, Parameters & parameters)

Writes out a model in distributed model format, meaning that each processor writes it's own local grid to a file in binary format. They can be combined, and or converted to ascii format using SPHERLSanal. This is for a tabulated equation of state model.

- $\leftarrow$  **sFileName** base name of the output files
- $\leftarrow procTop$

- $\leftarrow grid \\ \leftarrow time$
- $\leftarrow parameters$

References Parameters::dA, Parameters::dAlpha, Parameters::dAVThreshold, Time::dDeltat\_-nm1half, Time::dDeltat\_np1half, Grid::dLocalGridOld, Time::dt, DUMP\_VERSION, ProcTop::nCoords, Grid::nGlobalGridDims, Grid::nLocalGridDims, Grid::nNum1DZones, Grid::nNumGhostCells, Grid::nNumVars, ProcTop::nPeriodic, ProcTop::nProcDims, ProcTop::nRank, Time::nTimeStepIndex, Grid::nVariables, and Parameters::sEOSFileName.

Referenced by setMainFunctions().

#### 7.4.2.10 void setupLocalGrid (ProcTop & procTop, Grid & grid)

Determins size of local grids (Grid::nLocalGridDims) based on processor topology, and allocates memory for the local grids (Grid::dLocalGridNew, Grid::dLocalGridOld). It sets various other quantities aswell such as,

- the coordinates of all processors (ProcTop::nCoords)
- the offset for interface centered quantities (Grid::nCenIntOffset, which depends on zoning and boundary conditions
- the position the local grid is in relative to the global grid (Grid::nGlobalGridPositionLocalGrid).

#### Parameters:

- $\leftrightarrow procTop$  contains information about the processor topology
- $\leftrightarrow$  grid contains information about gird

References Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nCenIntOffset, ProcTop::nCoords, Grid::nD, Grid::nGlobalGridDims, Grid::nGlobalGridPositionLocalGrid, Grid::nLocalGridDims, Grid::nNum1DZones, Grid::nNumDims, Grid::nNumGhostCells, Grid::nNumIntVars, ProcTop::nNumProcs, Grid::nNumVars, ProcTop::nPeriodic, ProcTop::nProcDims, ProcTop::nRank, and Grid::nVariables.

Referenced by modelRead().

### 7.4.2.11 void updateLocalBoundaries (ProcTop & procTop, MessPass & messPass, Grid & grid)

Updates the boundaries of the local grids from the data in the local grids of other processors. It does this for all variables and updates to the old grid. It also has processor ProcTop::nRank=0 call average3DTo1DBoundariesOld which averages the 3D information into the 1D boundaries.

- $\leftarrow procTop$
- $\leftarrow messPass$
- $\leftrightarrow grid$

#### Todo

Shouldn't need MPI::COMM\_WORLD.Barrier() may want to test out removing this at some point as it might produce a bit of a speed up.

References average3DTo1DBoundariesOld(), Grid::dLocalGridNew, Grid::dLocalGridOld, ProcTop::nNeighborRanks, ProcTop::nNumNeighbors, ProcTop::nRank, Mess-Pass::requestRecv, MessPass::requestSend, MessPass::statusRecv, MessPass::statusSend, MessPass::typeRecvOldGrid, MessPass::typeSendNewGrid, and updateOldGrid().

Referenced by main().

### 7.4.2.12 void updateLocalBoundariesNewGrid (int nVar, ProcTop & procTop, MessPass & messPass, Grid & grid)

Updates the boundaries of the local grids from the data in the local grids of other processors. It does this for a specific variable specified by nVar and updates to the new grid. It also has processor ProcTop::nRank=0 call average3DTo1DBoundariesNew which averages the 3D information into the 1D boundaries for that specific variable.

#### Parameters:

- $\leftarrow procTop$
- $\leftarrow messPass$
- $\leftrightarrow grid$

#### Todo

May want to do some waiting on this message at some point before the end of the timestep, but it doesn't need to be done in this function. It might also be that this is built into the code by waiting at some other point. This is something that should be checked out at somepoint, perhaps once the preformance starts to be analyzed. I would think that if the send buffer was being modified before the send was completed, that there would be some errors poping up that would likely kill the program.

 $References\ average 3DTo 1DBoundaries New (),\ Grid:: dLocal Grid New,\ Proc Top:: nNeighbor Ranks,\ Proc Top:: nNum Neighbors,\ Proc Top:: nRank,\ Mess Pass:: request Recv,\ Mess Pass:: status Recv,\ Mess Pass:: type Recv New Var,\ and\ Mess Pass:: type Send New Var.$ 

 $Referenced \quad by \quad implicitSolve\_R(), \quad implicitSolve\_RT(), \quad implicitSolve\_RTP(), \quad main(), \\ updateLocalBoundaryVelocitiesNewGrid\_R(), \quad updateLocalBoundaryVelocitiesNewGrid\_RT(), \\ and \quad updateLocalBoundaryVelocitiesNewGrid \quad RTP().$ 

# 7.4.2.13 void updateLocalBoundaryVelocitiesNewGrid\_R (ProcTop & procTop, MessPass & messPass, Grid & grid)

Updates velocity boundaries of the new grid in a 1D calculations after the velocities have been newly calculated.

References Grid::nU, and updateLocalBoundariesNewGrid().

Referenced by setMainFunctions().

### 7.4.2.14 void updateLocalBoundaryVelocitiesNewGrid\_RT (ProcTop & procTop, MessPass & messPass, Grid & grid)

Updates velocity boundaries of the new grid in a 2D calculations after the velocities have been newly calculated.

References Grid::nU, Grid::nV, and updateLocalBoundariesNewGrid().

Referenced by setMainFunctions().

### 7.4.2.15 void updateLocalBoundaryVelocitiesNewGrid\_RTP (ProcTop & procTop, MessPass & messPass, Grid & grid)

Updates velocity boundaries of the new grid in a 3D calculations after the velocities have been newly calculated.

References Grid::nU, Grid::nV, Grid::nW, and updateLocalBoundariesNewGrid().

Referenced by setMainFunctions().

#### 7.4.2.16 void updateNewGridWithOld (Grid & grid, ProcTop & procTop)

Copies the contents of the old grid to the new grid including ghost cells.

#### Parameters:

- $\leftrightarrow grid$
- $\leftarrow procTop$

 $\label{localGridNew} References & Grid::dLocalGridNew, & Grid::dLocalGridOld, & Grid::nLocalGridDims, \\ Grid::nNumDims, & Grid::nNumGhostCells, & Grid::nNumIntVars, & Grid::nNumVars, & Proc-Top::nRank, and & Grid::nVariables. \\ \end{array}$ 

Referenced by main().

#### 7.4.2.17 void updateOldGrid (ProcTop & procTop, Grid & grid)

Updates the old grid with the new grid, not including boundaries.

#### Parameters:

- $\leftarrow procTop$
- $\leftrightarrow grid$

 $\label{lem:condition} References & Grid::dLocalGridNew, & Grid::dLocalGridOld, & Grid::nEndGhostUpdateExplicit, \\ Grid::nEndGhostUpdateImplicit, & Grid::nEndUpdateExplicit, & Grid::nEndUpdateImplicit, \\ Grid::nNumIntVars, & Grid::nNumVars, & Grid::nStartGhostUpdateExplicit, \\ Grid::nStartGhostUpdateImplicit, & Grid::nStartUpdateExplicit, \\ and & Grid::nStartUpdateImplicit. \\ \end{tabular}$ 

Referenced by updateLocalBoundaries().

### 7.5 dataMonitoring.cpp File Reference

```
#include <mpi.h>
#include <sstream>
#include <fstream>
#include <iostream>
#include <cmath>
#include <iomanip>
#include <string>
#include "watchzone.h"
#include "exception2.h"
#include "xmlFunctions.h"
#include "dataMonitoring.h"
#include "global.h"
#include "fileExists.h"
```

#### **Functions**

- void initWatchZones (XMLNode xParent, ProcTop &procTop, Grid &grid, Output &output, Parameters &parameters, Time &time)
- void writeWatchZones\_R\_GL (Output &output, Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void writeWatchZones\_R\_TEOS (Output &output, Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void writeWatchZones\_RT\_GL (Output &output, Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void writeWatchZones\_RT\_TEOS (Output &output, Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void writeWatchZones\_RTP\_GL (Output &output, Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void writeWatchZones\_RTP\_TEOS (Output &output, Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void finWatchZones (Output &output)

#### 7.5.1 Detailed Description

This file holds functions used for examining the grid data during execution. This includes initializing structures, handling watching zones during the execution of the program, opening files to write out the peak kinetic energy, etc.

#### 7.5.2 Function Documentation

#### 7.5.2.1 void finWatchZones (Output & output)

Closes the files opened for writting out the watchzones

#### Parameters:

 $\leftarrow output$ 

 $References\ Output :: of Watch Zone Files,\ and\ Output :: watch zone List.$ 

Referenced by fin().

### 7.5.2.2 void initWatchZones (XMLNode xParent, ProcTop & procTop, Grid & grid, Output & output, Parameters & parameters, Time & time)

Reads in watchzones set in configuration file "SPHERLS.xml". A list is created on each processor containing the watchzones on that processor's local grid. It also opens file streams for each watchzone and writes out a header.

#### Parameters:

- $\leftarrow xParent$
- $\leftarrow \mathit{proc}\,\mathsf{To}\,p$
- $\leftarrow grid$
- $\leftrightarrow output$
- $\leftarrow$  parameters
- $\leftarrow time$

References Parameters::bEOSGammaLaw, Parameters::dGamma, ProcTop::nCoords, Grid::nD, Grid::nGlobalGridDims, Grid::nLocalGridDims, Grid::nNum1DZones, Grid::nNumDims, Grid::nNumGhostCells, ProcTop::nNumProcs, ProcTop::nProcDims, ProcTop::nRank, Time::nTimeStepIndex, Output::ofWatchZoneFiles, Output::sBaseOutputFileName, and Output::watchzoneList.

Referenced by init().

# 7.5.2.3 void writeWatchZones\_R\_GL (Output & output, Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

Writes out the information for each watchzone specified in "SPHERLS.xml" in the case of a 1D gamma-law gas.

- $\leftrightarrow output$
- $\leftarrow \textit{grid}$
- $\leftarrow$  parameters

- $\leftarrow time$
- $\leftarrow procTop$

References Grid::dLocalGridOld, Parameters::dPi, Time::dt, Grid::nCenIntOffset, Grid::nD, Grid::nDM, Grid::nE, Grid::nP, Grid::nQ0, Grid::nR, Time::nTimeStepIndex, Grid::nU, Grid::nU0, Output::ofWatchZoneFiles, and Output::watchzoneList.

Referenced by setMainFunctions().

## 7.5.2.4 void writeWatchZones\_R\_TEOS (Output & output, Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

Writes out the information for each watchzone specified in "SPHERLS.xml" in the case of a 1D tabulated equation of state.

#### Parameters:

- $\leftrightarrow output$
- $\leftarrow grid$
- $\leftarrow$  parameters
- $\leftarrow time$
- $\leftarrow procTop$

References Grid::dLocalGridOld, Parameters::dPi, Time::dt, Grid::nCenIntOffset, Grid::nD, Grid::nDM, Grid::nE, Grid::nP, Grid::nQ0, Grid::nR, Grid::nT, Time::nTimeStepIndex, Grid::nU, Grid::nU0, Output::ofWatchZoneFiles, and Output::watchzoneList.

Referenced by setMainFunctions().

# 7.5.2.5 void writeWatchZones\_RT\_GL (Output & output, Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

Writes out the information for each watchzone specified in "SPHERLS.xml" in the case of a 2D gamma-law gas.

#### Parameters:

- $\leftrightarrow output$
- $\leftarrow qrid$
- $\leftarrow$  parameters
- $\leftarrow time$
- $\leftarrow procTop$

References Grid::dLocalGridOld, Parameters::dPi, Time::dt, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nE, Grid::nEddyVisc, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nR, Time::nTimeStepIndex, Parameters::nTypeTurbulanceMod, Grid::nU, Grid::nU0, Grid::nV, Output::ofWatchZoneFiles, and Output::watchzoneList.

Referenced by setMainFunctions().

### 7.5.2.6 void write Watch Zones RT TEOS (Output & output, Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

Writes out the information for each watchzone specified in "SPHERLS.xml" in the case of a 2D tabulated equation of state.

#### Parameters:

- $\leftrightarrow output$
- $\leftarrow grid$
- $\leftarrow \textit{parameters}$
- $\leftarrow time$
- $\leftarrow procTop$

References Grid::dLocalGridOld, Parameters::dPi, Time::dt, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nE, Grid::nEddyVisc, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nR, Grid::nT, Time::nTimeStepIndex, Parameters::nTypeTurbulanceMod, Grid::nU, Grid::nU0, Grid::nV, Output::ofWatchZoneFiles, and Output::watchzoneList.

Referenced by setMainFunctions().

### 7.5.2.7 void writeWatchZones\_RTP\_GL (Output & output, Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

Writes out the information for each watchzone specified in "SPHERLS.xml" in the case of a 3D gamma-law gas.

#### Parameters:

- $\leftrightarrow output$
- $\leftarrow \textit{grid}$
- $\leftarrow$  parameters
- $\leftarrow time$
- $\leftarrow procTop$

References Grid::dLocalGridOld, Parameters::dPi, Time::dt, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nE, Grid::nEddyVisc, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, Time::nTimeStepIndex, Parameters::nTypeTurbulanceMod, Grid::nU, Grid::nU0, Grid::nV, Grid::nW, Output::ofWatchZoneFiles, and Output::watchzoneList.

Referenced by setMainFunctions().

### 7.5.2.8 void writeWatchZones\_RTP\_TEOS (Output & output, Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

Writes out the information for each watchzone specified in "SPHERLS.xml" in the case of a 3D tabulated equation of state.

#### Parameters:

 $\leftrightarrow output$ 

- $\leftarrow grid$
- $\leftarrow \textit{parameters}$
- $\leftarrow \textit{time}$
- $\leftarrow procTop$

References Grid::dLocalGridOld, Parameters::dPi, Time::dt, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nE, Grid::nEddyVisc, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, Grid::nT, Time::nTimeStepIndex, Parameters::nTypeTurbulanceMod, Grid::nU, Grid::nU0, Grid::nV, Grid::nW, Output::ofWatchZoneFiles, and Output::watchzoneList.

Referenced by setMainFunctions().

### 7.6 dataMonitoring.h File Reference

```
#include <string>
#include "xmlParser.h"
#include "global.h"
```

#### Functions

- void initWatchZones (XMLNode xParent, ProcTop &procTop, Grid &grid, Output &output, Parameters &parameters, Time &time)
- void writeWatchZones\_R\_GL (Output &output, Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void writeWatchZones\_R\_TEOS (Output &output, Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void writeWatchZones\_RT\_GL (Output &output, Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void writeWatchZones\_RT\_TEOS (Output &output, Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void writeWatchZones\_RTP\_GL (Output &output, Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void writeWatchZones\_RTP\_TEOS (Output &output, Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void finWatchZones (Output &output)

#### 7.6.1 Detailed Description

Header file for dataMonitoring.cpp

#### 7.6.2 Function Documentation

#### 7.6.2.1 void finWatchZones (Output & output)

Closes the files opened for writting out the watchzones

#### Parameters:

 $\leftarrow output$ 

References Output::ofWatchZoneFiles, and Output::watchzoneList.

Referenced by fin().

# 7.6.2.2 void initWatchZones (XMLNode xParent, ProcTop & procTop, Grid & grid, Output & output, Parameters & parameters, Time & time)

Reads in watchzones set in configuration file "SPHERLS.xml". A list is created on each processor containing the watchzones on that processor's local grid. It also opens file streams for each watchzone and writes out a header.

#### Parameters:

- $\leftarrow xParent$
- $\leftarrow procTop$
- $\leftarrow grid$
- $\leftrightarrow output$
- $\leftarrow \textit{parameters}$
- $\leftarrow time$

 $\label{lem:condition} References\ Parameters::bEOSGammaLaw,\ Parameters::dGamma,\ ProcTop::nCoords,\ Grid::nD,\ Grid::nGlobalGridDims,\ Grid::nLocalGridDims,\ Grid::nNum1DZones,\ Grid::nNumDims,\ Grid::nNumGhostCells,\ ProcTop::nNumProcs,\ ProcTop::nProcDims,\ ProcTop::nRank,\ Time::nTimeStepIndex,\ Output::ofWatchZoneFiles,\ Output::sBaseOutputFileName,\ and\ Output::watchzoneList.$ 

Referenced by init().

## 7.6.2.3 void writeWatchZones\_R\_GL (Output & output, Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

Writes out the information for each watchzone specified in "SPHERLS.xml" in the case of a 1D gamma-law gas.

#### Parameters:

- $\leftrightarrow output$
- $\leftarrow grid$
- $\leftarrow parameters$
- $\leftarrow time$
- $\leftarrow procTop$

References Grid::dLocalGridOld, Parameters::dPi, Time::dt, Grid::nCenIntOffset, Grid::nD, Grid::nDM, Grid::nE, Grid::nP, Grid::nQ0, Grid::nR, Time::nTimeStepIndex, Grid::nU, Grid::nU0, Output::ofWatchZoneFiles, and Output::watchzoneList.

Referenced by setMainFunctions().

### 7.6.2.4 void writeWatchZones\_R\_TEOS (Output & output, Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

Writes out the information for each watchzone specified in "SPHERLS.xml" in the case of a 1D tabulated equation of state.

#### Parameters:

 $\leftrightarrow output$ 

- $\leftarrow grid$
- $\leftarrow \textit{parameters}$
- $\leftarrow time$
- $\leftarrow procTop$

References Grid::dLocalGridOld, Parameters::dPi, Time::dt, Grid::nCenIntOffset, Grid::nD, Grid::nDM, Grid::nE, Grid::nP, Grid::nQ0, Grid::nR, Grid::nT, Time::nTimeStepIndex, Grid::nU, Grid::nU0, Output::ofWatchZoneFiles, and Output::watchzoneList.

Referenced by setMainFunctions().

### 7.6.2.5 void writeWatchZones\_RT\_GL (Output & output, Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

Writes out the information for each watchzone specified in "SPHERLS.xml" in the case of a 2D gamma-law gas.

#### Parameters:

- $\leftrightarrow output$
- $\leftarrow grid$
- $\leftarrow \textit{parameters}$
- $\leftarrow time$
- $\leftarrow procTop$

References Grid::dLocalGridOld, Parameters::dPi, Time::dt, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nE, Grid::nEddyVisc, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nR, Time::nTimeStepIndex, Parameters::nTypeTurbulanceMod, Grid::nU, Grid::nU0, Grid::nV, Output::ofWatchZoneFiles, and Output::watchzoneList.

Referenced by setMainFunctions().

# 7.6.2.6 void writeWatchZones\_RT\_TEOS (Output & output, Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

Writes out the information for each watchzone specified in "SPHERLS.xml" in the case of a 2D tabulated equation of state.

#### Parameters:

- $\leftrightarrow \textit{output}$
- $\leftarrow qrid$
- $\leftarrow parameters$
- $\leftarrow time$
- $\leftarrow procTop$

References Grid::dLocalGridOld, Parameters::dPi, Time::dt, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nE, Grid::nEddyVisc, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nR, Grid::nT, Time::nTimeStepIndex, Parameters::nTypeTurbulanceMod, Grid::nU, Grid::nU0, Grid::nV, Output::ofWatchZoneFiles, and Output::watchzoneList.

Referenced by setMainFunctions().

### 7.6.2.7 void writeWatchZones\_RTP\_GL (Output & output, Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

Writes out the information for each watchzone specified in "SPHERLS.xml" in the case of a 3D gamma-law gas.

#### Parameters:

- $\leftrightarrow output$
- $\leftarrow grid$
- $\leftarrow$  parameters
- $\leftarrow \textit{time}$
- $\leftarrow procTop$

References Grid::dLocalGridOld, Parameters::dPi, Time::dt, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nE, Grid::nEddyVisc, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, Time::nTimeStepIndex, Parameters::nTypeTurbulanceMod, Grid::nU, Grid::nU0, Grid::nV, Grid::nW, Output::ofWatchZoneFiles, and Output::watchzoneList.

Referenced by setMainFunctions().

## 7.6.2.8 void writeWatchZones\_RTP\_TEOS (Output & output, Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

Writes out the information for each watchzone specified in "SPHERLS.xml" in the case of a 3D tabulated equation of state.

#### Parameters:

- $\leftrightarrow output$
- $\leftarrow grid$
- $\leftarrow \textit{parameters}$
- $\leftarrow time$
- $\leftarrow procTop$

References Grid::dLocalGridOld, Parameters::dPi, Time::dt, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nE, Grid::nEddyVisc, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, Grid::nT, Time::nTimeStepIndex, Parameters::nTypeTurbulanceMod, Grid::nU, Grid::nU0, Grid::nV, Grid::nW, Output::ofWatchZoneFiles, and Output::watchzoneList.

Referenced by setMainFunctions().

### 7.7 global.cpp File Reference

#include "global.h"

### 7.7.1 Detailed Description

Declares global variables used across files and functions. This file contains the constructors used to initialize the classes defined in global.h, and does little more than initilize the default values of various parameters.

### 7.8 global.h File Reference

```
#include <vector>
#include <mpi.h>
#include "watchzone.h"

#include "eos.h"

#include "petscksp.h"

#include <csignal>
#include <limits>
#include "profileData.h"

#include "procTop.h"

#include "time.h"
```

#### Classes

- class MessPass
- class Grid
- class Parameters
- class Output
- class Performance
- class Implicit
- class Functions
- class Global

#### **Defines**

- #define SIGNEGDEN 0
- #define SIGNEGENG 0
- #define SIGNEGTEMP 0
- #define SEDOV 0
- #define VISCOUS\_ENERGY\_EQ 1
- #define DUMP VERSION 1
- $\bullet$  #define DEBUG EQUATIONS 0
- #define DEDEM CLAMP 1

#### 7.8.1 Detailed Description

Header file for global.cpp.

This file contains definitions which are required throughout the program. The classes defined herein are used through out the program.

#### 7.8.2 Define Documentation

## 7.8.2.1 #define DEBUG EQUATIONS 0

If 1 will write out in the form of a profile file, all the horizontal maximum values of all terms in all equations.

Referenced by dImplicitEnergyFunction\_R(), dImplicitEnergyFunction\_RTP\_LES(), and dImplicitEnergyFunction RTP LES SB().

## 7.8.2.2 #define DEDEM CLAMP 1

If 1 a clamp on the DEDM gradient will be used to limit how large DE/DM becomes in the advection term in the energy equation.

# 7.8.2.3 #define DUMP\_VERSION 1

Sets the version of the dump file. Should be incremented if changes are made to the information that is printed out in a dump.

Referenced by modelRead(), modelWrite\_GL(), and modelWrite\_TEOS().

### 7.8.2.4 #define SEDOV 0

If 1 we are preforming the sedov test, which sets special boundary conditions, if 0 we use normal boundary conditions. It also handles artificial viscosity, and timestep slightly differently.

### 7.8.2.5 #define SIGNEGDEN 0

Raise signal on calculation of negative density if set to 1. Useful when debugging, it will stop the debugger at the location of the calculation of the negative density. If not 1, it will speed up calculation slightly and generate more useful output upon detection of negative densities. If 1 and not being run in the debugger, it likely won't generate any usefull output upon negative density, and wil simply abort the program.

## 7.8.2.6 #define SIGNEGENG 0

Raise signal on calculation of negative energy if set to 1, else don't rais a signal. Otherwise it will be handled through the normal exception method. This is useful when debugging, it will stop the debugger at the location of the calculation of the negative energy. If not 1, it will speed up calculation slightly and generate more useful output upon detection of negative energy. If 1 and not being run in the debugger, it likely won't generate any usefull output upon negative energies, and wil simply abort the program.

#### 7.8.2.7 #define SIGNEGTEMP 0

Raise signal on calculation of negative temperature if set to 1, else don't rais a signal. Otherwise it will be handled through the normal exception method. This is useful when debugging, it will stop

the debugger at the location of the calculation of the negative energy. If not 1, it will speed up calculation slightly and generate more useful output upon detection of negative energy. If 1 and not being run in the debugger, it likely won't generate any usefull output upon negative energies, and wil simply abort the program.

# 7.8.2.8 #define TRACKMAXSOLVERERROR 0

Report the error of the linear equation solver if set to 1, else don't. Not tracking the error reduces the calculations per iteration and will speed up running, however if there is question of weather the solver is working accurately this is very handy to turn on.

# 7.8.2.9 #define VISCOUS ENERGY EQ 1

If 1 will include viscosity in the energy equation. If 0 it won't. This normally should be set to 1

# 7.9 main.cpp File Reference

```
#include <mpi.h>
#include <sstream>
#include <string>
#include <fstream>
#include <cmath>
#include <vector>
#include <algorithm>
#include <iomanip>
#include <csignal>
#include <fenv.h>
#include "main.h"
#include "global.h"
#include "watchzone.h"
#include "exception2.h"
#include "xmlParser.h"
#include "xmlFunctions.h"
#include "dataManipulation.h"
#include "dataMonitoring.h"
#include "physEquations.h"
```

# **Functions**

- int main (int argc, char \*argv[])
- void signalHandler (int nSig)

# 7.9.1 Detailed Description

This file contains the main function which is the driver for SPHERLS.

## 7.9.2 Function Documentation

## 7.9.2.1 int main (int argc, char \* argv[])

Main driving function of SPHERLS.

#### Parameters:

- $\leftarrow$  argc number of arguments passed from the command line
- $\leftarrow$  argv array of character strings of size argc containing the arguments from the command line.

The flow of this function is as follows:

- Initilize program by calling init()
- Set function pointers by calling setMainFunctions()
- Update new grid with old grid by calling updateNewGridWithOld()
- Update boundaries of local grids
- Calculate the first time step by calling Functions::fpCalculateDeltat()
- Enter while loop until end time (Time::dEndTime) is reached, and for each interation of the loop:
  - Test to see if a model dump is needed (by checking Output::bDump and Output::nDumpFrequency), if so dump one by calling modelWrite()
  - Write out information for any watchzones present by calling writeWatchZones()
  - Write out information for peak kinetic energy per period by calling writePeakKE()
  - calculate time step by calling function pointed to by Functions::fpCalculateDeltat
- Calculate new velocities by calling the function pointed to by Functions::fpCalculateNewVelocities()
- Update velocities on new grid boundaries between processors by calling updateLocalBoundariesNewGrid() three times indicating the r-velocity (U),  $\theta$ -velocity (V) and the  $\phi$ -velocity (W).
- Calculate new grid velocities with Functions::fpCalculateNewGridVelocities().
- Calculate new radii with Functions::fpCalculateNewRadii().
- Update radii on new grid boundaries between processors by calling updateLocalBoundariesNewGrid() indicating radius is to be updated (R).
- Calculate new densities with Functions::fpCalculateNewDensities()
- Calculate new energies with Functions::fpCalculateNewEnergies()
- Update the old grid boundaries and centeres by calling updateLocalBoundaries()
- Calculating the next time step with Functions::fpCalculateDeltat()

Finish by dumping the last model computed

Parameters::bDEDM cut set, References Output::bDump, Output::bPrint, Time::dDelRho t Rho max, plicit::dAverageRHS, Implicit::dCurrentRelTError,  $Time::dDelT\_t\_T\_max,$ Time::dDeltat np1half, Time::dDelUmU0 t UmU0 max, Time::dDelV t V max, Output::dDumpFrequencyTime, Time::dDelW t W max, Time::dEndTime, Parameters::dMaxConvectiveVelocity, Implicit::dMaxErrorInRHS, put::dPrintFrequencyTime, Time::dt, Output::dTimeLastDump, Output::dTimeLastPrint, fin(), Functions::fpCalculateAveDensities, Functions::fpCalculateDeltat, Functions::fpCalculateNewAV, Functions::fpCalculateNewDensities, Functions::fpCalculateNewEddyVisc, Functions::fpCalculateNewEnergies, Functions::fpCalculateNewEOSVars, Functions::fpCalculateNewGridVelocities, Functions::fpCalculateNewRadii, Functions::fpCalculateNewVelocities, Functions::fpImplicitSolve, Functions::fpModelWrite, Functions::fpUpdateLocalBoundaryVelocitiesNewGrid, Functions::fpWriteWatchZones, Global::functions, Global::grid, Global::implicit, init(), Global::messPass, plicit::nCurrentNumIterations, Grid::nD, Grid::nDenAve, Output::nDumpFrequencyStep, Grid::nEddyVisc, Time::nEndTimeStep, Grid::nGamma, Implicit::nMaxNumSolverIterations, Implicit::nNumImplicitZones, Output::nNumTimeStepsSinceLastDump, Grid::nNumDims, Output::nNumTimeStepsSinceLastPrint, Grid::nP, Output::nPrintFrequencyStep, put::nPrintMode, Grid::nR, ProcTop::nRank, Grid::nT, Time::nTimeStepIndex, Grid::nU0, Global::output, Global::parameters, Global::performance, Global::procTop, Output::sBaseOutputFileName, Parameters::sDebugProfileOutput, setMainFunctions(), signal-Handler(), Global::time, updateLocalBoundaries(), updateLocalBoundariesNewGrid(), and updateNewGridWithOld().

## 7.9.2.2 void signalHandler (int nSig)

Used for catching signals.

Referenced by main().

# 7.10 main.h File Reference

#### **Functions**

- void signalHandler (int nSig)
- int main (int argc, char \*argv[])

# 7.10.1 Detailed Description

Header file for main.cpp

## 7.10.2 Function Documentation

### 7.10.2.1 int main (int argc, char \* argv[])

Main driving function of SPHERLS.

#### Parameters:

- $\leftarrow$  argc number of arguments passed from the command line
- $\leftarrow$  argv array of character strings of size argc containing the arguments from the command line.

The flow of this function is as follows:

- Initilize program by calling init()
- Set function pointers by calling setMainFunctions()
- Update new grid with old grid by calling updateNewGridWithOld()
- Update boundaries of local grids
- Calculate the first time step by calling Functions::fpCalculateDeltat()
- Enter while loop until end time (Time::dEndTime) is reached, and for each interation of the loop:
  - Test to see if a model dump is needed (by checking Output::bDump and Output::nDumpFrequency), if so dump one by calling modelWrite()
  - Write out information for any watchzones present by calling writeWatchZones()
  - Write out information for peak kinetic energy per period by calling writePeakKE()
  - calculate time step by calling function pointed to by Functions::fpCalculateDeltat
- Calculate new velocities by calling the function pointed to by Functions::fpCalculateNewVelocities()

- Update velocities on new grid boundaries between processors by calling updateLocalBoundariesNewGrid() three times indicating the r-velocity (U),  $\theta$ -velocity (V) and the  $\phi$ -velocity (W).
- Calculate new grid velocities with Functions::fpCalculateNewGridVelocities().
- Calculate new radii with Functions::fpCalculateNewRadii().
- Update radii on new grid boundaries between processors by calling updateLocalBoundariesNewGrid() indicating radius is to be updated (R).
- Calculate new densities with Functions::fpCalculateNewDensities()
- Calculate new energies with Functions::fpCalculateNewEnergies()
- Update the old grid boundaries and centeres by calling updateLocalBoundaries()
- Calculating the next time step with Functions::fpCalculateDeltat()

Finish by dumping the last model computed

Parameters::bDEDM cut set, References Output::bDump, Output::bPrint, Implicit::dAverageRHS, Implicit::dCurrentRelTError, Time::dDelRho t Rho max, Time::dDelT t T max, Time::dDeltat np1half, Time::dDelUmU0 t UmU0 max, Time::dDelV t V max, Time::dDelW t W max, Output::dDumpFrequencyTime, Time::dEndTime, Parameters::dMaxConvectiveVelocity, Implicit::dMaxErrorInRHS, Output::dPrintFrequencyTime, Time::dt, Output::dTimeLastDump, Output::dTimeLastPrint, fin(), Functions::fpCalculateAveDensities, Functions::fpCalculateDeltat, Functions::fpCalculateNewAV, Functions::fpCalculateNewDensities, Functions::fpCalculateNewEddyVisc, Functions::fpCalculateNewEnergies, Functions::fpCalculateNewEOSVars, Functions::fpCalculateNewGridVelocities, Functions::fpCalculateNewRadii, Functions::fpCalculateNewVelocities, Functions::fpImplicitSolve, Functions::fpModelWrite, Functions::fpUpdateLocalBoundaryVelocitiesNewGrid, Functions::fpWriteWatchZones, Global::implicit, Global::messPass, Global::functions, Global::grid, init(), plicit::nCurrentNumIterations, Grid::nD, Grid::nDenAve, Output::nDumpFrequencyStep, Grid::nEddyVisc, Time::nEndTimeStep, Grid::nGamma, Implicit::nMaxNumSolverIterations, Implicit::nNumImplicitZones, Output::nNumTimeStepsSinceLastDump, Grid::nNumDims. Output::nNumTimeStepsSinceLastPrint, Grid::nP, Output::nPrintFrequencyStep, put::nPrintMode, Grid::nR, ProcTop::nRank, Grid::nT, Time::nTimeStepIndex, Grid::nU0, Global::output, Global::parameters, Global::performance, Global::procTop, Output::sBaseOutputFileName, Parameters::sDebugProfileOutput, setMainFunctions(), Handler(), Global::time, updateLocalBoundaries(), updateLocalBoundariesNewGrid(), and updateNewGridWithOld().

## 7.10.2.2 void signal Handler (int nSig)

Used for catching signals.

Referenced by main().

# 7.11 physEquations.cpp File Reference

```
#include <cmath>
#include <sstream>
#include <signal.h>
#include "exception2.h"
#include "physEquations.h"
#include "dataManipulation.h"
#include "dataMonitoring.h"
#include "global.h"
#include <limits>
#include "profileData.h"
```

#### Functions

- void setMainFunctions (Functions &functions, ProcTop &procTop, Parameters &parameters, Grid &grid, Time &time, Implicit &implicit)
- void setInternalVarInf (Grid &grid, Parameters &parameters)
- void initInternalVars (Grid &grid, ProcTop &procTop, Parameters &parameters)
- void calNewVelocities\_R (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewVelocities\_R\_LES (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewVelocities\_RT (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewVelocities\_RT\_LES (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewVelocities\_RTP (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewVelocities\_RTP\_LES (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewU R (Grid &grid, Parameters &parameters, Time &time, ProcTop)
- void calNewU\_R\_LES (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewU RT (Grid &grid, Parameters &parameters, Time &time, ProcTop)
- void calNewU\_RT\_LES (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewU\_RTP (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewU\_RTP\_LES (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewV RT (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewV\_RT\_LES (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)

- void calNewV\_RTP (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewV\_RTP\_LES (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewW\_RTP (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewW\_RTP\_LES (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewU0\_R (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop, MessPass &messPass)
- void calNewU0\_RT (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop, MessPass &messPass)
- void calNewU0\_RTP (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop, MessPass &messPass)
- void calNewR (Grid &grid, Time &time)
- void calNewD R (Grid &grid, Parameters &parameters, Time &time, ProcTop)
- void calNewD RT (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewD\_RTP (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewE\_R\_AD (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewE\_R\_NA (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewE\_R\_NA\_LES (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewE\_RT\_AD (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewE\_RT\_NA (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewE\_RT\_NA\_LES (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewE\_RTP\_AD (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewE\_RTP\_NA (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewE\_RTP\_NA\_LES (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewDenave None (Grid &grid)
- void calNewDenave R (Grid &grid)
- void calNewDenave RT (Grid &grid)
- void calNewDenave RTP (Grid &grid)
- void calNewP GL (Grid &grid, Parameters &parameters)
- void calNewTPKappaGamma TEOS (Grid &grid, Parameters &parameters)
- void calNewPEKappaGamma TEOS (Grid &grid, Parameters &parameters)
- void calNewQ0\_R\_TEOS (Grid &grid, Parameters &parameters)
- void calNewQ0 R GL (Grid &grid, Parameters &parameters)
- void calNewQ0Q1 RT TEOS (Grid &grid, Parameters &parameters)
- void calNewQ0Q1 RT GL (Grid &grid, Parameters &parameters)
- void calNewQ0Q1Q2\_RTP\_TEOS (Grid &grid, Parameters &parameters)
- void calNewQ0Q1Q2\_RTP\_GL (Grid &grid, Parameters &parameters)
- void calNewEddyVisc None (Grid &grid, Parameters &parameters)

- void calNewEddyVisc R CN (Grid &grid, Parameters &parameters)
- void calNewEddyVisc RT CN (Grid &grid, Parameters &parameters)
- void calNewEddyVisc RTP CN (Grid &grid, Parameters &parameters)
- void calNewEddyVisc R SM (Grid &grid, Parameters &parameters)
- void calNewEddyVisc\_RT\_SM (Grid &grid, Parameters &parameters)
- void calNewEddyVisc RTP SM (Grid &grid, Parameters &parameters)
- void calOldDenave\_None (Grid &grid)
- void calOldDenave\_R (Grid &grid)
- void calOldDenave RT (Grid &grid)
- void calOldDenave RTP (Grid &grid)
- void calOldP GL (Grid &grid, Parameters &parameters)
- void calOldPEKappaGamma TEOS (Grid &grid, Parameters &parameters)
- void calOldQ0 R GL (Grid &grid, Parameters &parameters)
- void calOldQ0\_R\_TEOS (Grid &grid, Parameters &parameters)
- void calOldQ0Q1 RT GL (Grid &grid, Parameters &parameters)
- void calOldQ0Q1 RT TEOS (Grid &grid, Parameters &parameters)
- void calOldQ0Q1Q2 RTP GL (Grid &grid, Parameters &parameters)
- void calOldQ0Q1Q2 RTP TEOS (Grid &grid, Parameters &parameters)
- void calOldEddyVisc R CN (Grid &grid, Parameters &parameters)
- void calOldEddyVisc RT CN (Grid &grid, Parameters &parameters)
- void calOldEddyVisc RTP CN (Grid &grid, Parameters &parameters)
- void calOldEddyVisc R SM (Grid &grid, Parameters &parameters)
- void calOldEddyVisc RT SM (Grid &grid, Parameters &parameters)
- void calOldEddyVisc RTP SM (Grid &grid, Parameters &parameters)
- void calDelt\_R\_GL (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calDelt\_R\_TEOS (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calDelt\_RT\_GL (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calDelt\_RT\_TEOS (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calDelt\_RTP\_GL (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calDelt\_RTP\_TEOS (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calDelt\_CONST (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void implicitSolve\_None (Grid &grid, Implicit &implicit, Parameters &parameters, Time &time, ProcTop &procTop, MessPass &messPass, Functions &functions)
- void implicitSolve\_R (Grid &grid, Implicit &implicit, Parameters &parameters, Time &time, ProcTop &procTop, MessPass &messPass, Functions &functions)
- void implicitSolve\_RT (Grid &grid, Implicit &implicit, Parameters &parameters, Time &time, ProcTop &procTop, MessPass &messPass, Functions &functions)
- void implicitSolve\_RTP (Grid &grid, Implicit &implicit, Parameters &parameters, Time &time, ProcTop &procTop, MessPass &messPass, Functions &functions)
- double dImplicitEnergyFunction\_None (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)
- double dImplicitEnergyFunction\_R (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)

- double dImplicitEnergyFunction\_R\_SB (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)
- double dImplicitEnergyFunction\_RT (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)
- double dImplicitEnergyFunction\_RT\_SB (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)
- double dImplicitEnergyFunction\_RTP (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)
- double dImplicitEnergyFunction\_RTP\_SB (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)
- double dImplicitEnergyFunction\_R\_LES (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)
- double dImplicitEnergyFunction\_R\_LES\_SB (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)
- double dImplicitEnergyFunction\_RT\_LES (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)
- double dImplicitEnergyFunction\_RT\_LES\_SB (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)
- double dImplicitEnergyFunction\_RTP\_LES (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)
- double dImplicitEnergyFunction\_RTP\_LES\_SB (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)
- double dEOS GL (double dRho, double dE, Parameters parameters)
- void initDonorFracAndMaxConVel R GL (Grid &grid, Parameters &parameters)
- void initDonorFracAndMaxConVel R TEOS (Grid &grid, Parameters &parameters)
- void initDonorFracAndMaxConVel RT GL (Grid &grid, Parameters &parameters)
- void initDonorFracAndMaxConVel RT TEOS (Grid &grid, Parameters &parameters)
- void initDonorFracAndMaxConVel RTP GL (Grid &grid, Parameters &parameters)
- void initDonorFracAndMaxConVel RTP TEOS (Grid &grid, Parameters &parameters)

## 7.11.1 Detailed Description

This file is used to specify the functions which contain physics. This includes conservation equations, equation of state, etc.. It also sets function pointers for these functions, so that main() will know which functions to call. This implementation also allows the functions called to calculate, for example new densities, to be different depending on the processor. This allows one processor to handle the 1D region and other processors to handle a 3D region.

## 7.11.2 Function Documentation

# 7.11.2.1 void calDelt\_CONST (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function is used when a constant tie step is desired.

References Time::dConstTimeStep, Time::dDeltat\_n, Time::dDeltat\_nm1half, Time::dDeltat\_-np1half, Time::dt, ProcTop::nRank, and Time::nTimeStepIndex.

# 7.11.2.2 void calDelt\_R\_GL (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the time step by considering the sound crossing time in the radial direction only and is compatiable with a gamma law gass EOS.

#### Parameters:

- $\leftarrow$  grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- $\leftrightarrow$  time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology. This function uses ProcTop::nRank to pass messages.

References Time::dDelE t E max, Time::dDelRho t Rho max, Time::dDeltat -Time::dDeltat nm1half, Time::dDeltat np1half, Time::dDelUmU0 t UmU0 n, Time::dDelV t V max,  $Time::dDelW\_t\_W\_max,$ Parameters::dDonorCellMin, max, Parameters::dDonorCellMultiplier, Parameters::dGamma, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dMaxConvectiveVelocity, Time::dPerChange, Time::dt, Time::dTimeStepFactor, Grid::nCenIntOffset, Grid::nD, Grid::nDonorCellFrac, Grid::nE, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0, Grid::nR, ProcTop::nRank, Grid::nStartUpdateExplicit, Time::nTimeStepIndex, Grid::nU, and Grid::nU0. Referenced by setMainFunctions().

# 7.11.2.3 void calDelt\_R\_TEOS (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the time step by considering the sound crossing time in the radial direction only and is compatiable with a tabulated EOS.

### Parameters:

- $\leftarrow$  grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- $\leftrightarrow$  time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology. This function uses ProcTop::nRank to pass messages.

References Time::dDelRho t Rho max, Time::dDelT t T max, Time::dDeltat -Time::dDeltat nm1half, Time::dDeltat np1half, Time::dDelUmU0 t UmU0 max, Time::dDelV t V max, Time::dDelW t W max, Parameters::dDonorCellMin, rameters::dDonorCellMultiplier, Grid::dLocalGridNew, Grid::dLocalGridOld, Param-Time::dt, eters::dMaxConvectiveVelocity, Time::dPerChange, Time::dTimeStepFactor, Grid::nCenIntOffset, Grid∷nD, Grid::nDonorCellFrac, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGamma, Grid::nP, Grid::nQ0, Grid::nR, ProcTop::nRank, Grid::nStartUpdateExplicit, Grid::nT, Time::nTimeStepIndex, Grid::nU, and Grid::nU0.

# 7.11.2.4 void calDelt\_RT\_GL (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the time step by considering the sound crossing time in the radial and theta directions only and is compatiable with a gamma law gass EOS.

#### Parameters:

- $\leftarrow$  grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- $\leftrightarrow$  time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology. This function uses ProcTop::nRank to pass messages.

References Time::dDelE t E max, Time::dDelRho t Rho max, Time::dDeltat -Time::dDeltat nm1half, Time::dDeltat np1half, Time::dDelUmU0 t UmU0 max,  $\label{lem:delv_t_V_max} Time:: dDelW_t_W_max, \quad Parameters:: dDonor CellMin, \quad Parameters:$ ters::dDonorCellMultiplier, Parameters::dGamma, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dMaxConvectiveVelocity, Time::dPerChange, Time::dt, Time::dTimeStepFactor, Grid::nCenIntOffset, Grid::nD, Grid::nDonorCellFrac, Grid::nDTheta, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nR, ProcTop::nRank, Grid::nStartUpdateExplicit, Time::nTimeStepIndex, Grid::nU, Grid::nU0, and Grid::nV.

Referenced by setMainFunctions().

# 7.11.2.5 void calDelt\_RT\_TEOS (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the time step by considering the sound crossing time in the radial and theta directions and is compatiable with a tabulated EOS.

#### Parameters:

- ← grid contains the local grid, and will hold the newly updated densities
- ← parameters various parameters needed for the calculation
- $\leftrightarrow$  time contains time information, e.g. time step, current time etc.
- $\leftarrow$  procTop contains information about the processor topology. This function uses ProcTop::nRank to pass messages.

Time::dDelT t T max, References Time::dDelRho t Rho max, Time::dDeltat n. Time::dDeltat nm1half, Time::dDeltat np1half, Time::dDelUmU0 t - $Time::dDelV\_t\_V\_max,$ Time::dDelW t W max, UmU0 max, Parameters::dDonorCellMin, Parameters::dDonorCellMultiplier, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dMaxConvectiveVelocity, Time::dPerChange, Time::dt, Time::dTimeStepFactor, Grid::nCenIntOffset, Grid::nD, Grid::nDonorCellFrac, Grid::nDTheta, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGamma, Grid∷nP. Grid::nQ0, Grid::nQ1, Grid::nR, ProcTop::nRank, Grid::nStartUpdateExplicit, Grid∷nT, Time::nTimeStepIndex, Grid::nU, Grid::nU0, and Grid::nV.

# 7.11.2.6 void calDelt\_RTP\_GL (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the time step by considering the sound crossing time in the radial, theta and phi directions only and is compatiable with a gamma law gass EOS.

#### Parameters:

- ← grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- $\leftrightarrow$  time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology. This function uses ProcTop::nRank to pass messages.

References Time::dDelE\_t\_E\_max, Time::dDelRho\_t\_Rho\_max, Time::dDeltat\_n, Time::dDeltat\_nm1half, Time::dDeltat\_np1half, Time::dDelUmU0\_t\_UmU0\_max, Time::dDelV\_t\_V\_max, Time::dDelW\_t\_W\_max, Parameters::dDonorCellMin, Parameters::dDonorCellMultiplier, Parameters::dGamma, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dMaxConvectiveVelocity, Time::dPerChange, Time::dt, Time::dTimeStepFactor, Grid::nCenIntOffset, Grid::nD, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nE, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, ProcTop::nRank, Grid::nSinThetaIJK, Grid::nStartUpdateExplicit, Time::nTimeStepIndex, Grid::nU, Grid::nU0, Grid::nV, and Grid::nW.

Referenced by setMainFunctions().

# 7.11.2.7 void calDelt\_RTP\_TEOS (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the time step by considering the sound crossing time in the radial, theta and phi directions and is compatiable with a tabulated EOS.

#### Parameters:

- $\leftarrow$  grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- $\leftrightarrow$  time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology. This function uses ProcTop::nRank to pass messages.

References Time::dDelRho t Rho max, Time::dDelT t T max, Time::dDeltat -Time::dDeltat nm1half, Time::dDeltat np1half, Time::dDelUmU0 t UmU0 max, Time::dDelV t V max,  $Time::dDelW\_t\_W\_max,$ Parameters::dDonorCellMin, rameters::dDonorCellMultiplier, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dMaxConvectiveVelocity, Time::dPerChange, Time::dt, Time::dTimeStepFactor, Grid::nDonorCellFrac, Grid::nCenIntOffset, Grid::nD, Grid::nDPhi, Grid::nDTheta, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGamma, Grid::nP, Grid::nR, ProcTop::nRank, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nSinThetaIJK, Grid::nStartUpdateExplicit, Grid::nT, Time::nTimeStepIndex, Grid::nU, Grid::nU0, Grid::nV, and Grid::nW.

# 7.11.2.8 void calNewD\_R (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates new densities using terms in the radial direction only

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology, uses ProcTop::nRank when reporting negative densities

### **Boundary Conditions**

doesn't allow mass flux through outter interface

 $\label{lem:condition} References Time::dDeltat\_np1half, Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nD, Grid::nDonorCellFrac, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGlobalGridPositionLocalGrid, Grid::nNumGhostCells, Grid::nR, ProcTop::nRank, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, and Grid::nU0.$ 

Referenced by setMainFunctions().

# 7.11.2.9 void calNewD\_RT (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates new densities using terms in the radial and theta directions

### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- $\leftarrow$  time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology, uses ProcTop::nRank when reporting negative densities

#### **Boundary Conditions**

doesn't allow mass flux through outter interface

Time::dDeltat np1half, Grid::dLocalGridOld, References Grid::dLocalGridNew, Grid::nCenIntOffset, Grid::nD, Grid::nDCosThetaIJK, Grid::nDonorCellFrac, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, ProcTop::nRank, Grid∷nR, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0, and Grid::nV.

# 7.11.2.10 void calNewD\_RTP (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates new densities using terms in the radial, theta, and phi directions

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology, uses ProcTop::nRank when reporting negative densities

## **Boundary Conditions**

doesn't allow mass flux through outter interface

References Time::dDeltat np1half, Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid∷nD, Grid::nDCosThetaIJK, Grid::nDonorCellFrac, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid∷nDPhi, Grid::nDTheta, Grid::nGlobalGridPositionLocalGrid, Grid::nNumGhostCells, Grid∷nR, ProcTop::nRank, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0, Grid::nV, and Grid::nW.

Referenced by setMainFunctions().

## 7.11.2.11 void calNewDenave None (Grid & grid)

This function is a dumby funciton, and doesn't do anything. In the case of a 1D calculation the average density is undefined, and only the density is used. This is different from the case where the 1D region exsists on the rank 0 processor, but the grid as a whole is really 2D or 3D. In which case calNewDenave R should be used instead.

## Parameters:

 $\leftrightarrow grid$ 

Referenced by setMainFunctions().

#### 7.11.2.12 void cal New Denave R (Grid & grid)

This function calculates the horizontal average density in a 3\1D region. This really just copies the density from the particular radial zone into the averaged density variable. This way it can be used exactly the same way in the 1D region as it is in the 3D region. This is done using the density in the new grid, and places the result into the new grid.

#### Parameters:

 $\leftrightarrow$  grid supplies the information needed to calculate the horizontal density average, it also stores the calculated horizontally averaged density.

 $References \ Grid::dLocalGridNew, \ Grid::nDenAve, \ Grid::nEndGhostUpdateExplicit, \\ Grid::nEndUpdateExplicit, \ Grid::nStartGhostUpdateExplicit, \ and \ Grid::nStartUpdateExplicit.$ 

## 7.11.2.13 void calNewDenave RT (Grid & grid)

This function calculates the horizontal average density in a 2D region from the new grid density and stores the result in the new grid.

#### Parameters:

 $\leftrightarrow$  grid supplies the information needed to calculate the horizontal density average, it also stores the calculated horizontally averaged density.

 $\label{localGridNew} References & Grid::dLocalGridNew, & Grid::dLocalGridOld, & Grid::nCenIntOffset, \\ Grid::nD, & Grid::nDCosThetaIJK, & Grid::nDenAve, & Grid::nEndGhostUpdateExplicit, \\ Grid::nEndUpdateExplicit, & Grid::nR, & Grid::nStartGhostUpdateExplicit, \\ Grid::nStartUpdateExplicit. & Grid::nStartUpdateExplicit. \\ \end{tabular}$ 

Referenced by setMainFunctions().

## 7.11.2.14 void calNewDenave RTP (Grid & grid)

This function calculates the horizontal average density in a 3D region from the new grid density and stores the result in the new grid.

#### Parameters:

 $\leftrightarrow$  grid supplies the information needed to calculate the horizontal density average, it also stores the calculated horizontally averaged density.

 $\label{localGridNew} References Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nD, Grid::nDCosThetaIJK, Grid::nDenAve, Grid::nDPhi, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nR, Grid::nStartGhostUpdateExplicit, and Grid::nStartUpdateExplicit.$ 

Referenced by setMainFunctions().

# 7.11.2.15 void calNewE\_R\_AD (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates new adiabatic energies using terms in the radial direction.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$

 $References \quad Time::dDeltat\_np1half, \quad Grid::dLocalGridNew, \quad Grid::dLocalGridOld, \quad Parameters::dPi, \quad Grid::nCenIntOffset, \quad Grid::nD, \quad Grid::nDM, \quad Grid::nDonorCellFrac, \quad Grid::nE, \\ Grid::nEndGhostUpdateExplicit, \quad Grid::nEndUpdateExplicit, \quad Grid::nP, \quad Grid::nQ0, \quad Grid::nR, \\ ProcTop::nRank, \quad Grid::nStartGhostUpdateExplicit, \quad Grid::nStartUpdateExplicit, \quad Grid::nU, \quad and \\ Grid::nU0.$ 

# 7.11.2.16 void calNewE\_R\_NA (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates new non-adiabatic energies using terms in the radial direction and includes radiation diffusion terms.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- $\leftarrow$  time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nE][i+1][j][k] in calculation of  $E_{i+1/2,j,k}$  setting it equal to value at i.

### **Boundary Conditions**

 $grid.dLocalGridOld[grid.nDM][i+1][0][0] \ and \ grid.dLocalGridOld[grid.nE][i+1][j][k] \ missing in the calculation of upwind gradient in dA1. Using the centered gradient instead.$ 

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nT][i+1][0][0]

References Time::dDeltat\_np1half, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dSigma, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nE, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGlobalGridPositionLocalGrid, Grid::nKappa, Grid::nNumGhostCells, Grid::nP, Grid::nQ0, Grid::nR, ProcTop::nRank, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nT, Grid::nU, and Grid::nU0.

Referenced by setMainFunctions().

# 7.11.2.17 void calNewE\_R\_NA\_LES (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates new non-adiabatic energies using terms in the radial direction and includes radiation diffusion terms. It also includes the terms for including real viscosity, used in the LES.

### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nE][i+1][j][k] in calculation of  $E_{i+1/2,j,k}$  setting it equal to value at i.

## **Boundary Conditions**

grid.dLocalGridOld[grid.nDM][i+1][0][0] and grid.dLocalGridOld[grid.nE][i+1][j][k] missing in the calculation of upwind gradient in dA1. Using the centered gradient instead.

# **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nT][i+1][0][0]

### **Boundary Conditions**

missing energy outside the model, assuming it is the same as that in the last zone. That causes this term to be zero.

References Time::dDeltat\_np1half, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dSigma, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nE, Grid::nEddyVisc, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nKappa, Grid::nP, Grid::nQ0, Grid::nR, ProcTop::nRank, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, and Grid::nU0.

# 7.11.2.18 void calNewE\_RT\_AD (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates new adiabatic energies using terms in the radial and theta directions.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated densities
- ← parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$

### **Boundary Conditions**

grid.dLocalGridOld[grid.nE][i+1][i][k] is missing

#### **Boundary Conditions**

 $grid.dLocalGridOld[grid.nDM][i+1][0][0] \ \ and \ \ grid.dLocalGridOld[grid.nE][i+1][j][k] \ \ missing using inner gradient for both$ 

References Time::dDeltat np1half, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid∷nDTheta, Grid∷nE, Grid::nSinThetaIJK, Grid∷nP, Grid::nQ0, Grid∷nQ1, Grid∷nR, ProcTop::nRank, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0, and Grid::nV.

Referenced by setMainFunctions().

# 7.11.2.19 void calNewE\_RT\_NA (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates new non-adiabatic energies using terms in the radial and theta directions and includes radiation diffusion terms.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nE][i+1][j][k] in calculation of  $E_{i+1/2,j,k}$  setting it equal to value at i.

### **Boundary Conditions**

 $grid.dLocalGridOld[grid.nDM][i+1][0][0] \ and \ grid.dLocalGridOld[grid.nE][i+1][j][k] \ missing in the calculation of upwind gradient in dA1. Using the centered gradient instead.$ 

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nT][i+1][0][0] using flux equals  $2\sigma T^4$  at surface.

References Time::dDeltat\_np1half, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dSigma, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDTheta, Grid::nE, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nKappa, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nR, Proc-Top::nRank, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU, and Grid::nV.

Referenced by setMainFunctions().

# 7.11.2.20 void calNewE\_RT\_NA\_LES (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates new non-adiabatic energies using terms in the radial and theta directions and includes radiation diffusion terms. It also includes the terms for including real viscosity, used in the LES.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- $\leftarrow$  time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$

## **Boundary Conditions**

Missing  $\Delta M_r$  outside model using Parameters::dAlpha times  $\Delta M_r$  in the last zone instead.

#### **Boundary Conditions**

Setting energy at surface equal to energy in last zone.

#### **Boundary Conditions**

missing eddy viscosity outside the model setting it to zero

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nT][i+1][0][0] using flux equals  $2\sigma T^4$  at surface.

### **Boundary Conditions**

missing energy outside the model, assuming it is the same as that in the last zone. That causes this term to be zero.

References Parameters::dAlpha, Parameters::dAlphaExtra, Time::dDeltat\_np1half, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dPrt, Parameters::dSigma, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDTheta, Grid::nE, Grid::nEddyVisc, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nKappa, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nR, Proc-Top::nRank, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nT, Grid::nU, Grid::nU0, and Grid::nV.

Referenced by setMainFunctions().

# 7.11.2.21 void calNewE\_RTP\_AD (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates new adiabatic energies using terms in the radial, theta, and phi directions.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- $\leftarrow$  time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$

#### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nE][i+1][j][k] in calculation of  $E_{i+1/2,j,k}$  setting it equal to zero

### **Boundary Conditions**

grid.dLocalGridOld[grid.nDM][i+1][0][0] and grid.dLocalGridOld[grid.nE][i+1][j][k] missing in the calculation of upwind gradient in dA1.Using the centered gradient.

References Time::dDeltat\_np1half, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nE, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, ProcTop::nRank, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU, Grid::nV, and Grid::nW.

Referenced by setMainFunctions().

# 7.11.2.22 void calNewE\_RTP\_NA (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates new non-adiabatic energies using terms in the radial, theta, and phi directions and includes radiation diffusion terms.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated densities
- ← parameters various parameters needed for the calculation
- $\leftarrow$  time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nE][i+1][j][k] in calculation of  $E_{i+1/2,j,k}$  setting it equal to the value at i.

## **Boundary Conditions**

grid.dLocalGridOld[grid.nDM][i+1][0][0] and grid.dLocalGridOld[grid.nE][i+1][j][k] missing in the calculation of upwind gradient in dA1. Using the centered gradient instead.

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nT][i+1][0][0]

References Time::dDeltat\_np1half, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dSigma, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nE, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nKappa, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, ProcTop::nRank, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU, Grid::nU, Grid::nV, and Grid::nW.

Referenced by setMainFunctions().

# 7.11.2.23 void calNewE\_RTP\_NA\_LES (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates new non-adiabatic energies using terms in the radial, theta, and phi directions and includes radiation diffusion terms. It also includes the terms for including real viscosity, used in the LES.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- $\leftarrow$  time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$

## **Boundary Conditions**

Missing  $\Delta M_r$  outside model using Parameters::dAlpha times  $\Delta M_r$  in the last zone instead.

### **Boundary Conditions**

Missing W at i+1, assuming the same as at i

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nE][i+1][j][k] in calculation of  $E_{i+1/2,j,k}$  setting it equal to the value at i.

# **Boundary Conditions**

missing density outside model, setting it to zero

## **Boundary Conditions**

missing eddy viscosity outside the model setting it to zero

### **Boundary Conditions**

 $grid.dLocalGridOld[grid.nDM][i+1][0][0] \ and \ grid.dLocalGridOld[grid.nE][i+1][j][k] \ missing in the calculation of upwind gradient in dA1. Using the centered gradient instead.$ 

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nT][i+1][0][0]

References Parameters::dAlpha, Parameters::dAlphaExtra, Time::dDeltat\_np1half, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dPrt, Parameters::dSigma, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nE, Grid::nEddyVisc, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGlobalGridPositionLocalGrid, Grid::nKappa, Grid::nNumGhostCells, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, Proc-Top::nRank, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nT, Grid::nU, Grid::nU, Grid::nV, and Grid::nW.

Referenced by setMainFunctions().

#### 7.11.2.24 void calNewEddyVisc None (Grid & grid, Parameters & parameters)

This function is a empty function used as a place holder when no eddy viscosity model is being used.

## Parameters:

- $\leftrightarrow \textit{grid}$
- $\leftarrow$  parameters

Referenced by setMainFunctions().

## 7.11.2.25 void calNewEddyVisc R CN (Grid & grid, Parameters & parameters)

This function calculates the eddy viscosity using a constant times the zoning with only the radial terms.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the eddy viscosity.
- ← parameters contains parameters used in calculating the eddy viscosity.

 $\label{lem:references} References & Parameters:: dEddyViscosity, & Grid:: dLocalGridNew, & Parameters:: dMaxConvectiveVelocity, & Grid:: nCenIntOffset, & Grid:: nEddyVisc, \\ Grid:: nEndGhostUpdateExplicit, & Grid:: nEndUpdateExplicit, & Grid:: nR, \\ Grid:: nStartGhostUpdateExplicit, & and Grid:: nStartUpdateExplicit. \\ \\ \\$ 

# 7.11.2.26 void calNewEddyVisc\_R\_SM (Grid & grid, Parameters & parameters)

This function calculates the eddy viscosity with only the radial terms.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the eddy viscosity.
- ← parameters contains parameters used in calculating the eddy viscosity.

 $\label{lem:condition} References & Parameters:: dEddy Viscosity, & Grid:: dLocal Grid New, & Grid:: dLocal Grid Old, \\ Grid:: nCenInt Offset, & Grid:: nD, & Grid:: nEddy Visc, & Grid:: nEnd Ghost Update Explicit, \\ Grid:: nEnd Update Explicit, & Grid:: nR, & Grid:: nStart Ghost Update Explicit, \\ Grid:: nStart Update Explicit, & Grid:: nU. \\ \\$ 

## 7.11.2.27 void calNewEddyVisc RT CN (Grid & grid, Parameters & parameters)

This function calculates the eddy viscosity using a constant times the zoning with only the radial and theta terms.

#### Parameters:

- $\leftrightarrow$  *grid* supplies the input for calculating the eddy viscosity.
- ← parameters contains parameters used in calculating the eddy viscosity.

 $References \quad Parameters:: dEddy Viscosity, \quad Grid:: dLocal Grid New, \quad Grid:: dLocal Grid Old, \\ Parameters:: dMax Convective Velocity, \quad Grid:: nCenInt Offset, \quad Grid:: nDT heta, \\ Grid:: nEddy Visc, \quad Grid:: nEndGhost Update Explicit, \quad Grid:: nEndUpdate Explicit, \quad Grid:: nR, \\ Grid:: nStart Ghost Update Explicit, \quad and \quad Grid:: nStart Update Explicit. \\ \end{cases}$ 

Referenced by setMainFunctions().

#### 7.11.2.28 void calNewEddyVisc RT SM (Grid & grid, Parameters & parameters)

This function calculates the eddy viscosity with only the radial and theta terms.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the eddy viscosity.
- ← parameters contains parameters used in calculating the eddy viscosity.

 $\label{eq:references} \begin{array}{llll} References & Parameters::dEddyViscosity, & Grid::dLocalGridNew, & Grid::dLocalGridOld, \\ Grid::nCenIntOffset, & Grid::nD, & Grid::nDTheta, & Grid::nEddyVisc, \\ Grid::nEndGhostUpdateExplicit, & Grid::nEndUpdateExplicit, & Grid::nR, \\ Grid::nStartGhostUpdateExplicit, & Grid::nStartUpdateExplicit, & Grid::nU, & Grid::nU, & Grid::nU, & Grid::nU, & Grid::nU, & Grid::nU, & Grid::nV. \\ \end{array}$ 

# 7.11.2.29 void calNewEddyVisc\_RTP\_CN (Grid & grid, Parameters & parameters)

This function calculates the eddy viscosity using a constant times the zoning with only the radial, theta, and phi terms.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the eddy viscosity.
- ← parameters contains parameters used in calculating the eddy viscosity.

 $References \ Parameters:: dEddy Viscosity, \ Grid:: dLocalGridNew, \ Grid:: dLocalGridOld, \ Parameters:: dMaxConvectiveVelocity, \ Grid:: nCenIntOffset, \ Grid:: nDPhi, \ Grid:: nDTheta, \ Grid:: nEddy Visc, \ Grid:: nEndGhost UpdateExplicit, \ Grid:: nEndUpdateExplicit, \ Grid:: nR, \ Grid:: nSinThetaIJK, \ Grid:: nStartGhost UpdateExplicit, \ and \ Grid:: nStartUpdateExplicit.$ 

Referenced by setMainFunctions().

# 7.11.2.30 void calNewEddyVisc\_RTP\_SM (Grid & grid, Parameters & parameters)

This function calculates the eddy viscosity with only the radial, theta, and phi terms.

#### Parameters:

- $\leftrightarrow$  *grid* supplies the input for calculating the eddy viscosity.
- $\leftarrow$  parameters contains parameters used in calculating the eddy viscosity.

# **Boundary Conditions**

assuming that theta velocity is constant across surface

#### **Boundary Conditions**

assume phi velocity is constant across surface

References Parameters::dEddyViscosity, Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nCotThetaIJK, Grid::nD, Grid::nDPhi, Grid::nDTheta, Grid::nEddyVisc, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nR, Grid::nSinThetaIJK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU, Grid::nV, and Grid::nW.

Referenced by setMainFunctions().

## 7.11.2.31 void calNewP GL (Grid & grid, Parameters & parameters)

This function calculates the pressure. It is calculated using the new values of quantities and places the result in the new grid. It uses a gamma law gas give in dEOS\_GL to calculate the pressure.

#### Parameters:

 $\leftrightarrow$  *grid* supplies the input for calculating the pressure and also accepts the result of the pressure calculations.

← *parameters* contains parameters used in calculating the pressure, namely the adiabatic gamma that is used.

References dEOS\_GL(), Grid::dLocalGridNew, Grid::nD, Grid::nE, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nStartGhostUpdateExplicit, and Grid::nStartUpdateExplicit.

Referenced by setMainFunctions().

# 7.11.2.32 void calNewPEKappaGamma\_TEOS (Grid & grid, Parameters & parameters)

This function calculates the Energy, pressure and opacity of a cell. It calculates it using the new vaules of quantities and places the result in the new grid.

#### Parameters:

- $\leftrightarrow grid$  supplies the input for calculating the pressure and also accepts the result of the pressure calculation
- ← parameters contains parameters used in calculating the pressure.

References Grid::dLocalGridNew, Parameters::eosTable, eos::getPEKappaGamma(), Grid::nD, Grid::nE, Grid::nEndGhostUpdateImplicit, Grid::nEndUpdateImplicit, Grid::nGamma, Grid::nKappa, Grid::nP, Grid::nStartGhostUpdateImplicit, Grid::nStartUpdateImplicit, and Grid::nT.

Referenced by implicitSolve R(), implicitSolve RT(), and implicitSolve RTP().

## 7.11.2.33 void calNewQ0 R GL (Grid & grid, Parameters & parameters)

This funciton calculates the artificial viscosity of a cell. It calculates it using the new values of quantities and places the result in the new grid. It does this for the radial component of the viscosity only. It uses the sound speed derived from the adiabatic gamma given for the gamma law gas equation of state.

#### Parameters:

- $\leftrightarrow$  *grid* supplies the input for calculating the artificial viscosity and also accepts the result of the artificial viscosity calculation.
- $\leftarrow$  parameters contains parameters used when calculating the artificial viscosity, namely the adiabatic gamma.

References Parameters::dA, Parameters::dAVThreshold, Parameters::dGamma, Grid::dLocalGridNew, Grid::nCenIntOffset, Grid::nD, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0, Grid::nR, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, and Grid::nU.

Referenced by setMainFunctions().

## 7.11.2.34 void calNewQ0 R TEOS (Grid & grid, Parameters & parameters)

This function calculates the artificial viscosity of a cell. It calculates it using the old values of quantities and places the result in the old grid. It does this for the radial component of the viscosity only. It uses a sound speed derived from a tabulated equaiton of state for the calculation.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the artificial viscosity and also accepts the result of the artificial viscosity calculation
- ← parameters contains parameters used in calculating the artificial viscosity.

References Parameters::dA, Parameters::dAVThreshold, Grid::dLocalGridNew, Grid::nCenIntOffset, Grid::nD, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGamma, Grid::nP, Grid::nQ0, Grid::nR, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, and Grid::nU.

Referenced by setMainFunctions().

# 7.11.2.35 void calNewQ0Q1 RT GL (Grid & grid, Parameters & parameters)

This function calculates the artificial viscosity of a cell. It calculates it using the new values of quantities and places the result in the new grid. It does this for the radial and theta components of the viscosity. It uses the sound speed derived from the adiabatic gamma given for the gamma law gas equation of state.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the artificial viscosity and also accepts the result of the artificial viscosity calculations.
- ← *parameters* contains parameters used when calculating the artificial viscosity, namely the adiabatic gamma.

References Parameters::dA, Parameters::dAVThreshold, Parameters::dGamma, Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nD, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nV.

Referenced by setMainFunctions().

# 7.11.2.36 void calNewQ0Q1 RT TEOS (Grid & grid, Parameters & parameters)

This function calculates the artificial viscosity of a cell. It calculates it using the old values of quantities and places the result in the old grid. It does this for two component of the viscosity.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the artificial viscosity and also accepts the result of the artificial viscosity calculation
- ← parameters contains parameters used in calculating the artificial viscosity.

References Parameters::dA, Parameters::dAVThreshold, Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nD, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGamma, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, and Grid::nV.

# 7.11.2.37 void calNewQ0Q1Q2\_RTP\_GL (Grid & grid, Parameters & parameters)

This function calculates the artificial viscosity of a cell. It calculates it using the new values of quantities and places the result in the new grid. It does this for the radial, theta, and phi components of the viscosity. It uses the sound speed derived from the adiabatic gamma given for the gamma law gas equation of state.

#### Parameters:

- $\leftrightarrow$  *grid* supplies the input for calculating the artificial viscosity and also accepts the result of the artificial viscosity calculations.
- parameters contains parameters used when calculating the artificial viscosity, namely the adiabatic gamma.

References Parameters::dA, Parameters::dAVThreshold, Parameters::dGamma, Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid∷nD, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0. Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid∷nQ1, Grid::nQ2, Grid::nR, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nV, Grid∷nW.

Referenced by setMainFunctions().

# 7.11.2.38 void calNewQ0Q1Q2\_RTP\_TEOS (Grid & grid, Parameters & parameters)

This function calculates the artificial viscosity of a cell. It calculates it using the old values of quantities and places the result in the old grid. It does this for the three component of the viscosity.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the artificial viscosity and also accepts the result of the artificial viscosity calculation
- $\leftarrow$  parameters contains parameters used in calculating the artificial viscosity.

References Parameters::dA, Parameters::dAVThreshold, Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nD, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGamma, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nV, and Grid::nW.

Referenced by setMainFunctions().

#### 7.11.2.39 void calNewR (Grid & grid, Time & time)

This function calculates the radii, from the new radial grid velocities

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated radial velocities
- ← *time* contains time information, e.g. time step, current time etc.

References Time::dDeltat\_np1half, Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nR, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, and Grid::nU0.

Referenced by setMainFunctions().

# 7.11.2.40 void calNewTPKappaGamma\_TEOS (Grid & grid, Parameters & parameters)

This function calculates the Temperature, pressure and opacity of a cell. It calculates it using the new vaules of quantities and places the result in the new grid.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the pressure and also accepts the result of the pressure calculation
- $\leftarrow$  parameters contains parameters used in calculating the pressure.

References Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dTolerance, Parameters::eosTable, eos::getEAndDTDE(), eos::getPKappaGamma(), Grid::nD, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nE, Grid::nGamma, Grid::nKappa, Parameters::nMaxIterations, Grid∷nP, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, and Grid::nT.

Referenced by setMainFunctions().

# 7.11.2.41 void calNewU0\_R (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop, MessPass & messPass)

This function calculates the radial grid velocity, it does so by considering only the radial terms

#### Parameters:

- $\leftrightarrow grid$  contains the local grid, and will hold the newly updated radial grid velocities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology
- $\leftarrow messPass$

#### Todo

At some point I will likely want to make this function compatiable with a 3D domain decomposition instead of a purely radial domain decomposition.

#### **Boundary Conditions**

assuming density outside the star is 0 for the purposes of calculating the grid velocity. This boundary condition works, but is probably not strictly correct. The proper condition can be calculated by knowing that the mass conservation is independent of u0\_ip1half at the surface. This allows one to simply use the new densities and calculate the grid velocity from them.

References Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nCenIntOffset, ProcTop::nCoords, Grid::nD, Grid::nDonorCellFrac, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit,

 $\label{lem:procTop::nRadialNeighbors} ProcTop::nRadialNeighborNeighborIDs, ProcTop::nRadialNeighborRanks, ProcTop::nRank, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0, MessPass::typeRecvNewVar, and Mess-Pass::typeSendNewVar.$ 

Referenced by setMainFunctions().

# 7.11.2.42 void calNewU0\_RT (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop, MessPass & messPass)

This function calculates the radial grid velocity, and does it by only considering the radial and theta terms

#### Parameters:

- $\leftrightarrow grid$  contains the local grid, and will hold the newly updated radial grid velocities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology
- $\leftrightarrow messPass$  handles data needed for message passing

#### Todo

At some point I will likely want to make this function compatiable with a 3D domain decomposition instead of a purely radial domain decomposition.

## **Boundary Conditions**

grid.dLocalGridOld[grid.nD][i+1][j][k] is missing

 $\label{lem:condition} References\ Grid::dLocalGridNew,\ Grid::dLocalGridOld,\ Grid::nCenIntOffset,\ ProcTop::nCoords,\ Grid::nD,\ Grid::nDCosThetaIJK,\ Grid::nDonorCellFrac,\ Grid::nEndGhostUpdateExplicit,\ Grid::nEndUpdateExplicit,\ Grid::nLocalGridDims,\ Grid::nNumGhostCells,\ ProcTop::nNumRadialNeighbors,\ Grid::nR,\ ProcTop::nRadialNeighborNeighborIDs,\ ProcTop::nRadialNeighborRanks,\ ProcTop::nRank,\ Grid::nSinThetaIJp1halfK,\ Grid::nStartGhostUpdateExplicit,\ Grid::nStartUpdateExplicit,\ Grid::nU,\ Grid::nU,\ Grid::nV,\ MessPass::typeRecvNewVar,\ and\ MessPass::typeSendNewVar.$ 

Referenced by setMainFunctions().

# 7.11.2.43 void calNewU0\_RTP (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop, MessPass & messPass)

This function calculates the radial grid velocity, and does it by considering all radial, theta and phi terms

#### Parameters:

- $\leftrightarrow grid$  contains the local grid, and will hold the newly updated radial grid velocities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology
- $\leftrightarrow messPass$  handles data needed for message passing

#### Todo

At some point I will likely want to make this function compatiable with a 3D domain decomposition instead of a purely radial domain decomposition.

References Grid::dLocalGridNew, Grid::dLocalGridOld. Grid::nCenIntOffset, Proc-Top::nCoords, Grid::nD, Grid::nDCosThetaIJK, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nLocalGridDims, Grid::nNumGhostCells, ProcTop::nNumRadialNeighbors, Grid::nR, ProcTop::nRadialNeighborNeighborIDs, ProcTop::nRadialNeighborRanks, ProcTop::nRank, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid∷nU, Grid::nU0, Grid∷nV, Grid::nW, MessPass::typeRecvNewVar, and Pass::typeSendNewVar.

Referenced by setMainFunctions().

# 7.11.2.44 void calNewU\_R (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the radial velocity, and does it by only considering the radial terms.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated radial velocities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology

#### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nD][nICen+1][j][k] in calculation of  $\rho_{i+1/2}$ , setting it to 0.0

# **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nP][nICen+1][j][k] in calculation of  $S_1$ , setting it to -1.0\*grid.dLocalGridOld[grid.nP][nICen][j][k].

#### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nDM][nICen+1][0][0] in calculation of centered  $A_1$  gradient, setting it to zero.

## **Boundary Conditions**

 $\label{lem:missing grid.dLocalGridOld[grid.nU][i+1][j][k] and grid.dLocalGridOld[grid.nDM][nICen+1][0][0] in calculation of upwind gradient, when moving inward. Using centered gradient instead.}$ 

References Parameters::dAlpha, Parameters::dAlphaExtra, Time::dDeltat\_n, Parameters::dG, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nCenIntOffset, Grid::nD, Grid::nDM, Grid::nDonorCellFrac, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGlobalGridPositionLocalGrid, Grid::nM, Grid::nP, Grid::nQ0, Grid::nR, Grid::nStartGhostUpdateExplicit, Grid::nU, and Grid::nU0.

Referenced by calNewVelocities R().

# 7.11.2.45 void calNewU\_R\_LES (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the radial velocity, and does it by only considering the radial terms. It also includes the terms for including real viscosity, used in the LES.

#### Parameters:

- $\leftrightarrow grid$  contains the local grid, and will hold the newly updated radial velocities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nD][nICen+1][j][k] in calculation of  $\rho_{i+1/2}$ , setting it to 0.0

### **Boundary Conditions**

missing grid.dLocalGridOld[grid.nU][i+1][j][k] using velocity at i

### **Boundary Conditions**

Assuming eddy viscosity outside model is zero.

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nP][nICen+1][j][k] in calculation of  $S_1$ , setting it to -1.0\*grid.dLocalGridOld[grid.nP][nICen][j][k].

#### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nDM][nICen+1][0][0] in calculation of centered  $A_1$  gradient, setting it to zero.

#### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nU][i+1][j][k] and grid.dLocalGridOld[grid.nDM][nICen+1][0][0] in calculation of upwind gradient, when moving inward. Using centered gradient instead.

References Parameters::dAlpha, Parameters::dAlphaExtra, Time::dDeltat n, Pa-Grid:: dLocalGridNew,rameters::dG, Grid::dLocalGridOld, Parameters::dPi, Grid::nCenIntOffset, Grid∷nD, Grid::nDM, Grid::nDonorCellFrac, Grid::nEddyVisc, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nM, Grid::nP, Grid::nQ0, Grid::nR, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0.

Referenced by calNewVelocities\_R\_LES().

# 7.11.2.46 void calNewU\_RT (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the radial velocity, and does it by only considering the radial and theta terms.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated radial velocities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nD][nICen+1][j][k] in calculation of  $\rho_{i+1/2,j,k}$ , setting it to zero.

## **Boundary Conditions**

assuming theta velocity is constant across surface

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nDenAve][nICen+1][0][0] in calculation of  $\langle \rho \rangle_{i+1/2}$ , setting it to zero.

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nP][nICen+1][j][k] in calculation of  $S_1$ , setting it to -1.0\*dP\_ijk\_n.

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nDM][nICen+1][0][0] in calculation of centered  $A_1$  gradient, setting it to zero.

#### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nU][i+1][j][k] and grid.dLocalGridOld[grid.nDM][nICen+1][0][0] in calculation of upwind gradient, when moving inward. Using centered gradient instead.

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nDM][i+1][0][0] in calculation of  $S_1$  using Parameters::dAlpha \*grid.dLocalGridOld[grid.nDM][nICen][0][0] instead.

References Parameters::dAlpha, Parameters::dAlphaExtra, Time::dDeltat\_n, Parameters::dG, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDTheta, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nM, Grid::nP, Grid::nQ0, Grid::nR, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0, and Grid::nV.

Referenced by calNewVelocities RT().

# 7.11.2.47 void calNewU\_RT\_LES (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the radial velocity, and does it by only considering the radial and theta terms. It also includes the terms for including real viscosity, used in the LES.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated radial velocities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nDM][i+1][0][0] in calculation of  $S_1$  using Parameters::dAlpha \*grid.dLocalGridOld[grid.nDM][nICen][0][0] instead.

## **Boundary Conditions**

Missing density outside of surface, setting it to zero.

### **Boundary Conditions**

Missing density outside model, setting it to zero.

# **Boundary Conditions**

assuming theta and phi velocity same outside star as inside.

### **Boundary Conditions**

Assuming theta velocities are constant across surface.

## **Boundary Conditions**

assuming that \$V\$ at \$i+1\$ is equal to \$v\$ at \$i\$.

#### **Boundary Conditions**

Missing pressure outside surface setting it equal to negative pressure in the center of the first cell so that it will be zero at surface.

#### **Boundary Conditions**

assume viscosity is zero outside the star.

#### **Boundary Conditions**

Missing mass outside model, setting it to zero.

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nU][i+1][j][k] and grid.dLocalGridOld[grid.nDM][nICen+1][0][0] in calculation of upwind gradient, when moving inward. Using centered gradient instead.

References Parameters::dAlpha, Parameters::dAlphaExtra, Time::dDeltat n, ters::dG, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nCenIntOffset, Grid::nCotThetaIJK, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDTheta, Grid::nEddyVisc, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid∷nM. Grid::nQ0, Grid::nQ1, Grid∷nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0, and Grid::nV.

Referenced by calNewVelocities RT LES().

# 7.11.2.48 void calNewU\_RTP (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the radial velocity, and does it by including all radial, theta and phi terms.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated radial velocities
- $\leftarrow$  parameters various parameters needed for the calculation
- $\leftarrow$  time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology

### **Boundary Conditions**

missing grid.dLocalGridOld[grid.nU][i+1][j][k] in calculation of  $u_{i+1,j,k}$  setting  $u_{i+1,j,k}=u_{i+1/2,j,k}$ .

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nD][i+1][j][k] in calculation of  $\rho_{i+1/2,j,k}$ , setting it to zero.

### **Boundary Conditions**

assuming theta velocity is constant across the surface.

### **Boundary Conditions**

assuming phi velocity is constant across the surface.

#### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nDenAve][nICen+1][0][0] in calculation of  $\langle \rho \rangle_{i+1/2}$  setting it to zero.

#### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nDM][nICen+1][0][0] in calculation of centered  $A_1$  gradient, setting it equal to Parameters::dAlpha grid.dLocalGridOld[grid.nDM][nICen][0][0].

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nU][i+1][j][k] and grid.dLocalGridOld[grid.nDM][nICen+1][0][0] in calculation of upwind gradient, when moving inward. Using centered gradient instead.

## **Boundary Conditions**

$$\label{eq:missing_missing} \begin{split} & \text{Missing} \quad \text{grid.dLocalGridOld[grid.nDM][i+1][0][0]} \quad \text{in} \quad \text{calculation} \quad \text{of} \quad S_1 \quad \text{using} \\ & \text{Parameters::dAlpha} *& \text{grid.dLocalGridOld[grid.nDM][nICen][0][0]} \quad \text{instead.} \end{split}$$

References Parameters::dAlpha, Parameters::dAlphaExtra, Time::dDeltat\_n, Parameters::dG, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nM, Grid::nP, Grid::nQ0, Grid::nR, Grid::nSinThetaIJK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU, Grid::nV, and Grid::nW.

Referenced by calNewVelocities RTP().

# 7.11.2.49 void calNewU\_RTP\_LES (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the radial velocity, and does it by including all radial, theta and phi terms. It also includes the terms for including real viscosity, used in the LES.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated radial velocities
- $\leftarrow$  parameters various parameters needed for the calculation
- $\leftarrow$  time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nDM][i+1][0][0] in calculation of  $S_1$  using Parameters::dAlpha \*grid.dLocalGridOld[grid.nDM][nICen][0][0] instead.

# **Boundary Conditions**

Missing density outside of surface, setting it to zero.

### **Boundary Conditions**

Missing density outside model, setting it to zero.

### **Boundary Conditions**

assuming theta and phi velocity same outside star as inside.

#### **Boundary Conditions**

Assuming theta velocities are constant across surface.

## **Boundary Conditions**

assuming that V at i+1 is equal to v at i.

### **Boundary Conditions**

Missing pressure outside surface setting it equal to negative pressure in the center of the first cell so that it will be zero at surface.

#### **Boundary Conditions**

assume viscosity is zero outside the star.

### **Boundary Conditions**

Missing mass outside model, setting it to zero.

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nU][i+1][j][k] and grid.dLocalGridOld[grid.nDM][nICen+1][0][0] in calculation of upwind gradient, when moving inward. Using centered gradient instead.

References Parameters::dAlpha, Parameters::dAlphaExtra, Time::dDeltat\_n, Parameters::dG, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nCenIntOffset, Grid::nCotThetaIJK, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nEddyVisc, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGlobalGridPositionLocalGrid, Grid::nM, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU, Grid::nV, and Grid::nW.

Referenced by calNewVelocities\_RTP\_LES().

# 7.11.2.50 void calNewV\_RT (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the theta velocity, and does it by only considering the radial and theta terms.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated theta velocities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology

### **Boundary Conditions**

grid.dLocalGridOld[grid.nV][i+1][j+1][k] is missing

### **Boundary Conditions**

missing upwind gradient, using centred gradient instead

References  $Time::dDeltat_n$ , Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDTheta, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nR, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU, Grid::nU, Grid::nU, Grid::nU, Grid::nV.

Referenced by calNewVelocities RT().

# 7.11.2.51 void calNewV\_RT\_LES (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the theta velocity, and does it by only considering the radial and theta terms. It also includes the terms for including real viscosity, used in the LES.

- $\leftrightarrow grid$  contains the local grid, and will hold the newly updated theta velocities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology

### **Boundary Conditions**

Assuming density outside star is zero

### **Boundary Conditions**

Assuming theta velocity is constant across surface.

# **Boundary Conditions**

Assuming eddy viscosity is zero at surface.

References Time::dDeltat\_n, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nCenIntOffset, Grid::nCotThetaIJp1halfK, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDTheta, Grid::nEddyVisc, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU, Grid::nU, and Grid::nV.

Referenced by calNewVelocities RT LES().

# 7.11.2.52 void calNewV\_RTP (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the theta velocity, and does it by considering the radial, theta, and phi terms.

#### Parameters:

- $\leftrightarrow grid$  contains the local grid, and will hold the newly updated theta velocities
- ← parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology

# **Boundary Conditions**

Assuming theta and phi velocities are the same at the surface of the star as just inside the star.

#### **Boundary Conditions**

ussing cetnered gradient for upwind gradient outside star at surface.

References Time::dDeltat n, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nCotThetaIJp1halfK, Grid::nD, Grid::nDenAve, Grid::nCenIntOffset, Grid::nDM, Grid::nDonorCellFrac, Grid::nEndGhostUpdateExplicit, Grid::nDPhi, Grid::nDTheta, Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0, Grid::nV, and Grid::nW.

Referenced by calNewVelocities\_RTP().

# 7.11.2.53 void calNewV\_RTP\_LES (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the theta velocity, and does it by considering the radial, theta, and phi terms. It also includes the terms for including real viscosity, used in the LES.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated theta velocities
- ← parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology

### **Boundary Conditions**

Assuming density outside star is zero

# **Boundary Conditions**

Assuming theta velocity is constant across surface.

### **Boundary Conditions**

Assuming eddy viscosity is zero at surface.

References Time::dDeltat n, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi. Grid::nCenIntOffset, Grid::nCotThetaIJp1halfK, Grid∷nD, Grid∷nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nEddyVisc, Grid:: nEndGhostUpdateExplicit,Grid::nEndUpdateExplicit, Grid::nGlobalGridPositionLocalGrid, Grid::nNumGhostCells, Grid::nP, Grid::nQ0, Grid∷nQ1, Grid::nQ2, Grid∷nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0, Grid::nV, and Grid::nW.

Referenced by calNewVelocities RTP LES().

# 7.11.2.54 void calNewVelocities\_R (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function simply calls a function that calculate the radial velocity. Calls the function calNewU R to calculate radial velocity, including only radial terms.

### Parameters:

- $\leftrightarrow$  *grid* contains the local grid data and supplies the needed data to calculate the new velocities as well as holding the new velocities.
- $\leftarrow$  parameters contains parameters used in the calculation of the new velocities.
- ← time contains time step information, current time step, and current time
- $\leftarrow procTop$  contains processor topology information

References calNewU\_R().

Referenced by setMainFunctions().

# 7.11.2.55 void calNewVelocities R\_LES (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function simply calls a function that calculate the radial velocity. Calls the function calNewU R to calculate radial velocity, including only radial terms.

#### Parameters:

- $\leftarrow$  parameters contains parameters used in the calculation of the new velocities.
- $\leftarrow$  time contains time step information, current time step, and current time
- $\leftarrow procTop$  contains processor topology information

References calNewU\_R\_LES().

# 7.11.2.56 void calNewVelocities\_RT (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function simply calls two other functions that calculate the radial and theta velocities. Calls the two functions calNewU\_RT and calNewV\_RT to calculate radial and theta velocities, including both radial and theta terms.

#### Parameters:

- $\leftrightarrow$  *grid* contains the local grid data and supplies the needed data to calculate the new velocities as well as holding the new velocities.
- $\leftarrow$  parameters contains parameters used in the calculation of the new velocities.
- ← time contains time step information, current time step, and current time
- $\leftarrow procTop$  contains processor topology information

References  $calNewU_RT()$ , and  $calNewV_RT()$ .

Referenced by setMainFunctions().

# 7.11.2.57 void calNewVelocities RT\_LES (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function simply calls two other functions that calculate the radial and theta velocities. Calls the two functions calNewU\_RT and calNewV\_RT to calculate radial and theta velocities, including both radial and theta terms.

### Parameters:

- $\leftrightarrow$  grid contains the local grid data and supplies the needed data to calculate the new velocities as well as holding the new velocities.
- $\leftarrow$  parameters contains parameters used in the calculation of the new velocities.
- $\leftarrow time$  contains time step information, current time step, and current time
- $\leftarrow procTop$  contains processor topology information

References calNewU RT LES(), and calNewV RT LES().

Referenced by setMainFunctions().

# 7.11.2.58 void calNewVelocities\_RTP (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function simply calls three other functions that calculate the radial, theta and phi velocities. Calls the two functions calNewU\_RTP, calNewV\_RTP and calNewW\_RTP to calculate radial, theta, and phi velocities, including radial, theta, and phi terms.

#### Parameters:

- $\leftrightarrow$  *grid* contains the local grid data and supplies the needed data to calculate the new velocities as well as holding the new velocities.
- $\leftarrow$  parameters contains parameters used in the calculation of the new velocities.
- $\leftarrow$  time contains time step information, current time step, and current time
- $\leftarrow procTop$  contains processor topology information

References calNewU\_RTP(), calNewV\_RTP(), and calNewW\_RTP(). Referenced by setMainFunctions().

# 7.11.2.59 void calNewVelocities RTP LES (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function simply calls three other functions that calculate the radial, theta and phi velocities. Calls the two functions calNewU\_RTP, calNewV\_RTP and calNewW\_RTP to calculate radial, theta, and phi velocities, including radial, theta, and phi terms.

### Parameters:

- $\leftrightarrow$  grid contains the local grid data and supplies the needed data to calculate the new velocities as well as holding the new velocities.
- $\leftarrow$  parameters contains parameters used in the calculation of the new velocities.
- ← time contains time step information, current time step, and current time
- $\leftarrow procTop$  contains processor topology information

References calNewU\_RTP\_LES(), calNewV\_RTP\_LES(), and calNewW\_RTP\_LES(). Referenced by setMainFunctions().

# 7.11.2.60 void calNewW\_RTP (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the phi velocity, and does it by only considering the radial, theta, and phi terms.

- $\leftrightarrow grid$  contains the local grid, and will hold the newly updated theta velocities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology

### **Boundary Conditions**

missing grid.dLocalGridOld[grid.nW][i+1][j][k] assuming that the phi velocity at the outter most interface is the same as the phi velocity in the center of the zone.

# **Boundary Conditions**

missing grid.dLocalGridOld[grid.nW][i+1][j][k] in outter most zone. This is needed to calculate the upwind gradient for donnor cell. The centered gradient is used instead when moving in the negative direction.

References Time::dDeltat n, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nCenIntOffset, Grid::nCotThetaIJK, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR. Grid::nSinThetaIJK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0, Grid::nV, and Grid::nW.

Referenced by calNewVelocities RTP().

# 7.11.2.61 void calNewW\_RTP\_LES (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the phi velocity, and does it by only considering the radial, theta, and phi terms. It also includes the terms for including real viscosity, used in the LES.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated theta velocities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology

#### **Boundary Conditions**

assume theta and phi velocities are constant across surface

### **Boundary Conditions**

assume eddy viscosity is zero at surface

# **Boundary Conditions**

assume upwind gradient is the same as centered gradient across surface

Time::dDeltat n, References Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nCenIntOffset, Grid::nCotThetaIJK, Grid∷nD, Grid::nDenAve, Grid∷nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nEndGhostUpdateExplicit, Grid::nEddyVisc, Grid::nEndUpdateExplicit, Grid::nGlobalGridPositionLocalGrid, Grid::nNumGhostCells, Grid::nP, Grid::nQ0, Grid∷nQ1, Grid::nQ2, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0, Grid::nV, and Grid::nW.

Referenced by calNewVelocities RTP LES().

### 7.11.2.62 void calOldDenave None (Grid & grid)

This function is a dumby funciton, and doesn't do anything. In the case of a 1D calculation the average density is undefined, and only the density is used. This is different from the case where the 1D region exsists on the rank 0 processor, but the grid as a whole is really 2D or 3D. In which case calOldDenave R should be used instead.

### 7.11.2.63 void calOldDenave R (Grid & grid)

This function does nothing as the averaged density is not needed in 1D calculations.

#### Parameters:

 $\leftrightarrow$  *grid* supplies the information needed to calculate the horizontal density average, it also stores the calculated horizontally averaged density.

 $\label{lem:condition} References \ Grid::dLocalGridOld, \ Grid::nD, \ Grid::nDenAve, \ Grid::nEndGhostUpdateExplicit, \\ Grid::nEndGhostUpdateImplicit, \ Grid::nEndUpdateExplicit, \ Grid::nEndUpdateImplicit, \\ Grid::nStartGhostUpdateExplicit, \ Grid::nStartGhostUpdateImplicit, \\ Grid::nStartUpdateExplicit, \ and \ Grid::nStartUpdateImplicit.$ 

Referenced by initInternalVars().

### 7.11.2.64 void calOldDenave RT (Grid & grid)

This function calculates the horizontal average density in a 2D region. This function differs from calNewDenave\_RT in that it calculates the average density from the old grid density and stores the result in the old grid. While calNewDenave\_RT calculates the average density from the new grid density and places the result in the new grid.

### Parameters:

 $\leftrightarrow$  grid supplies the information needed to calculate the horizontal density average, it also stores the calculated horizontally averaged density.

References Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nD, Grid::nDCosThetaIJK, Grid::nDenAve, Grid::nEndGhostUpdateExplicit, Grid::nEndGhostUpdateImplicit, Grid::nEndUpdateExplicit, Grid::nEndUpdateImplicit, Grid::nStartGhostUpdateImplicit, Grid::nStartGhostUpdateImplicit, Grid::nStartUpdateExplicit, and Grid::nStartUpdateImplicit.

Referenced by initInternalVars().

### 7.11.2.65 void calOldDenave RTP (Grid & grid)

This function calculates the horizontal average density in a 3D region. This function differs from calNewDenave\_RTP in that it calculates the average density from the old grid density and stores the result in the old grid. While calNewDenave\_RTP calculates the average density from the new grid density and places the result in the new grid.

#### Parameters:

 $\leftrightarrow$  grid supplies the information needed to calculate the horizontal density average, it also stores the calculated horizontally averaged density.

References Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nD, Grid::nDCosThetaIJK, Grid::nDenAve, Grid::nDPhi, Grid::nEndGhostUpdateExplicit, Grid::nEndGhostUpdateImplicit, Grid::nEndUpdateExplicit, Grid::nEndUpdateImplicit, Grid::nStartGhostUpdateImplicit, Grid::nStartGhostUpdateImplicit, Grid::nStartUpdateExplicit, and Grid::nStartUpdateImplicit.

Referenced by initInternalVars().

# 7.11.2.66 void calOldEddyVisc R CN (Grid & grid, Parameters & parameters)

Calculates the eddy viscosity using a constant times the zoning including only the radial terms. It puts the result into the old grid. This function is used to initalize the eddy viscosity when the code begins execution.

 $References \quad Parameters:: dEddy Viscosity, \quad Grid:: dLocal Grid New, \quad Grid:: dLocal Grid Old, \\ Parameters:: dMax Convective Velocity, \quad Grid:: nCenInt Offset, \quad Grid:: nEddy Visc, \\ Grid:: nEnd Ghost Update Explicit, \quad Grid:: nEnd Update Explicit, \quad Grid:: nNum Ghost Cells, \quad Grid:: nR, \\ Grid:: nStart Ghost Update Explicit, \quad and \quad Grid:: nStart Update Explicit. \\ \end{cases}$ 

Referenced by initInternalVars().

# 7.11.2.67 void calOldEddyVisc R SM (Grid & grid, Parameters & parameters)

Calculates the eddy viscosity including only the radial terms. It puts the result into the old grid. This function is used to initalize the eddy viscosity when the code begins execution. It uses the Smagorinsky model for calculating the eddy viscosity.

#### Parameters:

- $\leftrightarrow$  *grid* supplies the input for calculating the eddy viscosity.
- ← parameters contains parameters used in calculating the eddy viscosity.

References Parameters::dEddyViscosity, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nD, Grid::nEddyVisc, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nNumGhostCells, Grid::nR, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, and Grid::nU.

Referenced by initInternalVars().

#### 7.11.2.68 void calOldEddyVisc RT CN (Grid & grid, Parameters & parameters)

Calculates the eddy viscosity using a constant times the zoning including only the radial and theta terms. It puts the result into the old grid. This function is used to initialize the eddy viscosity when the code begins execution.

# Parameters:

- $\leftrightarrow$  *grid* supplies the input for calculating the eddy viscosity.
- ← parameters contains parameters used in calculating the eddy viscosity.

 $References \ Parameters:: dEddy Viscosity, \ Grid:: dLocalGridNew, \ Grid:: dLocalGridOld, \ Parameters:: dMaxConvectiveVelocity, \ Grid:: nCenIntOffset, \ Grid:: nDTheta, \ Grid:: nEddy Visc, \ Grid:: nEndGhostUpdateExplicit, \ Grid:: nEndUpdateExplicit, \ Grid:: nNumGhostCells, \ Grid:: nR, \ Grid:: nStartGhostUpdateExplicit, \ and \ Grid:: nStartUpdateExplicit.$ 

Referenced by initInternalVars().

# 7.11.2.69 void calOldEddyVisc RT SM (Grid & grid, Parameters & parameters)

Calculates the eddy viscosity including only the radial and theta terms. It puts the result into the old grid. This function is used to initalize the eddy viscosity when the code begins execution. It uses the Smagorinsky model for calculating the eddy viscosity.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the eddy viscosity.
- ← parameters contains parameters used in calculating the eddy viscosity.

References Parameters::dEddyViscosity, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nD, Grid::nDTheta, Grid::nEddyVisc, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nNumGhostCells, Grid::nR, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU, and Grid::nV.

Referenced by initInternalVars().

# 7.11.2.70 void calOldEddyVisc\_RTP\_CN (Grid & grid, Parameters & parameters)

Calculates the eddy viscosity using a constant times the zoning including the radial, theta, and phi terms. It puts the result into the old grid. This function is used to initalize the eddy viscosity when the code begins execution.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the eddy viscosity.
- ← parameters contains parameters used in calculating the eddy viscosity.

 $\label{lem:convective} References \ Parameters:: dEddy Viscosity, \ Grid:: dLocalGridNew, \ Grid:: dLocalGridOld, \ Parameters:: dMaxConvectiveVelocity, \ Grid:: nCenInt Offset, \ Grid:: nDPhi, \ Grid:: nDTheta, \ Grid:: nEddy Visc, \ Grid:: nEndGhost Update Explicit, \ Grid:: nEndUpdate Explicit, \ Grid:: nStartGhost Update Explicit, \ and \ Grid:: nStart Update Explicit.$ 

Referenced by initInternalVars().

# 7.11.2.71 void calOldEddyVisc\_RTP\_SM (Grid & grid, Parameters & parameters)

Calculates the eddy viscosity including the radial, theta, and phi terms. It puts the result into the old grid. This function is used to initalize the eddy viscosity when the code begins execution. It uses the Smagorinsky model for calculating the eddy viscosity.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the eddy viscosity.
- ← parameters contains parameters used in calculating the eddy viscosity.

# **Boundary Conditions**

assuming that theta velocity is constant across surface

### **Boundary Conditions**

assume phi velocity is constant across surface

References Parameters::dEddyViscosity, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nCotThetaIJK, Grid::nD, Grid::nDPhi, Grid::nDTheta, Grid::nEddyVisc, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nNumGhostCells, Grid::nR, Grid::nSinThetaIJK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU, Grid::nV, and Grid::nW.

Referenced by initInternalVars().

# 7.11.2.72 void calOldP GL (Grid & grid, Parameters & parameters)

This function calculates the pressure using a gamma law gas, calculate by dEOS GL.

#### Parameters:

- $\leftrightarrow$  *grid* supplies the input for calculating the pressure and also accepts the results of the pressure calculations
- $\leftarrow$  parameters contains parameters used in calculating the pressure, namely the value of the adiabatic gamma

References dEOS\_GL(), Grid::dLocalGridOld, Grid::nD, Grid::nE, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nStartGhostUpdateExplicit, and Grid::nStartUpdateExplicit. Grid::nP,

Referenced by initInternalVars().

# 7.11.2.73 void calOldPEKappaGamma\_TEOS (Grid & grid, Parameters & parameters)

This function calculates the pressure, energy, opacity, and adiabatic index of a cell. It calculates it using the old vaules of quantities and places the result in the old grid. This function is used to initialize the internal variables pressure, energy and kappa, and is suitable for both 1D and 3D calculations.

#### Parameters:

- $\leftrightarrow grid$  supplies the input for calculating the pressure and also accepts the result of the pressure calculation
- $\leftarrow$  parameters contains parameters used in calculating the pressure.

References Grid::dLocalGridOld, Parameters::eosTable, eos::getPEKappaGamma(), Grid::nEndGhostUpdateExplicit, Grid::nEndGhostUpdateImplicit, Grid::nD, Grid::nE, Grid::nEndUpdateExplicit, Grid::nEndUpdateImplicit, Grid::nGamma, Grid::nKappa, Grid::nNumGhostCells, Grid::nP, Grid::nStartGhostUpdateExplicit, Grid::nStartGhostUpdateImplicit, Grid::nStartUpdateExplicit, Grid::nStartUpdateImplicit, and Grid::nT.

Referenced by initInternalVars().

# 7.11.2.74 void calOldQ0 R GL (Grid & grid, Parameters & parameters)

This function calculates the artificial viscosity of a cell. It calculates it using the old vaules of quantities and places the result in the old grid. It does this for the radial component of the viscosity only. This function is used when using a gamma law gas equation of state.

### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the artificial viscosity and also accepts the result of the artificial viscosity calculation
- ← parameters contains parameters used in calculating the artificial viscosity.

References Parameters::dA, Parameters::dAVThreshold, Parameters::dGamma, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nD, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0, Grid::nR, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, and Grid::nU.

Referenced by initInternalVars().

# $7.11.2.75 \quad {\rm void \; calOldQ0\_R\_TEOS \; (Grid \; \& \; \textit{grid}, \; \; Parameters \; \& \; \textit{parameters})}$

This function calculates the artificial viscosity of a cell. It calculates it using the old vaules of quantities and places the result in the old grid. It does this for 1D viscosity only.

#### Parameters:

- $\leftrightarrow grid$  supplies the input for calculating the pressure and also accepts the result of the pressure calculation
- ← parameters contains parameters used in calculating the artificial viscosity.

References Parameters::dA, Parameters::dAVThreshold, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nD, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGamma, Grid::nP, Grid::nQ0, Grid::nR, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, and Grid::nU.

Referenced by initInternalVars().

# 7.11.2.76 void calOldQ0Q1 RT GL (Grid & grid, Parameters & parameters)

This function calculates the artificial viscosity of a cell. It calculates it using the old vaules of quantities and places the result in the old grid. It does this for the two components of the viscosity. This function is used when using a gamma law gas equation of state.

### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the artificial viscosity and also accepts the result of the artificial viscosity calculation
- ← parameters contains parameters used in calculating the artificial viscosity.

References Parameters::dA, Parameters::dAVThreshold, Parameters::dGamma, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nD, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, and Grid::nV.

Referenced by initInternalVars().

# $7.11.2.77 \quad {\rm void \; calOldQ0Q1 \ \ \, RT \ \ \, TEOS \; (Grid \; \& \; \it{grid}, \; \; Parameters \; \& \; \it{parameters})}$

This function calculates the artificial viscosity of a cell. It calculates it using the old vaules of quantities and places the result in the old grid. It does this for two components of the viscosity. This function is used when using a tabulated equation of state.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the artificial viscosity and also accepts the result of the artificial viscosity calculation
- ← parameters contains parameters used in calculating the artificial viscosity.

Parameters::dA, Parameters::dAVThreshold, Grid::dLocalGridOld, References Grid::nCenIntOffset, Grid::nD, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit. Grid::nQ0, Grid::nSinThetaIJK, Grid::nGamma, Grid::nP, Grid::nQ1, Grid::nR, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, and Grid::nV.

Referenced by initInternalVars().

# 7.11.2.78 void calOldQ0Q1Q2 RTP GL (Grid & grid, Parameters & parameters)

This function calculates the artificial viscosity of a cell. It calculates it using the old vaules of quantities and places the result in the old grid. It does this for the three components of the viscosity. This function is used when using a gamma law gas equation of state.

#### Parameters:

- $\leftrightarrow grid$  supplies the input for calculating the artificial viscosity and also accepts the result of the artificial viscosity calculation
- ← parameters contains parameters used in calculating the artificial viscosity.

References Parameters::dA. Parameters::dAVThreshold. Parameters::dGamma. Grid::dLocalGridOld, Grid::nCenIntOffset, Grid∷nD, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid∷nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid∷nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nV, and Grid::nW.

Referenced by initInternalVars().

# 7.11.2.79 void calOldQ0Q1Q2\_RTP\_TEOS (Grid & grid, Parameters & parameters)

This function calculates the artificial viscosity of a cell. It calculates it using the old vaules of quantities and places the result in the old grid. It does this for the three components of the viscosity. This function is used when using a tabulated equation of state.

- $\leftrightarrow$  grid supplies the input for calculating the artificial viscosity and also accepts the result of the artificial viscosity calculation
- ← parameters contains parameters used in calculating the artificial viscosity.

References Parameters::dA, Parameters::dAVThreshold, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nD, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGamma, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nV, and Grid::nW.

Referenced by initInternalVars().

# 7.11.2.80 double dEOS\_GL (double dRho, double dE, Parameters parameters)

Calculates the pressure from the energy and density using a  $\gamma$ -law gas.

#### Parameters:

- $\leftarrow$  dRho the density of a cell
- $\leftarrow dE$  the energy of a cell
- $\leftarrow$  parameters contians various parameters, including  $\gamma$  needed to calculate the pressure.

#### Returns:

the pressure

This version of dEOS\_GL uses the same value of  $\gamma$  through out the model. The equation of state is given by  $\rho(\gamma - 1)E$ .

References Parameters::dGamma.

Referenced by calNewP GL(), and calOldP GL().

# 7.11.2.81 double dImplicitEnergyFunction\_None (Grid & grid, Parameters & parameters, Time & time, double dTemps[], int i, int j, int k)

This is an empty function, that isn't even called when no implicit solution is needed. This safe guards against future addition which may need to call an empty function when no implicit solve is being done.

Referenced by setMainFunctions().

# 7.11.2.82 double dImplicitEnergyFunction R (Grid & grid, Parameters & parameters, Time & time, double dTemps[], int i, int j, int k)

This function is used to determine the agreement of the updated values at n+1, with each other in the non-adiabatic energy equation. The \_R version of the function contains only the radial terms, and should be used for purely radial calculations. This function can also be used for calculating numerical deriviatives by varying the input temperatures.

- $\leftarrow grid$
- $\leftarrow$  parameters
- $\leftarrow time$
- $\leftarrow$  **dTemps** dTemps[0]=dT\_ijk\_np1 is the temperature at radial position (i, j, k) and time n+1,dTemps[1]=dT\_ip1jk\_np1 is the temperature at radial position (i+1, j, k) and time n+1, dTemps[2]=dT\_im1jk\_np1 is the temperature at radial position (i-1, j, k) and time n+1.

- $\leftarrow i$  is the radial index to evaluate the function at.
- $\leftarrow j$  is the theta index to evaluate the function at.
- $\leftarrow k$  is the phi index to evaluate the function at.

DEBUG EQUATIONS, References Time::dDeltat np1half, eos::dGetEnergy(), eos::dGetOpacity(), eos::dGetPressure(), Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dSigma, Parameters::eosTable, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nE, Grid::nGlobalGridPositionLocalGrid, Grid::nNumGhostCells, Grid::nQ0, Grid::nR, Grid::nT, Grid::nU, and Grid::nU0.

Referenced by setMainFunctions().

# 7.11.2.83 double dImplicitEnergyFunction\_R\_LES (Grid & grid, Parameters & parameters, Time & time, double dTemps[], int i, int j, int k)

This function is used to determine the agreement of the updated values at n+1, with each other in the non-adiabatic energy equation. The \_R version of the function contains only the radial terms, and should be used for purely radial calculations. This function can also be used for calculating numerical derivitatives by varying the input temperatures.

#### Parameters:

- $\leftarrow grid$
- $\leftarrow parameters$
- $\leftarrow time$
- $\leftarrow$  dTemps dTemps[0]=dT\_ijk\_np1 is the temperature at radial position (i,j,k) and time n+1,dTemps[1]=dT\_ip1jk\_np1 is the temperature at radial position (i+1,j,k) and time n+1, dTemps[2]=dT\_im1jk\_np1 is the temperature at radial position (i-1,j,k) and time n+1.
- $\leftarrow i$  is the radial index to evaluate the function at.
- $\leftarrow j$  is the theta index to evaluate the function at.
- $\leftarrow k$  is the phi index to evaluate the function at.

### Todo

this function should probably be turffed, the LES terms aren't needed in 1D. keeping it for now though.

References Time::dDeltat\_np1half, eos::dGetEnergy(), eos::dGetOpacity(), eos::dGetPressure(), Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dSigma, Parameters::eosTable, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nE, Grid::nEddyVisc, Grid::nGlobalGridPositionLocalGrid, Grid::nNumGhostCells, Grid::nQ0, Grid::nR, Grid::nT, Grid::nU, and Grid::nU0.

# 7.11.2.84 double dImplicitEnergyFunction\_R\_LES\_SB (Grid & grid, Parameters & parameters, Time & time, double $dTemps[\ ]$ , int i, int j, int k)

This function is used to determine the agreement of the updated values at n+1, with each other in the non-adiabatic energy equation. The \_R version of the function contains only the radial terms, and should be used for purely radial calculations. This function can also be used for

calculating numerical deriviatives by varying the input temperatures. This function differs from the version without the "\_SB" suffix (dImplicitEnergyFunction\_R)in that it is tailored to the surface boundary region.

#### Parameters:

- $\leftarrow qrid$
- $\leftarrow$  parameters
- $\leftarrow time$
- $\leftarrow$  dTemps dTemps[0]=dT\_ijk\_np1 is the temperature at radial position (i, j, k) and time n+1 and time n+1, dTemps[1]=dT\_im1jk\_np1 is the temperature at radial position (i-1, j, k) and time n+1.
- $\leftarrow i$  is the radial index to evaluate the function at.
- $\leftarrow j$  is the theta index to evaluate the function at.
- $\leftarrow k$  is the phi index to evaluate the function at.

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nE][i+1][j][k] in calculation of  $E_{i+1/2,j,k}$  setting it equal to value at i.

### **Boundary Conditions**

grid.dLocalGridOld[grid.nDM][i+1][0][0] and grid.dLocalGridOld[grid.nE][i+1][j][k] missing in the calculation of upwind gradient in dA1. Using the centered gradient instead.

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nT][i+1][0][0]

References Time::dDeltat\_np1half, eos::dGetEnergy(), eos::dGetOpacity(), eos::dGetPressure(), Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dSigma, Parameters::eosTable, Grid::nCenIntOffset, Grid::nD, Grid::nDM, Grid::nDonorCellFrac, Grid::nE, Grid::nEddyVisc, Grid::nQ0, Grid::nR, Grid::nT, Grid::nU, and Grid::nU0.

# 7.11.2.85 double dImplicitEnergyFunction R SB (Grid & grid, Parameters & parameters, Time & time, double dTemps[], int i, int j, int k)

This function is used to determine the agreement of the updated values at n+1, with each other in the non-adiabatic energy equation. The \_R version of the function contains only the radial terms, and should be used for purely radial calculations. This function can also be used for calculating numerical deriviatives by varying the input temperatures. This function differs from the version without the "\_SB" suffix (dImplicitEnergyFunction\_R)in that it is tailored to the surface boundary region.

- $\leftarrow grid$
- $\leftarrow$  parameters
- $\leftarrow time$
- $\leftarrow$  **dTemps** dTemps[0]=dT\_ijk\_np1 is the temperature at radial position (i, j, k) and time n+1 and time n+1, dTemps[1]=dT\_im1jk\_np1 is the temperature at radial position (i-1, j, k) and time n+1.

- $\leftarrow i$  is the radial index to evaluate the function at.
- $\leftarrow j$  is the theta index to evaluate the function at.
- $\leftarrow k$  is the phi index to evaluate the function at.

# **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nE][i+1][j][k] in calculation of  $E_{i+1/2,j,k}$  setting it equal to value at i.

### **Boundary Conditions**

A1 upwind set to zero as no material is flowing into the star

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nT][i+1][0][0]

References Time::dDeltat\_np1half, eos::dGetEnergy(), eos::dGetOpacity(), eos::dGetPressure(), Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dSigma, Parameters::eosTable, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nE, Grid::nGlobalGridPositionLocalGrid, Grid::nNumGhostCells, Grid::nQ0, Grid::nR, Grid::nT, Grid::nU, and Grid::nU0.

Referenced by setMainFunctions().

# 7.11.2.86 double dImplicitEnergyFunction\_RT (Grid & grid, Parameters & parameters, Time & time, double dTemps[], int i, int j, int k)

This function is used to determine the agreement of the updated values at n+1, with each other in the non-adiabatic energy equation. The \_RT version of the function contains only the radial and theta terms, and should be used for radial-theta calculations. This function can also be used for calculating numerical derivitatives by varying the input temperatures.

### Parameters:

- $\leftarrow grid$
- $\leftarrow$  parameters
- $\leftarrow time$
- ← dTemps dTemps[0]=dT\_ijk\_np1 is the temperature at radial position (i, j, k) and time n+1, dTemps[1]=dT\_ip1jk\_np1 is the temperature at radial position (i+1, j, k) and time n+1, dTemps[2]=dT\_im1jk\_np1 is the temperature at radial position (i-1, j, k) and time n+1, dTemps[3]=dT\_ijp1k\_np1 is the temperature at radial position (i, j+1, k) and time n+1, dTemps[4]=dT\_ijm1k\_np1 is the temperature at radial position (i, j-1, k) and time n+1.
- $\leftarrow i$  is the radial index to evaluate the function at.
- $\leftarrow j$  is the theta index to evaluate the function at.
- $\leftarrow k$  is the phi index to evaluate the function at.

References Time::dDeltat\_np1half, eos::dGetEnergy(), eos::dGetOpacity(), eos::dGetPressure(), Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dSigma, Parameters::eosTable, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDTheta, Grid::nE, Grid::nQ0, Grid::nQ1, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJF1halfK, Grid::nT, Grid::nU, Grid::nU0, and Grid::nV.

Referenced by setMainFunctions().

# 7.11.2.87 double dImplicitEnergyFunction RT\_LES (Grid & grid, Parameters & parameters, Time & time, double dTemps[], int i, int j, int k)

This function is used to determine the agreement of the updated values at n+1, with each other in the non-adiabatic energy equation. The \_RT version of the function contains only the radial and theta terms, and should be used for radial-theta calculations. This function can also be used for calculating numerical derivitatives by varying the input temperatures.

#### Parameters:

- $\leftarrow grid$
- $\leftarrow$  parameters
- $\leftarrow time$
- ← dTemps dTemps[0]=dT\_ijk\_np1 is the temperature at radial position (i, j, k) and time n+1, dTemps[1]=dT\_ip1jk\_np1 is the temperature at radial position (i+1, j, k) and time n+1, dTemps[2]=dT\_im1jk\_np1 is the temperature at radial position (i-1, j, k) and time n+1, dTemps[3]=dT\_ijp1k\_np1 is the temperature at radial position (i, j+1, k) and time n+1, dTemps[4]=dT\_ijm1k\_np1 is the temperature at radial position (i, j-1, k) and time n+1.
- $\leftarrow i$  is the radial index to evaluate the function at.
- $\leftarrow j$  is the theta index to evaluate the function at.
- $\leftarrow k$  is the phi index to evaluate the function at.

```
\begin{array}{lll} 5700 & CTEOS & if (grid.dLocalGridOld[grid.nM][nIInt][0][0]>=1.143732280336595e+33) \{ \\ dA1UpWindGrad=-1.796596699553508e-14; & dA1CenGrad=-1.796596699553508e-14; \\ 6100 & CTEOS & if (grid.dLocalGridOld[grid.nM][nIInt][0][0]>=1.143732392703405e+33) \{ \\ dA1UpWindGrad=-2.381754669392478e-14; & dA1CenGrad=-2.381754669392478e-14; \\ T6500 & CTEOS & if (grid.dLocalGridOld[grid.nM][nIInt][0][0]>=1.143732445236012e+33) \{ \\ dA1UpWindGrad=-3.795837002744412e-14; \\ dA1CenGrad=-3.795837002744412e-14; \} \\ \end{array}
```

References Time::dDeltat\_np1half, eos::dGetEnergy(), eos::dGetOpacity(), eos::dGetPressure(), Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dPrt, Parameters::dSigma, Parameters::eosTable, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDTheta, Grid::nE, Grid::nEddyVisc, Grid::nM, Grid::nQ0, Grid::nQ1, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nT, Grid::nU, Grid::nU0, and Grid::nV.

Referenced by setMainFunctions().

# 7.11.2.88 double dImplicitEnergyFunction\_RT\_LES\_SB (Grid & grid, Parameters & parameters, Time & time, double $dTemps[\ ]$ , int i, int j, int k)

This function is used to determine the agreement of the updated values at n+1, with each other in the non-adiabatic energy equation. The \_RT version of the function contains only the radial and theta terms, and should be used for radial-theta calculations. This function can also be used for calculating numerical deriviatives by varying the input temperatures. This function differs from the version without the "\_SB" suffix (dImplicitEnergyFunction\_RT)in that it is tailored to the surface boundary region.

#### Parameters:

 $\leftarrow grid$ 

- $\leftarrow parameters$
- $\leftarrow time$
- ← dTemps dTemps[0]=dT\_ijk\_np1 is the temperature at radial position (i, j, k) and time n+1, dTemps[1]=dT\_ip1jk\_np1 is the temperature at radial position (i+1, j, k) and time n+1, dTemps[2]=dT\_im1jk\_np1 is the temperature at radial position (i-1, j, k) and time n+1, dTemps[3]=dT\_ijp1k\_np1 is the temperature at radial position (i, j+1, k) and time n+1, dTemps[4]=dT\_ijm1k\_np1 is the temperature at radial position (i, j-1, k) and time n+1.
- $\leftarrow i$  is the radial index to evaluate the function at.
- $\leftarrow j$  is the theta index to evaluate the function at.
- $\leftarrow k$  is the phi index to evaluate the function at.

# **Boundary Conditions**

Missing  $\Delta M_r$  outside model using Parameters::dAlpha times  $\Delta M_r$  in the last zone instead.

# **Boundary Conditions**

missing density outside model assuming it is zero

# **Boundary Conditions**

missing desnity outside model assuming it is zero

### **Boundary Conditions**

assuming V at ip1half is the same as V at i

# **Boundary Conditions**

Assuming energy outside model is the same as the energy in the last zone inside the model.

### **Boundary Conditions**

Assuming energy outside model is the same as the energy in the last zone inside the model.

### **Boundary Conditions**

Using centered gradient for upwind gradient when motion is into the star at the surface

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nT][i+1][0][0] using flux equals  $2\sigma T^4$  at surface.

References Parameters::dAlpha, Parameters::dAlphaExtra, Time::dDeltat\_np1half, eos::dGetEnergy(), eos::dGetOpacity(), eos::dGetPressure(), Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dPrt, Parameters::dSigma, Parameters::eosTable, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDTheta, Grid::nE, Grid::nEddyVisc, Grid::nQ0, Grid::nQ1, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJF1halfK, Grid::nT, Grid::nU, Grid::nU0, and Grid::nV.

Referenced by setMainFunctions().

# 7.11.2.89 double dImplicitEnergyFunction\_RT\_SB (Grid & grid, Parameters & parameters, Time & time, double dTemps[], int i, int j, int k)

This function is used to determine the agreement of the updated values at n+1, with each other in the non-adiabatic energy equation. The \_RT version of the function contains only the radial and theta terms, and should be used for radial-theta calculations. This function can also be used for calculating numerical deriviatives by varying the input temperatures. This function differs from the version without the "\_SB" suffix (dImplicitEnergyFunction\_RT)in that it is tailored to the surface boundary region.

#### Parameters:

- $\leftarrow grid$
- $\leftarrow$  parameters
- $\leftarrow time$
- ← dTemps dTemps[0]=dT\_ijk\_np1 is the temperature at radial position (i, j, k) and time n+1, dTemps[1]=dT\_ip1jk\_np1 is the temperature at radial position (i+1, j, k) and time n+1, dTemps[2]=dT\_im1jk\_np1 is the temperature at radial position (i-1, j, k) and time n+1, dTemps[3]=dT\_ijp1k\_np1 is the temperature at radial position (i, j+1, k) and time n+1, dTemps[4]=dT\_ijm1k\_np1 is the temperature at radial position (i, j-1, k) and time n+1.
- $\leftarrow i$  is the radial index to evaluate the function at.
- $\leftarrow j$  is the theta index to evaluate the function at.
- $\leftarrow k$  is the phi index to evaluate the function at.

### **Boundary Conditions**

Using centered gradient for upwind gradient when motion is into the star at the surface

# **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nT][i+1][0][0] using flux equals  $2\sigma T^4$  at surface.

References Time::dDeltat\_np1half, eos::dGetEnergy(), eos::dGetOpacity(), eos::dGetPressure(), Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dSigma, Parameters::eosTable, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDTheta, Grid::nE, Grid::nQ0, Grid::nQ1, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJF1halfK, Grid::nT, Grid::nU, Grid::nU0, and Grid::nV.

Referenced by setMainFunctions().

# 7.11.2.90 double dImplicitEnergyFunction\_RTP (Grid & grid, Parameters & parameters, Time & time, double $dTemps[\ ]$ , int i, int j, int k)

This function is used to determine the agreement of the updated values at n+1, with each other in the non-adiabatic energy equation. The \_RTP version of the function contains terms for all three directions, and should be used for calculations involving all three directions. This function can also be used for calculating numerical deriviatives by varying the input temperatures. This function differs from the version without the "\_SB" suffix (dImplicitEnergyFunction\_RT)in that it is tailored to the surface boundary region.

#### Parameters:

 $\leftarrow grid$ 

- $\leftarrow parameters$
- $\leftarrow time$
- ←  $dTemps, dTemps[0] = dT_ijk_np1$  is the temperature at radial position (i, j, k) and time n+1,  $dTemps[1] = dT_ip1jk_np1$  is the temperature at radial position (i+1, j, k) and time n+1,  $dTemps[2] = dT_im1jk_np1$  is the temperature at radial position (i-1, j, k) and time n+1,  $dTemps[3] = dT_ip1k_np1$  is the temperature at radial position (i, j+1, k) and time n+1,  $dTemps[4] = dT_ip1k_np1$  is the temperature at radial position (i, j-1, k) and time n+1,  $dTemps[5] = dT_ipkp1_np1$  is the temperature at radial position (i, j, k+1) and time n+1,  $dTemps[6] = dT_ipkp1_np1$  is the temperature at radial position (i, j, k-1) and time n+1.
- $\leftarrow i$  is the radial index to evaluate the function at.
- $\leftarrow j$  is the theta index to evaluate the function at.
- $\leftarrow k$  is the phi index to evaluate the function at.

References Time::dDeltat\_np1half, eos::dGetEnergy(), eos::dGetOpacity(), eos::dGetPressure(), Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dSigma, Parameters::eosTable, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nE, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nT, Grid::nU, Grid::nU, Grid::nV, and Grid::nW.

Referenced by setMainFunctions().

# 7.11.2.91 double dImplicitEnergyFunction\_RTP\_LES (Grid & grid, Parameters & parameters, Time & time, double dTemps[], int i, int j, int k)

This function is used to determine the agreement of the updated values at n+1, with each other in the non-adiabatic energy equation. The \_RTP version of the function contains terms for all three directions, and should be used for calculations involving all three directions. This function can also be used for calculating numerical deriviatives by varying the input temperatures. This function differs from the version without the "\_SB" suffix (dImplicitEnergyFunction\_RT)in that it is tailored to the surface boundary region.

- $\leftarrow grid$
- $\leftarrow parameters$
- $\leftarrow time$
- ←  $dTemps, dTemps[0] = dT_ijk_np1$  is the temperature at radial position (i, j, k) and time n+1,  $dTemps[1] = dT_ip1jk_np1$  is the temperature at radial position (i+1, j, k) and time n+1,  $dTemps[2] = dT_im1jk_np1$  is the temperature at radial position (i-1, j, k) and time n+1,  $dTemps[3] = dT_ip1k_np1$  is the temperature at radial position (i, j+1, k) and time n+1,  $dTemps[4] = dT_ip1k_np1$  is the temperature at radial position (i, j-1, k) and time n+1,  $dTemps[5] = dT_ipkp1_np1$  is the temperature at radial position (i, j, k+1) and time n+1,  $dTemps[6] = dT_ipkp1_np1$  is the temperature at radial position (i, j, k-1) and time n+1.
- $\leftarrow i$  is the radial index to evaluate the function at.
- $\leftarrow j$  is the theta index to evaluate the function at.
- $\leftarrow k$  is the phi index to evaluate the function at.

References Time::dDeltat np1half, DEBUG EQUATIONS, eos::dGetEnergy(), eos::dGetOpacity(), eos::dGetPressure(), Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dPrt, Parameters::dSigma, Parameters::eosTable, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nE, Grid::nEddyVisc, Grid::nGlobalGridPositionLocalGrid, Grid::nM, Grid::nNumGhostCells, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nT, Grid::nU, Grid::nU0, Grid::nV, and Grid::nW.

Referenced by setMainFunctions().

# 7.11.2.92 double dImplicitEnergyFunction\_RTP\_LES\_SB (Grid & grid, Parameters & parameters, Time & time, double $dTemps[\ ]$ , int i, int j, int k)

This function is used to determine the agreement of the updated values at n+1, with each other in the non-adiabatic energy equation. The \_RTP version of the function contains terms for all three directions, and should be used for calculations involving all three directions. This function can also be used for calculating numerical deriviatives by varying the input temperatures. This function differs from the version without the "\_SB" suffix (dImplicitEnergyFunction\_RT)in that it is tailored to the surface boundary region.

### Parameters:

- $\leftarrow grid$
- $\leftarrow$  parameters
- $\leftarrow time$
- ← dTemps dTemps[0]=dT\_ijk\_np1 is the temperature at radial position (i, j, k) and time n+1, dTemps[1]=dT\_ip1jk\_np1 is the temperature at radial position (i+1, j, k) and time n+1, dTemps[2]=dT\_im1jk\_np1 is the temperature at radial position (i-1, j, k) and time n+1, dTemps[3]=dT\_ijp1k\_np1 is the temperature at radial position (i, j+1, k) and time n+1, dTemps[4]=dT\_ijm1k\_np1 is the temperature at radial position (i, j-1, k) and time n+1, dTemps[5]=dT\_ijkp1\_np1 is the temperature at radial position (i, j, k+1) and time n+1, dTemps[6]=dT\_ijkm1\_np1 is the temperature at radial position (i, j, k-1) and time n+1.
- $\leftarrow i$  is the radial index to evaluate the function at.
- $\leftarrow j$  is the theta index to evaluate the function at.
- $\leftarrow k$  is the phi index to evaluate the function at.

### **Boundary Conditions**

Missing  $\Delta M_r$  outside model using Parameters::dAlpha times  $\Delta M_r$  in the last zone instead.

# **Boundary Conditions**

missing density outside model assuming it is zero

### **Boundary Conditions**

missing desnity outside model assuming it is zero

### **Boundary Conditions**

assuming V at ip1half is the same as V at i

### **Boundary Conditions**

assuming W at ip1half is the same as W at i

### **Boundary Conditions**

Assuming energy outside model is the same as the energy in the last zone inside the model.

### **Boundary Conditions**

Assuming energy outside model is the same as the energy in the last zone inside the model.

# **Boundary Conditions**

A1 upwind set to zero as no material is flowing into the star

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nT][i+1][0][0] using flux equals  $2\sigma T^4$  at surface.

References Parameters::dAlpha, Parameters::dAlphaExtra, Time::dDeltat np1half, DEBUG EQUATIONS, eos::dGetEnergy(), eos::dGetOpacity(), eos::dGetPressure(), Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dPrt, Grid::nDenAve, rameters::dSigma, Parameters::eosTable, Grid::nCenIntOffset, Grid::nD, Grid::nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nE, Grid::nEddyVisc, Grid::nGlobalGridPositionLocalGrid, Grid::nM, Grid::nNumGhostCells, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nT, Grid::nU, Grid::nU0, Grid::nV, and Grid::nW.

Referenced by setMainFunctions().

# 7.11.2.93 double dImplicitEnergyFunction\_RTP\_SB (Grid & grid, Parameters & parameters, Time & time, double dTemps[], int i, int j, int k)

This function is used to determine the agreement of the updated values at n+1, with each other in the non-adiabatic energy equation. The \_RTP version of the function contains terms for all three directions, and should be used for calculations involving all three directions. This function can also be used for calculating numerical deriviatives by varying the input temperatures. This function differs from the version without the "\_SB" suffix (dImplicitEnergyFunction\_RT)in that it is tailored to the surface boundary region.

- $\leftarrow grid$
- $\leftarrow$  parameters
- $\leftarrow time$
- ← dTemps dTemps[0]=dT\_ijk\_np1 is the temperature at radial position (i, j, k) and time n+1, dTemps[1]=dT\_ip1jk\_np1 is the temperature at radial position (i+1, j, k) and time n+1, dTemps[2]=dT\_im1jk\_np1 is the temperature at radial position (i-1, j, k) and time n+1, dTemps[3]=dT\_ijp1k\_np1 is the temperature at radial position (i, j+1, k) and time n+1, dTemps[4]=dT\_ijm1k\_np1 is the temperature at radial position (i, j-1, k) and time n+1, dTemps[5]=dT\_ijkp1\_np1 is the temperature at radial position (i, j, k+1) and time n+1, dTemps[6]=dT\_ijkm1\_np1 is the temperature at radial position (i, j, k-1) and time n+1.

- $\leftarrow i$  is the radial index to evaluate the function at.
- $\leftarrow j$  is the theta index to evaluate the function at.
- $\leftarrow k$  is the phi index to evaluate the function at.

# **Boundary Conditions**

Using 
$$E_{i,j,k}^{n+1/2}$$
 for  $E_{i+1/2,j,k}^{n+1/2}$ 

### **Boundary Conditions**

Using centered gradient for upwind gradient when motion is into the star at the surface

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nT][i+1][0][0] using flux equals  $2\sigma T^4$  at surface.

References Time::dDeltat\_np1half, eos::dGetEnergy(), eos::dGetOpacity(), eos::dGetPressure(), Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dSigma, Parameters::eosTable, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nE, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nT, Grid::nU, Grid::nU, Grid::nV, and Grid::nW.

Referenced by setMainFunctions().

# 7.11.2.94 void implicitSolve\_None (Grid & grid, Implicit & implicit, Parameters & parameters, Time & time, ProcTop & procTop, MessPass & messPass, Functions & functions)

This is an empty function, to be called when no implicit solution is needed. This allows the same code in the main program to be executed wheather or not an implicit solution is being preformed by setting the function pointer to this function if there is no implicit solution required.

Referenced by setMainFunctions().

# 7.11.2.95 void implicitSolve\_R (Grid & grid, Implicit & implicit, Parameters & parameters, Time & time, ProcTop & procTop, MessPass & messPass, Functions & functions)

This function solves for temperature corrections based on derivatives of the radial non-adiabatic energy equation with respect to the new temperature. It then uses these derivatives as entries in the coeffecient matrix. The discrepancy in the balance of the energy equation with the new temperature, energy, pressure, and opacity are included as the right hand side of the system of equaitons. Solving this system of equaitons provides the corrections needed for the new temperature. This processes is then repeated until the corrections are small. At this point the new temperature is used to update the energy, pressure, and opacity in the new grid via the equaiton of state.

calNewPEKappaGamma TEOS(), References Implicit::dAverageRHS, Implicit::dCurrentRelTError, Implicit::dDerivativeStepFraction, Grid::dLocalGridNew, plicit::dMaxErrorInRHS, Implicit::dTolerance, Functions::fpImplicitEnergyFunction, Functions::fpImplicitEnergyFunction SB, Implicit::kspContext, Implicit::matCoeff, Implicit::nCurrentNumIterations, Implicit::nLocDer, Implicit::nLocFun, plicit::nMaxNumIterations, Implicit::nMaxNumSolverIterations, Implicit::nNumDerPerRow,

 $\label{lem:limbular} Implicit:::nNumRowsALocalSB, ProcTop::nRank, Grid::nT, Time::nTimeStepIndex, Implicit::nTypeDer, updateLocalBoundariesNewGrid(), Implicit::vecRHS, Implicit::vecscatTCorrections, Implicit::vecTCorrections, and Implicit::vecTCorrectionsLocal.$ 

Referenced by setMainFunctions().

# 7.11.2.96 void implicitSolve\_RT (Grid & grid, Implicit & implicit, Parameters & parameters, Time & time, ProcTop & procTop, MessPass & messPass, Functions & functions)

This function solves for temperature corrections based on derivatives of the radial-theta non-adiabatic energy equation with respect to the new temperature. It then uses these derivatives as entries in the coeffecient matrix. The discrepancy in the balance of the energy equation with the new temperature, energy, pressure, and opacity are included as the right hand side of the system of equaitons. Solving this system of equaitons provides the corrections needed for the new temperature. This processes is then repeated until the corrections are small. At this point the new temperature is used to update the energy, pressure, and opacity in the new grid via the equaiton of state.

calNewPEKappaGamma TEOS(), References Implicit::dAverageRHS, Implicit::dCurrentRelTError, Implicit::dDerivativeStepFraction, Grid::dLocalGridNew, Implicit::dMaxErrorInRHS, Implicit::dTolerance, Functions::fpImplicitEnergyFunction, Functions::fpImplicitEnergyFunction SB, Implicit::kspContext, Implicit::matCoeff, Implicit::nCurrentNumIterations, Implicit::nLocDer, Implicit::nLocFun, plicit::nMaxNumIterations, Implicit::nMaxNumSolverIterations, Implicit::nNumDerPerRow, Implicit::nNumRowsALocal, Implicit::nNumRowsALocalSB, ProcTop::nRank, Grid::nT, Time::nTimeStepIndex, Implicit::nTypeDer, updateLocalBoundariesNewGrid(), Implicit::vecRHS. Implicit::vecscatTCorrections, Implicit::vecTCorrections, Implicit::vecTCorrectionsLocal.

Referenced by setMainFunctions().

# 7.11.2.97 void implicitSolve\_RTP (Grid & grid, Implicit & implicit, Parameters & parameters, Time & time, ProcTop & procTop, MessPass & messPass, Functions & functions)

This function solves for temperature corrections based on derivatives of the radial-theta-phi non-adiabatic energy equation with respect to the new temperature. It then uses these derivatives as entries in the coeffecient matrix. The discrepancy in the balance of the energy equation with the new temperature, energy, pressure, and opacity are included as the right hand side of the system of equaitons. Solving this system of equaitons provides the corrections needed for the new temperature. This processes is then repeated until the corrections are small. At this point the new temperature is used to update the energy, pressure, and opacity in the new grid via the equaiton of state.

References calNewPEKappaGamma TEOS(), Implicit::dAverageRHS, Im-Implicit::dDerivativeStepFraction, Grid::dLocalGridNew, plicit::dCurrentRelTError, Implicit::dMaxErrorInRHS, Implicit::dTolerance, Functions::fpImplicitEnergyFunction, Functions::fpImplicitEnergyFunction SB, Implicit::kspContext, Implicit::matCoeff, Implicit::nCurrentNumIterations, Implicit::nLocDer, Implicit::nLocFun, Implicit::nMaxNumIterations, Implicit::nMaxNumSolverIterations, Implicit::nNumDerPerRow, Implicit::nNumRowsALocalSB, Implicit::nNumRowsALocal, ProcTop::nRank, Grid∷nT, Time::nTimeStepIndex, Implicit::nTypeDer, updateLocalBoundariesNewGrid(), Im $\label{limit} \begin{array}{lll} \textbf{plicit::} \textbf{vecRHS}, & \textbf{Implicit::} \textbf{vecTCorrections}, & \textbf{Implicit::} \textbf{vecTCorrections}, & \textbf{and} & \textbf{Implicit::} \textbf{vecTCorrectionsLocal}. \\ \end{array}$ 

Referenced by setMainFunctions().

# 7.11.2.98 void initDonorFracAndMaxConVel\_R\_GL (Grid & grid, Parameters & parameters)

Initializes the donor fraction, and the maximum convective velocity when starting a calculation. The donor fraction is used to determine the amount of upwinded donor cell to use in advection terms. The maximum convective velocity is used for calculation of constant eddy viscosity parameter. This version of the fuction is for 1D, gamma law calculations.

References Parameters::dDonorCellMin, Parameters::dDonorCellMultiplier, Parameters::dGamma, Grid::dLocalGridOld, Parameters::dMaxConvectiveVelocity, Grid::nCenIntOffset, Grid::nD, Grid::nDonorCellFrac, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0, Grid::nStartUpdateExplicit, Grid::nU, and Grid::nU0.

Referenced by initInternalVars().

# 7.11.2.99 void initDonorFracAndMaxConVel\_R\_TEOS (Grid & grid, Parameters & parameters)

Initializes the donor fraction, and the maximum convective velocity when starting a calculation. The donor fraction is used to determine the amount of upwinded donor cell to use in advection terms. The maximum convective velocity is used for calculation of constant eddy viscosity parameter. This version of the fuction is for 1D, tabulated equation of state calculations.

References Parameters::dDonorCellMin, Parameters::dDonorCellMultiplier, Grid::dLocalGridOld, Parameters::dMaxConvectiveVelocity, Grid::nCenIntOffset, Grid::nD, Grid::nDonorCellFrac, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGamma, Grid::nP, Grid::nQ0, Grid::nStartUpdateExplicit, Grid::nU, and Grid::nU0.

Referenced by initInternalVars().

# $7.11.2.100 \quad \text{void initDonorFracAndMaxConVel\_RT\_GL (Grid \& \textit{grid}, \text{ Parameters})} \\ \& \textit{parameters})$

Initializes the donor fraction, and the maximum convective velocity when starting a calculation. The donor fraction is used to determine the amount of upwinded donor cell to use in advection terms. The maximum convective velocity is used for calculation of constant eddy viscosity parameter. This version of the fuction is for 2D, gamma law calculations.

References Parameters::dDonorCellMin, Parameters::dDonorCellMultiplier, Parameters::dGamma, Grid::dLocalGridOld, Parameters::dMaxConvectiveVelocity, Grid::nCenIntOffset, Grid::nD, Grid::nDonorCellFrac, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0, and Grid::nV.

Referenced by initInternalVars().

# 7.11.2.101 void initDonorFracAndMaxConVel\_RT\_TEOS (Grid & grid, Parameters & parameters)

Initializes the donor fraction, and the maximum convective velocity when starting a calculation. The donor fraction is used to determine the amount of upwinded donor cell to use in advec-

tion terms. The maximum convective velocity is used for calculation of constant eddy viscosity parameter. This version of the fuction is for 2D, tabulated equation of state calculations.

References Parameters::dDonorCellMin, Parameters::dDonorCellMultiplier, Grid::dLocalGridOld, Parameters::dMaxConvectiveVelocity, Grid::nCenIntOffset, Grid::nD, Grid::nDonorCellFrac, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGamma, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0, and Grid::nV.

Referenced by initInternalVars().

# 7.11.2.102 void initDonorFracAndMaxConVel\_RTP\_GL (Grid & grid, Parameters & parameters)

Initializes the donor fraction, and the maximum convective velocity when starting a calculation. The donor fraction is used to determine the amount of upwinded donor cell to use in advection terms. The maximum convective velocity is used for calculation of constant eddy viscosity parameter. This version of the fuction is for 3D, gamma law calculations.

References Parameters::dDonorCellMin, Parameters::dDonorCellMultiplier, Parameters::dGamma, Grid::dLocalGridOld, Parameters::dMaxConvectiveVelocity, Grid::nCenIntOffset, Grid::nD, Grid::nDonorCellFrac, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0, Grid::nV, and Grid::nW.

Referenced by initInternalVars().

# 7.11.2.103 void initDonorFracAndMaxConVel\_RTP\_TEOS (Grid & grid, Parameters & parameters)

Initializes the donor fraction, and the maximum convective velocity when starting a calculation. The donor fraction is used to determine the amount of upwinded donor cell to use in advection terms. The maximum convective velocity is used for calculation of constant eddy viscosity parameter. This version of the fuction is for 3D, tabulated equation of state calculations.

References Parameters::dDonorCellMin, Parameters::dDonorCellMultiplier, Grid::dLocalGridOld, Parameters::dMaxConvectiveVelocity, Grid::nCenIntOffset, Grid::nD, Grid::nDonorCellFrac, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGamma, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0, Grid::nV, and Grid::nW.

Referenced by initInternalVars().

# 7.11.2.104 void initInternalVars (Grid & grid, ProcTop & procTop, Parameters & parameters)

This function function is used to set the initial values of the internal variables. While external variables are initialized from the starting model, internal variables are calculated at startup.

- $\leftrightarrow$  grid supplies information needed for initilizing internal variables as well as storing the initilized internal variables
- $\leftarrow procTop$  contians information about processor topology
- $\leftarrow$  parameters contains parameters used in initializing the internal variables.

### Warning:

 $\Delta\theta$ ,  $\Delta\phi$ ,  $\sin\theta_{i,j,k}$ ,  $\Delta\cos\theta_{i,j,k}$ , all don't have the first zone calculated. At the moment this is a ghost cell that doesn't matter, but it may become a problem if calculations require this quantity. This is an issue for quantities that aren't updated in time, as those that are will have boundary cells updated with periodic boundary conditions.

References Parameters::bEOSGammaLaw, calOldDenave R(), calOldDenave RT(), calOldDenave RTP(), calOldEddyVisc R CN(), calOldEddyVisc R SM(), calOldEddyVisc -RT CN(), calOldEddyVisc RT SM(), calOldEddyVisc RTP CN(), calOldEddyVisc RTP -SM(), calOldP GL(), calOldPEKappaGamma TEOS(), calOldQ0 R GL(), calOldQ0 R -TEOS(), calOldQ0Q1\_RT\_GL(), calOldQ0Q1\_RT\_TEOS(), calOldQ0Q1Q2\_RTP\_GL(), calOldQ0Q1Q2 RTP TEOS(), initDonorFracAndMaxConVel R -Grid::dLocalGridOld, GL(), initDonorFracAndMaxConVel R TEOS(), initDonorFracAndMaxConVel RT GL(), initDonorFracAndMaxConVel RT TEOS(), initDonorFracAndMaxConVel RTP GL(), initDonorFracAndMaxConVel RTP TEOS(), Grid::nCotThetaIJK, Grid::nCenIntOffset, Grid::nCotThetaIJp1halfK, Grid::nDCosThetaIJK, Grid::nDTheta, Grid∷nDPhi, Grid::nLocalGridDims, Grid::nNumDims, Grid::nNumGhostCells, Grid::nPhi, Proc-Top::nRank, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nTheta, and Parameters::nTypeTurbulanceMod.

Referenced by init().

# 7.11.2.105 void setInternalVarInf (Grid & grid, Parameters & parameters)

This function sets the information for internal variables. While external verabile information is derived from the starting model, internal variables infos are set in this function. In other words this function sets the values of Grid::nVariables.

#### Parameters:

- $\leftrightarrow$  grid supplies the information needed to calculate the horizontal density average, it also stores the calculated horizontally averaged density.
- ← *parameters* is used when setting variable infos, since one needs to know if the code is calculating using a gamma law gas, or a tabulated equation of state.

References Parameters::bEOSGammaLaw, Grid::nCotThetaIJK, Grid::nCotThetaIJp1halfK, Grid::nDCosThetaIJK, Grid::nDenAve, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid∷nE, Grid::nEddyVisc, Grid::nGamma, Grid::nKappa, Grid::nNumDims, Grid::nNumIntVars, Grid::nP, Grid::nQ0, Grid::nNumVars, Grid::nQ1, Grid::nQ2, Parameters::nTypeTurbulanceMod, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid∷nVariables.

Referenced by modelRead().

# 7.11.2.106 void setMainFunctions (Functions & functions, ProcTop & procTop, Parameters & parameters, Grid & grid, Time & time, Implicit & implicit)

Used to set the functions that main() uses to evolve the input model.

#### Parameters:

 $\rightarrow$  functions is of class Functions and is used to specify the functions called to calculate the evolution of the input model.

← *procTop* is of type ProcTop. ProcTop::nRank is used to set different functions based on processor rank. For instance processor rank 1 requires 1D versions of the equations.

- ← parameters is of class Parameters. It holds various constants and runtime parameters.
- ← *grid* of type Grid. This function requires the number of dimensions, specified by Grid::nNumDims.
- ← time of type Time. This function requires knowledge of the type of time setp being used, specified by Time::bVariableTimeStep.
- ← *implicit* of type Implicit. This function needs to know if there is an implicit region, specified when Implicit::nNumImplicitZones>0.

The functions are picked based on model geometry, and the physics requested or required by the input model, and the configuration file. The specific functions pointers that are set are described in the Functions class.

References Parameters::bAdiabatic, Parameters::bEOSGammaLaw, Time::bVariableTimeStep,  ${\tt calDelt\_CONST()}, \quad {\tt calDelt\_R\_GL()}, \quad {\tt calDelt\_R\_TEOS()}, \quad {\tt calDelt\_RT\_GL()}, \quad {\tt calDelt\_R\_CONST()}, \quad {\tt calDelt\_R\_GL()}, \quad {\tt calDelt\_R\_CONST()}, \quad {\tt calDelt\_R\_GL()}, \quad {\tt calDelt\_R\_CONST()}, \quad {\tt calDe$  $RT\_TEOS()$ ,  $calDelt\_RTP\_GL()$ ,  $calDelt\_RTP\_TEOS()$ , calNewD R(), calNewD RT(), calNewD RTP(), calNewDenave None(), calNewDenave R(),calNewDenave -RT(), calNewDenave RTP(), calNewE R AD(), calNewE R NA(), RT AD(),calNewE RT NA(), calNewE RT NA LES(), calNewE RTP  $calNewE_RTP \overline{NA} \overline{LES()},$ calNewE RTP NA(), AD(), calNewEddyVisc None(), calNewEddyVisc RT CN(), calNewEddyVisc RT SM(), calNewEddyVisc RTP CN(), calNewP GL(), calNewEddyVisc RTP SM(), calNewQ0 R GL(),calNewQ0 R calNewQ0Q1 RT GL(), calNewQ0Q1 RT TEOS(), calNewQ0Q1Q2 RTP -TEOS(), calNewTPKappaGamma TEOS(),  $calNewQ0Q1Q2\_RTP\_TEOS()$ , calNewR(), calNewU0 R(), calNewU0 RT(), calNewU0 RTP(), calNewVelocities R(), calNewVelocities calNewVelocities RT LES(), calNewVelocities RTP(), RT(),calNewVelocities -RTP LES(), dImplicitEnergyFunction None(), dImplicitEnergyFunction R(), dImplicitEnergyFunction R SB(), dImplicitEnergyFunction RT(), dImplicitEnergyFunction dImplicitEnergyFunction RT LES SB(), RT LES(),dImplicitEnergyFunction -RT SB(), dImplicitEnergyFunction RTP(), dImplicitEnergyFunction RTP LES(), dImplicitEnergyFunction RTP SB(), dImplicitEnergyFunction RTP LES SB(), tions::fpCalculateAveDensities, Functions::fpCalculateDeltat, Functions::fpCalculateNewAV, Functions::fpCalculateNewDensities, Functions::fpCalculateNewEddyVisc, Functions::fpCalculateNewEnergies, Functions::fpCalculateNewEOSVars, Functions::fpCalculateNewGridVelocities, Functions::fpCalculateNewRadii, Functions::fp Calculate New Velocities,Functions::fpImplicitEnergyFunction, Functions::fpImplicitEnergyFunction SB, Functions::fpImplicitSolve, Functions::fpModelWrite, Functions::fpUpdateLocalBoundaryVelocitiesNewGrid, Functions::fpWriteWatchZones, implicitSolve None(), implicitSolve R(),implicitSolve RT(), implicitSolve RTP(), modelWrite GL(), modelWrite TEOS(), Grid::nNumDims, Implicit::nNumImplicitZones, Proc-Top::nRank, Parameters::nTypeTurbulanceMod, updateLocalBoundaryVelocitiesNewGrid R(), updateLocalBoundaryVelocitiesNewGrid RT(), updateLocalBoundaryVelocitiesNewGrid - $RTP(), \ writeWatchZones \ R \ GL(), \ writeWatchZones \_R \_TEOS(), \ writeWatchZones \_RT \_-temperature - temperature - tempera$ GL(), writeWatchZones RT TEOS(), writeWatchZones RTP GL(), and writeWatchZones - $RTP\_TEOS()$ .

Referenced by main().

# 7.12 physEquations.h File Reference

#include "global.h"

# **Functions**

- void setMainFunctions (Functions &functions, ProcTop &procTop, Parameters &parameters, Grid &grid, Time &time, Implicit &implicit)
- void setInternalVarInf (Grid &grid, Parameters &parameters)
- void initInternalVars (Grid &grid, ProcTop &procTop, Parameters &parameters)
- void calNewVelocities\_R (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewVelocities\_R\_LES (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewVelocities\_RT (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewVelocities\_RT\_LES (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewVelocities\_RTP (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewVelocities\_RTP\_LES (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewU R (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewU\_R\_LES (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewU RT (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewU\_RT\_LES (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewU\_RTP (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewU\_RTP\_LES (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewV\_RT (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewV\_RT\_LES (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewV\_RTP (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewV\_RTP\_LES (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewW\_RTP (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewW\_RTP\_LES (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewU0\_R (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop, MessPass &messPass)
- void calNewU0\_RT (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop, MessPass &messPass)

• void calNewU0\_RTP (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop, MessPass &messPass)

- void calNewR (Grid &grid, Time &time)
- void calNewD R (Grid &grid, Parameters &parameters, Time &time, ProcTop)
- void calNewD\_RT (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewD\_RTP (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewE\_R\_AD (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewE\_R\_NA (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewE\_R\_NA\_LES (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewE\_RT\_AD (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewE\_RT\_NA (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewE\_RT\_NA\_LES (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewE\_RTP\_AD (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewE\_RTP\_NA (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewE\_RTP\_NA\_LES (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calNewDenave\_None (Grid &grid)
- void calNewDenave R (Grid &grid)
- void calNewDenave RT (Grid &grid)
- void calNewDenave RTP (Grid &grid)
- void calNewP\_GL (Grid &grid, Parameters &parameters)
- void calNewTPKappaGamma TEOS (Grid &grid, Parameters &parameters)
- void calNewPEKappaGamma TEOS (Grid &grid, Parameters &parameters)
- void calNewQ0 R GL (Grid &grid, Parameters &parameters)
- void calNewQ0 R TEOS (Grid &grid, Parameters &parameters)
- void calNewQ0Q1 RT GL (Grid &grid, Parameters &parameters)
- void calNewQ0Q1 RT TEOS (Grid &grid, Parameters &parameters)
- void calNewQ0Q1Q2\_RTP\_GL (Grid &grid, Parameters &parameters)
- void calNewQ0Q1Q2 RTP TEOS (Grid &grid, Parameters &parameters)
- void calNewEddyVisc None (Grid &grid, Parameters &parameters)
- void calNewEddyVisc R CN (Grid &grid, Parameters &parameters)
- void calNewEddyVisc RT CN (Grid &grid, Parameters &parameters)
- void calNewEddyVisc RTP CN (Grid &grid, Parameters &parameters)
- void calNewEddyVisc R SM (Grid &grid, Parameters &parameters)
- void calNewEddyVisc RT SM (Grid &grid, Parameters &parameters)
- void calNewEddyVisc RTP SM (Grid &grid, Parameters &parameters)
- void calOldDenave None (Grid &grid)
- void calOldDenave R (Grid &grid)
- void calOldDenave RT (Grid &grid)
- void calOldDenave RTP (Grid &grid)
- void calOldP GL (Grid &grid, Parameters &parameters)
- void calOldPEKappaGamma TEOS (Grid &grid, Parameters &parameters)

- void calOldQ0 R GL (Grid &grid, Parameters &parameters)
- void calOldQ0 R TEOS (Grid &grid, Parameters &parameters)
- void calOldQ0Q1 RT GL (Grid &grid, Parameters &parameters)
- void calOldQ0Q1 RT TEOS (Grid &grid, Parameters &parameters)
- $\bullet\,$ void cal<code>OldQ0Q1Q2\_RTP\_GL</code> (Grid &grid, Parameters &parameters)
- void calOldQ0Q1Q2\_RTP\_TEOS (Grid &grid, Parameters &parameters)
- void calOldEddyVisc R CN (Grid &grid, Parameters &parameters)
- void calOldEddyVisc\_RT\_CN (Grid &grid, Parameters &parameters)
- void calOldEddyVisc RTP CN (Grid &grid, Parameters &parameters)
- void calOldEddyVisc R SM (Grid &grid, Parameters &parameters)
- void calOldEddyVisc RT SM (Grid &grid, Parameters &parameters)
- void calOldEddyVisc RTP SM (Grid &grid, Parameters &parameters)
- void calDelt\_R\_GL (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calDelt\_R\_TEOS (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calDelt\_RT\_GL (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calDelt\_RT\_TEOS (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calDelt\_RTP\_GL (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calDelt\_RTP\_TEOS (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void calDelt\_CONST (Grid &grid, Parameters &parameters, Time &time, ProcTop &procTop)
- void implicitSolve\_None (Grid &grid, Implicit &implicit, Parameters &parameters, Time &time, ProcTop &procTop, MessPass &messPass, Functions &functions)
- void implicitSolve\_R (Grid &grid, Implicit &implicit, Parameters &parameters, Time &time, ProcTop &procTop, MessPass &messPass, Functions &functions)
- void implicitSolve\_RT (Grid &grid, Implicit &implicit, Parameters &parameters, Time &time, ProcTop &procTop, MessPass &messPass, Functions &functions)
- void implicitSolve\_RTP (Grid &grid, Implicit &implicit, Parameters &parameters, Time &time, ProcTop &procTop, MessPass &messPass, Functions &functions)
- double dImplicitEnergyFunction\_None (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)
- double dImplicitEnergyFunction\_R (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)
- double dImplicitEnergyFunction\_R\_SB (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)
- double dImplicitEnergyFunction\_RT (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)
- double dImplicitEnergyFunction\_RT\_SB (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)
- double dImplicitEnergyFunction\_RTP (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)
- double dImplicitEnergyFunction\_RTP\_SB (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)
- double dImplicitEnergyFunction\_R\_LES (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)
- double dImplicitEnergyFunction\_R\_LES\_SB (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)

• double dImplicitEnergyFunction\_RT\_LES (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)

- double dImplicitEnergyFunction\_RT\_LES\_SB (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)
- double dImplicitEnergyFunction\_RTP\_LES (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)
- double dImplicitEnergyFunction\_RTP\_LES\_SB (Grid &grid, Parameters &parameters, Time &time, double dTemps[], int i, int j, int k)
- double dEOS GL (double dRho, double dE, Parameters parameters)
- void initDonorFracAndMaxConVel R GL (Grid &grid, Parameters &parameters)
- void initDonorFracAndMaxConVel R TEOS (Grid &grid, Parameters &parameters)
- void initDonorFracAndMaxConVel RT GL (Grid &grid, Parameters &parameters)
- void initDonorFracAndMaxConVel\_RT\_TEOS (Grid &grid, Parameters &parameters)
- void initDonorFracAndMaxConVel\_RTP\_GL (Grid &grid, Parameters &parameters)
- void initDonorFracAndMaxConVel\_RTP\_TEOS (Grid &grid, Parameters &parameters)

# 7.12.1 Detailed Description

Header file for physEquations.cpp

### 7.12.2 Function Documentation

# 7.12.2.1 void calDelt\_CONST (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function is used when a constant tie step is desired.

References Time::dConstTimeStep, Time::dDeltat\_n, Time::dDeltat\_nm1half, Time::dDeltat\_-np1half, Time::dt, ProcTop::nRank, and Time::nTimeStepIndex.

Referenced by setMainFunctions().

# 7.12.2.2 void calDelt\_R\_GL (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the time step by considering the sound crossing time in the radial direction only and is compatiable with a gamma law gass EOS.

### Parameters:

- $\leftarrow$  grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- $\leftrightarrow$  time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology. This function uses ProcTop::nRank to pass messages.

 $\label{lem:continuous} Parameters::dDonorCellMultiplier, Parameters::dGamma, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dMaxConvectiveVelocity, Time::dPerChange, Time::dt, Time::dTimeStepFactor, Grid::nCenIntOffset, Grid::nD, Grid::nDonorCellFrac, Grid::nE, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nP, Grid::nQ0, Grid::nR, ProcTop::nRank, Grid::nStartUpdateExplicit, Time::nTimeStepIndex, Grid::nU, and Grid::nU0.$ 

Referenced by setMainFunctions().

# 7.12.2.3 void calDelt\_R\_TEOS (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the time step by considering the sound crossing time in the radial direction only and is compatiable with a tabulated EOS.

#### Parameters:

- ← grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- $\leftrightarrow$  time contains time information, e.g. time step, current time etc.
- ← *procTop* contains information about the processor topology. This function uses ProcTop::nRank to pass messages.

 $Time::dDelRho\_t\_Rho \ max,$ References Time::dDelT t T max, Time::dDeltat -Time::dDeltat nm1half, Time::dDeltat np1half, Time::dDelUmU0 t UmU0 max, Time::dDelV t V max,  $Time::dDelW_t_W_max,$ Parameters::dDonorCellMin, rameters::dDonorCellMultiplier, Grid::dLocalGridNew, Grid::dLocalGridOld, eters::dMaxConvectiveVelocity, Time::dPerChange, Time::dt, Time::dTimeStepFactor, Grid::nCenIntOffset. Grid∷nD, Grid::nDonorCellFrac, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGamma, Grid::nP, Grid::nQ0, Grid::nR, ProcTop::nRank, Grid::nStartUpdateExplicit, Grid::nT, Time::nTimeStepIndex, Grid::nU, and Grid::nU0.

Referenced by setMainFunctions().

# 7.12.2.4 void calDelt\_RT\_GL (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the time step by considering the sound crossing time in the radial and theta directions only and is compatiable with a gamma law gass EOS.

### Parameters:

- ← grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- $\leftrightarrow$  time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology. This function uses ProcTop::nRank to pass messages.

 $References \quad Time::dDelE\_t\_E\_max, \quad Time::dDelRho\_t\_Rho\_max, \quad Time::dDeltat\_n, \quad Time::dDeltat\_np1half, \quad Time::dDelUmU0\_t\_UmU0\_max, \quad Time::dDelV\_t\_V\_max, \quad Time::dDelW\_t\_W\_max, \quad Parameters::dDonorCellMin, \quad Parameters::dDonorCellMultiplier, \quad Parameters::dGamma, \quad Grid::dLocalGridNew, \quad Grid::dLocalGridOld, \quad Parameters::dMaxConvectiveVelocity, \quad Time::dPerChange, \quad Time::dt, \quad Time::dTimeStepFactor, \quad TimeStepFactor, \quad TimeSt$ 

Grid::nCenIntOffset, Grid::nD, Grid::nDonorCellFrac, Grid::nDTheta, Grid::nE, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nR, ProcTop::nRank, Grid::nStartUpdateExplicit, Time::nTimeStepIndex, Grid::nU, Grid::nU0, and Grid::nV.

Referenced by setMainFunctions().

# 7.12.2.5 void calDelt\_RT\_TEOS (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the time step by considering the sound crossing time in the radial and theta directions and is compatiable with a tabulated EOS.

#### Parameters:

- $\leftarrow$  grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- $\leftrightarrow$  time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology. This function uses ProcTop::nRank to pass messages.

References Time::dDelRho t Rho max, Time::dDelT t T max, Time::dDeltat -Time::dDeltat np1half, Time::dDeltat nm1half, Time::dDelUmU0 t -UmU0 max, Time::dDelV t V max, Time::dDelW t W max, Parameters::dDonorCellMin, Parameters::dDonorCellMultiplier, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dMaxConvectiveVelocity, Time::dPerChange, Time::dt, Time::dTimeStepFactor, Grid::nCenIntOffset, Grid::nD, Grid::nDonorCellFrac, Grid::nDTheta, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGamma, Grid::nP. Grid::nQ0, Grid::nQ1, Grid::nR, ProcTop::nRank, Grid::nStartUpdateExplicit, Grid∷nT, Time::nTimeStepIndex, Grid::nU, Grid::nU0, and Grid::nV.

Referenced by setMainFunctions().

# 7.12.2.6 void calDelt\_RTP\_GL (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the time step by considering the sound crossing time in the radial, theta and phi directions only and is compatiable with a gamma law gass EOS.

#### Parameters:

- ← grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- $\leftrightarrow$  time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology. This function uses ProcTop::nRank to pass messages.

 $\label{lem:condition} References & Time::dDelE\_t\_E\_max, & Time::dDelRho\_t\_Rho\_max, & Time::dDeltat\_n, & Time::dDeltat\_nm1half, & Time::dDeltat\_np1half, & Time::dDelUmU0\_t\_UmU0\_max, \\ Time::dDelV\_t\_V\_max, & Time::dDelW\_t\_W\_max, & Parameters::dDonorCellMin, & Parameters::dDonorCellMultiplier, & Parameters::dGamma, & Grid::dLocalGridNew, & Grid::dLocalGridOld, \\ Parameters::dMaxConvectiveVelocity, & Time::dPerChange, & Time::dt, & Time::dTimeStepFactor, \\ \end{array}$ 

Grid::nCenIntOffset, Grid::nD, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nE, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, ProcTop::nRank, Grid::nSinThetaIJK, Grid::nStartUpdateExplicit, Time::nTimeStepIndex, Grid::nU, Grid::nU0, Grid::nV, and Grid::nW.

Referenced by setMainFunctions().

# 7.12.2.7 void calDelt\_RTP\_TEOS (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the time step by considering the sound crossing time in the radial, theta and phi directions and is compatiable with a tabulated EOS.

# Parameters:

- $\leftarrow$  grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- $\leftrightarrow$  time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology. This function uses ProcTop::nRank to pass messages.

References Time::dDelRho t Rho max, Time::dDelT t T max, Time::dDeltat -Time::dDeltat nm1half, Time::dDeltat np1half, Time::dDelUmU0 t UmU0 max, Time::dDelV t V max, Time::dDelW t W max, Parameters::dDonorCellMin, Grid::dLocalGridNew, rameters::dDonorCellMultiplier, Grid::dLocalGridOld. Param-Time::dPerChange, Time::dt, eters::dMaxConvectiveVelocity, Time::dTimeStepFactor, Grid::nCenIntOffset, Grid∷nD, Grid::nDonorCellFrac, Grid∷nDPhi, Grid::nDTheta, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGamma, Grid::nP, Grid∷nQ1, Grid::nQ2, Grid::nR, ProcTop::nRank, Grid::nQ0,Grid::nSinThetaIJK, Grid::nStartUpdateExplicit, Grid::nT, Time::nTimeStepIndex, Grid::nU, Grid::nU0, Grid::nV, and Grid::nW.

Referenced by setMainFunctions().

# 7.12.2.8 void calNewD\_R (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates new densities using terms in the radial direction only

# Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology, uses ProcTop::nRank when reporting negative densities

### **Boundary Conditions**

doesn't allow mass flux through outter interface

References Time::dDeltat\_np1half, Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nD, Grid::nDonorCellFrac, Grid::nEndGhostUpdateExplicit,

$$\label{lem:grid::new} \begin{split} & Grid::nEndUpdateExplicit, & Grid::nGlobalGridPositionLocalGrid, & Grid::nNumGhostCells, \\ & Grid::nR, & ProcTop::nRank, & Grid::nStartGhostUpdateExplicit, & Grid::nStartUpdateExplicit, \\ & Grid::nU, \ and \ Grid::nU0. \end{split}$$

Referenced by setMainFunctions().

# 7.12.2.9 void calNewD\_RT (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates new densities using terms in the radial and theta directions

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology, uses ProcTop::nRank when reporting negative densities

### **Boundary Conditions**

doesn't allow mass flux through outter interface

References Time::dDeltat np1half, Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nD, Grid::nDCosThetaIJK, Grid::nDonorCellFrac, Grid::nEndGhostUpdateExplicit, ProcTop::nRank, Grid::nEndUpdateExplicit, Grid::nR, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0, and Grid::nV.

Referenced by setMainFunctions().

# 7.12.2.10 void calNewD\_RTP (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates new densities using terms in the radial, theta, and phi directions

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology, uses ProcTop::nRank when reporting negative densities

### **Boundary Conditions**

doesn't allow mass flux through outter interface

References Time::dDeltat np1half, Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nDCosThetaIJK, Grid::nDonorCellFrac, Grid::nD, Grid∷nDPhi, Grid::nDTheta, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGlobalGridPositionLocalGrid, Grid::nNumGhostCells, Grid::nR, ProcTop::nRank, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0, Grid::nV, and Grid::nW.

Referenced by setMainFunctions().

## 7.12.2.11 void calNewDenave None (Grid & grid)

This function is a dumby funciton, and doesn't do anything. In the case of a 1D calculation the average density is undefined, and only the density is used. This is different from the case where the 1D region exsists on the rank 0 processor, but the grid as a whole is really 2D or 3D. In which case calNewDenave R should be used instead.

#### Parameters:

 $\leftrightarrow grid$ 

Referenced by setMainFunctions().

## 7.12.2.12 void calNewDenave R (Grid & grid)

This function calculates the horizontal average density in a 3\1D region. This really just copies the density from the particular radial zone into the averaged density variable. This way it can be used exactly the same way in the 1D region as it is in the 3D region. This is done using the density in the new grid, and places the result into the new grid.

### Parameters:

 $\leftrightarrow$  grid supplies the information needed to calculate the horizontal density average, it also stores the calculated horizontally averaged density.

References Grid::dLocalGridNew, Grid::nD, Grid::nDenAve, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nStartGhostUpdateExplicit, and Grid::nStartUpdateExplicit.

Referenced by setMainFunctions().

### 7.12.2.13 void calNewDenave RT (Grid & grid)

This function calculates the horizontal average density in a 2D region from the new grid density and stores the result in the new grid.

#### Parameters:

 $\leftrightarrow$  grid supplies the information needed to calculate the horizontal density average, it also stores the calculated horizontally averaged density.

 $\label{lem:condition} References & Grid::dLocalGridNew, & Grid::dLocalGridOld, & Grid::nCenIntOffset, \\ Grid::nD, & Grid::nDCosThetaIJK, & Grid::nDenAve, & Grid::nEndGhostUpdateExplicit, \\ Grid::nEndUpdateExplicit, & Grid::nR, & Grid::nStartGhostUpdateExplicit, \\ Grid::nStartUpdateExplicit. & Grid::nStartUpdateExplicit. \\ \end{tabular}$ 

Referenced by setMainFunctions().

## 7.12.2.14 void calNewDenave RTP (Grid & grid)

This function calculates the horizontal average density in a 3D region from the new grid density and stores the result in the new grid.

#### Parameters:

 $\leftrightarrow$  grid supplies the information needed to calculate the horizontal density average, it also stores the calculated horizontally averaged density.

References Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nD, Grid::nDCosThetaIJK, Grid::nDenAve, Grid::nDPhi, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nStartGhostUpdateExplicit, and Grid::nStartUpdateExplicit.

Referenced by setMainFunctions().

# 7.12.2.15 void calNewE\_R\_AD (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates new adiabatic energies using terms in the radial direction.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$

References Time::dDeltat\_np1half, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nCenIntOffset, Grid::nD, Grid::nDM, Grid::nDonorCellFrac, Grid::nE, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nP, Grid::nQ0, Grid::nR, ProcTop::nRank, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, and Grid::nU0.

Referenced by setMainFunctions().

# 7.12.2.16 void calNewE\_R\_NA (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates new non-adiabatic energies using terms in the radial direction and includes radiation diffusion terms.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nE][i+1][j][k] in calculation of  $E_{i+1/2,j,k}$  setting it equal to value at i.

## **Boundary Conditions**

grid.dLocalGridOld[grid.nDM][i+1][0][0] and grid.dLocalGridOld[grid.nE][i+1][j][k] missing in the calculation of upwind gradient in dA1. Using the centered gradient instead.

#### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nT][i+1][0][0]

References Time::dDeltat\_np1half, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dSigma, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nE, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGlobalGridPositionLocalGrid, Grid::nKappa, Grid::nNumGhostCells, Grid::nP, Grid::nQ0, Grid::nR, ProcTop::nRank, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nT, Grid::nU, and Grid::nU0.

Referenced by setMainFunctions().

# 7.12.2.17 void calNewE\_R\_NA\_LES (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates new non-adiabatic energies using terms in the radial direction and includes radiation diffusion terms. It also includes the terms for including real viscosity, used in the LES.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nE][i+1][j][k] in calculation of  $E_{i+1/2,j,k}$  setting it equal to value at i.

## **Boundary Conditions**

 $grid.dLocalGridOld[grid.nDM][i+1][0][0] \ \ and \ \ grid.dLocalGridOld[grid.nE][i+1][j][k] \ \ missing in the calculation of upwind gradient in dA1. Using the centered gradient instead.$ 

#### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nT][i+1][0][0]

### **Boundary Conditions**

missing energy outside the model, assuming it is the same as that in the last zone. That causes this term to be zero.

 $References \quad Time::dDeltat\_np1half, \quad Grid::dLocalGridNew, \quad Grid::dLocalGridOld, \quad Parameters::dPi, \quad Parameters::dSigma, \quad Grid::nCenIntOffset, \quad Grid::nD, \quad Grid::nDenAve, \quad Grid::nDM, \quad Grid::nDonorCellFrac, \quad Grid::nE, \quad Grid::nEddyVisc, \quad Grid::nEndGhostUpdateExplicit, \quad Grid::nEndUpdateExplicit, \quad Grid::nP, \quad Grid::nP, \quad Grid::nQ0, \quad Grid::nR, \quad ProcTop::nRank, \quad Grid::nStartGhostUpdateExplicit, \quad Grid::nStartUpdateExplicit, \quad Grid::nT, \quad Grid::nU, \quad and \quad Grid::nU0.$ 

# 7.12.2.18 void calNewE\_RT\_AD (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates new adiabatic energies using terms in the radial and theta directions.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated densities
- ← parameters various parameters needed for the calculation
- ← time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$

## **Boundary Conditions**

grid.dLocalGridOld[grid.nE][i+1][j][k] is missing

## **Boundary Conditions**

 $grid.dLocalGridOld[grid.nDM][i+1][0][0] \ \ and \ \ grid.dLocalGridOld[grid.nE][i+1][j][k] \ \ missing using inner gradient for both$ 

References Time::dDeltat np1half, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid∷nE, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nDTheta, Grid::nR, Grid::nSinThetaIJK, Grid∷nP, Grid::nQ0, Grid::nQ1, ProcTop::nRank, Grid::nStartGhostUpdateExplicit,Grid::nStartUpdateExplicit, Grid::nSinThetaIJp1halfK, Grid::nU, Grid::nU0, and Grid::nV.

Referenced by setMainFunctions().

# 7.12.2.19 void calNewE\_RT\_NA (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates new non-adiabatic energies using terms in the radial and theta directions and includes radiation diffusion terms.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated densities
- ← parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nE][i+1][j][k] in calculation of  $E_{i+1/2,j,k}$  setting it equal to value at i.

### **Boundary Conditions**

 $grid.dLocalGridOld[grid.nDM][i+1][0][0] \ and \ grid.dLocalGridOld[grid.nE][i+1][j][k] \ missing in the calculation of upwind gradient in dA1. Using the centered gradient instead.$ 

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nT][i+1][0][0] using flux equals  $2\sigma T^4$  at surface.

References Time::dDeltat\_np1half, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dSigma, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve,

Grid::nDM, Grid::nDonorCellFrac, Grid::nDTheta, Grid::nE, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nRappa, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nR, Proc-Top::nRank, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU, Grid::nU, and Grid::nV.

Referenced by setMainFunctions().

# 7.12.2.20 void calNewE\_RT\_NA\_LES (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates new non-adiabatic energies using terms in the radial and theta directions and includes radiation diffusion terms. It also includes the terms for including real viscosity, used in the LES.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- $\leftarrow$  time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$

## **Boundary Conditions**

Missing  $\Delta M_r$  outside model using Parameters::dAlpha times  $\Delta M_r$  in the last zone instead.

## **Boundary Conditions**

Setting energy at surface equal to energy in last zone.

## **Boundary Conditions**

missing eddy viscosity outside the model setting it to zero

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nT][i+1][0][0] using flux equals  $2\sigma T^4$  at surface.

### **Boundary Conditions**

missing energy outside the model, assuming it is the same as that in the last zone. That causes this term to be zero.

References Parameters::dAlpha, Parameters::dAlphaExtra, Time::dDeltat\_np1half, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dPrt, Parameters::dSigma, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDTheta, Grid::nE, Grid::nEddyVisc, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nKappa, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nR, Proc-Top::nRank, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nT, Grid::nU, Grid::nU0, and Grid::nV.

Referenced by setMainFunctions().

# 7.12.2.21 void calNewE\_RTP\_AD (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates new adiabatic energies using terms in the radial, theta, and phi directions.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated densities
- ← parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nE][i+1][j][k] in calculation of  $E_{i+1/2,j,k}$  setting it equal to zero.

## **Boundary Conditions**

 $grid.dLocalGridOld[grid.nDM][i+1][0][0] \ \ and \ \ grid.dLocalGridOld[grid.nE][i+1][j][k] \ \ missing in the calculation of upwind gradient in dA1. Using the centered gradient.$ 

References Time::dDeltat\_np1half, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nE, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, ProcTop::nRank, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU, Grid::nV, and Grid::nW.

Referenced by setMainFunctions().

# 7.12.2.22 void calNewE\_RTP\_NA (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates new non-adiabatic energies using terms in the radial, theta, and phi directions and includes radiation diffusion terms.

## Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated densities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nE][i+1][j][k] in calculation of  $E_{i+1/2,j,k}$  setting it equal to the value at i.

## **Boundary Conditions**

 $grid.dLocalGridOld[grid.nDM][i+1][0][0] \ and \ grid.dLocalGridOld[grid.nE][i+1][j][k] \ missing in the calculation of upwind gradient in dA1. Using the centered gradient instead.$ 

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nT][i+1][0][0]

References Time::dDeltat\_np1half, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dSigma, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nE, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nKappa, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, ProcTop::nRank, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU, Grid::nU, Grid::nV, and Grid::nW.

Referenced by setMainFunctions().

# 7.12.2.23 void calNewE\_RTP\_NA\_LES (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates new non-adiabatic energies using terms in the radial, theta, and phi directions and includes radiation diffusion terms. It also includes the terms for including real viscosity, used in the LES.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated densities
- ← parameters various parameters needed for the calculation
- ← time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$

## **Boundary Conditions**

Missing  $\Delta M_r$  outside model using Parameters::dAlpha times  $\Delta M_r$  in the last zone instead.

### **Boundary Conditions**

Missing W at i+1, assuming the same as at i

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nE][i+1][j][k] in calculation of  $E_{i+1/2,j,k}$  setting it equal to the value at i.

## **Boundary Conditions**

missing density outside model, setting it to zero

## **Boundary Conditions**

missing eddy viscosity outside the model setting it to zero

## **Boundary Conditions**

grid.dLocalGridOld[grid.nDM][i+1][0][0] and grid.dLocalGridOld[grid.nE][i+1][j][k] missing in the calculation of upwind gradient in dA1. Using the centered gradient instead.

#### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nT][i+1][0][0]

References Parameters::dAlpha, Parameters::dAlphaExtra, Time::dDeltat np1half, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dPrt, Parameters ters::dSigma, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nE, Grid::nEddyVisc, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGlobalGridPositionLocalGrid, Grid::nKappa, Grid::nNumGhostCells, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, Proc-Top::nRank, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nT, Grid::nU, Grid::nU0, Grid::nV, and Grid::nW.

Referenced by setMainFunctions().

## 7.12.2.24 void calNewEddyVisc None (Grid & grid, Parameters & parameters)

This function is a empty function used as a place holder when no eddy viscosity model is being used.

#### Parameters:

- $\leftrightarrow qrid$
- $\leftarrow$  parameters

Referenced by setMainFunctions().

## 7.12.2.25 void calNewEddyVisc R CN (Grid & grid, Parameters & parameters)

This function calculates the eddy viscosity using a constant times the zoning with only the radial terms.

### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the eddy viscosity.
- ← parameters contains parameters used in calculating the eddy viscosity.

 $\label{lem:references} References & Parameters:: dEddy Viscosity, & Grid:: dLocal Grid New, & Parameters:: dMax Convective Velocity, & Grid:: nCenInt Offset, & Grid:: nEddy Visc, Grid:: nEndGhost Update Explicit, & Grid:: nEndUpdate Explicit, & Grid:: nR, Grid:: nStart Ghost Update Explicit, and Grid:: nStart Update Explicit. \\ \\$ 

## 7.12.2.26 void calNewEddyVisc R SM (Grid & grid, Parameters & parameters)

This function calculates the eddy viscosity with only the radial terms.

## Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the eddy viscosity.
- ← parameters contains parameters used in calculating the eddy viscosity.

References Parameters::dEddyViscosity, Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nD, Grid::nEddyVisc, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nR, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, and Grid::nU.

## 7.12.2.27 void calNewEddyVisc RT CN (Grid & grid, Parameters & parameters)

This function calculates the eddy viscosity using a constant times the zoning with only the radial and theta terms.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the eddy viscosity.
- ← parameters contains parameters used in calculating the eddy viscosity.

 $\label{lem:convective} References & Parameters:: dEddy Viscosity, & Grid:: dLocalGridNew, & Grid:: dLocalGridOld, \\ Parameters:: dMaxConvectiveVelocity, & Grid:: nCenIntOffset, & Grid:: nDTheta, \\ Grid:: nEddy Visc, & Grid:: nEndGhost Update Explicit, & Grid:: nEndUpdate Explicit, & Grid:: nR, \\ Grid:: nStartGhost Update Explicit, & and & Grid:: nStartUpdate Explicit. \\ \end{cases}$ 

Referenced by setMainFunctions().

## 7.12.2.28 void calNewEddyVisc RT SM (Grid & grid, Parameters & parameters)

This function calculates the eddy viscosity with only the radial and theta terms.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the eddy viscosity.
- ← parameters contains parameters used in calculating the eddy viscosity.

 $\label{eq:references} \begin{array}{llll} References & Parameters::dEddyViscosity, & Grid::dLocalGridNew, & Grid::dLocalGridOld, \\ Grid::nCenIntOffset, & Grid::nD, & Grid::nDTheta, & Grid::nEddyVisc, \\ Grid::nEndGhostUpdateExplicit, & Grid::nEndUpdateExplicit, & Grid::nR, \\ Grid::nStartGhostUpdateExplicit, & Grid::nStartUpdateExplicit, & Grid::nU, & Grid::nU, & Grid::nU, & Grid::nU, & Grid::nU, & Grid::nV. \\ \end{array}$ 

Referenced by setMainFunctions().

# 7.12.2.29 void calNewEddyVisc\_RTP\_CN (Grid & grid, Parameters & parameters)

This function calculates the eddy viscosity using a constant times the zoning with only the radial, theta, and phi terms.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the eddy viscosity.
- ← parameters contains parameters used in calculating the eddy viscosity.

References Parameters::dEddyViscosity, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dMaxConvectiveVelocity, Grid::nCenIntOffset, Grid::nDPhi, Grid::nDTheta, Grid::nEddyVisc, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nR, Grid::nSinThetaIJK, Grid::nStartGhostUpdateExplicit, and Grid::nStartUpdateExplicit.

Referenced by setMainFunctions().

# 7.12.2.30 void calNewEddyVisc\_RTP\_SM (Grid & grid, Parameters & parameters)

This function calculates the eddy viscosity with only the radial, theta, and phi terms.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the eddy viscosity.
- ← parameters contains parameters used in calculating the eddy viscosity.

## **Boundary Conditions**

assuming that theta velocity is constant across surface

## **Boundary Conditions**

assume phi velocity is constant across surface

References Parameters::dEddyViscosity, Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nCotThetaIJK, Grid::nD, Grid::nDPhi, Grid::nDTheta, Grid::nEddyVisc, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nR, Grid::nSinThetaIJK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU, Grid::nV, and Grid::nW.

Referenced by setMainFunctions().

## 7.12.2.31 void calNewP GL (Grid & grid, Parameters & parameters)

This function calculates the pressure. It is calculated using the new values of quantities and places the result in the new grid. It uses a gamma law gas give in dEOS GL to calculate the pressure.

#### Parameters:

- $\leftrightarrow$  *grid* supplies the input for calculating the pressure and also accepts the result of the pressure calculations.
- ← parameters contains parameters used in calculating the pressure, namely the adiabatic gamma that is used.

Referenced by setMainFunctions().

# $7.12.2.32 \quad \text{void calNewPEKappaGamma\_TEOS (Grid \& \textit{grid}, Parameters \& \textit{parameters})}$

This function calculates the Energy, pressure and opacity of a cell. It calculates it using the new vaules of quantities and places the result in the new grid.

#### Parameters:

 $\leftrightarrow grid$  supplies the input for calculating the pressure and also accepts the result of the pressure calculation

← parameters contains parameters used in calculating the pressure.

References Grid::dLocalGridNew, Parameters::eosTable, eos::getPEKappaGamma(), Grid::nD, Grid::nE, Grid::nEndGhostUpdateImplicit, Grid::nEndUpdateImplicit, Grid::nGamma, Grid::nKappa, Grid::nP, Grid::nStartGhostUpdateImplicit, Grid::nStartUpdateImplicit, and Grid::nT.

Referenced by implicitSolve R(), implicitSolve RT(), and implicitSolve RTP().

## 7.12.2.33 void calNewQ0 R GL (Grid & grid, Parameters & parameters)

This funciton calculates the artificial viscosity of a cell. It calculates it using the new values of quantities and places the result in the new grid. It does this for the radial component of the viscosity only. It uses the sound speed derived from the adiabatic gamma given for the gamma law gas equation of state.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the artificial viscosity and also accepts the result of the artificial viscosity calculation.
- $\leftarrow$  parameters contains parameters used when calculating the artificial viscosity, namely the adiabatic gamma.

 $\label{lem:condition} References & Parameters:: dA, & Parameters:: dAVThreshold, & Parameters:: dGamma, Grid:: dLocalGridNew, & Grid:: nCenIntOffset, & Grid:: nD, & Grid:: nEndGhostUpdateExplicit, Grid:: nEndUpdateExplicit, Grid:: nP, & Grid:: nQ0, & Grid:: nR, & Grid:: nStartGhostUpdateExplicit, Grid:: nStartUpdateExplicit, and Grid:: nU.$ 

Referenced by setMainFunctions().

## 7.12.2.34 void calNewQ0 R TEOS (Grid & grid, Parameters & parameters)

This function calculates the artificial viscosity of a cell. It calculates it using the old values of quantities and places the result in the old grid. It does this for the radial component of the viscosity only. It uses a sound speed derived from a tabulated equaiton of state for the calculation.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the artificial viscosity and also accepts the result of the artificial viscosity calculation
- ← parameters contains parameters used in calculating the artificial viscosity.

References Parameters::dA, Parameters::dAVThreshold, Grid::dLocalGridNew, Grid::nCenIntOffset, Grid::nD, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGamma, Grid::nP, Grid::nQ0, Grid::nR, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, and Grid::nU.

Referenced by setMainFunctions().

### 7.12.2.35 void calNewQ0Q1 RT GL (Grid & grid, Parameters & parameters)

This function calculates the artificial viscosity of a cell. It calculates it using the new values of quantities and places the result in the new grid. It does this for the radial and theta components

of the viscosity. It uses the sound speed derived from the adiabatic gamma given for the gamma law gas equation of state.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the artificial viscosity and also accepts the result of the artificial viscosity calculations.
- ← parameters contains parameters used when calculating the artificial viscosity, namely the adiabatic gamma.

 $\label{lem:references} References & Parameters::dA, & Parameters::dAVThreshold, & Parameters::dGamma, Grid::dLocalGridNew, & Grid::dLocalGridOld, & Grid::nCenIntOffset, & Grid::nD, Grid::nEndGhostUpdateExplicit, & Grid::nEndUpdateExplicit, & Grid::nP, & Grid::nQ0, & Grid::nQ1, Grid::nR, & Grid::nSinThetaIJK, & Grid::nSinThetaIJp1halfK, & Grid::nStartGhostUpdateExplicit, Grid::nV.$ 

Referenced by setMainFunctions().

## 7.12.2.36 void calNewQ0Q1 RT TEOS (Grid & grid, Parameters & parameters)

This function calculates the artificial viscosity of a cell. It calculates it using the old values of quantities and places the result in the old grid. It does this for two component of the viscosity.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the artificial viscosity and also accepts the result of the artificial viscosity calculation
- ← parameters contains parameters used in calculating the artificial viscosity.

References Parameters::dA, Parameters::dAVThreshold, Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nD, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGamma, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, and Grid::nV.

Referenced by setMainFunctions().

# 7.12.2.37 void calNewQ0Q1Q2\_RTP\_GL (Grid & grid, Parameters & parameters)

This function calculates the artificial viscosity of a cell. It calculates it using the new values of quantities and places the result in the new grid. It does this for the radial, theta, and phi components of the viscosity. It uses the sound speed derived from the adiabatic gamma given for the gamma law gas equation of state.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the artificial viscosity and also accepts the result of the artificial viscosity calculations.
- $\leftarrow$  parameters contains parameters used when calculating the artificial viscosity, namely the adiabatic gamma.

References Parameters::dA, Parameters::dAVThreshold, Parameters::dGamma, Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nD,

 $\label{eq:Grid::nEndGhostUpdateExplicit} Grid::nEndUpdateExplicit, & Grid::nP, & Grid::nQ0, \\ Grid::nQ1, & Grid::nQ2, & Grid::nR, & Grid::nSinThetaIJK, & Grid::nSinThetaIJp1halfK, \\ Grid::nStartGhostUpdateExplicit, & Grid::nStartUpdateExplicit, & Grid::nU, & Grid::nV, & and \\ Grid::nW. & Grid::nV, & Gri$ 

Referenced by setMainFunctions().

# 7.12.2.38 void calNewQ0Q1Q2\_RTP\_TEOS (Grid & grid, Parameters & parameters)

This function calculates the artificial viscosity of a cell. It calculates it using the old values of quantities and places the result in the old grid. It does this for the three component of the viscosity.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the artificial viscosity and also accepts the result of the artificial viscosity calculation
- ← parameters contains parameters used in calculating the artificial viscosity.

References Parameters::dA, Parameters::dAVThreshold, Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nD, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGamma, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nV, and Grid::nW.

Referenced by setMainFunctions().

## 7.12.2.39 void calNewR (Grid & grid, Time & time)

This function calculates the radii, from the new radial grid velocities

### Parameters:

- $\leftrightarrow grid$  contains the local grid, and will hold the newly updated radial velocities
- $\leftarrow$  time contains time information, e.g. time step, current time etc.

References Time::dDeltat\_np1half, Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nR, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, and Grid::nU0.

Referenced by setMainFunctions().

# 7.12.2.40 void calNewTPKappaGamma\_TEOS (Grid & grid, Parameters & parameters)

This function calculates the Temperature, pressure and opacity of a cell. It calculates it using the new vaules of quantities and places the result in the new grid.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the pressure and also accepts the result of the pressure calculation
- $\leftarrow$  parameters contains parameters used in calculating the pressure.

References Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dTolerance, Parameters::eosTable, eos::getEAndDTDE(), eos::getPKappaGamma(), Grid::nD, Grid::nE, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGamma, Grid::nKappa, Parameters::nMaxIterations, Grid∷nP, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, and Grid::nT.

Referenced by setMainFunctions().

# 7.12.2.41 void calNewU0\_R (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop, MessPass & messPass)

This function calculates the radial grid velocity, it does so by considering only the radial terms

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated radial grid velocities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology
- $\leftarrow messPass$

#### Todo

At some point I will likely want to make this function compatiable with a 3D domain decomposition instead of a purely radial domain decomposition.

## **Boundary Conditions**

assuming density outside the star is 0 for the purposes of calculating the grid velocity. This boundary condition works, but is probably not strictly correct. The proper condition can be calculated by knowing that the mass conservation is independent of u0\_ip1half at the surface. This allows one to simply use the new densities and calculate the grid velocity from them.

 $\label{lem:conder} References\ Grid::dLocalGridNew,\ Grid::dLocalGridOld,\ Grid::nCenIntOffset,\ ProcTop::nCoords,\ Grid::nD,\ Grid::nDonorCellFrac,\ Grid::nEndGhostUpdateExplicit,\ Grid::nEndUpdateExplicit,\ ProcTop::nNumRadialNeighbors,\ Grid::nR,\ ProcTop::nRadialNeighborNeighborIDs,\ ProcTop::nRadialNeighborRanks,\ ProcTop::nRank,\ Grid::nStartGhostUpdateExplicit,\ Grid::nU,\ Grid::nU0,\ MessPass::typeRecvNewVar,\ and\ Mess-Pass::typeSendNewVar.$ 

Referenced by setMainFunctions().

# 7.12.2.42 void calNewU0\_RT (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop, MessPass & messPass)

This function calculates the radial grid velocity, and does it by only considering the radial and theta terms

### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated radial grid velocities
- $\leftarrow$  parameters various parameters needed for the calculation
- $\leftarrow$  time contains time information, e.g. time step, current time etc.

- $\leftarrow procTop$  contains information about the processor topology
- $\leftrightarrow messPass$  handles data needed for message passing

### Todo

At some point I will likely want to make this function compatiable with a 3D domain decomposition instead of a purely radial domain decomposition.

## **Boundary Conditions**

```
grid.dLocalGridOld[grid.nD][i+1][j][k] is missing
```

 $\label{lem:conder} References\ Grid::dLocalGridNew,\ Grid::dLocalGridOld,\ Grid::nCenIntOffset,\ ProcTop::nCoords,\ Grid::nD,\ Grid::nDCosThetaIJK,\ Grid::nDonorCellFrac,\ Grid::nEndGhostUpdateExplicit,\ Grid::nEndUpdateExplicit,\ Grid::nLocalGridDims,\ Grid::nNumGhostCells,\ ProcTop::nNumRadialNeighbors,\ Grid::nR,\ ProcTop::nRadialNeighborNeighborIDs,\ ProcTop::nRadialNeighborRanks,\ ProcTop::nRank,\ Grid::nSinThetaIJp1halfK,\ Grid::nStartGhostUpdateExplicit,\ Grid::nStartUpdateExplicit,\ Grid::nU,\ Grid::nU,\ Grid::nV,\ MessPass::typeRecvNewVar,\ and\ MessPass::typeSendNewVar.$ 

Referenced by setMainFunctions().

# 7.12.2.43 void calNewU0\_RTP (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop, MessPass & messPass)

This function calculates the radial grid velocity, and does it by considering all radial, theta and phi terms

#### Parameters:

- $\leftrightarrow grid$  contains the local grid, and will hold the newly updated radial grid velocities
- $\leftarrow \textit{parameters} \,\,$  various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology
- $\leftrightarrow messPass$  handles data needed for message passing

## Todo

At some point I will likely want to make this function compatiable with a 3D domain decomposition instead of a purely radial domain decomposition.

References Grid::dLocalGridNew, Grid::dLocalGridOld, Grid::nCenIntOffset, Top::nCoords, Grid::nD, Grid::nDCosThetaIJK, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nLocalGridDims, Grid::nNumGhostCells, ProcTop::nNumRadialNeighbors, Grid::nR, ProcTop::nRadialNeighborNeighborIDs, ProcTop::nRadialNeighborRanks, ProcTop::nRank, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid∷nU. Grid::nU0, Grid∷nV, Grid::nW, MessPass::typeRecvNewVar, Pass::typeSendNewVar.

Referenced by setMainFunctions().

# 7.12.2.44 void calNewU\_R (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the radial velocity, and does it by only considering the radial terms.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated radial velocities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nD][nICen+1][j][k] in calculation of  $\rho_{i+1/2}$ , setting it to 0.0

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nP][nICen+1][j][k] in calculation of  $S_1$ , setting it to -1.0\*grid.dLocalGridOld[grid.nP][nICen][j][k].

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nDM][nICen+1][0][0] in calculation of centered  $A_1$  gradient, setting it to zero.

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nU][i+1][j][k] and grid.dLocalGridOld[grid.nDM][nICen+1][0][0] in calculation of upwind gradient, when moving inward. Using centered gradient instead.

References Parameters::dAlpha, Parameters::dAlphaExtra, Time::dDeltat\_n, Parameters::dG, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nCenIntOffset, Grid::nD, Grid::nDM, Grid::nDonorCellFrac, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGlobalGridPositionLocalGrid, Grid::nM, Grid::nP, Grid::nQ0, Grid::nR, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, and Grid::nU0.

Referenced by calNewVelocities R().

# 7.12.2.45 void calNewU\_R\_LES (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the radial velocity, and does it by only considering the radial terms. It also includes the terms for including real viscosity, used in the LES.

#### Parameters:

- $\leftrightarrow grid$  contains the local grid, and will hold the newly updated radial velocities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology

#### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nD][nICen+1][j][k] in calculation of  $\rho_{i+1/2}$ , setting it to 0.0

## **Boundary Conditions**

missing grid.dLocalGridOld[grid.nU][i+1][j][k] using velocity at i

## **Boundary Conditions**

Assuming eddy viscosity outside model is zero.

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nP][nICen+1][j][k] in calculation of  $S_1$ , setting it to -1.0\*grid.dLocalGridOld[grid.nP][nICen][j][k].

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nDM][nICen+1][0][0] in calculation of centered  $A_1$  gradient, setting it to zero.

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nU][i+1][j][k] and grid.dLocalGridOld[grid.nDM][nICen+1][0][0] in calculation of upwind gradient, when moving inward. Using centered gradient instead.

References Parameters::dAlpha, Parameters::dAlphaExtra, Time::dDeltat n, Parameters::dG, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nDonorCellFrac, Grid::nCenIntOffset, Grid::nD, Grid::nDM, Grid::nEddyVisc, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nM, Grid::nP, Grid::nQ0, Grid∷nR, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid∷nU0.

Referenced by calNewVelocities R LES().

# 7.12.2.46 void calNewU\_RT (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the radial velocity, and does it by only considering the radial and theta terms.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated radial velocities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nD][nICen+1][j][k] in calculation of  $\rho_{i+1/2,j,k}$ , setting it to zero.

## **Boundary Conditions**

assuming theta velocity is constant across surface

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nDenAve][nICen+1][0][0] in calculation of  $\langle \rho \rangle_{i+1/2}$ , setting it to zero.

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nP][nICen+1][j][k] in calculation of  $S_1$ , setting it to -1.0\*dP\_ijk\_n.

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nDM][nICen+1][0][0] in calculation of centered  $A_1$  gradient, setting it to zero.

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nU][i+1][j][k] and grid.dLocalGridOld[grid.nDM][nICen+1][0][0] in calculation of upwind gradient, when moving inward. Using centered gradient instead.

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nDM][i+1][0][0] in calculation of  $S_1$  using Parameters::dAlpha \*grid.dLocalGridOld[grid.nDM][nICen][0][0] instead.

References Parameters::dAlpha, Parameters::dAlphaExtra, Time::dDeltat\_n, Parameters::dG, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDTheta, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nM, Grid::nP, Grid::nQ0, Grid::nR, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0, and Grid::nV.

Referenced by calNewVelocities RT().

# 7.12.2.47 void calNewU\_RT\_LES (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the radial velocity, and does it by only considering the radial and theta terms. It also includes the terms for including real viscosity, used in the LES.

### Parameters:

- $\leftrightarrow grid$  contains the local grid, and will hold the newly updated radial velocities
- ← parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology

## **Boundary Conditions**

$$\label{eq:missing_missing} \begin{split} & \text{Missing} \quad \text{grid.dLocalGridOld[grid.nDM][i+1][0][0]} \quad \text{in} \quad \text{calculation} \quad \text{of} \quad S_1 \quad \text{using} \\ & \text{Parameters::dAlpha} *& \text{grid.dLocalGridOld[grid.nDM][nICen][0][0]} \quad \text{instead.} \end{split}$$

## **Boundary Conditions**

Missing density outside of surface, setting it to zero.

## **Boundary Conditions**

Missing density outside model, setting it to zero.

## **Boundary Conditions**

assuming theta and phi velocity same outside star as inside.

## **Boundary Conditions**

Assuming theta velocities are constant across surface.

## **Boundary Conditions**

assuming that \$V\$ at \$i+1\$ is equal to \$v\$ at \$i\$.

## **Boundary Conditions**

Missing pressure outside surface setting it equal to negative pressure in the center of the first cell so that it will be zero at surface.

## **Boundary Conditions**

assume viscosity is zero outside the star.

## **Boundary Conditions**

Missing mass outside model, setting it to zero.

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nU][i+1][j][k] and grid.dLocalGridOld[grid.nDM][nICen+1][0][0] in calculation of upwind gradient, when moving inward. Using centered gradient instead.

References Parameters::dAlpha, Parameters::dAlphaExtra, Time::dDeltat n, ters::dG, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nCenIntOffset, Grid::nCotThetaIJK, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDTheta, Grid::nEddyVisc, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid∷nM, Grid∷nP, Grid∷nQ0, Grid::nQ1, Grid∷nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0, and Grid::nV.

Referenced by calNewVelocities RT LES().

# 7.12.2.48 void calNewU\_RTP (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the radial velocity, and does it by including all radial, theta and phi terms.

#### Parameters:

- $\leftrightarrow grid$  contains the local grid, and will hold the newly updated radial velocities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology

## **Boundary Conditions**

missing grid.dLocalGridOld[grid.nU][i+1][j][k] in calculation of  $u_{i+1,j,k}$  setting  $u_{i+1,j,k}=u_{i+1/2,j,k}$ .

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nD][i+1][j][k] in calculation of  $\rho_{i+1/2,j,k}$ , setting it to zero.

## **Boundary Conditions**

assuming theta velocity is constant across the surface.

## **Boundary Conditions**

assuming phi velocity is constant across the surface.

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nDenAve][nICen+1][0][0] in calculation of  $\langle \rho \rangle_{i+1/2}$  setting it to zero

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nDM][nICen+1][0][0] in calculation of centered  $A_1$  gradient, setting it equal to Parameters::dAlpha grid.dLocalGridOld[grid.nDM][nICen][0][0].

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nU][i+1][j][k] and grid.dLocalGridOld[grid.nDM][nICen+1][0][0] in calculation of upwind gradient, when moving inward. Using centered gradient instead.

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nDM][i+1][0][0] in calculation of  $S_1$  using Parameters::dAlpha \*grid.dLocalGridOld[grid.nDM][nICen][0][0] instead.

References Parameters::dAlpha, Parameters::dAlphaExtra, Time::dDeltat\_n, Parameters::dG, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nM, Grid::nP, Grid::nQ0, Grid::nR, Grid::nSinThetaIJK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU, Grid::nV, and Grid::nW.

Referenced by calNewVelocities RTP().

# 7.12.2.49 void calNewU\_RTP\_LES (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the radial velocity, and does it by including all radial, theta and phi terms. It also includes the terms for including real viscosity, used in the LES.

## Parameters:

- $\leftrightarrow grid$  contains the local grid, and will hold the newly updated radial velocities
- $\leftarrow$  parameters various parameters needed for the calculation

- $\leftarrow$  time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nDM][i+1][0][0] in calculation of  $S_1$  using Parameters::dAlpha \*grid.dLocalGridOld[grid.nDM][nICen][0][0] instead.

## **Boundary Conditions**

Missing density outside of surface, setting it to zero.

## **Boundary Conditions**

Missing density outside model, setting it to zero.

## **Boundary Conditions**

assuming theta and phi velocity same outside star as inside.

## **Boundary Conditions**

Assuming theta velocities are constant across surface.

## **Boundary Conditions**

assuming that \$V\$ at \$i+1\$ is equal to \$v\$ at \$i\$.

#### **Boundary Conditions**

Missing pressure outside surface setting it equal to negative pressure in the center of the first cell so that it will be zero at surface.

## **Boundary Conditions**

assume viscosity is zero outside the star.

## **Boundary Conditions**

Missing mass outside model, setting it to zero.

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nU][i+1][j][k] and grid.dLocalGridOld[grid.nDM][nICen+1][0][0] in calculation of upwind gradient, when moving inward. Using centered gradient instead.

References Parameters::dAlpha, Parameters::dAlphaExtra, Time::dDeltat\_n, Parameters::dG, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nCenIntOffset, Grid::nCotThetaIJK, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nEddyVisc, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGlobalGridPositionLocalGrid, Grid::nM, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU0, Grid::nV, and Grid::nW.

Referenced by calNewVelocities RTP LES().

# 7.12.2.50 void calNewV\_RT (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the theta velocity, and does it by only considering the radial and theta terms.

## Parameters:

- $\leftrightarrow grid$  contains the local grid, and will hold the newly updated theta velocities
- $\leftarrow$  parameters various parameters needed for the calculation
- $\leftarrow$  time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology

## **Boundary Conditions**

grid.dLocalGridOld[grid.nV][i+1][j+1][k] is missing

## **Boundary Conditions**

missing upwind gradient, using centred gradient instead

References Time::dDeltat\_n, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDTheta, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nR, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU, Grid::nU, and Grid::nV.

Referenced by calNewVelocities RT().

# 7.12.2.51 void calNewV\_RT\_LES (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the theta velocity, and does it by only considering the radial and theta terms. It also includes the terms for including real viscosity, used in the LES.

## Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated theta velocities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology

### **Boundary Conditions**

Assuming density outside star is zero

## **Boundary Conditions**

Assuming theta velocity is constant across surface.

#### **Boundary Conditions**

Assuming eddy viscosity is zero at surface.

References Time::dDeltat\_n, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nCenIntOffset, Grid::nCotThetaIJp1halfK, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDTheta, Grid::nEddyVisc, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU, and Grid::nV.

Referenced by calNewVelocities RT LES().

# 7.12.2.52 void calNewV\_RTP (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the theta velocity, and does it by considering the radial, theta, and phi terms.

#### Parameters:

- $\leftrightarrow grid$  contains the local grid, and will hold the newly updated theta velocities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology

## **Boundary Conditions**

Assuming theta and phi velocities are the same at the surface of the star as just inside the star

## **Boundary Conditions**

ussing cetnered gradient for upwind gradient outside star at surface.

References Time::dDeltat n, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nCotThetaIJp1halfK, Grid::nD, Grid::nDenAve, Grid::nCenIntOffset, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0, Grid∷nQ1, Grid::nQ2, Grid::nR, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0, Grid::nV, and Grid::nW.

Referenced by calNewVelocities RTP().

# 7.12.2.53 void calNewV\_RTP\_LES (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the theta velocity, and does it by considering the radial, theta, and phi terms. It also includes the terms for including real viscosity, used in the LES.

#### Parameters:

- $\leftrightarrow grid$  contains the local grid, and will hold the newly updated theta velocities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← *time* contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology

## **Boundary Conditions**

Assuming density outside star is zero

## **Boundary Conditions**

Assuming theta velocity is constant across surface.

## **Boundary Conditions**

Assuming eddy viscosity is zero at surface.

Grid::dLocalGridNew, Grid::dLocalGridOld, References Time::dDeltat n, Pa-Grid::nCenIntOffset, rameters::dPi, Grid::nCotThetaIJp1halfK, Grid∷nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nEddvVisc, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGlobalGridPositionLocalGrid, Grid::nNumGhostCells, Grid::nP, Grid::nQ0, Grid::nQ2, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid∷nQ1, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0, Grid::nV, and Grid::nW.

Referenced by calNewVelocities\_RTP\_LES().

# 7.12.2.54 void calNewVelocities\_R (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function simply calls a function that calculate the radial velocity. Calls the function calNewU R to calculate radial velocity, including only radial terms.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid data and supplies the needed data to calculate the new velocities as well as holding the new velocities.
- $\leftarrow$  parameters contains parameters used in the calculation of the new velocities.
- $\leftarrow$  time contains time step information, current time step, and current time
- $\leftarrow procTop$  contains processor topology information

References calNewU R().

Referenced by setMainFunctions().

# 7.12.2.55 void calNewVelocities $R_LES$ (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function simply calls a function that calculate the radial velocity. Calls the function calNewU R to calculate radial velocity, including only radial terms.

### Parameters:

- $\leftrightarrow$  grid contains the local grid data and supplies the needed data to calculate the new velocities as well as holding the new velocities.
- $\leftarrow$  parameters contains parameters used in the calculation of the new velocities.
- $\leftarrow$  time contains time step information, current time step, and current time
- $\leftarrow procTop$  contains processor topology information

References calNewU R LES().

# 7.12.2.56 void calNewVelocities\_RT (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function simply calls two other functions that calculate the radial and theta velocities. Calls the two functions calNewU\_RT and calNewV\_RT to calculate radial and theta velocities, including both radial and theta terms.

#### Parameters:

- $\leftarrow$  parameters contains parameters used in the calculation of the new velocities.
- $\leftarrow$  time contains time step information, current time step, and current time
- ← procTop contains processor topology information

References calNewU\_RT(), and calNewV\_RT().

Referenced by setMainFunctions().

# 7.12.2.57 void calNewVelocities RT LES (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function simply calls two other functions that calculate the radial and theta velocities. Calls the two functions calNewU\_RT and calNewV\_RT to calculate radial and theta velocities, including both radial and theta terms.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid data and supplies the needed data to calculate the new velocities as well as holding the new velocities.
- $\leftarrow$  parameters contains parameters used in the calculation of the new velocities.
- ← time contains time step information, current time step, and current time
- $\leftarrow procTop$  contains processor topology information

References calNewU RT LES(), and calNewV RT LES().

Referenced by setMainFunctions().

# 7.12.2.58 void calNewVelocities\_RTP (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function simply calls three other functions that calculate the radial, theta and phi velocities. Calls the two functions calNewU\_RTP, calNewV\_RTP and calNewW\_RTP to calculate radial, theta, and phi velocities, including radial, theta, and phi terms.

### Parameters:

- $\leftrightarrow$  grid contains the local grid data and supplies the needed data to calculate the new velocities as well as holding the new velocities.
- ← parameters contains parameters used in the calculation of the new velocities.
- ← time contains time step information, current time step, and current time

 $\leftarrow procTop$  contains processor topology information

References calNewU\_RTP(), calNewV\_RTP(), and calNewW\_RTP(). Referenced by setMainFunctions().

# 7.12.2.59 void calNewVelocities RTP LES (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function simply calls three other functions that calculate the radial, theta and phi velocities. Calls the two functions calNewU\_RTP, calNewV\_RTP and calNewW\_RTP to calculate radial, theta, and phi velocities, including radial, theta, and phi terms.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid data and supplies the needed data to calculate the new velocities as well as holding the new velocities.
- $\leftarrow$  parameters contains parameters used in the calculation of the new velocities.
- $\leftarrow$  time contains time step information, current time step, and current time
- $\leftarrow procTop$  contains processor topology information

References calNewU\_RTP\_LES(), calNewV\_RTP\_LES(), and calNewW\_RTP\_LES(). Referenced by setMainFunctions().

# 7.12.2.60 void calNewW\_RTP (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the phi velocity, and does it by only considering the radial, theta, and phi terms.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated theta velocities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology

## **Boundary Conditions**

missing grid.dLocalGridOld[grid.nW][i+1][j][k] assuming that the phi velocity at the outter most interface is the same as the phi velocity in the center of the zone.

### **Boundary Conditions**

missing grid.dLocalGridOld[grid.nW][i+1][j][k] in outter most zone. This is needed to calculate the upwind gradient for donnor cell. The centered gradient is used instead when moving in the negative direction.

References Time::dDeltat\_n, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nCenIntOffset, Grid::nCotThetaIJK, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nEndGhostUpdateExplicit,

Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, Grid::nSinThetaIJK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0, Grid::nV, and Grid::nW.

Referenced by calNewVelocities RTP().

# 7.12.2.61 void calNewW\_RTP\_LES (Grid & grid, Parameters & parameters, Time & time, ProcTop & procTop)

This function calculates the phi velocity, and does it by only considering the radial, theta, and phi terms. It also includes the terms for including real viscosity, used in the LES.

#### Parameters:

- $\leftrightarrow$  grid contains the local grid, and will hold the newly updated theta velocities
- $\leftarrow$  parameters various parameters needed for the calculation
- ← time contains time information, e.g. time step, current time etc.
- $\leftarrow procTop$  contains information about the processor topology

## **Boundary Conditions**

assume theta and phi velocities are constant across surface

## **Boundary Conditions**

assume eddy viscosity is zero at surface

#### **Boundary Conditions**

assume upwind gradient is the same as centered gradient across surface

References Time::dDeltat n, Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Grid::nCenIntOffset, Grid::nCotThetaIJK, Grid::nD, Grid::nDenAve, Grid∷nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nEddyVisc, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGlobalGridPositionLocalGrid, Grid::nNumGhostCells, Grid::nP, Grid::nQ0, Grid∷nQ1, Grid::nQ2, Grid∷nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0, Grid::nV, and Grid::nW.

Referenced by calNewVelocities RTP LES().

### 7.12.2.62 void calOldDenave None (Grid & grid)

This function is a dumby function, and doesn't do anything. In the case of a 1D calculation the average density is undefined, and only the density is used. This is different from the case where the 1D region exsists on the rank 0 processor, but the grid as a whole is really 2D or 3D. In which case calOldDenave R should be used instead.

### 7.12.2.63 void calOldDenave R (Grid & grid)

This function does nothing as the averaged density is not needed in 1D calculations.

#### Parameters:

 $\leftrightarrow$  *grid* supplies the information needed to calculate the horizontal density average, it also stores the calculated horizontally averaged density.

 $\label{lem:condition} References \ Grid::dLocalGridOld, \ Grid::nD, \ Grid::nDenAve, \ Grid::nEndGhostUpdateExplicit, \ Grid::nEndUpdateExplicit, \ Grid::nEndUpdateExplicit, \ Grid::nEndUpdateExplicit, \ Grid::nStartGhostUpdateExplicit, \ Grid::nStartGhostUpdateImplicit, \ Grid::nStartUpdateExplicit, \ and \ Grid::nStartUpdateImplicit.$ 

Referenced by initInternalVars().

## 7.12.2.64 void calOldDenave RT (Grid & grid)

This function calculates the horizontal average density in a 2D region. This function differs from calNewDenave\_RT in that it calculates the average density from the old grid density and stores the result in the old grid. While calNewDenave\_RT calculates the average density from the new grid density and places the result in the new grid.

#### Parameters:

 $\leftrightarrow$  grid supplies the information needed to calculate the horizontal density average, it also stores the calculated horizontally averaged density.

References Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nD, Grid::nDCosThetaIJK, Grid::nDenAve, Grid::nEndGhostUpdateExplicit, Grid::nEndGhostUpdateImplicit, Grid::nEndUpdateExplicit, Grid::nEndUpdateImplicit, Grid::nStartGhostUpdateExplicit, Grid::nStartGhostUpdateImplicit, Grid::nStartUpdateExplicit, and Grid::nStartUpdateImplicit.

Referenced by initInternalVars().

#### 7.12.2.65 void calOldDenave RTP (Grid & grid)

This function calculates the horizontal average density in a 3D region. This function differs from calNewDenave\_RTP in that it calculates the average density from the old grid density and stores the result in the old grid. While calNewDenave\_RTP calculates the average density from the new grid density and places the result in the new grid.

## Parameters:

 $\leftrightarrow$  grid supplies the information needed to calculate the horizontal density average, it also stores the calculated horizontally averaged density.

Referenced by initInternalVars().

### 7.12.2.66 void calOldEddyVisc R CN (Grid & grid, Parameters & parameters)

Calculates the eddy viscosity using a constant times the zoning including only the radial terms. It puts the result into the old grid. This function is used to initalize the eddy viscosity when the

code begins execution.

 $\label{lem:convective} References & Parameters:: dEddy Viscosity, & Grid:: dLocalGridNew, & Grid:: dLocalGridOld, \\ Parameters:: dMaxConvectiveVelocity, & Grid:: nCenIntOffset, & Grid:: nEddy Visc, \\ Grid:: nEndGhost Update Explicit, & Grid:: nEndUpdate Explicit, & Grid:: nNumGhost Cells, & Grid:: nR, \\ Grid:: nStartGhost Update Explicit, & and & Grid:: nStartUpdate Explicit. \\ \end{cases}$ 

Referenced by initInternalVars().

## 7.12.2.67 void calOldEddyVisc R SM (Grid & grid, Parameters & parameters)

Calculates the eddy viscosity including only the radial terms. It puts the result into the old grid. This function is used to initalize the eddy viscosity when the code begins execution. It uses the Smagorinsky model for calculating the eddy viscosity.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the eddy viscosity.
- ← parameters contains parameters used in calculating the eddy viscosity.

 $References \quad Parameters:: dEddy Viscosity, \quad Grid:: dLocal Grid Old, \quad Grid:: nCenInt Offset, \\ Grid:: nD, \quad Grid:: nEddy Visc, \quad Grid:: nEndGhost Update Explicit, \quad Grid:: nEndUpdate Explicit, \\ Grid:: nNumGhost Cells, \quad Grid:: nR, \quad Grid:: nStart Ghost Update Explicit, \\ Grid:: nU.$ 

Referenced by initInternalVars().

## 7.12.2.68 void calOldEddyVisc RT CN (Grid & grid, Parameters & parameters)

Calculates the eddy viscosity using a constant times the zoning including only the radial and theta terms. It puts the result into the old grid. This function is used to initialize the eddy viscosity when the code begins execution.

#### Parameters:

- $\leftrightarrow$  *grid* supplies the input for calculating the eddy viscosity.
- ← parameters contains parameters used in calculating the eddy viscosity.

 $References \ Parameters:: dEddy Viscosity, \ Grid:: dLocalGridNew, \ Grid:: dLocalGridOld, \ Parameters:: dMaxConvectiveVelocity, \ Grid:: nCenIntOffset, \ Grid:: nDTheta, \ Grid:: nEddy Visc, \ Grid:: nEndGhostUpdateExplicit, \ Grid:: nEndUpdateExplicit, \ Grid:: nNumGhostCells, \ Grid:: nR, \ Grid:: nStartGhostUpdateExplicit, \ and \ Grid:: nStartUpdateExplicit.$ 

Referenced by initInternalVars().

## 7.12.2.69 void calOldEddyVisc RT SM (Grid & grid, Parameters & parameters)

Calculates the eddy viscosity including only the radial and theta terms. It puts the result into the old grid. This function is used to initalize the eddy viscosity when the code begins execution. It uses the Smagorinsky model for calculating the eddy viscosity.

#### Parameters:

 $\leftrightarrow$  grid supplies the input for calculating the eddy viscosity.

← parameters contains parameters used in calculating the eddy viscosity.

References Parameters::dEddyViscosity, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nD, Grid::nDTheta, Grid::nEddyVisc, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nNumGhostCells, Grid::nR, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU, and Grid::nV.

Referenced by initInternalVars().

# 7.12.2.70 void calOldEddyVisc\_RTP\_CN (Grid & grid, Parameters & parameters)

Calculates the eddy viscosity using a constant times the zoning including the radial, theta, and phi terms. It puts the result into the old grid. This function is used to initalize the eddy viscosity when the code begins execution.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the eddy viscosity.
- $\leftarrow$  parameters contains parameters used in calculating the eddy viscosity.

 $\label{lem:reconstruction} References & Parameters:: dEddy Viscosity, & Grid:: dLocalGridNew, & Grid:: dLocalGridOld, & Parameters:: dMaxConvectiveVelocity, & Grid:: nCenIntOffset, & Grid:: nDPhi, & Grid:: nDTheta, & Grid:: nEddy Visc, & Grid:: nEndGhostUpdateExplicit, & Grid:: nEndUpdateExplicit, & Grid:: nStartGhostUpdateExplicit, & Grid:: nStartGhostUpdateExplicit, & Grid:: nStartUpdateExplicit.$ 

Referenced by initInternalVars().

## 7.12.2.71 void calOldEddyVisc\_RTP\_SM (Grid & grid, Parameters & parameters)

Calculates the eddy viscosity including the radial, theta, and phi terms. It puts the result into the old grid. This function is used to initalize the eddy viscosity when the code begins execution. It uses the Smagorinsky model for calculating the eddy viscosity.

#### Parameters:

- $\leftrightarrow$  *grid* supplies the input for calculating the eddy viscosity.
- ← parameters contains parameters used in calculating the eddy viscosity.

## **Boundary Conditions**

assuming that theta velocity is constant across surface

#### **Boundary Conditions**

assume phi velocity is constant across surface

References Parameters::dEddyViscosity, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nCotThetaIJK, Grid::nD, Grid::nDPhi, Grid::nDTheta, Grid::nEddyVisc, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nNumGhostCells, Grid::nR, Grid::nSinThetaIJK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU, Grid::nV, and Grid::nW.

Referenced by initInternalVars().

## 7.12.2.72 void calOldP GL (Grid & grid, Parameters & parameters)

This function calculates the pressure using a gamma law gas, calculate by dEOS\_GL.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the pressure and also accepts the results of the pressure calculations
- $\leftarrow$  parameters contains parameters used in calculating the pressure, namely the value of the adiabatic gamma

References dEOS\_GL(), Grid::dLocalGridOld, Grid::nD, Grid::nE, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nStartGhostUpdateExplicit, and Grid::nStartUpdateExplicit. Grid::nP,

Referenced by initInternalVars().

# 7.12.2.73 void calOldPEKappaGamma\_TEOS (Grid & grid, Parameters & parameters)

This function calculates the pressure, energy, opacity, and adiabatic index of a cell. It calculates it using the old vaules of quantities and places the result in the old grid. This function is used to initialize the internal variables pressure, energy and kappa, and is suitable for both 1D and 3D calculations.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the pressure and also accepts the result of the pressure calculation
- ← parameters contains parameters used in calculating the pressure.

References Grid::dLocalGridOld, Parameters::eosTable, eos::getPEKappaGamma(), Grid::nEndGhostUpdateExplicit, Grid::nEndGhostUpdateImplicit, Grid::nD, Grid∷nE, Grid::nEndUpdateExplicit, Grid::nEndUpdateImplicit, Grid::nGamma, Grid::nKappa, Grid::nNumGhostCells, Grid::nP, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nStartUpdateImplicit, Grid::nStartGhostUpdateImplicit, and Grid::nT.

Referenced by initInternalVars().

## 7.12.2.74 void calOldQ0 R GL (Grid & grid, Parameters & parameters)

This function calculates the artificial viscosity of a cell. It calculates it using the old vaules of quantities and places the result in the old grid. It does this for the radial component of the viscosity only. This function is used when using a gamma law gas equation of state.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the artificial viscosity and also accepts the result of the artificial viscosity calculation
- ← parameters contains parameters used in calculating the artificial viscosity.

References Parameters::dA, Parameters::dAVThreshold, Parameters::dGamma, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nD, Grid::nEndGhostUpdateExplicit,

Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0, Grid::nR, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, and Grid::nU.

Referenced by initInternalVars().

## 7.12.2.75 void calOldQ0 R TEOS (Grid & grid, Parameters & parameters)

This function calculates the artificial viscosity of a cell. It calculates it using the old vaules of quantities and places the result in the old grid. It does this for 1D viscosity only.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the pressure and also accepts the result of the pressure calculation
- ← parameters contains parameters used in calculating the artificial viscosity.

References Parameters::dA, Parameters::dAVThreshold, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nD, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGamma, Grid::nP, Grid::nQ0, Grid::nR, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, and Grid::nU.

Referenced by initInternalVars().

## 7.12.2.76 void calOldQ0Q1 RT GL (Grid & grid, Parameters & parameters)

This function calculates the artificial viscosity of a cell. It calculates it using the old vaules of quantities and places the result in the old grid. It does this for the two components of the viscosity. This function is used when using a gamma law gas equation of state.

## Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the artificial viscosity and also accepts the result of the artificial viscosity calculation
- ← parameters contains parameters used in calculating the artificial viscosity.

 $\label{lem:references} References & Parameters::dA, & Parameters::dAVThreshold, & Parameters::dGamma, Grid::dLocalGridOld, & Grid::nCenIntOffset, & Grid::nD, & Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, & Grid::nP, & Grid::nQ1, & Grid::nR, & Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, & Grid::nStartGhostUpdateExplicit, & Grid::nStartUpdateExplicit, Grid::nU, and & Grid::nV.$ 

Referenced by initInternalVars().

## 7.12.2.77 void calOldQ0Q1 RT TEOS (Grid & grid, Parameters & parameters)

This function calculates the artificial viscosity of a cell. It calculates it using the old vaules of quantities and places the result in the old grid. It does this for two components of the viscosity. This function is used when using a tabulated equation of state.

#### Parameters:

 $\leftrightarrow$  grid supplies the input for calculating the artificial viscosity and also accepts the result of the artificial viscosity calculation

← parameters contains parameters used in calculating the artificial viscosity.

References Parameters::dA, Parameters::dAVThreshold, Grid::dLocalGridOld, Grid::nEndUpdateExplicit, Grid::nCenIntOffset, Grid::nD, Grid::nEndGhostUpdateExplicit, Grid::nGamma, Grid::nP, Grid∷nQ0, Grid::nQ1, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, and Grid::nV.

Referenced by initInternalVars().

## 7.12.2.78 void calOldQ0Q1Q2 RTP GL (Grid & grid, Parameters & parameters)

This function calculates the artificial viscosity of a cell. It calculates it using the old vaules of quantities and places the result in the old grid. It does this for the three components of the viscosity. This function is used when using a gamma law gas equation of state.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the artificial viscosity and also accepts the result of the artificial viscosity calculation
- ← parameters contains parameters used in calculating the artificial viscosity.

References Parameters::dA, Parameters::dAVThreshold, Parameters::dGamma, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid∷nD, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0, Grid∷nQ1, Grid::nQ2, Grid::nR, Grid::nSinThetaIJp1halfK, Grid::nSinThetaIJK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nV, and Grid::nW.

Referenced by initInternalVars().

# 7.12.2.79 void calOldQ0Q1Q2\_RTP\_TEOS (Grid & grid, Parameters & parameters)

This function calculates the artificial viscosity of a cell. It calculates it using the old vaules of quantities and places the result in the old grid. It does this for the three components of the viscosity. This function is used when using a tabulated equation of state.

#### Parameters:

- $\leftrightarrow$  grid supplies the input for calculating the artificial viscosity and also accepts the result of the artificial viscosity calculation
- ← parameters contains parameters used in calculating the artificial viscosity.

References Parameters::dA, Parameters::dAVThreshold, Grid::dLocalGridOld, Grid::nCenIntOffset, Grid::nD, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGamma, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nStartGhostUpdateExplicit, Grid::nStartUpdateExplicit, Grid::nU, Grid::nV, and Grid::nW.

Referenced by initInternalVars().

#### 7.12.2.80 double dEOS GL (double dRho, double dE, Parameters parameters)

Calculates the pressure from the energy and density using a  $\gamma$ -law gas.

#### Parameters:

- $\leftarrow dRho$  the density of a cell
- $\leftarrow dE$  the energy of a cell
- $\leftarrow$  parameters contians various parameters, including  $\gamma$  needed to calculate the pressure.

#### Returns:

the pressure

This version of dEOS\_GL uses the same value of  $\gamma$  through out the model. The equation of state is given by  $\rho(\gamma - 1)E$ .

References Parameters::dGamma.

Referenced by calNewP GL(), and calOldP GL().

# 7.12.2.81 double dImplicitEnergyFunction\_None (Grid & grid, Parameters & parameters, Time & time, double dTemps[], int i, int j, int k)

This is an empty function, that isn't even called when no implicit solution is needed. This safe guards against future addition which may need to call an empty function when no implicit solve is being done.

Referenced by setMainFunctions().

# 7.12.2.82 double dImplicitEnergyFunction\_R (Grid & grid, Parameters & parameters, Time & time, double dTemps[], int i, int j, int k)

This function is used to determine the agreement of the updated values at n+1, with each other in the non-adiabatic energy equation. The \_R version of the function contains only the radial terms, and should be used for purely radial calculations. This function can also be used for calculating numerical deriviatives by varying the input temperatures.

#### Parameters:

- $\leftarrow grid$
- $\leftarrow parameters$
- $\leftarrow time$
- $\leftarrow$  dTemps dTemps[0]=dT\_ijk\_np1 is the temperature at radial position (i,j,k) and time n+1,dTemps[1]=dT\_ip1jk\_np1 is the temperature at radial position (i+1,j,k) and time n+1, dTemps[2]=dT\_im1jk\_np1 is the temperature at radial position (i-1,j,k) and time n+1.
- $\leftarrow i$  is the radial index to evaluate the function at.
- $\leftarrow j$  is the theta index to evaluate the function at.
- $\leftarrow k$  is the phi index to evaluate the function at.

References Time::dDeltat np1half, DEBUG EQUATIONS, eos::dGetEnergy(), eos::dGetOpacity(), eos::dGetPressure(), Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dSigma, Parameters::dPi, Parameters::eosTable, Grid::nCenIntOffset, Grid∷nD. Grid::nDenAve, Grid::nDonorCellFrac, Grid∷nDM, Grid::nE. Grid::nGlobalGridPositionLocalGrid, Grid::nNumGhostCells, Grid::nQ0, Grid::nR, Grid::nT, Grid::nU, and Grid::nU0.

Referenced by setMainFunctions().

# 7.12.2.83 double dImplicitEnergyFunction\_R\_LES (Grid & grid, Parameters & parameters, Time & time, double $dTemps[\ ]$ , int i, int j, int k)

This function is used to determine the agreement of the updated values at n+1, with each other in the non-adiabatic energy equation. The \_R version of the function contains only the radial terms, and should be used for purely radial calculations. This function can also be used for calculating numerical derivitatives by varying the input temperatures.

#### Parameters:

- $\leftarrow grid$
- $\leftarrow$  parameters
- $\leftarrow time$
- $\leftarrow$  **dTemps** dTemps[0]=dT\_ijk\_np1 is the temperature at radial position (i, j, k) and time n+1,dTemps[1]=dT\_ip1jk\_np1 is the temperature at radial position (i+1, j, k) and time n+1, dTemps[2]=dT\_im1jk\_np1 is the temperature at radial position (i-1, j, k) and time n+1.
- $\leftarrow i$  is the radial index to evaluate the function at.
- $\leftarrow j$  is the theta index to evaluate the function at.
- $\leftarrow k$  is the phi index to evaluate the function at.

### Todo

this function should probably be turffed, the LES terms aren't needed in 1D. keeping it for now though.

References Time::dDeltat\_np1half, eos::dGetEnergy(), eos::dGetOpacity(), eos::dGetPressure(), Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dSigma, Parameters::eosTable, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nE, Grid::nEddyVisc, Grid::nGlobalGridPositionLocalGrid, Grid::nNumGhostCells, Grid::nQ0, Grid::nR, Grid::nT, Grid::nU, and Grid::nU0.

# 7.12.2.84 double dImplicitEnergyFunction\_R\_LES\_SB (Grid & grid, Parameters & parameters, Time & time, double $dTemps[\ ]$ , int i, int j, int k)

This function is used to determine the agreement of the updated values at n+1, with each other in the non-adiabatic energy equation. The \_R version of the function contains only the radial terms, and should be used for purely radial calculations. This function can also be used for calculating numerical derivitatives by varying the input temperatures. This function differs from the version without the "\_SB" suffix (dImplicitEnergyFunction\_R)in that it is tailored to the surface boundary region.

## Parameters:

- $\leftarrow \textit{grid}$
- $\leftarrow$  parameters
- $\leftarrow time$
- $\leftarrow$  **dTemps** dTemps[0]=dT\_ijk\_np1 is the temperature at radial position (i, j, k) and time n+1 and time n+1, dTemps[1]=dT\_im1jk\_np1 is the temperature at radial position (i-1, j, k) and time n+1.
- $\leftarrow i$  is the radial index to evaluate the function at.

- $\leftarrow j$  is the theta index to evaluate the function at.
- $\leftarrow k$  is the phi index to evaluate the function at.

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nE][i+1][j][k] in calculation of  $E_{i+1/2,j,k}$  setting it equal to value at i.

## **Boundary Conditions**

grid.dLocalGridOld[grid.nDM][i+1][0][0] and grid.dLocalGridOld[grid.nE][i+1][j][k] missing in the calculation of upwind gradient in dA1. Using the centered gradient instead.

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nT][i+1][0][0]

References Time::dDeltat\_np1half, eos::dGetEnergy(), eos::dGetOpacity(), eos::dGetPressure(), Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dSigma, Parameters::eosTable, Grid::nCenIntOffset, Grid::nD, Grid::nDM, Grid::nDonorCellFrac, Grid::nE, Grid::nEddyVisc, Grid::nQ0, Grid::nR, Grid::nT, Grid::nU, and Grid::nU0.

# 7.12.2.85 double dImplicitEnergyFunction R SB (Grid & grid, Parameters & parameters, Time & time, double dTemps[], int i, int j, int k)

This function is used to determine the agreement of the updated values at n+1, with each other in the non-adiabatic energy equation. The \_R version of the function contains only the radial terms, and should be used for purely radial calculations. This function can also be used for calculating numerical derivitatives by varying the input temperatures. This function differs from the version without the "\_SB" suffix (dImplicitEnergyFunction\_R)in that it is tailored to the surface boundary region.

## Parameters:

- $\leftarrow qrid$
- $\leftarrow parameters$
- $\leftarrow time$
- $\leftarrow$  dTemps dTemps[0]=dT\_ijk\_np1 is the temperature at radial position (i, j, k) and time n+1 and time n+1, dTemps[1]=dT\_im1jk\_np1 is the temperature at radial position (i-1, j, k) and time n+1.
- $\leftarrow i$  is the radial index to evaluate the function at.
- $\leftarrow j$  is the theta index to evaluate the function at.
- $\leftarrow k$  is the phi index to evaluate the function at.

## **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nE][i+1][j][k] in calculation of  $E_{i+1/2,j,k}$  setting it equal to value at i.

## **Boundary Conditions**

All upwind set to zero as no material is flowing into the star

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nT][i+1][0][0]

References Time::dDeltat\_np1half, eos::dGetEnergy(), eos::dGetOpacity(), eos::dGetPressure(), Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dSigma, Parameters::eosTable, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nE, Grid::nGlobalGridPositionLocalGrid, Grid::nNumGhostCells, Grid::nQ0, Grid::nR, Grid::nT, Grid::nU, and Grid::nU0.

Referenced by setMainFunctions().

# 7.12.2.86 double dImplicitEnergyFunction\_RT (Grid & grid, Parameters & parameters, Time & time, double dTemps[], int i, int j, int k)

This function is used to determine the agreement of the updated values at n+1, with each other in the non-adiabatic energy equation. The \_RT version of the function contains only the radial and theta terms, and should be used for radial-theta calculations. This function can also be used for calculating numerical derivitives by varying the input temperatures.

### Parameters:

- $\leftarrow grid$
- $\leftarrow$  parameters
- $\leftarrow time$
- ← dTemps dTemps[0]=dT\_ijk\_np1 is the temperature at radial position (i, j, k) and time n+1, dTemps[1]=dT\_ip1jk\_np1 is the temperature at radial position (i+1, j, k) and time n+1, dTemps[2]=dT\_im1jk\_np1 is the temperature at radial position (i-1, j, k) and time n+1, dTemps[3]=dT\_ijp1k\_np1 is the temperature at radial position (i, j+1, k) and time n+1, dTemps[4]=dT\_ijm1k\_np1 is the temperature at radial position (i, j-1, k) and time n+1.
- $\leftarrow i$  is the radial index to evaluate the function at.
- $\leftarrow j$  is the theta index to evaluate the function at.
- $\leftarrow k$  is the phi index to evaluate the function at.

References Time::dDeltat\_np1half, eos::dGetEnergy(), eos::dGetOpacity(), eos::dGetPressure(), Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dSigma, Parameters::eosTable, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDTheta, Grid::nE, Grid::nQ0, Grid::nQ1, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJF1halfK, Grid::nT, Grid::nU, Grid::nU0, and Grid::nV.

Referenced by setMainFunctions().

# 7.12.2.87 double dImplicitEnergyFunction RT\_LES (Grid & grid, Parameters & parameters, Time & time, double dTemps[], int i, int j, int k)

This function is used to determine the agreement of the updated values at n+1, with each other in the non-adiabatic energy equation. The \_RT version of the function contains only the radial and theta terms, and should be used for radial-theta calculations. This function can also be used for calculating numerical deriviatives by varying the input temperatures.

### Parameters:

 $\leftarrow grid$ 

- $\leftarrow$  parameters
- $\leftarrow time$
- ← dTemps dTemps[0]=dT\_ijk\_np1 is the temperature at radial position (i, j, k) and time n+1, dTemps[1]=dT\_ip1jk\_np1 is the temperature at radial position (i+1, j, k) and time n+1, dTemps[2]=dT\_im1jk\_np1 is the temperature at radial position (i-1, j, k) and time n+1, dTemps[3]=dT\_ijp1k\_np1 is the temperature at radial position (i, j+1, k) and time n+1, dTemps[4]=dT\_ijm1k\_np1 is the temperature at radial position (i, j-1, k) and time n+1.
- $\leftarrow i$  is the radial index to evaluate the function at.
- $\leftarrow j$  is the theta index to evaluate the function at.
- $\leftarrow k$  is the phi index to evaluate the function at.

```
\begin{array}{lll} 5700 & CTEOS & if (grid.dLocalGridOld[grid.nM][nIInt][0][0]>=1.143732280336595e+33) \{ \\ dA1UpWindGrad=-1.796596699553508e-14; & dA1CenGrad=-1.796596699553508e-14; \\ 6100 & CTEOS & if (grid.dLocalGridOld[grid.nM][nIInt][0][0]>=1.143732392703405e+33) \{ \\ dA1UpWindGrad=-2.381754669392478e-14; & dA1CenGrad=-2.381754669392478e-14; \\ T6500 & CTEOS & if (grid.dLocalGridOld[grid.nM][nIInt][0][0]>=1.143732445236012e+33) \{ \\ dA1UpWindGrad=-3.795837002744412e-14; \\ dA1CenGrad=-3.795837002744412e-14; \} \\ \end{array}
```

References Time::dDeltat\_np1half, eos::dGetEnergy(), eos::dGetOpacity(), eos::dGetPressure(), Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dPrt, Parameters::dSigma, Parameters::eosTable, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDTheta, Grid::nE, Grid::nEddyVisc, Grid::nM, Grid::nQ0, Grid::nQ1, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nT, Grid::nU, Grid::nU0, and Grid::nV.

Referenced by setMainFunctions().

# 7.12.2.88 double dImplicitEnergyFunction\_RT\_LES\_SB (Grid & grid, Parameters & parameters, Time & time, double $dTemps[\ ]$ , int i, int j, int k)

This function is used to determine the agreement of the updated values at n+1, with each other in the non-adiabatic energy equation. The \_RT version of the function contains only the radial and theta terms, and should be used for radial-theta calculations. This function can also be used for calculating numerical deriviatives by varying the input temperatures. This function differs from the version without the "\_SB" suffix (dImplicitEnergyFunction\_RT)in that it is tailored to the surface boundary region.

### Parameters:

- $\leftarrow grid$
- $\leftarrow$  parameters
- $\leftarrow time$
- ← dTemps dTemps[0]=dT\_ijk\_np1 is the temperature at radial position (i, j, k) and time n+1, dTemps[1]=dT\_ip1jk\_np1 is the temperature at radial position (i+1, j, k) and time n+1, dTemps[2]=dT\_im1jk\_np1 is the temperature at radial position (i-1, j, k) and time n+1, dTemps[3]=dT\_ijp1k\_np1 is the temperature at radial position (i, j+1, k) and time n+1, dTemps[4]=dT\_ijm1k\_np1 is the temperature at radial position (i, j-1, k) and time n+1.
- $\leftarrow i$  is the radial index to evaluate the function at.

- $\leftarrow j$  is the theta index to evaluate the function at.
- $\leftarrow k$  is the phi index to evaluate the function at.

### **Boundary Conditions**

Missing  $\Delta M_r$  outside model using Parameters::dAlpha times  $\Delta M_r$  in the last zone instead.

### **Boundary Conditions**

missing density outside model assuming it is zero

### **Boundary Conditions**

missing desnity outside model assuming it is zero

### **Boundary Conditions**

assuming V at ip1half is the same as V at i

### **Boundary Conditions**

Assuming energy outside model is the same as the energy in the last zone inside the model.

### **Boundary Conditions**

Assuming energy outside model is the same as the energy in the last zone inside the model.

### **Boundary Conditions**

Using centered gradient for upwind gradient when motion is into the star at the surface

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nT][i+1][0][0] using flux equals  $2\sigma T^4$  at surface.

References Parameters::dAlpha, Parameters::dAlphaExtra, Time::dDeltat\_np1half, eos::dGetEnergy(), eos::dGetOpacity(), eos::dGetPressure(), Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dPrt, Parameters::dSigma, Parameters::eosTable, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDTheta, Grid::nE, Grid::nEddyVisc, Grid::nQ0, Grid::nQ1, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJF1halfK, Grid::nT, Grid::nU, Grid::nU0, and Grid::nV.

Referenced by setMainFunctions().

# 7.12.2.89 double dImplicitEnergyFunction\_RT\_SB (Grid & grid, Parameters & parameters, Time & time, double $dTemps[\ ]$ , int i, int j, int k)

This function is used to determine the agreement of the updated values at n+1, with each other in the non-adiabatic energy equation. The \_RT version of the function contains only the radial and theta terms, and should be used for radial-theta calculations. This function can also be used for calculating numerical deriviatives by varying the input temperatures. This function differs from the version without the "\_SB" suffix (dImplicitEnergyFunction\_RT)in that it is tailored to the surface boundary region.

### Parameters:

 $\leftarrow grid$ 

- $\leftarrow parameters$
- $\leftarrow time$
- ← dTemps dTemps[0]=dT\_ijk\_np1 is the temperature at radial position (i, j, k) and time n+1, dTemps[1]=dT\_ip1jk\_np1 is the temperature at radial position (i+1, j, k) and time n+1, dTemps[2]=dT\_im1jk\_np1 is the temperature at radial position (i-1, j, k) and time n+1, dTemps[3]=dT\_ijp1k\_np1 is the temperature at radial position (i, j+1, k) and time n+1, dTemps[4]=dT\_ijm1k\_np1 is the temperature at radial position (i, j-1, k) and time n+1.
- $\leftarrow i$  is the radial index to evaluate the function at.
- $\leftarrow j$  is the theta index to evaluate the function at.
- $\leftarrow k$  is the phi index to evaluate the function at.

### **Boundary Conditions**

Using centered gradient for upwind gradient when motion is into the star at the surface

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nT][i+1][0][0] using flux equals  $2\sigma T^4$  at surface.

References Time::dDeltat\_np1half, eos::dGetEnergy(), eos::dGetOpacity(), eos::dGetPressure(), Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dSigma, Parameters::eosTable, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDTheta, Grid::nE, Grid::nQ0, Grid::nQ1, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJF1halfK, Grid::nT, Grid::nU, Grid::nU0, and Grid::nV.

Referenced by setMainFunctions().

## 7.12.2.90 double dImplicitEnergyFunction RTP (Grid & grid, Parameters & parameters, Time & time, double $dTemps[\ ]$ , int i, int j, int k)

This function is used to determine the agreement of the updated values at n+1, with each other in the non-adiabatic energy equation. The \_RTP version of the function contains terms for all three directions, and should be used for calculations involving all three directions. This function can also be used for calculating numerical deriviatives by varying the input temperatures. This function differs from the version without the "\_SB" suffix (dImplicitEnergyFunction\_RT)in that it is tailored to the surface boundary region.

### Parameters:

- $\leftarrow grid$
- $\leftarrow$  parameters
- $\leftarrow time$
- ←  $dTemps, dTemps[0] = dT_ijk_np1$  is the temperature at radial position (i, j, k) and time n+1,  $dTemps[1] = dT_ip1jk_np1$  is the temperature at radial position (i+1, j, k) and time n+1,  $dTemps[2] = dT_im1jk_np1$  is the temperature at radial position (i-1, j, k) and time n+1,  $dTemps[3] = dT_ip1k_np1$  is the temperature at radial position (i, j+1, k) and time n+1,  $dTemps[4] = dT_ip1k_np1$  is the temperature at radial position (i, j-1, k) and time n+1,  $dTemps[5] = dT_ipkp1_np1$  is the temperature at radial position (i, j, k+1) and time n+1,  $dTemps[6] = dT_ipkp1_np1$  is the temperature at radial position (i, j, k-1) and time n+1.
- $\leftarrow i$  is the radial index to evaluate the function at.

- $\leftarrow j$  is the theta index to evaluate the function at.
- $\leftarrow k$  is the phi index to evaluate the function at.

References Time::dDeltat\_np1half, eos::dGetEnergy(), eos::dGetOpacity(), eos::dGetPressure(), Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dSigma, Parameters::eosTable, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nE, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nT, Grid::nU, Grid::nU, Grid::nV, and Grid::nW.

Referenced by setMainFunctions().

# 7.12.2.91 double dImplicitEnergyFunction\_RTP\_LES (Grid & grid, Parameters & parameters, Time & time, double dTemps[], int i, int j, int k)

This function is used to determine the agreement of the updated values at n+1, with each other in the non-adiabatic energy equation. The \_RTP version of the function contains terms for all three directions, and should be used for calculations involving all three directions. This function can also be used for calculating numerical deriviatives by varying the input temperatures. This function differs from the version without the "\_SB" suffix (dImplicitEnergyFunction\_RT)in that it is tailored to the surface boundary region.

#### Parameters:

- $\leftarrow grid$
- $\leftarrow$  parameters
- $\leftarrow time$
- ←  $dTemps, dTemps[0] = dT_ijk_np1$  is the temperature at radial position (i, j, k) and time n+1,  $dTemps[1] = dT_ip1jk_np1$  is the temperature at radial position (i+1, j, k) and time n+1,  $dTemps[2] = dT_im1jk_np1$  is the temperature at radial position (i-1, j, k) and time n+1,  $dTemps[3] = dT_ip1k_np1$  is the temperature at radial position (i, j+1, k) and time n+1,  $dTemps[4] = dT_ip1k_np1$  is the temperature at radial position (i, j-1, k) and time n+1,  $dTemps[5] = dT_ipkp1_np1$  is the temperature at radial position (i, j, k+1) and time n+1,  $dTemps[6] = dT_ipkp1_np1$  is the temperature at radial position (i, j, k-1) and time n+1.
- $\leftarrow i$  is the radial index to evaluate the function at.
- $\leftarrow i$  is the theta index to evaluate the function at.
- $\leftarrow k$  is the phi index to evaluate the function at.

References Time::dDeltat np1half, DEBUG EQUATIONS, eos::dGetEnergy(), eos::dGetOpacity(), eos::dGetPressure(), Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dPrt, Parameters::dSigma, Parameters::eosTable, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nE, Grid::nEddyVisc, Grid::nGlobalGridPositionLocalGrid, Grid::nM, Grid::nNumGhostCells, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nT, Grid::nU, Grid::nU0, Grid::nV, and Grid::nW.

Referenced by setMainFunctions().

# 7.12.2.92 double dImplicitEnergyFunction\_RTP\_LES\_SB (Grid & grid, Parameters & parameters, Time & time, double $dTemps[\ ]$ , int i, int j, int k)

This function is used to determine the agreement of the updated values at n+1, with each other in the non-adiabatic energy equation. The \_RTP version of the function contains terms for all three directions, and should be used for calculations involving all three directions. This function can also be used for calculating numerical deriviatives by varying the input temperatures. This function differs from the version without the "\_SB" suffix (dImplicitEnergyFunction\_RT)in that it is tailored to the surface boundary region.

#### Parameters:

- $\leftarrow grid$
- $\leftarrow$  parameters
- $\leftarrow time$
- ← dTemps dTemps[0]=dT\_ijk\_np1 is the temperature at radial position (i, j, k) and time n+1, dTemps[1]=dT\_ip1jk\_np1 is the temperature at radial position (i+1, j, k) and time n+1, dTemps[2]=dT\_im1jk\_np1 is the temperature at radial position (i-1, j, k) and time n+1, dTemps[3]=dT\_ijp1k\_np1 is the temperature at radial position (i, j+1, k) and time n+1, dTemps[4]=dT\_ijm1k\_np1 is the temperature at radial position (i, j-1, k) and time n+1, dTemps[5]=dT\_ijkp1\_np1 is the temperature at radial position (i, j, k+1) and time n+1, dTemps[6]=dT\_ijkm1\_np1 is the temperature at radial position (i, j, k-1) and time n+1.
- $\leftarrow i$  is the radial index to evaluate the function at.
- $\leftarrow j$  is the theta index to evaluate the function at.
- $\leftarrow k$  is the phi index to evaluate the function at.

### **Boundary Conditions**

Missing  $\Delta M_r$  outside model using Parameters::dAlpha times  $\Delta M_r$  in the last zone instead.

### **Boundary Conditions**

missing density outside model assuming it is zero

### **Boundary Conditions**

missing desnity outside model assuming it is zero

### **Boundary Conditions**

assuming V at ip1half is the same as V at i

### **Boundary Conditions**

assuming W at ip1half is the same as W at i

### **Boundary Conditions**

Assuming energy outside model is the same as the energy in the last zone inside the model.

### **Boundary Conditions**

Assuming energy outside model is the same as the energy in the last zone inside the model.

### **Boundary Conditions**

A1 upwind set to zero as no material is flowing into the star

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nT][i+1][0][0] using flux equals  $2\sigma T^4$  at surface.

References Parameters::dAlpha, Parameters::dAlphaExtra, Time::dDeltat np1half, DEBUG EQUATIONS, eos::dGetEnergy(), eos::dGetOpacity(), eos::dGetPressure(), Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dPrt. rameters::dSigma, Parameters::eosTable, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nE, Grid::nEddyVisc, Grid::nGlobalGridPositionLocalGrid, Grid::nM, Grid::nNumGhostCells, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nT, Grid::nU, Grid::nU0, Grid::nV, and Grid::nW.

Referenced by setMainFunctions().

# 7.12.2.93 double dImplicitEnergyFunction\_RTP\_SB (Grid & grid, Parameters & parameters, Time & time, double dTemps[], int i, int j, int k)

This function is used to determine the agreement of the updated values at n+1, with each other in the non-adiabatic energy equation. The \_RTP version of the function contains terms for all three directions, and should be used for calculations involving all three directions. This function can also be used for calculating numerical deriviatives by varying the input temperatures. This function differs from the version without the "\_SB" suffix (dImplicitEnergyFunction\_RT)in that it is tailored to the surface boundary region.

### Parameters:

- $\leftarrow grid$
- $\leftarrow$  parameters
- $\leftarrow time$
- ← dTemps dTemps[0]=dT\_ijk\_np1 is the temperature at radial position (i, j, k) and time n+1, dTemps[1]=dT\_ip1jk\_np1 is the temperature at radial position (i+1, j, k) and time n+1, dTemps[2]=dT\_im1jk\_np1 is the temperature at radial position (i-1, j, k) and time n+1, dTemps[3]=dT\_ijp1k\_np1 is the temperature at radial position (i, j+1, k) and time n+1, dTemps[4]=dT\_ijm1k\_np1 is the temperature at radial position (i, j-1, k) and time n+1, dTemps[5]=dT\_ijkp1\_np1 is the temperature at radial position (i, j, k+1) and time n+1, dTemps[6]=dT\_ijkm1\_np1 is the temperature at radial position (i, j, k-1) and time n+1.
- $\leftarrow i$  is the radial index to evaluate the function at.
- $\leftarrow j$  is the theta index to evaluate the function at.
- $\leftarrow k$  is the phi index to evaluate the function at.

### **Boundary Conditions**

Using 
$$E_{i,j,k}^{n+1/2}$$
 for  $E_{i+1/2,j,k}^{n+1/2}$ 

### **Boundary Conditions**

Using centered gradient for upwind gradient when motion is into the star at the surface

### **Boundary Conditions**

Missing grid.dLocalGridOld[grid.nT][i+1][0][0] using flux equals  $2\sigma T^4$  at surface.

References Time::dDeltat\_np1half, eos::dGetEnergy(), eos::dGetOpacity(), eos::dGetPressure(), Grid::dLocalGridNew, Grid::dLocalGridOld, Parameters::dPi, Parameters::dSigma, Parameters::eosTable, Grid::nCenIntOffset, Grid::nD, Grid::nDenAve, Grid::nDM, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid::nE, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nR, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nT, Grid::nU, Grid::nU, Grid::nV, and Grid::nW.

Referenced by setMainFunctions().

# 7.12.2.94 void implicitSolve\_None (Grid & grid, Implicit & implicit, Parameters & parameters, Time & time, ProcTop & procTop, MessPass & messPass, Functions & functions)

This is an empty function, to be called when no implicit solution is needed. This allows the same code in the main program to be executed wheather or not an implicit solution is being preformed by setting the function pointer to this function if there is no implicit solution required.

Referenced by setMainFunctions().

# 7.12.2.95 void implicitSolve\_R (Grid & grid, Implicit & implicit, Parameters & parameters, Time & time, ProcTop & procTop, MessPass & messPass, Functions & functions)

This function solves for temperature corrections based on derivatives of the radial non-adiabatic energy equation with respect to the new temperature. It then uses these derivatives as entries in the coeffecient matrix. The discrepancy in the balance of the energy equation with the new temperature, energy, pressure, and opacity are included as the right hand side of the system of equaitons. Solving this system of equaitons provides the corrections needed for the new temperature. This processes is then repeated until the corrections are small. At this point the new temperature is used to update the energy, pressure, and opacity in the new grid via the equaiton of state.

Im-References calNewPEKappaGamma TEOS(), Implicit::dAverageRHS, plicit::dCurrentRelTError, Implicit::dDerivativeStepFraction, Grid::dLocalGridNew, plicit::dMaxErrorInRHS, Implicit::dTolerance, Functions::fpImplicitEnergyFunction, Functions::fpImplicitEnergyFunction SB, Implicit::kspContext, Implicit::matCoeff, Implicit::nLocDer, Implicit::nCurrentNumIterations, Implicit::nLocFun, plicit::nMaxNumIterations, Implicit::nMaxNumSolverIterations, Implicit::nNumDerPerRow, Implicit::nNumRowsALocal, Implicit::nNumRowsALocalSB, ProcTop::nRank, Grid∷nT, Time::nTimeStepIndex, updateLocalBoundariesNewGrid(), Implicit::nTypeDer, Implicit::vecRHS, Implicit::vecscatTCorrections, Implicit::vecTCorrections, and Implicit::vecTCorrectionsLocal.

Referenced by setMainFunctions().

# 7.12.2.96 void implicitSolve\_RT (Grid & grid, Implicit & implicit, Parameters & parameters, Time & time, ProcTop & procTop, MessPass & messPass, Functions & functions)

This function solves for temperature corrections based on derivatives of the radial-theta non-adiabatic energy equation with respect to the new temperature. It then uses these derivatives

as entries in the coeffecient matrix. The discrepancy in the balance of the energy equation with the new temperature, energy, pressure, and opacity are included as the right hand side of the system of equaitons. Solving this system of equaitons provides the corrections needed for the new temperature. This processes is then repeated until the corrections are small. At this point the new temperature is used to update the energy, pressure, and opacity in the new grid via the equaiton of state.

References calNewPEKappaGamma TEOS(), Implicit::dAverageRHS, Implicit::dCurrentRelTError, Implicit::dDerivativeStepFraction, Grid::dLocalGridNew, Implicit::dMaxErrorInRHS, Implicit::dTolerance, Functions::fpImplicitEnergyFunction, Functions::fpImplicitEnergyFunction SB, Implicit::kspContext, Implicit::matCoeff. Implicit::nCurrentNumIterations, Implicit::nLocDer, Implicit::nLocFun, plicit::nMaxNumIterations, Implicit::nMaxNumSolverIterations, Implicit::nNumDerPerRow, Implicit::nNumRowsALocalSB, ProcTop::nRank, Implicit::nNumRowsALocal, Grid∷nT, Time::nTimeStepIndex, Implicit::nTypeDer, updateLocalBoundariesNewGrid(), Implicit::vecRHS, Implicit::vecscatTCorrections, Implicit::vecTCorrections, Implicit::vecTCorrectionsLocal.

Referenced by setMainFunctions().

# 7.12.2.97 void implicitSolve\_RTP (Grid & grid, Implicit & implicit, Parameters & parameters, Time & time, ProcTop & procTop, MessPass & messPass, Functions & functions)

This function solves for temperature corrections based on derivatives of the radial-theta-phi non-adiabatic energy equation with respect to the new temperature. It then uses these derivatives as entries in the coeffecient matrix. The discrepancy in the balance of the energy equation with the new temperature, energy, pressure, and opacity are included as the right hand side of the system of equaitons. Solving this system of equaitons provides the corrections needed for the new temperature. This processes is then repeated until the corrections are small. At this point the new temperature is used to update the energy, pressure, and opacity in the new grid via the equaiton of state.

References calNewPEKappaGamma TEOS(), Implicit::dAverageRHS, Tmplicit::dCurrentRelTError, Implicit::dDerivativeStepFraction, Grid::dLocalGridNew, Implicit::dMaxErrorInRHS, Implicit::dTolerance, Functions::fpImplicitEnergyFunction, Functions::fpImplicitEnergyFunction SB, Implicit::kspContext, Implicit::matCoeff, Implicit::nCurrentNumIterations, Implicit::nLocFun, Implicit::nLocDer, Implicit::nMaxNumIterations, Implicit::nMaxNumSolverIterations, Implicit::nNumDerPerRow, ProcTop::nRank, Implicit::nNumRowsALocal, Implicit::nNumRowsALocalSB, Grid∷nT, Time::nTimeStepIndex, Implicit::nTypeDer, updateLocalBoundariesNewGrid(), Implicit::vecRHS, Implicit::vecscatTCorrections, Implicit::vecTCorrections, and Implicit::vecTCorrectionsLocal.

Referenced by setMainFunctions().

# $7.12.2.98 \quad \text{void initDonorFracAndMaxConVel\_R\_GL (Grid \& \textit{grid}, \text{ Parameters \& parameters)}}$

Initializes the donor fraction, and the maximum convective velocity when starting a calculation. The donor fraction is used to determine the amount of upwinded donor cell to use in advection terms. The maximum convective velocity is used for calculation of constant eddy viscosity parameter. This version of the fuction is for 1D, gamma law calculations.

References Parameters::dDonorCellMin, Parameters::dDonorCellMultiplier, Parameters

 $ters::dGamma,\ Grid::dLocalGridOld,\ Parameters::dMaxConvectiveVelocity,\ Grid::nCenIntOffset,\ Grid::nD,\ Grid::nDonorCellFrac,\ Grid::nEndGhostUpdateExplicit,\ Grid::nEndUpdateExplicit,\ Grid::nP,\ Grid::nQ0,\ Grid::nStartUpdateExplicit,\ Grid::nU,\ and\ Grid::nU0.$ 

Referenced by initInternalVars().

## 7.12.2.99 void initDonorFracAndMaxConVel\_R\_TEOS (Grid & grid, Parameters & parameters)

Initializes the donor fraction, and the maximum convective velocity when starting a calculation. The donor fraction is used to determine the amount of upwinded donor cell to use in advection terms. The maximum convective velocity is used for calculation of constant eddy viscosity parameter. This version of the fuction is for 1D, tabulated equation of state calculations.

References Parameters::dDonorCellMin, Parameters::dDonorCellMultiplier, Grid::dLocalGridOld, Parameters::dMaxConvectiveVelocity, Grid::nCenIntOffset, Grid::nD, Grid::nDonorCellFrac, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGamma, Grid::nP, Grid::nQ0, Grid::nStartUpdateExplicit, Grid::nU, and Grid::nU0.

Referenced by initInternalVars().

# 7.12.2.100 void initDonorFracAndMaxConVel\_RT\_GL (Grid & grid, Parameters & parameters)

Initializes the donor fraction, and the maximum convective velocity when starting a calculation. The donor fraction is used to determine the amount of upwinded donor cell to use in advection terms. The maximum convective velocity is used for calculation of constant eddy viscosity parameter. This version of the fuction is for 2D, gamma law calculations.

References Parameters:: dDonor CellMin, Parameters:: dDonor CellMultiplier, Parameters:: dGamma, Grid:: dLocalGridOld, Parameters:: dMaxConvectiveVelocity, Grid:: nCenIntOffset, Grid:: nD, Grid:: nDonor CellFrac, Grid:: nEndGhostUpdateExplicit, Grid:: nEndUpdateExplicit, Grid:: nP, Grid:: nQ0, Grid:: nQ1, Grid:: nStartUpdateExplicit, Grid:: nU, Grid:: nU0, and Grid:: nV.

Referenced by initInternalVars().

# $7.12.2.101 \quad \text{void initDonorFracAndMaxConVel\_RT\_TEOS (Grid \& \textit{grid}, \\ Parameters \& \textit{parameters})}$

Initializes the donor fraction, and the maximum convective velocity when starting a calculation. The donor fraction is used to determine the amount of upwinded donor cell to use in advection terms. The maximum convective velocity is used for calculation of constant eddy viscosity parameter. This version of the fuction is for 2D, tabulated equation of state calculations.

References Parameters::dDonorCellMin, Parameters::dDonorCellMultiplier, Grid::dLocalGridOld, Parameters::dMaxConvectiveVelocity, Grid::nCenIntOffset, Grid::nD, Grid::nDonorCellFrac, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGamma, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0, and Grid::nV.

Referenced by initInternalVars().

## 7.12.2.102 void initDonorFracAndMaxConVel\_RTP\_GL (Grid & grid, Parameters & parameters)

Initializes the donor fraction, and the maximum convective velocity when starting a calculation. The donor fraction is used to determine the amount of upwinded donor cell to use in advection terms. The maximum convective velocity is used for calculation of constant eddy viscosity parameter. This version of the fuction is for 3D, gamma law calculations.

References Parameters::dDonorCellMin, Parameters::dDonorCellMultiplier, Parameters::dGamma, Grid::dLocalGridOld, Parameters::dMaxConvectiveVelocity, Grid::nCenIntOffset, Grid::nD, Grid::nDonorCellFrac, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0, Grid::nV, and Grid::nW.

Referenced by initInternalVars().

## 7.12.2.103 void initDonorFracAndMaxConVel\_RTP\_TEOS (Grid & grid, Parameters & parameters)

Initializes the donor fraction, and the maximum convective velocity when starting a calculation. The donor fraction is used to determine the amount of upwinded donor cell to use in advection terms. The maximum convective velocity is used for calculation of constant eddy viscosity parameter. This version of the fuction is for 3D, tabulated equation of state calculations.

References Parameters::dDonorCellMin, Parameters::dDonorCellMultiplier, Grid::dLocalGridOld, Parameters::dMaxConvectiveVelocity, Grid::nCenIntOffset, Grid::nD, Grid::nDonorCellFrac, Grid::nEndGhostUpdateExplicit, Grid::nEndUpdateExplicit, Grid::nGamma, Grid::nP, Grid::nQ0, Grid::nQ1, Grid::nQ2, Grid::nStartUpdateExplicit, Grid::nU, Grid::nU0, Grid::nV, and Grid::nW.

Referenced by initInternalVars().

## 7.12.2.104 void initInternalVars (Grid & grid, ProcTop & procTop, Parameters & parameters)

This function function is used to set the initial values of the internal variables. While external variables are initialized from the starting model, internal variables are calculated at startup.

#### Parameters:

- $\leftrightarrow$  grid supplies information needed for initilizing internal variables as well as storing the initilized internal variables
- $\leftarrow procTop$  contians information about processor topology
- ← parameters contains parameters used in initializing the internal variables.

### Warning:

 $\Delta\theta$ ,  $\Delta\phi$ ,  $\sin\theta_{i,j,k}$ ,  $\Delta\cos\theta_{i,j,k}$ , all don't have the first zone calculated. At the moment this is a ghost cell that doesn't matter, but it may become a problem if calculations require this quantity. This is an issue for quantities that aren't updated in time, as those that are will have boundary cells updated with periodic boundary conditions.

References Parameters::bEOSGammaLaw,  $calOldDenave\_R()$ ,  $calOldDenave\_RT()$ ,  $calOldDenave\_RT()$ , calOldEddyVisc R CN(), calOldEddyVisc R SM(), calOldEddyVisc -

RT CN(), calOldEddyVisc RT SM(), calOldEddyVisc RTP CN(), calOldEddyVisc RTP -SM(), calOldP GL(), calOldPEKappaGamma TEOS(), calOldQ0 R GL(), calOldQ0 R -TEOS(), calOldQ0Q1 RT GL(), calOldQ0Q1 RT TEOS(), calOldQ0Q1Q2 RTP GL(), calOldQ0Q1Q2 RTP TEOS(), Grid::dLocalGridOld, initDonorFracAndMaxConVel R -GL(), initDonorFracAndMaxConVel R TEOS(), initDonorFracAndMaxConVel RT GL(), initDonorFracAndMaxConVel RT TEOS(), initDonorFracAndMaxConVel RTP GL(), initDonorFracAndMaxConVel RTP TEOS(), Grid::nCenIntOffset, Grid::nCotThetaIJK, Grid::nCotThetaIJp1halfK, Grid::nDCosThetaIJK, Grid::nDPhi, Grid::nDTheta, Grid::nLocalGridDims, Grid::nNumDims, Grid::nNumGhostCells, Grid::nPhi, Proc-Top::nRank, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Grid::nTheta, and Parameters::nTypeTurbulanceMod.

Referenced by init().

### 7.12.2.105 void setInternalVarInf (Grid & grid, Parameters & parameters)

This function sets the information for internal variables. While external verabile information is derived from the starting model, internal variables infos are set in this function. In other words this function sets the values of Grid::nVariables.

### Parameters:

- $\leftrightarrow$  grid supplies the information needed to calculate the horizontal density average, it also stores the calculated horizontally averaged density.
- ← *parameters* is used when setting variable infos, since one needs to know if the code is calculating using a gamma law gas, or a tabulated equation of state.

References Parameters::bEOSGammaLaw, Grid::nCotThetaIJK, Grid::nCotThetaIJp1halfK, Grid::nDCosThetaIJK, Grid::nDenAve, Grid::nDonorCellFrac, Grid::nDPhi, Grid::nDTheta, Grid∷nE, Grid::nEddyVisc, Grid::nGamma, Grid::nKappa, Grid::nNumDims, Grid::nNumVars, Grid::nNumIntVars, Grid::nQ1, Grid::nP, Grid::nQ0, Grid∷nQ2, Grid::nSinThetaIJK, Grid::nSinThetaIJp1halfK, Parameters::nTypeTurbulanceMod, Grid::nVariables.

Referenced by modelRead().

# 7.12.2.106 void setMainFunctions (Functions & functions, ProcTop & procTop, Parameters & parameters, Grid & grid, Time & time, Implicit & implicit)

Used to set the functions that main() uses to evolve the input model.

### Parameters:

- $\rightarrow$  functions is of class Functions and is used to specify the functions called to calculate the evolution of the input model.
- ← *procTop* is of type ProcTop. ProcTop::nRank is used to set different functions based on processor rank. For instance processor rank 1 requires 1D versions of the equations.
- ← parameters is of class Parameters. It holds various constants and runtime parameters.
- ← *grid* of type Grid. This function requires the number of dimensions, specified by Grid::nNumDims.
- ← *time* of type Time. This function requires knowledge of the type of time setp being used, specified by Time::bVariableTimeStep.

← *implicit* of type Implicit. This function needs to know if there is an implicit region, specified when Implicit::nNumImplicitZones>0.

The functions are picked based on model geometry, and the physics requested or required by the input model, and the configuration file. The specific functions pointers that are set are described in the Functions class.

References Parameters::bAdiabatic, Parameters::bEOSGammaLaw, Time::bVariableTimeStep, calDelt CONST(), calDelt R GL(), calDelt R TEOS(), calDelt RT GL(), calDelt - $RT\_TEOS()$ ,  $calDelt\_RTP\_GL()$ ,  $calDelt\_RTP\_TEOS()$ ,  $calNewD\_R()$ , calNewD -RT(),calNewD RTP(), calNewDenave None(), calNewDenave R(), calNewDenave calNewDenave RTP(), calNewE R AD(), calNewE R NA(), RT(),calNewE calNewE RT NA(), RT AD(),calNewE RT NA LES(), calNewE RTP calNewEddyVisc None(), calNewE RTP NA LES(), AD(), calNewE RTP NA(), calNewEddyVisc RT SM(), calNewEddyVisc RT CN(), calNewEddyVisc RTP CN(), calNewEddyVisc RTP SM(), calNewP GL(), calNewQ0 R GL(),calNewQ0 R calNewQ0Q1 RT GL(), calNewQ0Q1 RT TEOS(), calNewQ0Q1Q2 RTP calNewQ0Q1Q2 RTP TEOS(), calNewR(), calNewTPKappaGamma TEOS(), GL(),calNewU0 R(), calNewU0 RT(), calNewU0 RTP(), calNewVelocities R(), calNewVelocities calNewVelocities RT LES(), calNewVelocities RTP(), calNewVelocities -RT(), RTP LES(), dImplicitEnergyFunction None(), dImplicitEnergyFunction R(), dImplicitEnergyFunction R SB(), dImplicitEnergyFunction RT(), dImplicitEnergyFunction -RT LES(), dImplicitEnergyFunction RT LES SB(), dImplicitEnergyFunction -RT SB(), dImplicitEnergyFunction RTP(), dImplicitEnergyFunction RTP LES(), dImplicitEnergyFunction RTP LES SB(), dImplicitEnergyFunction RTP SB(), tions::fpCalculateAveDensities, Functions::fpCalculateDeltat, Functions::fpCalculateNewAV, Functions::fpCalculateNewDensities, Functions::fpCalculateNewEddyVisc, Functions::fpCalculateNewEOSVars, tions::fpCalculateNewEnergies, Functions::fpCalculateNewGridVelocities, Functions::fpCalculateNewRadii, Functions::fpCalculateNewVelocities, Functions::fpImplicitEnergyFunction, Functions::fpImplicitEnergyFunction SB, Functions::fpImplicitSolve, Functions::fpModelWrite, Functions::fpUpdateLocalBoundaryVelocitiesNewGrid, Functions::fpWriteWatchZones, implicitSolve None(), implicitSolve R(),implicitSolve RT(), implicitSolve RTP(), modelWrite GL(), modelWrite TEOS(), Grid::nNumDims, Implicit::nNumImplicitZones, Proc-Top::nRank, Parameters::nTypeTurbulanceMod, updateLocalBoundaryVelocitiesNewGrid R(), updateLocalBoundaryVelocitiesNewGrid RT(), updateLocalBoundaryVelocitiesNewGrid -RTP(), writeWatchZones R GL(), writeWatchZones R TEOS(), writeWatchZones RT -GL(), writeWatchZones RT TEOS(), writeWatchZones RTP GL(), and writeWatchZones -RTP TEOS().

Referenced by main().

## 7.13 procTop.cpp File Reference

```
#include "procTop.h"
#include <cstring>
```

### 7.13.1 Detailed Description

Implementation file for the ProcTop class

## 7.14 procTop.h File Reference

### Classes

• class ProcTop

## 7.14.1 Detailed Description

Header file for the  ${\bf ProcTop}$  class

## 7.15 profile Data.h File Reference

```
#include <string>
#include <vector>
#include <map>
#include <limits>
#include "time.h"
#include "procTop.h"
#include <fstream>
```

### Classes

 $\bullet$  class **profileData** 

### 7.15.1 Detailed Description

Header file for keepMax::cpp

## 7.16 time.cpp File Reference

```
#include "time.h"
#include <limits>
```

### 7.16.1 Detailed Description

Implementation file for the Time class

## 7.17 time.h File Reference

## Classes

• class Time

## 7.17.1 Detailed Description

Header file for the  ${\bf ProcTop}$  class

## 7.18 userguide.h File Reference

## 7.18.1 Detailed Description

Contains the text for the "Using and Modifying SPHERIS" section of this manual.

## 7.19 watchzone.cpp File Reference

```
#include "watchzone.h"
#include "exception2.h"
#include <sstream>
```

### 7.19.1 Detailed Description

This file holds the implementation of the watchzone class.

## 7.20 watchzone.h File Reference

```
#include <string>
#include <fstream>
```

### Classes

• class WatchZone

### 7.20.1 Detailed Description

This file holds the definition of the watchzone class.

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