

# Carmen–MHD Code Manual

(version 1.0)

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# Chapter 1

## Adaptive Multiresolution MHD Simulation

### 1.1 CARMEN–MHD code

CARMEN code was firstly developed by Olivier Roussel in his Ph.D dissertation to perform the simulation of the Advection-diffusion, Burgers-diffusion, Flame front, Flame ball, Flame-curl interaction and Navier-Stokes equations with the finite volume method in the context of the adaptive multiresolution analysis for cell-averages [17]. The code was extended to the resistive and ideal MHD equations in order to take advantage of the wavelet based multiresolution algorithm. Its development was done in such a way that the code structure was exclusively for MHD simulations, and CARMEN code became CARMEN–MHD code [6].

The bidimensional MHD solver implementation in the uniform mesh started with the MHD–FV code in *C* language, which was developed in Gomes' Master thesis [5] sit on top on a revised version of Bastien's MHD code in FORTRAN [2]. The MHD–FV code could perform the first order accuracy MHD finite volume simulation in two dimensions, with the Harten-Lax-Van Leer (HLL) and Harten-Lax-Van Leer-Discontinuities (HLLD) Riemann solvers to compute the fluxes [14, 16].

The bidimensional ideal MHD model implementation in CARMEN–MHD have been done since 2012 [5, 4, 7]. The first implementation came with the HLL and HLLD Riemann solvers, mixed divergence cleaning, second order accuracy in time and first order accuracy in space. The MHD solver was coupled to the pre existing adaptive multiresolution algorithm.

After that, during Gomes' Ph.D, some modifications have been made to improve the CPU time and boundary conditions, reconstruct the variables, fix MHD waves evaluation, divergence cleaning spacial fix and more [12]. Subsequently, the three-dimensional model, finite volume approach for MHD, wavelet coefficient normalization improvement and artificial diffusion source terms were implemented [10, 9]. These modification improved the mesh adaptivity to the solution and the comparisons of CPU time, allowing the three dimensional simulations and ensuring numerical stability. There was also a bug fixed in the original CARMEN code finite volume three dimensional approach implementation, which was shifting the solution's computational domain.

The resistive MHD model in two and three dimensions is the earlier implementation coupled to the adaptive algorithm [8, 11]. The time step was updated to the resistive case and the resistivity scalar function can be constant or vary in space, allowing the simulation of a variety of physical phenomena.

CARMEN–MHD code is available for download in GitHub (<https://github.com/waveletApplications/carmenMHD>). If you want to cite CARMEN–MHD code, you can refer any of the following publications:

- GOMES, Anna Karina Fontes. Simulação numérica de um modelo magneto-hidrodinâmico multidimensional no contexto da multirresolução adaptativa por médias celulares. 2017. PhD thesis, Instituto Nacional de Pesquisas Espaciais (INPE), São José dos Campos.
- GOMES, Anna Karina Fontes et al. An adaptive multiresolution method for ideal magnetohydrodynamics using divergence cleaning with parabolic-hyperbolic correction. *Applied Numerical Mathematics*, v. 95, p. 199-213, 2015.
- DOMINGUES, Margarete O. et al. Extended generalized Lagrangian multipliers for magnetohydrodynamics using adaptive multiresolution methods. In: ESAIM: Proceedings. EDP Sciences, 2013. p. 95-107.

- GOMES, Anna Karina Fontes. Análise multirresolução adaptativa no contexto da resolução numérica de um modelo de magnetohidrodinâmica ideal. 2012. Master thesis, Instituto Nacional de Pesquisas Espaciais (INPE), São José dos Campos.

## 1.2 GLM–MHD equations

The resistive MHD equations with the mixed divergence cleaning are given by

$$\frac{\partial \rho}{\partial t} + \nabla \cdot (\rho \mathbf{u}) = 0, \quad (1.1a)$$

$$\frac{\partial \mathcal{E}}{\partial t} + \nabla \cdot \left[ \left( \mathcal{E} + p + \frac{|\mathbf{B}|^2}{2} \right) \mathbf{u} - \mathbf{B}(\mathbf{u} \cdot \mathbf{B}) \right] = \nabla \cdot [\mathbf{B} \times \eta (\nabla \times \mathbf{B})], \quad (1.1b)$$

$$\frac{\partial (\rho \mathbf{u})}{\partial t} + \nabla \cdot \left[ \rho \mathbf{u}' \mathbf{u} + \mathbf{I} \left( p + \frac{|\mathbf{B}|^2}{2} \right) - \mathbf{B}' \mathbf{B} \right] = \mathbf{0}, \quad (1.1c)$$

$$\frac{\partial \mathbf{B}}{\partial t} + \nabla \cdot [\mathbf{u}' \mathbf{B} - \mathbf{B}' \mathbf{u}] = -\nabla \times (\eta \nabla \times \mathbf{B}), \quad (1.1d)$$

where  $\rho$  is the density,  $\mathbf{u}$  the fluid velocity,  $\mathbf{B}$  the magnetic field,  $\eta$  the resistivity, and  $\mathbf{I}$  the  $3 \times 3$  identity matrix. The total energy density  $\mathcal{E}$  is given by

$$\mathcal{E} = \frac{p}{\gamma - 1} + \frac{\rho \mathbf{u}}{2} + \frac{\mathbf{B}}{2}, \quad (1.2)$$

where  $p$  is the pressure and  $\gamma$  the adiabatic constant. The constraint  $\nabla \cdot \mathbf{B} = 0$  is enforced by the parabolic-hyperbolic (or mixed) divergence cleaning proposed by Dedner *et al.* in [1] in combination with the correction proposed by Mignone and Tzeferacos in [15].

The mixed correction adds a differential operator  $D = \frac{1}{c_h^2} \frac{\partial}{\partial t} + \frac{1}{c_p^2}$  to the  $\nabla \cdot \mathbf{B} = 0$  equation, resulting in a new MHD model composed by the density, energy density and momentum equations, along with the additional equations

$$\frac{\partial \mathbf{B}}{\partial t} + \nabla \cdot (\mathbf{u} \mathbf{B} - \mathbf{B} \mathbf{u} + \psi \mathbf{I}) = 0, \quad \frac{\partial \psi}{\partial t} + c_h^2 \nabla \cdot \mathbf{B} = -\frac{c_h^2}{c_p^2} \psi \quad (1.3)$$

where  $\psi$  is a scalar function,  $c_p$  and  $c_h$  are the parabolic and hyperbolic constants, respectively.

## 1.3 Adaptive Multiresolution

The adaptivity of the computational mesh is performed by the adaptive multiresolution (MR) approach, which is based on an adaptive cell average approach as discussed in [13, 3]. The main idea of MR is to decompose the data into several levels of refinement. The refinement levels are associated to a multiresolution mesh structure that creates dyadic embedded cell meshes, that have different numbers of cells according to the level they belong to. The idea of adaptivity starts from the wavelet coefficients, which can measure the local regularity of the data according to a given threshold parameter  $\varepsilon^\ell = \varepsilon(\varepsilon^0, \ell)$ , where  $\ell$  denotes the cell scale level and  $\varepsilon^0$  is the initial threshold parameter. When the wavelet coefficients are larger than  $\varepsilon^\ell$ , the computational mesh needs to be more refined locally; otherwise the mesh can remain coarser. This methodology allows the computational mesh to be more refined only where it is required. In the context of our work, our data is an MHD solution represented by a set of cell averages (mesh) [12].

## 1.4 Development team

As part of efforts for developing the Space Science studies at the Brazilian Institute for Space Research (INPE), some numerical codes has been improved to extend fluid dynamic solutions to the magnetohydrodynamics context. So, the CARMEN–MHD code has been maintained by Anna Karina Fontes Gomes<sup>1</sup>, Muller Moreira Souza Lopes<sup>2</sup> and Margarete Oliveira Domingues<sup>3</sup>, members of the *Wavelet and Applications* research group in this institute at

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## 1.6 License

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<sup>4</sup><https://www.gnu.org/licenses/gpl-3.0.en.html>



## Chapter 2

# How to use CARMEN–MHD

In this chapter we present the details about CARMEN-MHD code initialization files and algorithm. We also provide the information about its compilation and run.

### 2.1 Running the code

To compile the code you need to install the  $g^{++}$  compiler. We have used CARMEN–MHD code in Ubuntu 14.04 with  $g^{++}$  version 4.8.4 without any problem. You can try another compiler, but we cannot ensure its operation as we did not try it before.

With the proper compiler installed, open the terminal and type `make`. If you make any changes and want to recompile the code you should type `make clean` before `make` to ensure that every file of the code will be compiled again. This process will create an executable file named `carmen`, by which you can run the code. Just type `./carmen` in the terminal and the simulation will begin.

When the simulation is finished, it will create output files named `Average.vtk`, `Mesh.dat`, `Integral.dat` and `carmen.prf`. The `Average.vtk` file contains the cell-average values of the numerical solution. It can be visualized in `VisIt`<sup>1</sup> or `Paraview`<sup>2</sup>. The `Mesh.dat` file contains the adaptive mesh of the simulation, represented by the coordinates of the center of the cells and the level it belongs to. The scripts need for the visualization of these files is available in `github`<sup>3</sup>. The `Integral.dat` file contains the values of some quantities, as global energy and number of iterations, over time. You can have information about the behavior of these quantities as the time varies. Finally, the `carmen.prf` contained the simulation information, as CPU time, memory compression and some parameters that are set in the initialization.

### 2.2 Code operation

The CARMEN–MHD code algorithm is described by Figure 2.1. The algorithm is basically divided in five sections. At first, we have the *Initialization*, where the initial condition and parameters are read and the initial mesh created. This step occurs once during the simulation. The next steps *Time Evolution*, *Rebuild mesh* and *Output* occur as long as the final time is not reached. In time evolution, the fluxes are calculated and the MHD variables are evolved in time. After that, the stability of the solution is checked. If it is stable, the divergence cleaning correction is applied to the variables. Otherwise, the simulation is aborted. The mesh is updated according to the new values of the variables and adapted again. The evolved variables and the new adapted mesh is written in `Average.vtk` and `Mesh.dat`, respectively. The last step is *Finish*, in which the memory is freed.

---

<sup>1</sup><https://visit.llnl.gov/>

<sup>2</sup><https://www.paraview.org/>

<sup>3</sup><https://github.com/waveletApplications/carmenMHD>

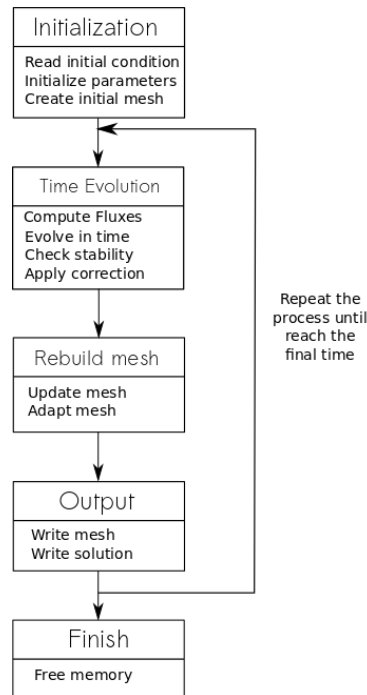


Figure 2.1: Flow chart of the CARMEN-MHD algorithm.

For the initialization of the code, you need to define the initial conditions and parameters to perform the simulation. These settings has to be make in `carmen.ini` and `carmen.par` files. In 2.1 we present an example of a `carmen.ini` file, given by the initial condition for the Kelvin-Helmholtz instability. The primitive variables values of the MHD model are defined and then assigned to their respective conservative variables in the vector of quantities  $Q[i]$ , where  $i \in \{1, \dots, 9\}$ , considering  $\rho$ ,  $\rho u_x$ ,  $\rho u_y$ ,  $\rho u_z$ ,  $\mathcal{E}$ ,  $\psi$ ,  $B_x$ ,  $B_y$  and  $B_z$ . An example `carmen.par` file is presented in 2.2. The file is divided in sections, where you can find the parameters related to each one. The parameters are described below.

### 1 - Time Integration

**StepNb** It is related to the accuracy order of Runge-Kutta time evolution scheme. We use 2 for second order.

**SchemeNb** It is related to the Riemann solver choice. We set 1 for HLL and 2 for HLLD.

**PhysicalTime** Final time of the simulation.

**CFL** Courant number related to the CFL condition.

### 2 - Solved Equations

**EquationType** CARMEN-MHD code only works if it is set to be 7, which means MHD equations.

### 3 - Geometry

**Dimension** It can be choose 2 for two dimensions and 3 for three dimensions.

**XMin** It is the inferior limit of the domain. You choose 1, 2 or 3 inside the brackets to set the limits in  $x$ ,  $y$  or  $z$  direction, respectively.

**XMax** It is the superior limit of the domain. You choose 1, 2 or 3 inside the brackets to set the limits in  $x$ ,  $y$  or  $z$  direction, respectively.



Listing 2.1: Example of a carmen.ini file.

---

```
// — Initial condition Q[i] in function of (x,y,z) —

// Pi constant
real PI = 4.0*atan(double(1.0));

// Conservative Variables
//rho, rho*ux, rho*uy, rho*uz, E, psi, Bx, By, Bz

real rho,E,ux,uy,uz,psi,Bx,By,Bz,p;

rho = 1.0;
ux = 5*(tanh(20*(y+0.5)) - (tanh(20*(y-0.5)) + 1));
uy = 0.25*sin(2*PI*x)*(exp(-100*(y+0.5)*(y+0.5)) - exp(-100*(y-0.5)*(y-0.5)));
uz = 0.;
p = 50.0;
psi = 0.;
Bx = 1.0;
By = 0.0;
Bz = 0.;
E = (p/(Gamma-1.0)) + rho*0.5*(ux*ux + uy*uy + uz*uz) + 0.5*(Bx*Bx + By*By +Bz*Bz);

Q[1] = rho;
Q[2] = rho*ux;
Q[3] = rho*uy;
Q[4] = rho*uz;
Q[5] = E;
Q[6] = psi;
Q[7] = Bx;
Q[8] = By;
Q[9] = Bz;
```

---

**CMin** It is the boundary condition on the inferior limit of the domain. You choose 1, 2 or 3 inside the brackets to set the limits in  $x$ ,  $y$  or  $z$  direction, respectively. You assign 2 for Neumann and 3 for periodic conditions.

**CMax** It is the boundary condition on the superior limit of the domain. You choose 1, 2 or 3 inside the brackets to set the limits in  $x$ ,  $y$  or  $z$  direction, respectively. You assign 2 for Neumann and 3 for periodic conditions.

#### 4 - Multiresolution

**Multiresolution** If it is `true`, the multiresolution approach is used. Otherwise, you choose a full regular mesh.

**ScaleNb** It is the number of scale, related to the maximum refinement of the mesh, which is  $2^{\text{Dimension} * \text{ScaleNb}}$ .

**Tolerance** This is the value of the threshold parameter (only for multiresolution).

#### 5 - Physics

**Gamma** The adiabatic constant.

**Resistivity** If it is `true`, the resistivity approach is used. Otherwise, you choose the ideal MHD model.

**eta** Resistivity constant (only for resistive MHD)

#### Divergence cleaning

**cr** Parameter related to the mixed divergence cleaning. We usually choose  $cr = 0.4$ .

#### Others

**ThresholdNorm** If it is 0, the threshold parameter remains the same for every refinement level. If it is 1, the threshold parameter changes according to the level of refinement.

Depending on the initial condition you want to input, the parameters have to be adjusted. If you want to use the resistive model with variable resistivity, you can define the scalar function  $\eta$  in `carmen.eta` file. In 2.3 we show an example file used for the magnetic reconnection simulation. In this case, we assign the function to the variable `Res`, which depends on the independent variables  $x$  and  $y$ . This can be extended to three dimensions by adding the variable  $z$ . You can define a function that is more adequate to the problem of your interest. The next chapters are describing the classes, files and variables of the code in details. This documentation can help you with the understanding of the code routines and more.

Listing 2.2: Example of a carmen.par file.

---

```
// Carmen parameter file
// Generated by Carmen Editor
// Kelvin–Helmholtz instability in two dimensions

// 1) Time integration -----
StepNb = 2;
SchemeNb = 2;
PhysicalTime = 0.2;
CFL = 4.00000e-01;

// 2) Solved equations -----
EquationType = 7;

// 3) Geometry -----
Dimension = 2;

XMin[1] = 0.000000e+00;
XMax[1] = 1.000000e+00;
XMin[2] = -1.000000e+00;
XMax[2] = 1.000000e+00;

CMin[1] = 2;
CMax[1] = 2;
CMin[2] = 2;
CMax[2] = 2;

// 4) Multiresolution -----
Multiresolution = true;
ScaleNb = 9;
Tolerance = 1.000000e-01;
// 5) Physics -----
Gamma = 1.4;
Resistivity = true;
eta = 0.02;

// 6) Divergence Cleaning -----
cr = 0.4;

// 7) Others -----
ThresholdNorm = 0;
```

---

Listing 2.3: Example of a carmen.ini file.

---

```
// — Magnetic Resistivity —  
// If Resistivity = True, edit this  
  
if (fabs(x) <= 0.05)  
{  
    if (fabs(y) <= 0.2)  
        Res = eta * (cos(pi*x/0.1) + 1.) * (cos(pi*y/0.4) + 1) / 4.;  
}  
else  
    Res = 0.;
```

---

# Chapter 3

## Class Index

### 3.1 Class List

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

<a href="#">Cell</a>	An object <a href="#">Cell</a> contains all the informations of a cell for both multiresolution and finite volume computations . . . . .	17
<a href="#">FineMesh</a>	An object <a href="#">FineMesh</a> is a regular fine mesh, used for finite volume computations. It is not used for multiresolution computations . . . . .	56
<a href="#">Matrix</a>	Standard class for every matrix in Carmen . . . . .	89
<a href="#">Node</a>	An object <a href="#">Node</a> is an element of a graded tree structure, used for multiresolution computations. Its contains the following informations: . . . . .	110
<a href="#">PrintGrid</a>	An object <a href="#">PrintGrid</a> is a special regular grid created to write tree-structured data into an output file	146
<a href="#">TimeAverageGrid</a>	Time Average Grid . . . . .	162
<a href="#">Timer</a>	An object <a href="#">Timer</a> gives information on the CPU time of long-time computations . . . . .	169
<a href="#">Vector</a>	Standard class for every vector in Carmen . . . . .	174



# Chapter 4

## File Index

### 4.1 File List

Here is a list of all files with brief descriptions:

<a href="#">AdaptTimeStep.cpp</a>	This function computes the time step for the next iteration . . . . .	203
<a href="#">ArtificialViscosity.cpp</a>	Computes the Laplacian terms for density, energy and momentum equations. It helps with the stability . . . . .	204
<a href="#">Backup.cpp</a>	Backup the last simulation . . . . .	206
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<a href="#">Cell.cpp</a>	Constructor and distructor of the cell class. Also computes the cell-averages of the MHD variables . . . . .	278
<a href="#">Cell.h</a>	. . . . .	278
<a href="#">ComputedTolerance.cpp</a>	Adapt trreshold parameter or use it fixed . . . . .	278
<a href="#">CPUTimeRef.cpp</a>	Compute the reference CPU time with the finite volume solver. The output is the CPU time for 1 iteration . . . . .	279
<a href="#">DigitNumber.cpp</a>	This function returns the number of digits of an integer . . . . .	281
<a href="#">DivCleaning.cpp</a>	. . . . .	282
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<a href="#">Flux.cpp</a>	Computes the numerical fluxes HLL and HLLD . . . . .	286
<a href="#">FluxCorrection.cpp</a>	Computes the mixed correction in the numerical fluxes (Dedner, 2002) . . . . .	287
<a href="#">GetBoundaryCells.cpp</a>	Computes the cells C1, C2, C3, C4 in function of the cells Cell1, Cell2, Cell3, Cell4 taking into account boundary conditions . . . . .	289

<a href="#">InitAverage.cpp</a>	Fill the variables vector with the initial condition . . . . .	297
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<a href="#">Parameters.h</a>	This header contains all parameters as global variables . . . . .	350
<a href="#">Performance.cpp</a>	Simulation information . . . . .	367
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<a href="#">Source.cpp</a>	Computes the source terms of the system . . . . .	394
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---

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## Chapter 5

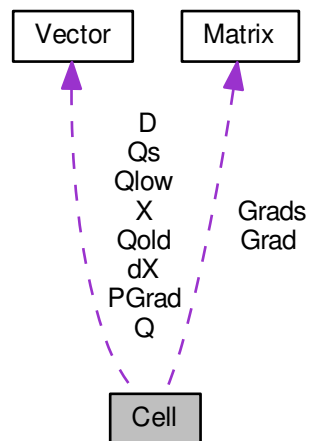
# Class Documentation

### 5.1 Cell Class Reference

An object `Cell` contains all the informations of a cell for both multiresolution and finite volume computations.

```
#include <Cell.h>
```

Collaboration diagram for `Cell`:



#### Public Member Functions

- `Cell ()`  
*Cell constructor.*
- `~Cell ()`  
*Cell distructor.*
- `void operator= (const Cell &C)`  
*Defines the cell operators. It is possible to obtain the cell values as:*
- `void setSize (const int AxisNo, const real UserSize)`  
*Sets the size of the cell in the direction AxisNo to UserSize. Example:*

- void `setSize` (const `Vector` &UserSize)
 

*Sets the size of the cell in every direction to the vector UserSize . Example:*
- void `setCenter` (const int AxisNo, const `real` UserX)
 

*Sets the coordinate of the cell-center in the direction AxisNo to UserX. Example:*
- void `setCenter` (const `Vector` &UserX)
 

*Sets the position of the cell-center to the vector UserX. Example:*
- void `setAverage` (const int QuantityNo, const `real` UserAverage)
 

*Sets the cell-average of the quantity QuantityNo to UserAverage. Example:*
- void `setAverage` (const `Vector` &UserAverage)
 

*Sets all the cell-average quantities to the vector UserAverage. Example:*
- void `setAverageZero` ()
 

*Sets all the cell-average values to zero.*
- void `setTempAverage` (const int QuantityNo, const `real` UserAverage)
 

*Identical to `setAverage` (int QuantityNo, real UserAverage), but for the vector of the temporary cell-average values.*
- void `setTempAverage` (const `Vector` &UserAverage)
 

*Identical to void `setAverage` (const `Vector`& UserAverage), but for the vector of the temporary cell-average values.*
- void `setTempAverageZero` ()
 

*Sets all the temporary cell-average values to zero.*
- void `setLowAverage` (const int QuantityNo, const `real` UserAverage)
 

*Identical to `setAverage` (int QuantityNo, real UserAverage), but for the vector of the cell-average values with low precision in the Runge-Kutta-Fehlberg method.*
- void `setLowAverage` (const `Vector` &UserAverage)
 

*Identical to void `setAverage` (const `Vector`& UserAverage), void `setAverage` (const `Vector`& UserAverage), but for the vector of the cell-average values with low precision in the Runge-Kutta-Fehlberg method.*
- void `setOldAverage` (const int QuantityNo, const `real` UserAverage)
 

*Identical to `setAverage` (int QuantityNo, real UserAverage), but for the vector of the old cell-average values.*
- void `setOldAverage` (const `Vector` &UserAverage)
 

*Identical to void `setAverage` (const `Vector`& UserAverage), but for the vector of the cell-average values.*
- void `setDivergence` (const int QuantityNo, const `real` UserAverage)
 

*Identical to void `setAverage` (int QuantityNo, real UserAverage), but for the divergence vector.*
- void `setDivergence` (const `Vector` &UserAverage)
 

*Identical to void `setAverage` (const `Vector`& UserAverage), but for the divergence vector.*
- void `setPsiGrad` (const int Dimension, const `real` UserAverage)
 

*Identical to void `setAverage` (int QuantityNo, real UserAverage), but for the gradient of psi vector.*
- void `setPsiGrad` (const `Vector` &UserAverage)
 

*Identical to void `setAverage` (int QuantityNo, real UserAverage), but for the gradient of psi vector.*
- void `setRes` (const `real` UserAverage)
 

*Set resistivity.*
- void `setDivergenceZero` ()
 

*Sets all the components of the divergence vector to zero.*
- void `setGradient` (const int i, const int j, const `real` UserAverage)
 

*Sets the component no. i, j of the quantity gradient to UserAverage.*
- void `setTempGradient` (const int i, const int j, const `real` UserAverage)
 

*Identical to the previous one for the temporary values. Does not work for MHD!*
- void `setGradient` (const `Matrix` &UserAverage)
 

*Sets the quantity gradient to the matrix UserAverage. Does not work for MHD! Example:*
- void `setTempGradient` (const `Matrix` &UserAverage)
 

*identical to the previous one for the temporary values.*
- void `setGradientZero` ()
 

*Sets all the components of the velocity gradient to zero.*
- bool `isInsideBoundary` () const

- Returns true if the cell is inside the boundary.*

  - `bool isInFluid ()` const
- Returns true if the cell is inside the fluid.*

  - `real size (const int AxisNo)` const
- Returns the cell size in the direction AxisNo.*

  - `Vector size ()` const
- Returns the vector containing the cell size in every direction.*

  - `real center (const int AxisNo)` const
- Returns the component no. AxisNo of the cell-center position.*

  - `Vector center ()` const
- Returns the cell-center position vector.*

  - `real average (const int QuantityNo)` const
- Returns the component no. QuantityNo of the cell-average value.*

  - `Vector average ()` const
- Returns the cell-average value vector.*

  - `real tempAverage (const int QuantityNo)` const
- Returns the component no. QuantityNo of the temporary cell-average value.*

  - `Vector tempAverage ()` const
- Returns the temporary cell-average value vector.*

  - `real lowAverage (const int QuantityNo)` const
- Returns the component no. QuantityNo of the cell-average value with low precision in the Runge-Kutta-Fehlberg method.*

  - `Vector lowAverage ()` const
- Returns the cell-average vector with low precision in the Runge-Kutta-Fehlberg method.*

  - `real oldAverage (const int QuantityNo)` const
- Returns the component no. QuantityNo of the old cell-average values.*

  - `Vector oldAverage ()` const
- Returns the old cell-average values.*

  - `real divergence (const int QuantityNo)` const
- Returns the component no. QuantityNo of the divergence vector.*

  - `Vector divergence ()` const
- Returns the divergence vector.*

  - `real PsiGrad (const int Dimension)` const
- Returns the component of PsiGrad vector.*

  - `Vector PsiGrad ()` const
- Returns the PsiGrad vector.*

  - `real gradient (const int i, const int j)` const
- Returns the component no. i, j of the velocity gradient. Does not work for MHD!*

  - `real tempGradient (const int i, const int j)` const
- Identical to the previous one for the temporary values.*

  - `Matrix gradient ()` const
- Returns the velocity gradient in matrix form.*

  - `Matrix tempGradient ()` const
- Identical to the previous one for the temporary values.*

  - `real density ()` const
- Returns the cell-average density.*

  - `real tempDensity ()` const
- Identical to the previous one for the temporary values.*

  - `real psi ()` const
- Returns the cell-average density.*

  - `real tempPsi ()` const

- Identical to the previous one for the temporary values.*
- **real etaConst** () const  
*Returns the cell-average resistivity.*
  - **real pressure** () const  
*Returns the cell-average pressure.*
  - **real tempPressure** () const  
*Identical to the previous one for the temporary values.*
  - **real oldPressure** () const  
*Identical to the previous one for the values at the instant n-1.*
  - **real temperature** () const  
*Returns the cell-average temperature. Does not work for MHD!*
  - **real tempTemperature** () const  
*Identical to the previous one for the temporary values. Does not work for MHD!*
  - **real concentration** () const  
*Returns the cell-average concentration of the limiting reactant. Does not work for MHD!*
  - **real energy** () const  
*Returns the cell-average energy per unit of volume.*
  - **real tempEnergy** () const  
*Identical to the previous one for the temporary values.*
  - **real magField** (const int AxisNo) const  
*Returns the component no. AxisNo of the cell-average magnetic field.*
  - **real tempMagField** (const int AxisNo) const  
*Identical to the previous one for the temporary values.*
  - **Vector magField** () const  
*Returns the cell-average magnetic field vector.*
  - **Vector tempMagField** () const  
*Identical to the previous one for the temporary values.*
  - **real velocity** (const int AxisNo) const  
*Returns the component no. AxisNo of the cell-average velocity.*
  - **real tempVelocity** (const int AxisNo) const  
*Identical to the previous one for the temporary values.*
  - **Vector velocity** () const  
*Returns the cell-average velocity vector.*
  - **Vector tempVelocity** () const  
*Identical to the previous one for the temporary values.*
  - **real speedOfSound** () const  
*Returns the cell-average speed of sound.*
  - **real entropy** () const  
*Returns the entropy ( $p/\rho^\Gamma$ ). Does not work for MHD!*
  - **real fastSpeed** (const int AxisNo) const  
*Returns the fast speed vector.*
  - **real volume** () const  
*Returns the volume of the cell (length in 1D, area in 2D, volume in 3D).*
  - **bool isOverflow** () const  
*Return true if one of the cell-average quantities is greater than the maximum. This usually means the computation is numerically unstable.*

## Public Attributes

- [Vector X](#)
- [Vector dX](#)
- [Vector Q](#)
- [Vector Qs](#)
- [Vector Qlow](#)
- [Vector Qold](#)
- [Vector D](#)
- [real Res](#)
- [Vector PGrad](#)
- [Matrix Grad](#)
- [Matrix Grads](#)

### 5.1.1 Detailed Description

An object [Cell](#) contains all the informations of a cell for both multiresolution and finite volume computations.

### 5.1.2 Constructor & Destructor Documentation

#### 5.1.2.1 [Cell::Cell \( \)](#)

[Cell](#) constructor.

```

34         :
35         X(Dimension),
36         dX(Dimension),
37         Q(QuantityNb),
38         Qs(QuantityNb),
39         Qlow((!ConstantTimeStep && StepNb > 2) ||
TimeAdaptivity)? QuantityNb:0),
40         Qold((UseBoundaryRegions)? QuantityNb:0),
41         D(QuantityNb),
42         Res(),
43         PGrad(Dimension),
44         Grad((EquationType==6)? Dimension:0, ((
EquationType==6)? QuantityNb:0),
45         Grads((EquationType==6) && SchemeNb > 5)?
Dimension:0, ((EquationType==6) && SchemeNb > 5)?
QuantityNb:0)
46 {
47     // Empty constructor
48     ;
49 }
```

#### 5.1.2.2 [Cell::~~Cell \( \)](#)

[Cell](#) distructor.

```

62 {
63     // Empty distructor
64 }
```

### 5.1.3 Member Function Documentation

#### 5.1.3.1 `real Cell::average ( const int QuantityNo ) const` `[inline]`

Returns the component no. *QuantityNo* of the cell-average value.

## Parameters

<i>QuantityNo</i>	Number of MHD variables = 9.
-------------------	------------------------------

## Returns

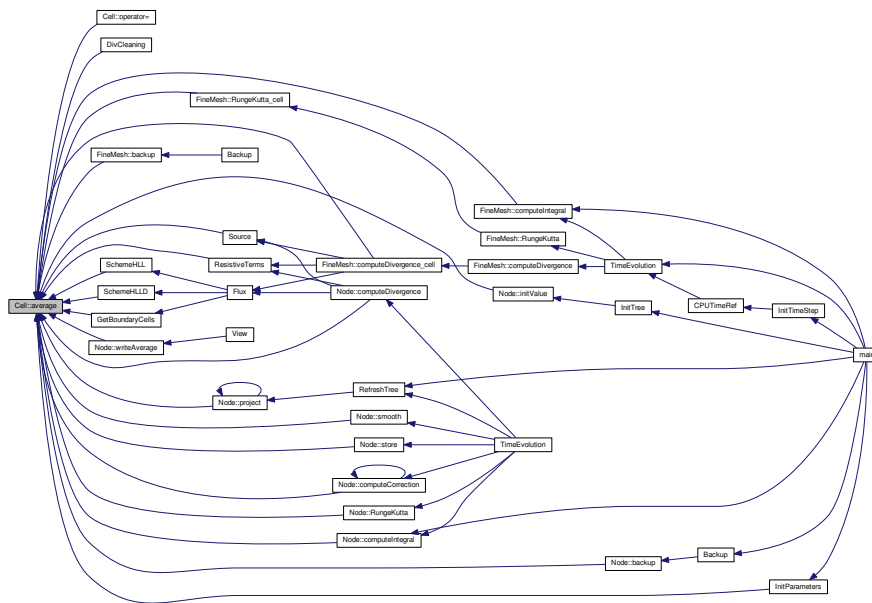
real

```

1129 {
1130     return Q.value(QuantityNo);
1131 }

```

Here is the caller graph for this function:



### 5.1.3.2 Vector Cell::average ( ) const [inline]

Returns the cell-average value vector.

## Returns

Vector

```

1137 {
1138     return Q;
1139 }

```

Here is the caller graph for this function:





```
5.1.3.3 real Cell::center ( const int AxisNo ) const [inline]
```

Returns the component no. *AxisNo* of the cell-center position.

## Parameters

<i>AxisNo</i>	Space direction the function is computed.
---------------	-------------------------------------------

## Returns

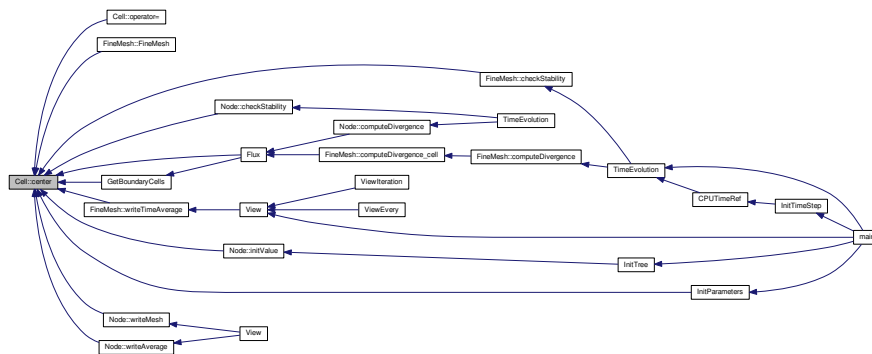
real

```

1113 {
1114     return X.value(AxisNo);
1115 }

```

Here is the caller graph for this function:



#### 5.1.3.4 Vector Cell::center ( ) const [inline]

Returns the cell-center position vector.

## Returns

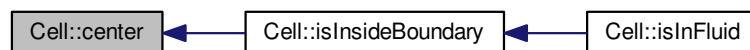
Vector

```

1121 {
1122     return X;
1123 }

```

Here is the caller graph for this function:



#### 5.1.3.5 real Cell::concentration ( ) const [inline]

Returns the cell-average concentration of the limiting reactant. Does not work for MHD!



## Returns

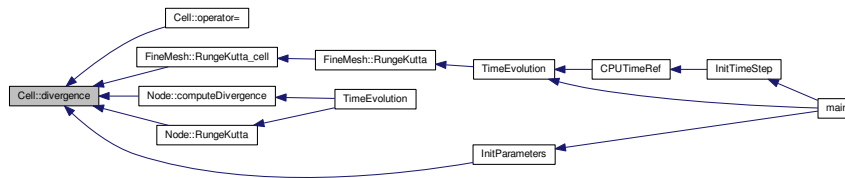
real

```

1193 {
1194     return D.value(QuantityNo);
1195 }

```

Here is the caller graph for this function:



### 5.1.3.8 Vector Cell::divergence ( ) const [inline]

Returns the divergence vector.

## Returns

Vector

```

1201 {
1202     return D;
1203 }

```

### 5.1.3.9 real Cell::energy ( ) const [inline]

Returns the cell-average energy per unit of volume.

## Returns

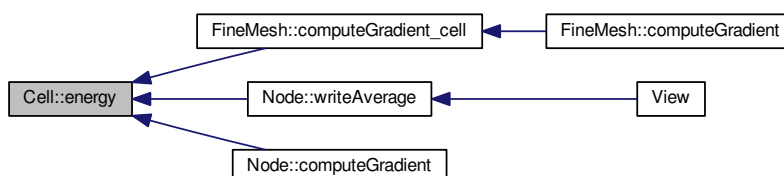
real

```

1308 {
1309     return Q.value(5);
1310 }

```

Here is the caller graph for this function:



**5.1.3.10** `real Cell::entropy( ) const [inline]`

Returns the entropy ( $p/\rho^\Gamma$ ). Does not work for MHD!

**Returns**

real

```
1380 {
1381     return pressure()*exp(-Gamma*log(density()));
1382 }
```

**5.1.3.11** `real Cell::etaConst( ) const [inline]`

Returns the cell-average resistivity.

**Returns**

real

```
1299 {
1300     return Res;
1301 }
```

**5.1.3.12** `real Cell::fastSpeed( const int AxisNo ) const`

Returns the fast speed vector.

Computes the fast magnetoacoustic wave at each direction.

**Parameters**

<i>AxisNo</i>	
---------------	--

**Returns**

real

**Parameters**

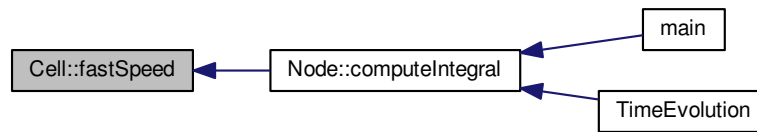
<i>AxisNo</i>	Number of axis of interest. 1: x-direction, 2: y-direction, 3: z-direction.
---------------	-----------------------------------------------------------------------------

**Returns**

double

```
341 {
342     real result;
343     real a = Gamma*pressure()/density();
344     real b = (magField()*magField())/density();
345     real bk= (magField(AxisNo)*magField(AxisNo))/density();
346
347
348     result = sqrt(.5*(a+b+sqrt((a+b)*(a+b) - 4.0*a*bk)));
349
350     return result;
351 }
```

Here is the caller graph for this function:



#### 5.1.3.13 `real Cell::gradient ( const int i, const int j ) const [inline]`

Returns the component no.  $i, j$  of the velocity gradient. Does not work for MHD!

##### Parameters

$i$	
$j$	

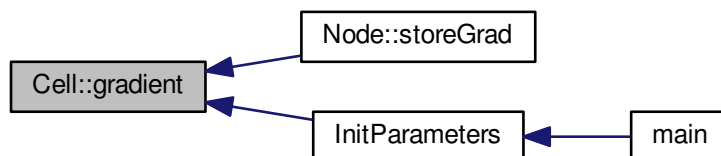
##### Returns

real

```

1226 {
1227     return Grad.value(i, j);
1228 }
  
```

Here is the caller graph for this function:



#### 5.1.3.14 `Matrix Cell::gradient ( ) const [inline]`

Returns the velocity gradient in matrix form.

##### Returns

Matrix

```

1246 {
1247     return Grad;
1248 }
  
```

## 5.1.3.15 bool Cell::isInFluid ( ) const

Returns true if the cell is inside the fluid.

## Returns

bool

brief Returns true if the cell is inside the boundary

return bool

```
488 {
489     return (!isInsideBoundary());
490 }
```

## 5.1.3.16 bool Cell::isInsideBoundary ( ) const

Returns true if the cell is inside the boundary.

## Returns

bool

```
445 {
446     int ei = 1;
447     int ej = (Dimension > 1) ? 1:0;
448     int ek = (Dimension > 2) ? 1:0;
449
450     int i, j, k;
451
452     Vector Edge(Dimension);
453
454     bool result=false;
455
456     // Loop: if one edge of the cell is in the boundary, return true
457     for (i=-ei; i <= ei; i+=2)
458     for (j=-ej; j <= ej; j+=2)
459     for (k=-ek; k <= ek; k+=2)
460     {
461         Edge.setValue(1,center(1) + i*0.5*size(1));
462
463         if (Dimension > 1)
464             Edge.setValue(2,center(2) + j*0.5*size(2));
465         if (Dimension > 2)
466             Edge.setValue(3,center(3) + k*0.5*size(3));
467
468         if (BoundaryRegion(Edge) != 0) result = true;
469     }
470
471     return result;
472
473 }
```

Here is the caller graph for this function:



### 5.1.3.17 bool Cell::isOverflow ( ) const

Return true if one of the cell-average quantities is greater than the maximum. This usually means the computation is numerically unstable.

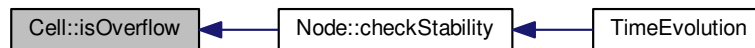
#### Returns

bool

```

383 {
384     // --- Local variables ---
385
386     int n;                // Counter on the quantities
387
388     // --- If one of the values is overflow, return true ---
389
390     for (n = 1; n <= QuantityNb; n++)
391         if (average().isNaN()) return true;
392
393     return false;
394 }
```

Here is the caller graph for this function:



### 5.1.3.18 real Cell::lowAverage ( const int *QuantityNo* ) const [inline]

Returns the component no. *QuantityNo* of the cell-average value with low precision in the Runge-Kutta-Fehlberg method.

#### Parameters

<i>QuantityNo</i>	Number of MHD variables = 9.
-------------------	------------------------------

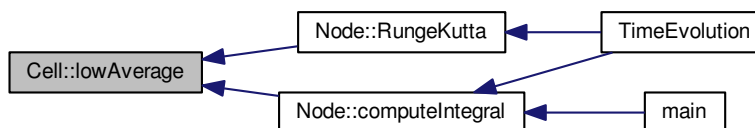
#### Returns

real

```

1161 {
1162     return Qlow.value(QuantityNo);
1163 }
```

Here is the caller graph for this function:





## 5.1.3.19 Vector Cell::lowAverage ( ) const [inline]

Returns the cell-average vector with low precision in the Runge-Kutta-Fehlberg method.

## Returns

Vector

```
1169 {
1170     return Qlow;
1171 }
```

## 5.1.3.20 real Cell::magField ( const int AxisNo ) const [inline]

Returns the component no. *AxisNo* of the cell-average magnetic field.

## Parameters

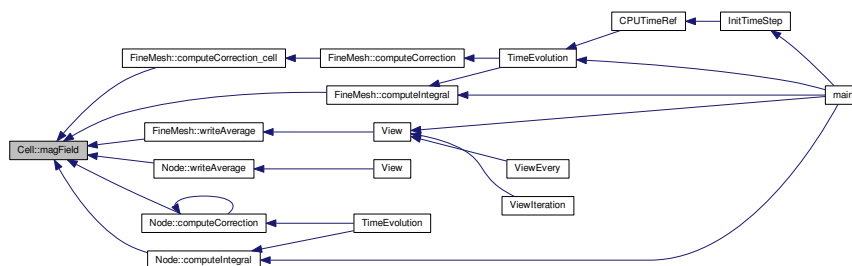
<i>AxisNo</i>
---------------

## Returns

real

```
1337 {
1338     return Q.value(6+AxisNo);
1339 }
```

Here is the caller graph for this function:



## 5.1.3.21 Vector Cell::magField ( ) const

Returns the cell-average magnetic field vector.

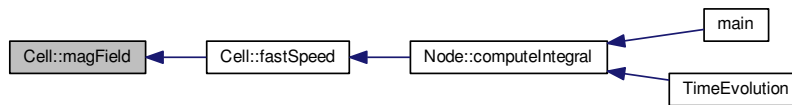
Computes the magnetic field. Allocates the magnetic field initial condition  $Q[i+6]$  to its components.

## Returns

Vector

```
244 {
245     // Local variables
246     Vector B(3);
247     int i;
248     for (i=1; i<=3; i++)
249         B.setValue(i, Q.value(i+6));
250     return B;
251 }
252
253
254 }
```

Here is the caller graph for this function:



### 5.1.3.22 `real Cell::oldAverage ( const int QuantityNo ) const [inline]`

Returns the component no. *QuantityNo* of the old cell-average values.

Parameters

<i>QuantityNo</i>	Number of MHD variables = 9.
-------------------	------------------------------

Returns

`real`

```

1177 {
1178     return Qold.value(QuantityNo);
1179 }
  
```

Here is the caller graph for this function:



### 5.1.3.23 `Vector Cell::oldAverage ( ) const [inline]`

Returns the old cell-average values.

Returns

`Vector`

```

1185 {
1186     return Qold;
1187 }
  
```

### 5.1.3.24 `real Cell::oldPressure ( ) const`

Identical to the previous one for the values at the instant  $n-1$ .

Computes the pressure of the fluid at the instant  $n-1$ . This value is useful for time integration computations.

## Returns

real  
double

```

152 {
153     // Conservative quantities;
154
155     real rho, rhoE;
156     Vector rhoV(3), B(3);
157
158     // Get conservative quantities
159
160     rho = Qold.value(1);
161
162     for (int i=1 ; i<=3 ; i++){
163         rhoV.setValue(i, Qold.value(i+1));
164         B.setValue(i, Qold.value(i+6));
165     }
166     rhoE = Qold.value(5);
167
168     // Return pressure
169
170     return (Gamma-1.)*(rhoE - .5*(rhoV*rhoV)/rho - .5*(B*B));
171
172 }

```

Here is the caller graph for this function:



## 5.1.3.25 void Cell::operator=( const Cell &amp; C )

Defines the cell operators. It is possible to obtain the cell values as:

- Center coordinates
- Size
- Average
- Temporary average
- Divergence
- Gradient of Psi

## Parameters

<i>C</i>	Cell of interest
----------	------------------

## Returns

void

```

417 {
418     // local cell becomes equal to cell C
419
420     setCenter(C.center());
421     setSize(C.size());
422     setAverage(C.average());
423     setTempAverage(C.tempAverage());
424     setDivergence(C.divergence());
425     setPsiGrad(C.PsiGrad());
426
427     return;
428
429 }

```

### 5.1.3.26 real Cell::pressure ( ) const

Returns the cell-average pressure.

Computes the pressure of the fluid. The pressure is computed in terms of the energy density, velocity and magnetic field. It is given by  $(\Gamma - 1) \cdot (\rho E - .5 \cdot (\rho v \cdot \rho v) / \rho - .5 \cdot (B \cdot B))$ ; return double.

Returns

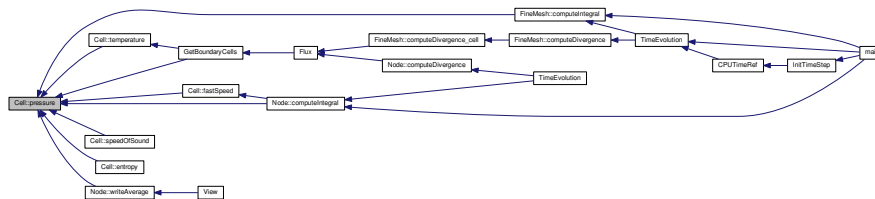
real

```

85 {
86     // Conservative quantities;
87
88     real rho, rhoE;
89     Vector rhoV(3), B(3);
90
91     // Get conservative quantities
92
93     rho = Q.value(1);
94
95     for (int i=1 ; i<=3; i++){
96         B.setValue(i, Q.value(i+6));
97         rhoV.setValue(i, Q.value(i+1));
98     }
99     rhoE = Q.value(5);
100
101     // Return pressure
102
103     return (Gamma-1.)*(rhoE - .5*(rhoV*rhoV)/rho - .5*(B*B));
104
105 }
106 }

```

Here is the caller graph for this function:



### 5.1.3.27 real Cell::psi ( ) const [inline]

Returns the cell-average density.

Returns

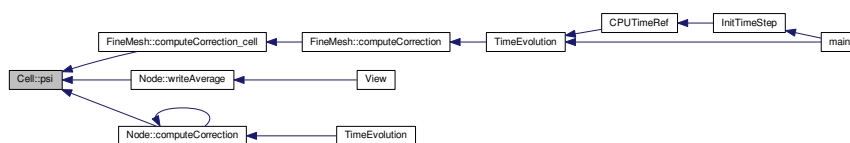
real

```

1281 {
1282     return Q.value(6);
1283 }

```

Here is the caller graph for this function:



5.1.3.28 `real Cell::PsiGrad ( const int Dimension ) const` `[inline]`

Returns the component of PsiGrad vector.

## Parameters

<i>Dimension</i>
------------------

## Returns

real

```
1209 {
1210     return PGrad.value(Dimension);
1211 }
```

Here is the caller graph for this function:

5.1.3.29 `Vector Cell::PsiGrad ( ) const` `[inline]`

Returns the PsiGrad vector.

## Returns

Vector

```
1217 {
1218     return PGrad;
1219 }
```

5.1.3.30 `void Cell::setAverage ( const int QuantityNo, const real UserAverage )` `[inline]`

Sets the cell-average of the quantity *QuantityNo* to *UserAverage*. Example:

```
#include "Carmen.h"
QuantityNb = 2;
real T=0.,Y=1.;
Cell c;
c.setAverage(1,T);
c.setAverage(2,Y);
```



## Returns

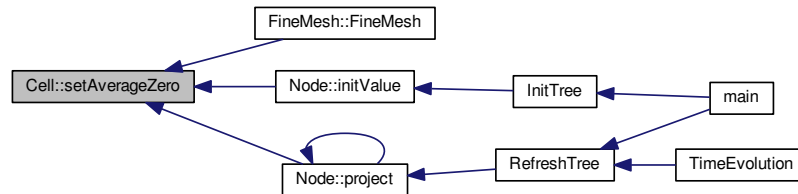
void

```

938 {
939     Q.setZero();
940 }

```

Here is the caller graph for this function:



### 5.1.333 void Cell::setCenter ( const int *AxisNo*, const real *UserX* ) [inline]

Sets the coordinate of the cell-center in the direction *AxisNo* to *UserX*. Example:

```

#include "Carmen.h"
Dimension = 3;
real x=1., y=0., z=0.;
Cell c;
c.setCenter(1,x);
c.setCenter(2,y);
c.setCenter(3,z);

```

Parameters

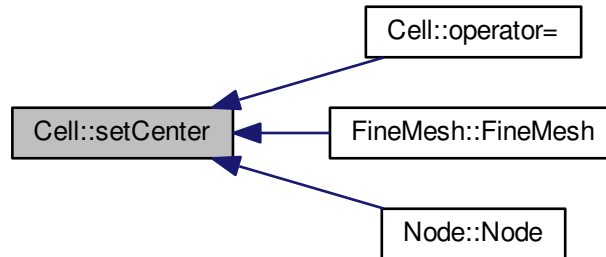
<i>AxisNo</i>	Number of axis of interest. 1: x-direction, 2: y-direction, 3: z-direction.
<i>UserX</i>	Center of the cell.

Returns

```
void
```

```
906 {
907     X.setValue(AxisNo, UserX);
908 }
```

Here is the caller graph for this function:



#### 5.1.334 void Cell::setCenter ( const Vector & UserX ) [inline]

Sets the position of the cell-center to the vector *UserX*. Example:

```
#include "Carmen.h"
Dimension = 3;
Vector V(1.,0.,0.);
Cell c;
c.setCenter(V);
```

Parameters

<i>UserX</i>	Center of the cell.
--------------	---------------------

Returns

```
void
```

```
914 {
915     X = UserX;
916 }
```

#### 5.1.335 void Cell::setDivergence ( const int QuantityNo, const real UserAverage ) [inline]

Identical to void [setAverage](#) (int QuantityNo, real UserAverage), but for the divergence vector.



## Parameters

<i>QuantityNo</i>	Number of MHD variables
<i>UserAverage</i>	Divergence values

## Returns

void

```

1020 {
1021     D.setValue(QuantityNo, UserAverage);
1022 }

```

Here is the caller graph for this function:



### 5.1.3.36 void Cell::setDivergence ( const Vector & UserAverage ) [inline]

Identical to void [setAverage](#) (const Vector& UserAverage), but for the divergence vector.

## Parameters

<i>UserAverage</i>	Divergence values
--------------------	-------------------

## Returns

void

```

1036 {
1037     D = UserAverage;
1038 }

```

### 5.1.3.37 void Cell::setDivergenceZero ( ) [inline]

Sets all the components of the divergence vector to zero.

## Returns

void

```

1044 {
1045     D.setZero();
1046 }

```

### 5.1.3.38 void Cell::setGradient ( const int i, const int j, const real UserAverage ) [inline]

Sets the component no. *i, j* of the quantity gradient to *UserAverage*.

Does not work for MHD!

Example:

```
#include "Carmen.h"
```

```

QuantityNb = 2;
real Gxx=0., Gxy=1., Gyx=1., Gyy=0.;
Cell c;
c.setGradient(1,1,Gxx);
c.setGradient(1,2,Gxy);
c.setGradient(2,1,Gyx);
c.setGradient(2,2,Gyy);

```

Parameters

<i>i</i>	
<i>j</i>	
<i>UserAverage</i>	Gradient value

Returns

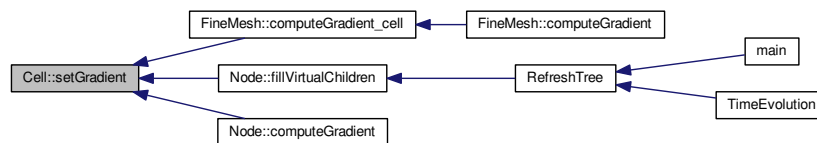
void

```

1052 {
1053     Grad.setValue(i, j, UserAverage);
1054 }

```

Here is the caller graph for this function:



### 5.1.3.39 void Cell::setGradient ( const Matrix & UserAverage ) [inline]

Sets the quantity gradient to the matrix *UserAverage*. Does not work for MHD! Example:

```
#include "Carmen.h"
```

```
QuantityNb = 5;
```

```
Dimension = 3;
```

```
Matrix G(3,5);
```

- Cell c
  - ...
- ```
c.SetGradient(G);
```

Parameters

|                    |                |
|--------------------|----------------|
| <i>UserAverage</i> | Gradient value |
|--------------------|----------------|

Returns

void

```

1068 {
1069     Grad = UserAverage;
1070 }

```

## 5.1.3.40 void Cell::setGradientZero ( ) [inline]

Sets all the components of the velocity gradient to zero.

## Returns

void

```
1085 {
1086     Grad.setZero();
1087 }
```

5.1.3.41 void Cell::setLowAverage ( const int *QuantityNo*, const real *UserAverage* ) [inline]

Identical to [setAverage](#) (int *QuantityNo*, real *UserAverage*), but for the vector of the cell-average values with low precision in the Runge-Kutta-Fehlberg method.

## Parameters

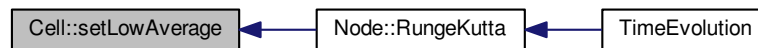
|                    |                         |
|--------------------|-------------------------|
| <i>QuantityNo</i>  | Number of MHD variables |
| <i>UserAverage</i> | Average value           |

## Returns

void

```
970 {
971     Qlow.setValue(QuantityNo, UserAverage);
972 }
```

Here is the caller graph for this function:

5.1.3.42 void Cell::setLowAverage ( const Vector & *UserAverage* ) [inline]

Identical to void [setAverage](#) (const [Vector](#)& *UserAverage*), void [setAverage](#) (const [Vector](#)& *UserAverage*), but for the vector of the cell-average values with low precision in the Runge-Kutta-Fehlberg method.

## Parameters

|                    |               |
|--------------------|---------------|
| <i>UserAverage</i> | Average value |
|--------------------|---------------|

## Returns

void

```
978 {
979     Qlow = UserAverage;
980 }
```

5.1.3.43 void Cell::setOldAverage ( const int *QuantityNo*, const real *UserAverage* ) [inline]

Identical to [setAverage](#) (int *QuantityNo*, real *UserAverage*), but for the vector of the old cell-average values.

## Parameters

|                    |                         |
|--------------------|-------------------------|
| <i>QuantityNo</i>  | Number of MHD variables |
| <i>UserAverage</i> | Average value           |

## Returns

void

```

986 {
987     Qold.setValue(QuantityNo, UserAverage);
988 }
```

Here is the caller graph for this function:



#### 5.1.3.44 void Cell::setOldAverage ( const Vector & UserAverage ) [inline]

Identical to void [setAverage](#) (const Vector& UserAverage), but for the vector of the cell-average values.

## Parameters

|                    |               |
|--------------------|---------------|
| <i>UserAverage</i> | Average value |
|--------------------|---------------|

## Returns

void

```

994 {
995     Qold = UserAverage;
996 }
```

#### 5.1.3.45 void Cell::setPsiGrad ( const int Dimension, const real UserAverage ) [inline]

Identical to void [setAverage](#) (int QuantityNo, real UserAverage), but for the gradient of psi vector.

## Parameters

|                    |                         |
|--------------------|-------------------------|
| <i>Dimension</i>   | Dimension of the system |
| <i>UserAverage</i> | Gradient of psi value   |

## Returns

void

```

1003 {
1004     PGrad.setValue(Dimension, UserAverage);
1005 }
```

Here is the caller graph for this function:



#### 5.1.3.46 void Cell::setPsiGrad ( const Vector & UserAverage ) [inline]

Identical to void `setAverage` (int QuantityNo, real UserAverage), but for the gradient of psi vector.

##### Parameters

|                    |                       |
|--------------------|-----------------------|
| <i>UserAverage</i> | Gradient of psi value |
|--------------------|-----------------------|

##### Returns

void

```

1028 {
1029     PGrad = UserAverage;
1030 }
  
```

#### 5.1.3.47 void Cell::setRes ( const real UserAverage ) [inline]

Set resistivity.

##### Parameters

|                    |                    |
|--------------------|--------------------|
| <i>UserAverage</i> | Resistivity values |
|--------------------|--------------------|

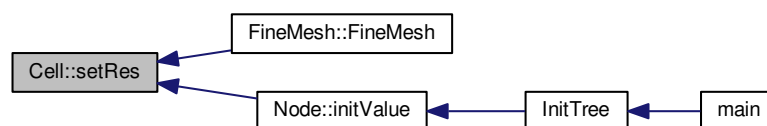
##### Returns

void

```

1012 {
1013     Res = UserAverage;
1014 }
  
```

Here is the caller graph for this function:



#### 5.1.3.48 void Cell::setSize ( const int *AxisNo*, const real *UserSize* ) [inline]

Sets the size of the cell in the direction *AxisNo* to *UserSize*. Example:

```
#include "Carmen.h"
Dimension = 2;
real dx=1., dy=2.;
Cell c;
c.setSize(1, dx);
c.setSize(2, dy);
```

Parameters

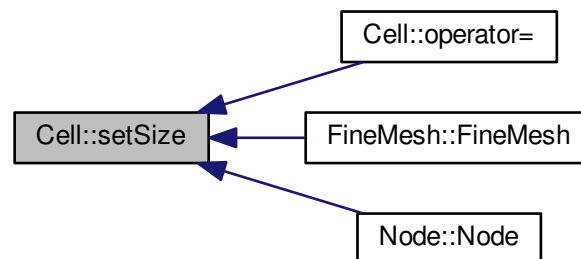
|                 |                                                                             |
|-----------------|-----------------------------------------------------------------------------|
| <i>AxisNo</i>   | Number of axis of interest. 1: x-direction, 2: y-direction, 3: z-direction. |
| <i>UserSize</i> | Size of the cell.                                                           |

Returns

void

```
889 {
890     dx.setValue(AxisNo, UserSize);
891 }
```

Here is the caller graph for this function:



#### 5.1.3.49 void Cell::setSize ( const Vector & *UserSize* ) [inline]

Sets the size of the cell in every direction to the vector *UserSize*. Example:

```
#include "Carmen.h"
Dimension = 2;
Vector W(1.,2.);
Cell c;
c.setSize(W);
```

## Parameters

|                 |                           |
|-----------------|---------------------------|
| <i>UserSize</i> | Size of the cell (vector) |
|-----------------|---------------------------|

## Returns

void

```
898 {
899     dx = UserSize;
900 }
```

### 5.1.3.50 void Cell::setTempAverage ( const int *QuantityNo*, const real *UserAverage* ) [inline]

Identical to [setAverage](#) (int *QuantityNo*, real *UserAverage*), but for the vector of the temporary cell-average values.

## Parameters

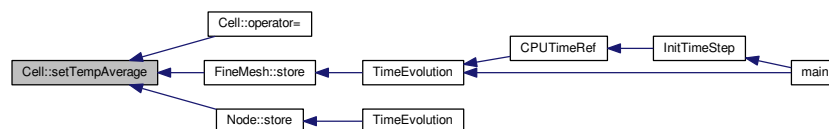
|                    |                         |
|--------------------|-------------------------|
| <i>QuantityNo</i>  | Number of MHD variables |
| <i>UserAverage</i> | Average value           |

## Returns

void

```
946 {
947     Qs.setValue(QuantityNo, UserAverage);
948 }
```

Here is the caller graph for this function:



### 5.1.3.51 void Cell::setTempAverage ( const Vector & *UserAverage* ) [inline]

Identical to void [setAverage](#) (const [Vector](#)& *UserAverage*), but for the vector of the temporary cell-average values.

## Parameters

|                    |               |
|--------------------|---------------|
| <i>UserAverage</i> | Average value |
|--------------------|---------------|

## Returns

void

```
954 {
955     Qs = UserAverage;
956 }
```

### 5.1.3.52 void Cell::setTempAverageZero ( ) [inline]

Sets all the temporary cell-average values to zero.

#### Returns

void

```
962 {
963     Qs.setZero();
964 }
```

### 5.1.3.53 void Cell::setTempGradient ( const int *i*, const int *j*, const real *UserAverage* ) [inline]

Identical to the previous one for the temporary values. Does not work for MHD!

#### Parameters

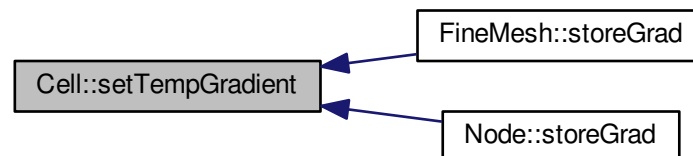
|                    |                |
|--------------------|----------------|
| <i>i</i>           |                |
| <i>j</i>           |                |
| <i>UserAverage</i> | Gradient value |

#### Returns

void

```
1060 {
1061     Grads.setValue(i, j, UserAverage);
1062 }
```

Here is the caller graph for this function:



### 5.1.3.54 void Cell::setTempGradient ( const Matrix & *UserAverage* ) [inline]

identical to the previous one for the temporary values.

#### Parameters

|                    |                |
|--------------------|----------------|
| <i>UserAverage</i> | Gradient value |
|--------------------|----------------|

#### Returns

void

```
1076 {
1077     Grads = UserAverage;
1078 }
```



---

5.1.3.55 `real Cell::size ( const int AxisNo ) const` `[inline]`

Returns the cell size in the direction *AxisNo*.



5.1.3.57 `real Cell::speedOfSound ( ) const [inline]`

Returns the cell-average speed of sound.

## Returns

real

```
1372 {
1373     return sqrt(Gamma*pressure()/density());
1374 }
```

5.1.3.58 `real Cell::tempAverage ( const int QuantityNo ) const [inline]`

Returns the component no. *QuantityNo* of the temporary cell-average value.

## Parameters

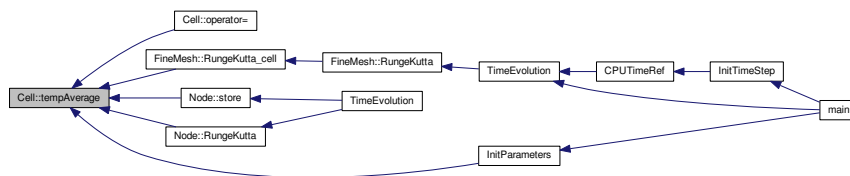
|                   |                              |
|-------------------|------------------------------|
| <i>QuantityNo</i> | Number of MHD variables = 9. |
|-------------------|------------------------------|

## Returns

real

```
1145 {
1146     return Qs.value(QuantityNo);
1147 }
```

Here is the caller graph for this function:

5.1.3.59 `Vector Cell::tempAverage ( ) const [inline]`

Returns the temporary cell-average value vector.

## Returns

Vector

```
1153 {
1154     return Qs;
1155 }
```

5.1.3.60 `real Cell::tempDensity ( ) const [inline]`

Identical to the previous one for the temporary values.

**Returns**

real

```

1272 {
1273     return Qs.value(1);
1274 }

```

Here is the caller graph for this function:

**5.1.3.61 real Cell::tempEnergy ( ) const [inline]**

Identical to the previous one for the temporary values.

**Returns**

real

```

1317 {
1318     return Qs.value(5);
1319 }

```

**5.1.3.62 real Cell::temperature ( ) const**

Returns the cell-average temperature. Does not work for MHD!

Computes the temperature. Not useful for MHD computations.

**Returns**real  
double

```

185 {
186
187     if(!ComputeTemp) exit(1);
188
189     // Conservative quantities;
190     real rho, p, T;
191
192     // Get conservative quantities
193     rho = density();
194     p = pressure();
195     T = p/rho;
196
197     // Return temperature
198
199     return T;
200 }

```

Here is the caller graph for this function:



### 5.1.3.63 `real Cell::tempGradient ( const int i, const int j ) const [inline]`

Identical to the previous one for the temporary values.

#### Parameters

|          |  |
|----------|--|
| <i>i</i> |  |
| <i>j</i> |  |

#### Returns

real

```

1236 {
1237     return Grads.value(i, j);
1238 }
  
```

### 5.1.3.64 `Matrix Cell::tempGradient ( ) const [inline]`

Identical to the previous one for the temporary values.

#### Returns

Matrix

```

1255 {
1256     return Grads;
1257 }
  
```

### 5.1.3.65 `real Cell::tempMagField ( const int AxisNo ) const [inline]`

Identical to the previous one for the temporary values.

#### Parameters

|               |  |
|---------------|--|
| <i>AxisNo</i> |  |
|---------------|--|

#### Returns

real

```

1346 {
1347     return Qs.value(6+AxisNo);
1348 }
  
```

### 5.1.3.66 Vector Cell::tempMagField ( ) const

Identical to the previous one for the temporary values.

Computes temporary the magnetic field. This value is useful for time integration computations.

#### Returns

Vector

```

267 {
268
269     // Local variables
270
271     Vector B(3);
272     int i;
273
274     for (i=1; i<=3; i++)
275         B.setValue(i,Qs.value(i+6));
276
277     return B;
278 }
```

### 5.1.3.67 real Cell::tempPressure ( ) const

Identical to the previous one for the temporary values.

Computes the temporary pressure of the fluid. This computation is needed for the Runge-Kutta time integration.

#### Returns

real  
double

```

118 {
119     // Conservative quantities;
120
121     real rho, rhoE;
122     Vector rhoV(3), B(3);
123
124     // Get conservative quantities
125
126     rho = Qs.value(1);
127
128     for (int i=1 ; i<=3 ; i++){
129         rhoV.setValue(i,Qs.value(i+1));
130         B.setValue(i, Qs.value(i+6));
131     }
132
133     rhoE = Qs.value(5);
134
135     // Return pressure
136
137     return (Gamma-1.)*(rhoE - .5*(rhoV*rhoV)/rho - .5*(B*B));
138
139 }
```

Here is the caller graph for this function:



**5.1.3.68** `real Cell::tempPsi( ) const` `[inline]`

Identical to the previous one for the temporary values.

**Returns**

real

```
1290 {
1291     return Qs.value(6);
1292 }
```

**5.1.3.69** `real Cell::tempTemperature ( ) const`

Identical to the previous one for the temporary values. Does not work for MHD!

Computes the temporary temperature. Not useful for MHD computations.

**Returns**

real  
double

```
213 {
214
215     if(!ComputeTemp) exit(1);
216
217     // Conservative quantities;
218     real rho, p, T;
219     T = Qs.value(1);
220
221     // Get conservative quantities
222     rho = tempDensity();
223     p = tempPressure();
224     T = Gamma*Ma*Ma*p/rho;
225
226
227     // Return temperature
228
229     return T;
230 }
```

**5.1.3.70** `real Cell::tempVelocity ( const int AxisNo ) const` `[inline]`

Identical to the previous one for the temporary values.

**Parameters**

|               |  |
|---------------|--|
| <i>AxisNo</i> |  |
|---------------|--|

**Returns**

real

```
1364 {
1365     return Qs.value(1+AxisNo)/Qs.value(1);
1366 }
```

**5.1.3.71** `Vector Cell::tempVelocity ( ) const`

Identical to the previous one for the temporary values.

Computes the temporary velocity of the fluid. This value is useful for time integration computations.

## Returns

Vector

```

315 {
316
317     // Local variables
318
319     Vector V(3);
320     int i;
321
322     for (i=1; i<=3; i++)
323         V.setValue(i,Qs.value(i+1)/Qs.value(1));
324
325     return V;
326 }

```

## 5.1.3.72 real Cell::velocity ( const int AxisNo ) const [inline]

Returns the component no. *AxisNo* of the cell-average velocity.

## Parameters

|               |
|---------------|
| <i>AxisNo</i> |
|---------------|

## Returns

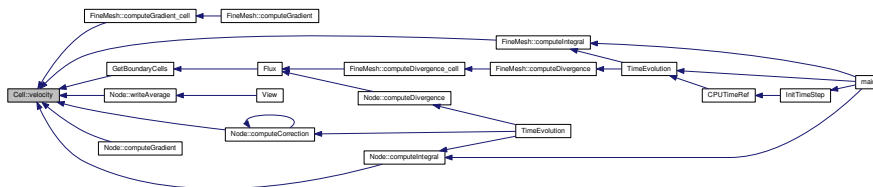
real

```

1355 {
1356     return Q.value(1+AxisNo)/Q.value(1);
1357 }

```

Here is the caller graph for this function:



## 5.1.3.73 Vector Cell::velocity ( ) const

Returns the cell-average velocity vector.

Computes the velocity of the fluid. Allocates the velocity initial condition  $Q[i+6]$  to its components.

## Returns

Vector

```

291 {
292
293     // Local variables
294
295     Vector V(3);
296     int i;
297
298     for (i=1; i<=3; i++)
299         V.setValue(i,Q.value(i+1)/Q.value(1));
300
301     return V;
302 }

```



### 5.1.3.74 `real Cell::volume ( ) const`

Returns the volume of the cell (length in 1D, area in 2D, volume in 3D).

Computes the volume of a cell.

#### Returns

`real`  
`double`

```

364 {
365     int AxisNo = 1;
366     real result = 1.;
367
368     for (AxisNo = 1; AxisNo <= Dimension; AxisNo++)
369         result *= size(AxisNo);
370
371     return result;
372
373 }
```

## 5.1.4 Member Data Documentation

### 5.1.4.1 `Vector Cell::D`

Divergence vector. Its dimension is *QuantityNb*.

### 5.1.4.2 `Vector Cell::dX`

`Cell` size in each direction. Its dimension is *Dimension*.

### 5.1.4.3 `Matrix Cell::Grad`

Quantity gradient. Only necessary for a Navier-Stokes computation. Not working!

### 5.1.4.4 `Matrix Cell::Grads`

Temporary quantity gradient. Only necessary for a Navier-Stokes computation. Not working!

### 5.1.4.5 `Vector Cell::PGrad`

Gradient of psi scalar function. Its dimension is *Dimension*.

### 5.1.4.6 `Vector Cell::Q`

`Vector` containing the cell-average values. Its dimension is *QuantityNb*.

### 5.1.4.7 `Vector Cell::Qlow`

This vector is used to store the cell-averages computed with the N-stage Runge-Kutta-Fehlberg N(N+1) method.

### 5.1.4.8 `Vector Cell::Qold`

Cell-average values at the instant *n-1*.

#### 5.1.4.9 Vector Cell::Qs

Temporary cell-average values. This vector is used to store the intermediary value in a multi-step Runge-Kutta or McCormack time integration.

#### 5.1.4.10 real Cell::Res

Resistivity scalar function.

#### 5.1.4.11 Vector Cell::X

Position of the cell center. Its dimension is *Dimension*.

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

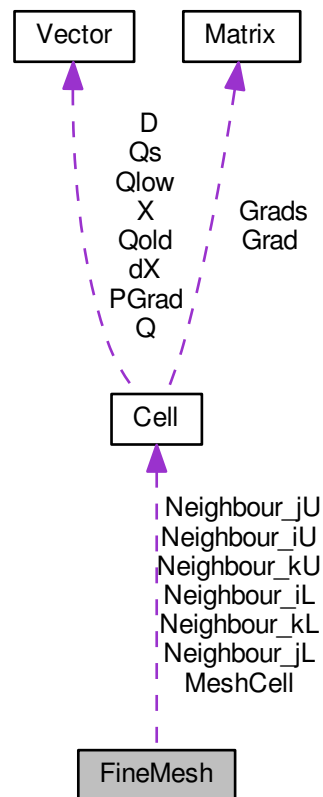
- [Cell.h](#)
- [Cell.cpp](#)

## 5.2 FineMesh Class Reference

An object [FineMesh](#) is a regular fine mesh, used for finite volume computations. It is not used for multiresolution computations.

```
#include <FineMesh.h>
```

Collaboration diagram for FineMesh:



## Public Member Functions

- [FineMesh](#) ()
  - Constructor of [FineMesh](#) class Generates a regular fine mesh containing  $2 \times (\text{Dimension} \times \text{ScaleNb})$  cells. The cell-averages are initialized from the initial condition contained in file `carmen.ini`.*
- [~FineMesh](#) ()
  - Destructor the regular fine mesh.*
- void [store](#) ()
  - Stores cell-average values into temporary cell-average values.*
- void [storeGrad](#) ()
  - Stores gradient values into temporary gradient values.*
- void [computeDivergence](#) (int)
  - Computes the divergence vector with the space discretization scheme.*
- void [computeDivergence\\_cell](#) (int)
  - Computes one [Cell](#) Divergence.*
- void [RungeKutta\\_cell](#) (int)
  - Computes one Runge-Kutta step.*
- void [RungeKutta](#) (int)
  - Computes the Runge-Kutta step.*
- void [computeIntegral](#) ()

- Computes integral values like e.g. flame velocity, global error, etc.*

  - void [computeCorrection](#) (int)
    - Computes divergence cleaning source term (only for MHD).*
  - void [computeCorrection\\_cell](#) (int)
    - Computes divergence cleaning source term (only for MHD) at one cell.*
  - void [computeGradient](#) (int)
    - Computes velocity gradient (only for Navier-Stokes). one cell.*
  - void [computeGradient\\_cell](#) (int)
    - Computes velocity gradient (only for Navier-Stokes). each cells.*
  - void [computeTimeAverage](#) ()
    - Computes the time-average value in every cell.*
  - void [checkStability](#) () const
    - Checks if the computation is numerically unstable, i.e. if one of the cell-averages is overflow. In case of numerical instability, the computation is stopped and a message appears.*
  - void [writeHeader](#) (const char \*FileName) const
    - Write header for GNUplot, Data Explorer, TecPLot and VTK into file FileName.*
  - void [writeAverage](#) (const char \*FileName)
    - Write cell-averages for GNUplot, Data Explorer, TecPLot and VTK into file FileName.*
  - void [writeTimeAverage](#) (const char \*FileName) const
    - Write time-averages into file FileName.*
  - void [backup](#) ()
    - Backs up the tree structure and the cell-averages into a file carmen.bak. In further computations, the data can be recovered using **Restore()**.*
  - void [restore](#) ()
    - Restores the tree structure and the cell-averages from the file carmen.bak. This file was created by the method **Backup()**.*

## Public Attributes

- Cell \*\*\* [Neighbour\\_iL](#)
- Cell \*\*\* [Neighbour\\_iU](#)
- Cell \*\*\* [Neighbour\\_jL](#)
- Cell \*\*\* [Neighbour\\_jU](#)
- Cell \*\*\* [Neighbour\\_kL](#)
- Cell \*\*\* [Neighbour\\_kU](#)
- Cell \* [MeshCell](#)

### 5.2.1 Detailed Description

An object [FineMesh](#) is a regular fine mesh, used for finite volume computations. It is not used for multiresolution computations.

It contains an array of cells *\*MeshCell*.

NOTE: for reasons of simplicity, only periodic and Neuman boundary conditions have been implemented.

### 5.2.2 Constructor & Destructor Documentation

#### 5.2.2.1 [FineMesh::FineMesh](#) ( )

Constructor of [FineMesh](#) class Generates a regular fine mesh containing  $2^{**}(Dimension*ScaleNb)$  cells. The cell-averages are initialized from the initial condition contained in file *carmen.ini*.

!!DEBUG

```

34 {
35     // --- Local variables ---
36
37     int n=0, i=0, j=0, k=0;           // position numbers
38
39     real x=0., y=0., z=0.; // positions
40     real dx=0., dy=0., dz=0.; // space steps
41
42     // --- Create an array of 2^(ScaleNb*Dimension) cells ---
43
44     MeshCell = new Cell[(1<<(ScaleNb*Dimension))];
45
46 //Parallel
47
48 #if defined PARMPI
49     Neighbour_iL = new Cell**[NeighbourNb];
50     Neighbour_iU = new Cell**[NeighbourNb];
51     Neighbour_jL = new Cell**[NeighbourNb];
52     Neighbour_jU = new Cell**[NeighbourNb];
53     Neighbour_kL = new Cell**[NeighbourNb];
54     Neighbour_kU = new Cell**[NeighbourNb];
55
56     for (i=0;i<NeighbourNb;i++) {
57         Neighbour_iL[i] = new Cell*[one_D];
58         Neighbour_iU[i] = new Cell*[one_D];
59         Neighbour_jL[i] = new Cell*[one_D];
60         Neighbour_jU[i] = new Cell*[one_D];
61         Neighbour_kL[i] = new Cell*[one_D];
62         Neighbour_kU[i] = new Cell*[one_D];
63     }
64
65     for (i=0;i<NeighbourNb;i++)
66         for (j=0;j<one_D;j++) {
67             Neighbour_iL[i][j] = new Cell[two_D];
68             Neighbour_iU[i][j] = new Cell[two_D];
69             Neighbour_jL[i][j] = new Cell[two_D];
70             Neighbour_jU[i][j] = new Cell[two_D];
71             Neighbour_kL[i][j] = new Cell[two_D];
72             Neighbour_kU[i][j] = new Cell[two_D];
73         }
74 #endif
75
76
77
78     // --- Create a time-average grid ---
79
80     if (TimeAveraging)
81         MyTimeAverageGrid = new TimeAverageGrid(ScaleNb);
82
83     // --- Compute dx, dy, dz ---
84
85     dx = (XMax[1]-XMin[1])/(1<<ScaleNb);
86     if (Dimension > 1) dy = (XMax[2]-XMin[2])/(1<<ScaleNb);
87     if (Dimension > 2) dz = (XMax[3]-XMin[3])/(1<<ScaleNb);
88
89     // --- Loop on all cells ---
90
91     for (n = 0; n < 1<<(ScaleNb*Dimension); n++)
92     {
93         // -- Compute i, j, k --
94
95         i = n%(1<<ScaleNb);
96         if (Dimension > 1) j = (n%(1<<(2*ScaleNb)))/(1<<ScaleNb);
97         if (Dimension > 2) k = n/(1<<(2*ScaleNb));
98         // -- Compute x, y, z --
99
100        x = XMin[1] + (i+.5)*dx;
101        if (Dimension > 1) y = XMin[2] + (j+.5)*dy;
102        if (Dimension > 2) z = XMin[3] + (k+.5)*dz;
103
104        // -- Set position --
105
106        cell(n)->setCenter(1,x);
107        if (Dimension > 1) cell(n)->setCenter(2,y);
108        if (Dimension > 2) cell(n)->setCenter(3,z);
109
110        // -- Set size --
111
112        cell(n)->setSize(1,dx);
113        if (Dimension > 1) cell(n)->setSize(2,dy);
114        if (Dimension > 2) cell(n)->setSize(3,dz);
115    }
116    // --- End loop on all cells ---
117
118 //Parallel
119
120 #if defined PARMPI

```

```

121
122     for (i=0;i<one_D;i++)
123         for (j=0;j<two_D;j++)
124             for (k=0;k<NeighbourNb;k++) {
125                 Neighbour_iL[k][i][j].setSize(1,dx);
126                 Neighbour_iU[k][i][j].setSize(1,dx);
127                 Neighbour_jL[k][i][j].setSize(1,dx);
128                 Neighbour_jU[k][i][j].setSize(1,dx);
129                 Neighbour_kL[k][i][j].setSize(1,dx);
130                 Neighbour_kU[k][i][j].setSize(1,dx);
131
132                 if (Dimension > 1) {
133                     Neighbour_iL[k][i][j].setSize(2,dy);
134                     Neighbour_iU[k][i][j].setSize(2,dy);
135                     Neighbour_jL[k][i][j].setSize(2,dy);
136                     Neighbour_jU[k][i][j].setSize(2,dy);
137                     Neighbour_kL[k][i][j].setSize(2,dy);
138                     Neighbour_kU[k][i][j].setSize(2,dy);
139                 }
140
141                 if (Dimension > 2) {
142                     Neighbour_iL[k][i][j].setSize(3,dz);
143                     Neighbour_iU[k][i][j].setSize(3,dz);
144                     Neighbour_jL[k][i][j].setSize(3,dz);
145                     Neighbour_jU[k][i][j].setSize(3,dz);
146                     Neighbour_kL[k][i][j].setSize(3,dz);
147                     Neighbour_kU[k][i][j].setSize(3,dz);
148                 }
149             }
150
151 #endif
152
153
154
155     // -- Set initial cell-average value --
156
157
158     /*
159     printf("\nRecovery: %d",Recovery);
160     printf("\nUseBackup: %d",UseBackup);
161     printf("\nComputeCPUTimeRef: %d\n",ComputeCPUTimeRef);
162 */
163
164
165     if (Recovery && UseBackup && !ComputeCPUTimeRef) {
166         // printf("\nRestore!\n");
167         restore();
168     }
169     else
170     {
171         for (n = 0; n < 1<<(ScaleNb*Dimension); n++)
172         {
173             cell(n)->setAverageZero();
174
175             if (UseBoundaryRegions && BoundaryRegion(cell(n)->center()) !=
176 0)
177             {
178                 x = cell(n)->center(1);
179                 y = (Dimension > 1)? cell(n)->center(2): 0.;
180                 z = (Dimension > 2)? cell(n)->center(3): 0.;
181                 cell(n)->setAverage(InitAverage(x,y,z));
182             }
183             else
184             {
185                 switch (Dimension)
186                 {
187                     case 1:
188                         for (i=0;i<=1;i++)
189                             cell(n)->setAverage( cell(n)->average()+.5*
190 InitAverage(
191                                 cell(n)->center(1)+(i-0.5)*cell(n)->size(1) ) );
192                         break;
193                     case 2:
194                         if(IcNb){
195                             for (i=0;i<=1;i++)
196                                 for (j=0;j<=1;j++){
197                                     cell(n)->setAverage( cell(n)->average()+.25*
198 InitAverage(
199                                         cell(n)->center(1)+(i-0.5)*cell(n)->size(1),
200                                         cell(n)->center(2)+(j-0.5)*cell(n)->size(2) ) );
201                                     cell(n)->setRes(InitResistivity(cell(n)->center(1)
202 ,cell(n)->center(2)));
203                                 }
204                             }else{
205                                 cell(n)->setAverage(InitAverage(cell(n)->center(1),cell(n)
206 ->center(2)));

```

```

204         cell(n)->setRes(InitResistivity(cell(n)->center(1),cell(n)
->center(2)));
205     }
206     break;
207
208     case 3:
209         if(IcNb){
210             for (i=0;i<=1;i++)
211                 for (j=0;j<=1;j++)
212                     for (k=0;k<=1;k++){
213                         cell(n)->setAverage( cell(n)->average()+.125*
InitAverage(
214                             cell(n)->center(1)+(i-0.5)*cell(n)->size(1),
215                             cell(n)->center(2)+(j-0.5)*cell(n)->size(2),
216                             cell(n)->center(3)+(k-0.5)*cell(n)->size(3) ) );
217                         cell(n)->setRes(InitResistivity(cell(n)->
center(1),cell(n)->center(2),cell(n)->center(3)));
218                     }
219                 }else{
220                     cell(n)->setAverage(InitAverage(cell(n)->center(1),cell(n)
->center(2),cell(n)->center(3)));
221                     cell(n)->setRes(InitResistivity(cell(n)->center(1),cell(n)
->center(2),cell(n)->center(3)));
222                 }
223             }
224         break;
225     };
226 }
227 }
228 }
229
230 #if defined PARMPI
231     //Important moment: Exchange boundary (neighbour) cells before start computation (1st iteration)
232
233     CPUEXchange(this, SendQ);
234     if (MPIRecvType == 1) MPI_Waitall(4*Dimension,req,st); //Send quantity number
one (code name "Q")
235
236     if (EquationType==6) {
237         CPUEXchange(this, SendGrad);
238         if (MPIRecvType == 1) MPI_Waitall(4*Dimension,req,st); //Send gradient
239     }
240 #endif
241
242 }

```

### 5.2.2.2 FineMesh::~FineMesh ( )

Destructor the regular fine mesh.

```

251 {
252     // --- Delete pointers to cells ---
253
254     delete[] MeshCell;
255
256     #if defined PARMPI
257         int i,j;
258
259         for (i=0;i<NeighbourNb;i++)
260             for (j=0;j<one_D;j++){
261                 delete[] Neighbour_iL[i][j];
262                 delete[] Neighbour_iU[i][j];
263                 delete[] Neighbour_jL[i][j];
264                 delete[] Neighbour_jU[i][j];
265                 delete[] Neighbour_kL[i][j];
266                 delete[] Neighbour_kU[i][j];
267             }
268
269         for (i=0;i<NeighbourNb;i++) {
270             delete[] Neighbour_iL[i];
271             delete[] Neighbour_iU[i];
272             delete[] Neighbour_jL[i];
273             delete[] Neighbour_jU[i];
274             delete[] Neighbour_kL[i];
275             delete[] Neighbour_kU[i];
276         }
277
278         delete[] Neighbour_iL;
279         delete[] Neighbour_iU;
280         delete[] Neighbour_jL;
281         delete[] Neighbour_jU;
282         delete[] Neighbour_kL;

```

```

283     delete[] Neighbour_kU;
284 #endif
285
286
287     if (TimeAveraging)
288         delete MyTimeAverageGrid;
289 }

```

## 5.2.3 Member Function Documentation

### 5.2.3.1 void FineMesh::backup ( )

Backs up the tree structure and the cell-averages into a file *carmen.bak*. In further computations, the data can be recovered using **Restore()**.

#### Returns

void

```

1742 {
1743     int n=0;                // Cell number
1744     FILE* output;         // Output file
1745     int QuantityNo;       // Counter on quantities
1746
1747     // --- Init ---
1748
1749     output = fopen(BackupName,"w");
1750
1751     // --- Backup data on cells ---
1752
1753     fprintf(output, "Backup at iteration %i, physical time %e\n",
1754             IterationNo, ElapsedTime);
1755     for (n = 0; n < 1<<(ScaleNb*Dimension); n++)
1756         for (QuantityNo=1; QuantityNo <= QuantityNb; QuantityNo++)
1757             fprintf(output, FORMAT, cell(n)->average(QuantityNo));
1758     fclose(output);
1759 }

```

Here is the caller graph for this function:



### 5.2.3.2 void FineMesh::checkStability ( ) const

Checks if the computation is numerically unstable, i.e. if one of the cell-averages is overflow. In case of numerical instability, the computation is stopped and a message appears.

#### Returns

void

```

928 {
929     // --- Local variables ---
930
931     int n=0,iaux;          // cell number
932     real x=0.,y=0.,z=0.; // Real position
933     iaux = 0;

```

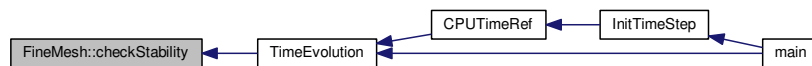


```

934 // --- Loop on all cells ---
935
936 for (n = 0; n < 1<<(ScaleNb*Dimension); n++)
937 {
938     // --- Compute x, y, z ---
939
940     x = cell(n)->center(1);
941     if (Dimension > 1) y = cell(n)->center(2);
942     if (Dimension > 2) z = cell(n)->center(3);
943
944     // --- Test if one cell is overflow ---
945
946     if (cell(n)->isOverflow())
947     {
948         iaux=system("echo Unstable computation.>> carmen.prf");
949         if (Cluster == 0) iaux=system("echo carmen: unstable computation. >> OUTPUT");
950         cout << "carmen: instability detected at iteration no. "<<
IterationNo <<"\n";
951         cout << "carmen: position ("<< x << ", "<<y<< ", "<<z<<")\n";
952         cout << "carmen: abort execution.\n";
953         exit(1);
954     }
955 }
956 // --- End loop on all cells ---
957 }

```

Here is the caller graph for this function:



### 5.2.3.3 void FineMesh::computeCorrection ( int mode )

Computes divergence cleaning source term (only for MHD).

#### Parameters

|            |                                                                   |
|------------|-------------------------------------------------------------------|
| <i>int</i> | It can be zero or one. Associated to the time integration scheme. |
|------------|-------------------------------------------------------------------|

#### Returns

void

```

541 {
542     int i, j, k, d;
543     // --- Loops for internal cells
544     if (mode==0) {
545         if (Dimension==1)
546             for (i=NeighbourNb; i<(1<<ScaleNb)-NeighbourNb; i++) {
547                 j=0; k=0;
548                 d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
549                 computeCorrection_cell(d);
550             }
551         if (Dimension==2)
552             for (i=NeighbourNb; i<(1<<ScaleNb)-NeighbourNb; i++)
553                 for (j=NeighbourNb; j<(1<<ScaleNb)-NeighbourNb; j++) {
554                     k=0;
555                     d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
556                     computeCorrection_cell(d);
557                 }
558         if (Dimension==3)
559             for (i=NeighbourNb; i<(1<<ScaleNb)-NeighbourNb; i++)
560                 for (j=NeighbourNb; j<(1<<ScaleNb)-NeighbourNb; j++)
561                     for (k=NeighbourNb; k<(1<<ScaleNb)-NeighbourNb; k++) {
562                         d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
563                         computeCorrection_cell(d);
564                     }
565     }
566 }

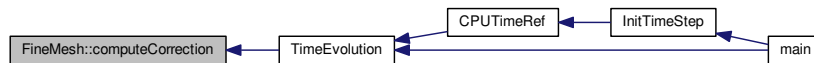
```

```

567         }
568     }
569
570 // --- loop for neighbour cells
571 if (mode==1) {
572
573     if (Dimension==1)
574         for (i=0;i<(1<<ScaleNb);i++)
575             if (i<NeighbourNb || i>=(1<<ScaleNb)-NeighbourNb) {
576                 j=0; k=0;
577                 d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
578                 computeCorrection_cell(d);
579             }
580
581     if (Dimension==2)
582         for (i=0;i<(1<<ScaleNb);i++)
583             for (j=0;j<one_D;j++)
584                 if (i<NeighbourNb || j<NeighbourNb ||
585                     i>=(1<<ScaleNb)-NeighbourNb || j>=(1<<
586                     ScaleNb)-NeighbourNb) {
587                     k=0;
588                     d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
589                     computeCorrection_cell(d);
590                 }
591
592     if (Dimension==3)
593         for (i=0;i<(1<<ScaleNb);i++)
594             for (j=0;j<one_D;j++)
595                 for (k=0;k<two_D;k++)
596                     if (i<NeighbourNb || j<NeighbourNb || k<
597                         NeighbourNb ||
598                             i>=(1<<ScaleNb)-NeighbourNb || j>=(1<<
599                             ScaleNb)-NeighbourNb || k>=(1<<ScaleNb)-NeighbourNb) {
600                             d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
601                             computeCorrection_cell(d);
602                         }
603 }
604 /*
605 int n;
606 for (n = 0; n < 1<<(ScaleNb*Dimension); n++) computeDivergence_cell(n);*/
607 }

```

Here is the caller graph for this function:



### 5.2.3.4 void FineMesh::computeCorrection\_cell ( int n )

Computes divergence cleaning source term (only for MHD) at one cell.

#### Parameters

|            |                                                                   |
|------------|-------------------------------------------------------------------|
| <i>int</i> | It can be zero or one. Associated to the time integration scheme. |
|------------|-------------------------------------------------------------------|

#### Returns

void

```

508 {
509     // --- Local variables ---
510     real rho=0., psi=0.; // Variables density and psi
511     int q=0; // Counters
512     real Bx=0.; // Magnetic field
513
514     // --- Computation ---
515
516
517     if(DivClean==1) // EGLM
518     {

```

```

519
520     rho = cell(n)->density();
521     psi = cell(n)->psi();
522
523     for (q=1; q <= 3; q++) //GLM
524     {
525         Bx = cell(n)->magField(q);
526         cell(n)->setAverage(q+1, cell(n)->average(q+1) -
TimeStep*Bx*Bdivergence/(ch*ch));
527     }
528
529     cell(n)->setAverage(5, cell(n)->average(5) - TimeStep*
PsiGrad);
530     cell(n)->setAverage(6,psi*exp(-(cr*ch*TimeStep/
SpaceStep)));
531
532     }else if(DivClean==2)
533     {
534         psi = cell(n)->psi();
535         cell(n)->setAverage(6,psi*exp(-(cr*ch*TimeStep/
SpaceStep)));
536     }
537
538
539 }

```

Here is the caller graph for this function:



### 5.2.3.5 void FineMesh::computeDivergence ( int mode )

Computes the divergence vector with the space discretization scheme.

#### Parameters

|            |                                                                   |
|------------|-------------------------------------------------------------------|
| <i>int</i> | It can be zero or one. Associated to the time integration scheme. |
|------------|-------------------------------------------------------------------|

#### Returns

void

```

427 {
428     int i,j,k,d;
429
430
431     // --- Loops for internal cells
432     if (mode==0)
433     {
434         if (Dimension==1)
435             for (i=2*NeighbourNb;i<(1<<<ScaleNb)-2*NeighbourNb;i++)
436             {
437                 j=0; k=0;
438                 d=i + (1<<<ScaleNb)*(j + (1<<<ScaleNb)*k);
439                 computeDivergence_cell(d);
440             }
441
442         if (Dimension==2)
443             for (i=2*NeighbourNb;i<(1<<<ScaleNb)-2*NeighbourNb;i++)
444                 for (j=2*NeighbourNb;j<(1<<<ScaleNb)-2*NeighbourNb;j++)
445                 {
446                     k=0;
447                     d=i + (1<<<ScaleNb)*(j + (1<<<ScaleNb)*k);
448                     computeDivergence_cell(d);
449                 }
450
451         if (Dimension==3)
452             for (i=2*NeighbourNb;i<(1<<<ScaleNb)-2*NeighbourNb;i++)
453                 for (j=2*NeighbourNb;j<(1<<<ScaleNb)-2*NeighbourNb;j++)
454                     for (k=2*NeighbourNb;k<(1<<<ScaleNb)-2*

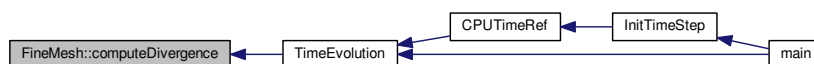
```

```

    NeighbourNb;k++) {
456         d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
457         computeDivergence_cell(d);
458     }
459 }
460
461 // --- loop for neighbour cells
462 if (mode==1) {
463
464     if (Dimension==1)
465         for (i=0;i<(1<<ScaleNb);i++)
466             if (i<2*NeighbourNb || i>=(1<<ScaleNb)-2*NeighbourNb) {
467                 j=0; k=0;
468                 d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
469                 computeDivergence_cell(d);
470             }
471
472     if (Dimension==2)
473         for (i=0;i<(1<<ScaleNb);i++)
474             for (j=0;j<one_D;j++)
475                 if (i<2*NeighbourNb || j<2*NeighbourNb ||
476                     i>=(1<<ScaleNb)-2*NeighbourNb || j>=(1<<
ScaleNb)-2*NeighbourNb) {
477                     k=0;
478                     d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
479                     computeDivergence_cell(d);
480                 }
481
482     if (Dimension==3)
483         for (i=0;i<(1<<ScaleNb);i++)
484             for (j=0;j<one_D;j++)
485                 for (k=0;k<two_D;k++)
486                     if (i<2*NeighbourNb || j<2*NeighbourNb || k<2*
NeighbourNb ||
487                         i>=(1<<ScaleNb)-2*NeighbourNb || j>=(1<<
ScaleNb)-2*NeighbourNb || k>=(1<<ScaleNb)-2*
NeighbourNb) {
488                         d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
489                         computeDivergence_cell(d);
490                     }
491 }
492 }
493
494 /*
495     int n;
496     for (n = 0; n < 1<<(ScaleNb*Dimension); n++) computeDivergence_cell(n);*/
497 }

```

Here is the caller graph for this function:



### 5.2.3.6 void FineMesh::computeDivergence\_cell ( int n )

Computes one [Cell](#) Divergence.

#### Parameters

|            |                                                                   |
|------------|-------------------------------------------------------------------|
| <i>int</i> | It can be zero or one. Associated to the time integration scheme. |
|------------|-------------------------------------------------------------------|

#### Returns

void

2D resistive part of the model added to the Flux

```
335 {
```

```

336 // --- Local variables ---
337
338 int i=0, j=0, k=0; // position numbers
339 Vector FluxIn, FluxOut; // ingoing and outgoing fluxes
340 real divCor=0.;
341
342 // --- Loop on all cells ---
343
344 // --- Only in fluid region ---
345
346 if (!UseBoundaryRegions || BoundaryRegion(cell(n)->center())==0)
347 {
348 // --- Compute source term --
349
350 cell(n)->setDivergence(Source(*cell(n)));
351
352 // -- Compute i, j, k --
353
354 i = n%(1<<ScaleNb);
355 if (Dimension > 1) j = (n%(1<<(2*ScaleNb)))/(1<<ScaleNb);
356 if (Dimension > 2) k = n/(1<<(2*ScaleNb));
357
358 // Add flux in x-direction
359
360 FluxIn = Flux( *cell(i-2, j, k), *cell(i-1, j, k), *cell(i, j, k), *cell(i+1, j, k), 1 );
361 divCor = -auxvar;
362 FluxOut = Flux( *cell(i-1, j, k), *cell(i, j, k), *cell(i+1, j, k), *cell(i+2, j, k), 1 );
363 divCor += auxvar;
364
365
366
367 if(Resistivity){
368 FluxIn = FluxIn - ResistiveTerms(*cell(i,j,k) , *cell(i-1,j,k), *cell(i,j-1,k)
369 , *cell(i,j,k-1) , 1);
369 FluxOut = FluxOut - ResistiveTerms(*cell(i+1,j,k), *cell(i,j,k) , *cell(i+1,j-1,
k), *cell(i+1,j,k-1), 1);
370 }
371
372 cell(n)->setDivergence( cell(n)->divergence() + (FluxIn - FluxOut)/(cell(n)->
size(1)));
373
374 // Variables \grad(psi) and \div(B) to evaluate GLM and EGLM divergence cleaning
375 PsiGrad = cell(n)->average(7)*(FluxOut.value(7) - FluxIn.
value(7) - divCor)/(cell(n)->size(1));
376 Bdivergence += (FluxOut.value(6) - FluxIn.value(6))/(cell(n)->
size(1));
377
378 // Add flux in y-direction
379
380 if (Dimension > 1)
381 {
382 FluxIn = Flux( *cell(i, j-2, k), *cell(i, j-1, k), *cell(i, j, k), *cell(i, j+1, k), 2 )
;
383 divCor = -auxvar;
384 FluxOut = Flux( *cell(i, j-1, k), *cell(i, j, k), *cell(i, j+1, k), *cell(i, j+2, k), 2 )
;
385 divCor += auxvar;
386
387 if(Resistivity){
388 FluxIn = FluxIn - ResistiveTerms(*cell(i,j,k) , *cell(i-1,j,k) , *cell(i,
j-1,k), *cell(i,j,k-1) , 2);
389 FluxOut = FluxOut - ResistiveTerms(*cell(i,j+1,k), *cell(i-1,j+1,k), *cell(i,
j,k) , *cell(i,j+1,k-1), 2);
390 }
391
392 cell(n)->setDivergence( cell(n)->divergence() + (FluxIn - FluxOut)/(cell(n)->
size(2)) );
393
394 // Variables \grad(psi) and \div(B) to evaluate GLM and EGLM divergence cleaning
395 PsiGrad += cell(n)->average(8)*(FluxOut.value(8) - FluxIn.
value(8) - divCor)/(cell(n)->size(2));
396 Bdivergence += (FluxOut.value(6) - FluxIn.value(6))/(cell(n)->
size(2));
397 }
398
399 // Add flux in z-direction
400
401 if (Dimension > 2)
402 {
403 FluxIn = Flux( *cell(i, j, k-2), *cell(i, j, k-1), *cell(i, j, k), *cell(i, j, k+1), 3 )
;
404 divCor = -auxvar;
405 FluxOut = Flux( *cell(i, j, k-1), *cell(i, j, k), *cell(i, j, k+1), *cell(i, j, k+2), 3 )
;
406 divCor += auxvar;
407
408 if(Resistivity){
409 FluxIn = FluxIn - ResistiveTerms(*cell(i,j,k) , *cell(i-1,j,k) , *cell(i,

```

```

j-1,k) , *cell(i,j,k-1), 3);
410     FluxOut = FluxOut - ResistiveTerms(*cell(i,j,k+1), *cell(i-1,j,k+1), *cell(i,
j-1,k+1), *cell(i,j,k) , 3);
411 }
412
413     cell(n)->setDivergence( cell(n)->divergence() + (FluxIn - FluxOut)/(cell(n)->
size(3)) );
414
415     // Variables \grad(psi) and \div(B) to evaluate GLM and EGLM divergence cleaning
416     PsiGrad += cell(n)->average(9)*(FluxOut.value(9) - FluxIn.
value(9) - divCor)/(cell(n)->size(3));
417     Bdivergence += (FluxOut.value(6) - FluxIn.value(6))/(cell(n)->
size(3));
418 }
419 }
420
421 // --- End loop on all cells ---
422 }

```

Here is the caller graph for this function:



### 5.2.3.7 void FineMesh::computeGradient ( int mode )

Computes velocity gradient (only for Navier-Stokes). one cell.

#### Parameters

|            |                                                                   |
|------------|-------------------------------------------------------------------|
| <i>int</i> | It can be zero or one. Associated to the time integration scheme. |
|------------|-------------------------------------------------------------------|

#### Returns

void

```

690 {
691     int i,j,k,d;
692     // --- Loops for internal cells
693     if (mode==0) {
694     if (Dimension==1)
695         for (i=NeighbourNb;i<(1<<ScaleNb)-NeighbourNb;i++) {
696             j=0; k=0;
697             d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
698             computeGradient_cell(d);
699         }
700
701     if (Dimension==2)
702         for (i=NeighbourNb;i<(1<<ScaleNb)-NeighbourNb;i++)
703             for (j=NeighbourNb;j<(1<<ScaleNb)-NeighbourNb;j++) {
704                 k=0;
705                 d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
706                 computeGradient_cell(d);
707             }
708
709
710     if (Dimension==3)
711         for (i=NeighbourNb;i<(1<<ScaleNb)-NeighbourNb;i++)
712             for (j=NeighbourNb;j<(1<<ScaleNb)-NeighbourNb;j++)
713                 for (k=NeighbourNb;k<(1<<ScaleNb)-NeighbourNb;k++) {
714                     d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
715                     computeGradient_cell(d);
716                 }
717     }
718
719 // --- loop for neighbour cells
720 if (mode==1) {
721
722 if (Dimension==1)
723     for (i=0;i<(1<<ScaleNb);i++)
724     if (i<NeighbourNb || i>=(1<<ScaleNb)-NeighbourNb) {
725         j=0; k=0;

```

```

726             d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
727             computeGradient_cell(d);
728         }
729
730     if (Dimension==2)
731         for (i=0;i<(1<<ScaleNb);i++)
732             for (j=0;j<one_D;j++)
733                 if (i<NeighbourNb || j<NeighbourNb ||
734                     i>=(1<<ScaleNb)-NeighbourNb || j>=(1<<
ScaleNb)-NeighbourNb) {
735                     k=0;
736                     d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
737                     computeGradient_cell(d);
738                 }
739
740     if (Dimension==3)
741         for (i=0;i<(1<<ScaleNb);i++)
742             for (j=0;j<one_D;j++)
743                 for (k=0;k<two_D;k++)
744                     if (i<NeighbourNb || j<NeighbourNb || k<
NeighbourNb ||
745                         i>=(1<<ScaleNb)-NeighbourNb || j>=(1<<
ScaleNb)-NeighbourNb || k>=(1<<ScaleNb)-NeighbourNb) {
746                             d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
747                             computeGradient_cell(d);
748                         }
749     }
750 /*
751     int n;
752     for (n = 0; n < 1<<(ScaleNb*Dimension); n++) computeDivergence_cell(n);*/
753 }

```

### 5.2.3.8 void FineMesh::computeGradient\_cell ( int n )

Computes velocity gradient (only for Navier-Stokes). each cells.

#### Parameters

|            |                                                                   |
|------------|-------------------------------------------------------------------|
| <i>int</i> | It can be zero or one. Associated to the time integration scheme. |
|------------|-------------------------------------------------------------------|

#### Returns

void

```

615 {
616     // --- Local variables ---
617     int i=0, j=0, k=0;           // Counter on children
618     real V1=0., V2=0.;         // Velocities
619     real dx=0.;                // Distance between the centers of the neighbour cells
620     real dxV=0.;               // Correction of dx for the computation of GradV close to solid walls
621     // Cell size
622     real rho1=0., rho2=0.;     // Densities
623     real rhoE1=0., rhoE2=0.;  // Energies
624     int p=0, q=0;              // Counters on dimension (between 0 and Dimension)
625     int ei=0, ej=0, ek=0;     // 1 if this direction is chosen, 0 elsewhere
626
627     real result;
628     result = 0.;
629
630     // --- Recursion ---
631     if (EquationType != 6)
632     {
633         cout << "FineMesh.cpp: In method `void FineMesh::computeGradient()':\n";
634         cout << "FineMesh.cpp: EquationType not equal to 6 \n";
635         cout << "carmen: *** [FineMesh.o] Execution error\n";
636         cout << "carmen: abort execution.\n";
637         exit(1);
638     }
639
640     // --- Loop on all cells ---
641
642     // Only in the fluid
643     if (BoundaryRegion(cell(n)->center()) == 0)
644     {
645         i = n%(1<<ScaleNb);
646         if (Dimension > 1) j = (n%(1<<(2*ScaleNb)))/(1<<ScaleNb);
647         if (Dimension > 2) k = n/(1<<(2*ScaleNb));
648

```

```

649     for (p=1; p <= Dimension; p++)
650     {
651         ei = (p==1)? 1:0;
652         ej = (p==2)? 1:0;
653         ek = (p==3)? 1:0;
654
655         dx = cell(i,j,k)->size(p);
656         dx *= 2.;
657
658         // dxV = correction on dx for the computation of GradV close to solid walls
659
660         if (BoundaryRegion(cell(i+ei,j+ej,k+ek)->center()) > 3 ||
661             BoundaryRegion(cell(i-ei,j-ej,k-ek)->center()) > 3 )
662             dxV = 0.75*dx;
663         else
664             dxV = dx;
665
666         rho1 = cell(i+ei,j+ej,k+ek)->density();
667         rho2 = cell(i-ei,j-ej,k-ek)->density();
668
669         cell(n)->setGradient(p, 1, (rho1-rho2)/dx);
670
671         for (q=1; q <= Dimension; q++)
672         {
673             V1=cell(i+ei,j+ej,k+ek)->velocity(q);
674             V2=cell(i-ei,j-ej,k-ek)->velocity(q);
675             result = (V1-V2)/dx;
676             cell(n)->setGradient(p, q+1, (V1-V2)/dxV);
677         }
678
679         rhoE1 = cell(i+ei,j+ej,k+ek)->energy();
680         rhoE2 = cell(i-ei,j-ej,k-ek)->energy();
681
682         cell(n)->setGradient(p, Dimension+2, (rhoE1-rhoE2)/dx);
683     }
684 }
685
686 }

```

Here is the caller graph for this function:



### 5.2.3.9 void FineMesh::computeIntegral ( )

Computes integral values like e.g. flame velocity, global error, etc.

#### Returns

void

```

966 {
967     // --- Local variables ---
968
969     int n=0;           // cell number
970     int  AxisNo;      // Counter on dimension
971     real dx=0., dy=0., dz=0.; // Cell size
972     Vector  Center(Dimension); // local center of the flame ball
973     real VelocityMax=0.; // local maximum of the velocity
974     real divB=0;
975     Vector  GradDensity(Dimension); // gradient of density
976     Vector  GradPressure(Dimension); // gradient of pressure
977     real B1=0., B2=0.; // Left and right magnetic field cells
978     real modB=0.;
979     real MaxSpeed;
980     real QuantityNo=0.;
981
982     int  ei=0, ej=0, ek=0; // 1 if this direction is chosen, 0 elsewhere

```



```

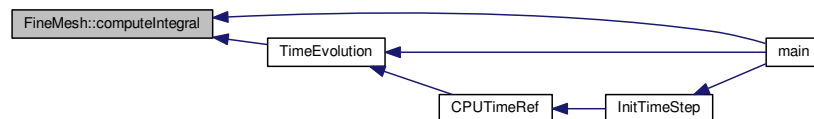
983     int     i=0, j=0, k=0;         // Counter on children
984
985     // --- Init ---
986
987     // Init integral values
988
989     FlameVelocity      = 0.;
990     GlobalMomentum     = 0.;
991     GlobalEnergy       = 0.;
992     GlobalEnstrophy    = 0.;
993     ExactMomentum     = 0.;
994     ExactEnergy        = 0.;
995
996     GlobalReactionRate = 0.;
997     AverageRadius      = 0.;
998     ReactionRateMax    = 0.;
999
1000     for (AxisNo=1; AxisNo <= Dimension; AxisNo++)
1001         Center.setValue(AxisNo,XCenter[AxisNo]);
1002
1003     ErrorMax           = 0.;
1004     ErrorMid           = 0.;
1005     ErrorL2            = 0.;
1006     ErrorNb            = 0.;
1007
1008     RKFEError         = 0.;
1009
1010     Eigenvalue         = 0.;
1011     QuantityMax.setZero();
1012
1013     IntVorticity=0.;
1014     IntDensity=0.;
1015     IntMomentum.setZero();
1016     BaroclinicEffect=0.;
1017
1018 // --- Loop on all cells ---
1019
1020     for (n = 0; n < 1<<(ScaleNb*Dimension); n++)
1021     {
1022         i = n%(1<<ScaleNb);
1023         if (Dimension > 1) j = (n%(1<<(2*ScaleNb)))/(1<<ScaleNb);
1024         if (Dimension > 2) k = n/(1<<(2*ScaleNb));
1025
1026         dx = cell(n)->size(1);
1027         dy = (Dimension > 1) ? cell(n)->size(2) : 1.;
1028         dz = (Dimension > 2) ? cell(n)->size(3) : 1.;
1029
1030         // --- Compute the global momentum, global energy and global enstrophy ---
1031
1032         GlobalMomentum += cell(n)->average(2)*dx*dy*dz;
1033         GlobalEnergy   += .5*(cell(n)->magField()*cell(n)->
magField() + cell(n)->density()*cell(n)->velocity()*cell(n)->
velocity()) + cell(n)->pressure()/(Gamma-1.0);
1034         //GlobalEnergy += .5*(cell(n)->density()*cell(n)->velocity()*cell(n)->velocity());
1035         GlobalEnergy   *= dx*dy*dz;
1036         Helicity       += (cell(n)->magField(2)*cell(n)->
velocity(3) - cell(n)->magField(3)*cell(n)->velocity(2))*cell(n)->
magField(1) +
1037             (cell(n)->magField(3)*cell(n)->velocity(1) - cell(n)->
magField(1)*cell(n)->velocity(3))*cell(n)->magField(2) +
1038             (cell(n)->magField(1)*cell(n)->velocity(2) - cell(n)->
magField(2)*cell(n)->velocity(1))*cell(n)->magField(3);
1039         Helicity       *= 2*dx*dy*dz;
1040
1041         // --- Compute maximum of the conservative quantities ---
1042
1043         for (QuantityNo=1; QuantityNo <=QuantityNb; QuantityNo++)
1044         {
1045             if ( QuantityMax.value(QuantityNo) < fabs(cell(n)->average(QuantityNo)) )
1046                 QuantityMax.setValue(QuantityNo, fabs(cell(n)->average(QuantityNo))
);
1047         }
1048
1049         // --- Compute the maximal eigenvalue ---
1050
1051         VelocityMax = 0.;
1052         MaxSpeed    = 0.;
1053         for (AxisNo=1; AxisNo <= Dimension; AxisNo++){
1054             VelocityMax = Max( VelocityMax, fabs(cell(n)->velocity(AxisNo)));
1055             MaxSpeed    = Max( MaxSpeed , fabs(cell(n)->fastSpeed(AxisNo)));
1056         }
1057
1058         VelocityMax += MaxSpeed;
1059
1060         EigenvalueMax = Max( EigenvalueMax, VelocityMax);
1061
1062         for (AxisNo = 1; AxisNo <= Dimension; AxisNo ++)
```

```

1063     {
1064
1065         ei = (AxisNo == 1)? 1:0;
1066         ej = (AxisNo == 2)? 1:0;
1067         ek = (AxisNo == 3)? 1:0;
1068
1069         dx = cell(n)->size(AxisNo);
1070         // dx *= 2.;
1071
1072         B1=cell(i+ei, j+ej, k+ek)->magField(AxisNo);
1073         B2=cell(i-ei, j-ej, k-ek)->magField(AxisNo);
1074         modB += (B1 + B2)/dx;
1075         divB += (B1-B2)/dx;
1076     }
1077     modB += 1.120e-13;
1078     DIVBMax = Max(DIVBMax, 0.5*Abs(divB));
1079     DIVB = DIVBMax/modB;
1080     break;
1081 }
1082 }
1083
1084 // --- End loop on all cells ---
1085
1086 ReduceIntegralValues();
1087 }

```

Here is the caller graph for this function:



### 5.2.3.10 void FineMesh::computeTimeAverage ( )

Computes the time-average value in every cell.

#### Returns

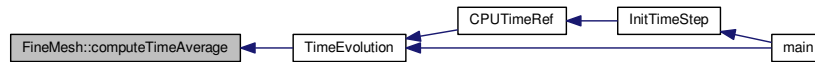
void

```

1804 {
1805     // Local variables
1806
1807     int i=0, j=0, k=0, n=0; // Counters on directions
1808
1809     // Start this procedure when the physical time is larger than StartTimeAveraging
1810
1811     if (TimeStep*IterationNo <= StartTimeAveraging)
1812         return;
1813
1814     // Update time-average values with values in FineMesh
1815
1816     for (n = 0; n < 1<<(ScaleNb*Dimension); n++)
1817     {
1818         i = n%(1<<ScaleNb);
1819         if (Dimension > 1) j = (n%(1<<(2*ScaleNb)))/(1<<ScaleNb);
1820         if (Dimension > 2) k = n/(1<<(2*ScaleNb));
1821
1822         MyTimeAverageGrid->updateValue(i, j, k, cell(i, j, k)->average());
1823     }
1824
1825     // Update the number of samples (Warning: currently only for constant time step !!)
1826     MyTimeAverageGrid->updateSample();
1827 }

```

Here is the caller graph for this function:



### 5.2.3.11 void FineMesh::restore ( )

Restores the tree structure and the cell-averages from the file *carmen.bak*. This file was created by the method [Backup\(\)](#).

Returns

void

```

1769 {
1770     int    n,iaux;    // Cell number
1771     int    QuantityNo; // Counter on quantities
1772     FILE*  input;    // Input file
1773     real   buf;      // Buffer
1774     //      char   line[1024];
1775
1776     // --- Init ---
1777
1778     input = fopen(BackupName,"r");
1779
1780     // When there is no back-up file, return
1781     if (!input) return;
1782
1783     // --- Restore data on cells ---
1784
1785     //fgets(line, 1024, input);
1786     for (n = 0; n < 1<<(ScaleNb*Dimension); n++)
1787     for (QuantityNo=1; QuantityNo <= QuantityNb; QuantityNo++)
1788     {
1789         iaux=fscanf(input, BACKUP_FILE_FORMAT, &buf);
1790         cell(n)->setAverage(QuantityNo, buf);
1791     }
1792
1793     fclose(input);
1794     return;
1795 }
  
```

Here is the caller graph for this function:



### 5.2.3.12 void FineMesh::RungeKutta ( int mode )

Computes the Runge-Kutta step.

## Parameters

|            |                                                                   |
|------------|-------------------------------------------------------------------|
| <i>int</i> | It can be zero or one. Associated to the time integration scheme. |
|------------|-------------------------------------------------------------------|

## Returns

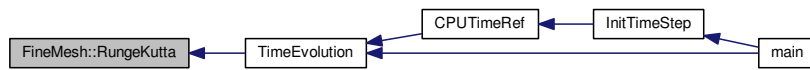
void

```

807                                     {
808     int i,j,k,d;
809
810     // --- Loops for internal cells
811
812     if (mode==0) {
813     if (Dimension==1)
814         for (i=NeighbourNb;i<(1<<ScaleNb)-NeighbourNb;i++) {
815             j=0; k=0;
816             d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
817             RungeKutta_cell(d);
818         }
819
820     if (Dimension==2)
821         for (i=NeighbourNb;i<(1<<ScaleNb)-NeighbourNb;i++)
822             for (j=NeighbourNb;j<(1<<ScaleNb)-NeighbourNb;j++) {
823                 k=0;
824                 d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
825                 RungeKutta_cell(d);
826             }
827
828
829     if (Dimension==3)
830         for (i=NeighbourNb;i<(1<<ScaleNb)-NeighbourNb;i++)
831             for (j=NeighbourNb;j<(1<<ScaleNb)-NeighbourNb;j++)
832                 for (k=NeighbourNb;k<(1<<ScaleNb)-NeighbourNb;k++) {
833                     d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
834                     RungeKutta_cell(d);
835                 }
836     }
837
838     // --- loop for neighbour cells
839     if (mode==1) {
840
841     if (Dimension==1)
842         for (i=0;i<(1<<ScaleNb);i++)
843             if (i<NeighbourNb || i>=(1<<ScaleNb)-NeighbourNb) {
844                 j=0; k=0;
845                 d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
846                 RungeKutta_cell(d);
847             }
848
849     if (Dimension==2)
850         for (i=0;i<(1<<ScaleNb);i++)
851             for (j=0;j<one_D;j++)
852                 if (i<NeighbourNb || j<NeighbourNb ||
853                     i>=(1<<ScaleNb)-NeighbourNb || j>=(1<<
854     ScaleNb)-NeighbourNb) {
855                         k=0;
856                         d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
857                         RungeKutta_cell(d);
858                     }
859
860     if (Dimension==3)
861         for (i=0;i<(1<<ScaleNb);i++)
862             for (j=0;j<one_D;j++)
863                 for (k=0;k<two_D;k++)
864                     if (i<NeighbourNb || j<NeighbourNb || k<
865     NeighbourNb ||
866                         i>=(1<<ScaleNb)-NeighbourNb || j>=(1<<
867     ScaleNb)-NeighbourNb || k>=(1<<ScaleNb)-NeighbourNb) {
868                             d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
869                             RungeKutta_cell(d);
870                         }
871
872     }
873
874     // int n;
875     // for (n = 0; n < 1<<(ScaleNb*Dimension); n++) RungeKutta_cell(n);
876 }

```

Here is the caller graph for this function:



### 5.2.3.13 void FineMesh::RungeKutta\_cell ( int n )

Computes one Runge-Kutta step.

#### Parameters

|            |                                                                   |
|------------|-------------------------------------------------------------------|
| <i>int</i> | It can be zero or one. Associated to the time integration scheme. |
|------------|-------------------------------------------------------------------|

#### Returns

void

```

764 {
765     // --- Local variables ---
766     real c1=0., c2=0., c3=0.;          // Runge-Kutta coefficients
767
768     Vector Q(QuantityNb), Qs(QuantityNb), D(QuantityNb);          //
769     // Cell-average, temporary cell-average and divergence
770
771     // --- Loop on all cells ---
772     if (!UseBoundaryRegions || BoundaryRegion(cell(n)->center()) == 0)
773     {
774         switch(StepNo)
775         {
776             case 1:
777                 c1 = 1.; c2 = 0.; c3 = 1.;
778                 break;
779             case 2:
780                 if (StepNb == 2) {c1 = .5; c2 = .5; c3 = .5; }
781                 if (StepNb == 3) {c1 = .75; c2 = .25; c3 = .25; }
782                 break;
783             case 3:
784                 if (StepNb == 3) {c1 = 1./3.; c2 = 2.*c1; c3 = c2;}
785                 break;
786         };
787
788         // --- Runge-Kutta step ---
789
790         Q = cell(n)->average();
791         Qs = cell(n)->tempAverage();
792         D = cell(n)->divergence();
793
794         cell(n)->setAverage( c1*Qs + c2*Q + (c3 * TimeStep)*D );
795     }
796
797 }
  
```

Here is the caller graph for this function:



### 5.2.3.14 void FineMesh::store ( )

Stores cell-average values into temporary cell-average values.

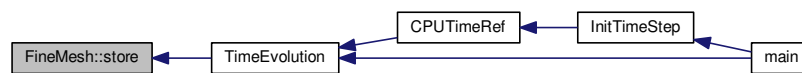
#### Returns

void

```

883 {
884     // --- Local variables ---
885
886     int     n=0;                // cell number
887
888     for (n = 0; n < 1<<(ScaleNb*Dimension); n++)
889     {
890         if (UseBoundaryRegions)
891         {
892             if (IterationNo == 1)
893                 cell(n)->setOldAverage(cell(n)->average());
894             else
895                 cell(n)->setOldAverage(cell(n)->tempAverage());
896         }
897
898         cell(n)->setTempAverage(cell(n)->average());
899     }
900 }
```

Here is the caller graph for this function:



### 5.2.3.15 void FineMesh::storeGrad ( )

Stores gradient values into temporary gradient values.

#### Returns

void

```

911 {
912     // --- Local variables ---
913
914     int     n=0;                // cell number
915
916     for (n = 0; n < 1<<(ScaleNb*Dimension); n++)
917         cell(n)->setTempGradient(cell(n)->gradient());
918 }
```

### 5.2.3.16 void FineMesh::writeAverage ( const char \* FileName )

Write cell-averages for GNUplot, Data Explorer, TecPlot and VTK into file *FileName*.

#### Parameters

| <i>FileName</i> | Name of the file to write. |
|-----------------|----------------------------|
|-----------------|----------------------------|

## Returns

void

```

1264 {
1265     // --- Local variables ---
1266
1267     int i=0,j=0,k=0, n=0;           // Coordinates
1268     FILE *output;                 // pointer to output file
1269
1270     //real x=0., y=0., z=0., t=0.;
1271
1272     // --- Open file ---
1273
1274     if ((output = fopen(FileName,"a")) != NULL)
1275     {
1276         // --- Eventually coarsen grid
1277         if (PrintMoreScales == -1)
1278         {
1279             coarsen();
1280             ScaleNb--;
1281         }
1282
1283         // --- Loop on all cells ---
1284
1285         for (n=0; n < (1<<(Dimension*ScaleNb)); n++)
1286         {
1287
1288             // -- Compute i, j, k --
1289
1290             // For Gnuplot and DX, loop order: for i... {for j... {for k...} }
1291             if (PostProcessing != 3)
1292             {
1293                 switch(Dimension)
1294                 {
1295                     case 1:
1296                         i = n;
1297                         j = k = 0;
1298                         break;
1299
1300                     case 2:
1301                         j = n%(1<<ScaleNb);
1302                         i = n/(1<<ScaleNb);
1303                         k = 0;
1304                         break;
1305
1306                     case 3:
1307                         k = n%(1<<ScaleNb);
1308                         j = (n%(1<<(2*ScaleNb)))/(1<<ScaleNb);
1309                         i = n/(1<<(2*ScaleNb));
1310                         break;
1311                 };
1312             }
1313             else
1314             {
1315                 // For Tecplot, loop order: for k... {for j... {for i...} }
1316                 i = n%(1<<ScaleNb);
1317                 if (Dimension > 1) j = (n%(1<<(2*ScaleNb)))/(1<<
ScaleNb);
1318                 if (Dimension > 2) k = n/(1<<(2*ScaleNb));
1319             }
1320
1321             if(PostProcessing == 4)
1322             {
1323                 fprintf(output, "\n\nSCALARS eta float\nLOOKUP_TABLE default\n");
1324                 for (n=0; n < (1<<(Dimension*ScaleNb)); n++){
1325                     switch(Dimension)
1326                     {
1327                         case 1:
1328                             i = n;
1329                             j = k = 0;
1330                             break;
1331
1332                         case 2:
1333                             j = n%(1<<ScaleNb);
1334                             i = n/(1<<ScaleNb);
1335                             k = 0;
1336                             break;
1337
1338                         case 3:
1339                             k = n%(1<<ScaleNb);
1340                             j = (n%(1<<(2*ScaleNb)))/(1<<ScaleNb);

```

```

1341         i = n/(1<<(2*ScaleNb));
1342         break;
1343     };
1344     FileWrite(output, FORMAT, cell(i,j,k)->etaConst());
1345     fprintf(output, "\n");
1346 }
1347
1348 fprintf(output, "\n\nSCALARS Density float\nLOOKUP_TABLE default\n");
1349 for (n=0; n < (1<<(Dimension*ScaleNb)); n++){
1350     switch(Dimension)
1351     {
1352     case 1:
1353         i = n;
1354         j = k = 0;
1355         break;
1356
1357     case 2:
1358         j = n%(1<<ScaleNb);
1359         i = n/(1<<ScaleNb);
1360         k = 0;
1361         break;
1362
1363     case 3:
1364         k = n%(1<<ScaleNb);
1365         j = (n%(1<<(2*ScaleNb)))/(1<<ScaleNb);
1366         i = n/(1<<(2*ScaleNb));
1367         break;
1368     };
1369     FileWrite(output, FORMAT, cell(i,j,k)->density());
1370     fprintf(output, "\n");
1371 }
1372
1373 fprintf(output, "\n\nSCALARS Pressure float\nLOOKUP_TABLE default\n");
1374 for (n=0; n < (1<<(Dimension*ScaleNb)); n++){
1375     switch(Dimension)
1376     {
1377     case 1:
1378         i = n;
1379         j = k = 0;
1380         break;
1381
1382     case 2:
1383         j = n%(1<<ScaleNb);
1384         i = n/(1<<ScaleNb);
1385         k = 0;
1386         break;
1387
1388     case 3:
1389         k = n%(1<<ScaleNb);
1390         j = (n%(1<<(2*ScaleNb)))/(1<<ScaleNb);
1391         i = n/(1<<(2*ScaleNb));
1392         break;
1393     };
1394     FileWrite(output, FORMAT, cell(i,j,k)->pressure());
1395     fprintf(output, "\n");
1396 }
1397
1398 fprintf(output, "\n\nSCALARS Energy float\nLOOKUP_TABLE default\n");
1399 for (n=0; n < (1<<(Dimension*ScaleNb)); n++){
1400     switch(Dimension)
1401     {
1402     case 1:
1403         i = n;
1404         j = k = 0;
1405         break;
1406
1407     case 2:
1408         j = n%(1<<ScaleNb);
1409         i = n/(1<<ScaleNb);
1410         k = 0;
1411         break;
1412
1413     case 3:
1414         k = n%(1<<ScaleNb);
1415         j = (n%(1<<(2*ScaleNb)))/(1<<
ScaleNb);
1416         i = n/(1<<(2*ScaleNb));
1417         break;
1418     };
1419     FileWrite(output, FORMAT, cell(i,j,k)->energy());
1420     fprintf(output, "\n");
1421 }
1422
1423 fprintf(output, "\n\nSCALARS Vx float\nLOOKUP_TABLE default\n");
1424 for (n=0; n < (1<<(Dimension*ScaleNb)); n++){
1425     switch(Dimension)
1426     {

```



```

1427         case 1:
1428             i = n;
1429             j = k = 0;
1430             break;
1431
1432         case 2:
1433             j = n%(1<<ScaleNb);
1434             i = n/(1<<ScaleNb);
1435             k = 0;
1436             break;
1437
1438         case 3:
1439             k = n%(1<<ScaleNb);
1440             j = (n%(1<<(2*ScaleNb)))/(1<<ScaleNb);
1441             i = n/(1<<(2*ScaleNb));
1442             break;
1443     };
1444     FileWrite(output, FORMAT, cell(i,j,k)->velocity(1));
1445     fprintf(output, "\n");
1446 }
1447
1448 fprintf(output, "\n\nSCALARS Vy float\nLOOKUP_TABLE default\n");
1449 for (n=0; n < (1<<(Dimension*ScaleNb)); n++){
1450     switch(Dimension)
1451     {
1452         case 1:
1453             i = n;
1454             j = k = 0;
1455             break;
1456
1457         case 2:
1458             j = n%(1<<ScaleNb);
1459             i = n/(1<<ScaleNb);
1460             k = 0;
1461             break;
1462
1463         case 3:
1464             k = n%(1<<ScaleNb);
1465             j = (n%(1<<(2*ScaleNb)))/(1<<
ScaleNb);
1466             i = n/(1<<(2*ScaleNb));
1467             break;
1468     };
1469     FileWrite(output, FORMAT, cell(i,j,k)->velocity(2));
1470     fprintf(output, "\n");
1471 }
1472
1473 fprintf(output, "\n\nSCALARS Vz float\nLOOKUP_TABLE default\n");
1474 for (n=0; n < (1<<(Dimension*ScaleNb)); n++){
1475     switch(Dimension)
1476     {
1477         case 1:
1478             i = n;
1479             j = k = 0;
1480             break;
1481
1482         case 2:
1483             j = n%(1<<ScaleNb);
1484             i = n/(1<<ScaleNb);
1485             k = 0;
1486             break;
1487
1488         case 3:
1489             k = n%(1<<ScaleNb);
1490             j = (n%(1<<(2*ScaleNb)))/(1<<
ScaleNb);
1491             i = n/(1<<(2*ScaleNb));
1492             break;
1493     };
1494     FileWrite(output, FORMAT, cell(i,j,k)->velocity(3));
1495     fprintf(output, "\n");
1496 }
1497
1498 fprintf(output, "\n\nSCALARS Bx float\nLOOKUP_TABLE default\n");
1499 for (n=0; n < (1<<(Dimension*ScaleNb)); n++){
1500     switch(Dimension)
1501     {
1502         case 1:
1503             i = n;
1504             j = k = 0;
1505             break;
1506
1507         case 2:
1508             j = n%(1<<ScaleNb);
1509             i = n/(1<<ScaleNb);
1510             k = 0;
1511             break;

```

```

1512
1513
1514         case 3:
1515             k = n%(1<<ScaleNb);
1516                 j = (n%(1<<(2*ScaleNb)))/(1<<
ScaleNb);
1517             i = n/(1<<(2*ScaleNb));
1518             break;
1519         };
1520         FileWrite(output, FORMAT, cell(i,j,k)->magField(1));
1521         fprintf(output, "\n");
1522     }
1523     fprintf(output, "\n\nSCALARS By float\nLOOKUP_TABLE default\n");
1524     for (n=0; n < (1<<(Dimension*ScaleNb)); n++){
1525         switch(Dimension)
1526         {
1527             case 1:
1528                 i = n;
1529                 j = k = 0;
1530                 break;
1531             case 2:
1532                 j = n%(1<<ScaleNb);
1533                 i = n/(1<<ScaleNb);
1534                 k = 0;
1535                 break;
1536             case 3:
1537                 k = n%(1<<ScaleNb);
1538                 j = (n%(1<<(2*ScaleNb)))/(1<<
ScaleNb);
1539                 i = n/(1<<(2*ScaleNb));
1540                 break;
1541         };
1542         FileWrite(output, FORMAT, cell(i,j,k)->magField(2));
1543         fprintf(output, "\n");
1544     }
1545     fprintf(output, "\n\nSCALARS Bz float\nLOOKUP_TABLE default\n");
1546     for (n=0; n < (1<<(Dimension*ScaleNb)); n++){
1547         switch(Dimension)
1548         {
1549             case 1:
1550                 i = n;
1551                 j = k = 0;
1552                 break;
1553             case 2:
1554                 j = n%(1<<ScaleNb);
1555                 i = n/(1<<ScaleNb);
1556                 k = 0;
1557                 break;
1558             case 3:
1559                 k = n%(1<<ScaleNb);
1560                 j = (n%(1<<(2*ScaleNb)))/(1<<
ScaleNb);
1561                 i = n/(1<<(2*ScaleNb));
1562                 break;
1563         };
1564         FileWrite(output, FORMAT, cell(i,j,k)->magField(3));
1565         fprintf(output, "\n");
1566     }
1567     fprintf(output, "\n\nSCALARS DivB float\nLOOKUP_TABLE default\n");
1568     for (n=0; n < (1<<(Dimension*ScaleNb)); n++){
1569         switch(Dimension)
1570         {
1571             case 1:
1572                 i = n;
1573                 j = k = 0;
1574                 break;
1575             case 2:
1576                 j = n%(1<<ScaleNb);
1577                 i = n/(1<<ScaleNb);
1578                 k = 0;
1579                 break;
1580             case 3:
1581                 k = n%(1<<ScaleNb);
1582                 j = (n%(1<<(2*ScaleNb)))/(1<<ScaleNb);
1583                 i = n/(1<<(2*ScaleNb));
1584                 break;
1585         };
1586         real divB=0, B1=0., B2=0., dx=0.;
1587         int ei=0,ej=0,ek=0;
1588     }
1589
1590
1591
1592
1593
1594
1595

```

```

1596         for (int AxisNo = 1; AxisNo <= Dimension; AxisNo ++ )
1597         {
1598
1599             ei = (AxisNo == 1)? 1:0;
1600             ej = (AxisNo == 2)? 1:0;
1601             ek = (AxisNo == 3)? 1:0;
1602
1603             dx = cell(i,j,k)->size(AxisNo);
1604             dx *= 2.;
1605
1606             B1 = cell(i+ei, j+ej, k+ek)->magField(AxisNo);
1607             B2 = cell(i-ei, j-ej, k-ek)->magField(AxisNo);
1608
1609             divB += (B1-B2)/dx;
1610         }
1611         FileWrite(output, FORMAT, divB);
1612         fprintf(output, "\n");
1613     }
1614
1615     fprintf(output, "\n\nVECTORS Velocity float\n");
1616     for (n=0; n < (1<<(Dimension*ScaleNb)); n++){
1617         switch(Dimension)
1618         {
1619             case 1:
1620                 i = n;
1621                 j = k = 0;
1622                 break;
1623
1624             case 2:
1625                 j = n%(1<<ScaleNb);
1626                 i = n/(1<<ScaleNb);
1627                 k = 0;
1628                 break;
1629
1630             case 3:
1631                 k = n%(1<<ScaleNb);
1632                 j = (n%(1<<(2*ScaleNb)))/(1<<
1633         ScaleNb);
1634                 i = n/(1<<(2*ScaleNb));
1635                 break;
1636         };
1637
1638         FileWrite(output, FORMAT, cell(i,j,k)->velocity(1));
1639         FileWrite(output, FORMAT, cell(i,j,k)->velocity(2));
1640         FileWrite(output, FORMAT, cell(i,j,k)->velocity(3));
1641     }
1642
1643     fprintf(output, "\n\nVECTORS MagField float\n");
1644     for (n=0; n < (1<<(Dimension*ScaleNb)); n++){
1645         switch(Dimension)
1646         {
1647             case 1:
1648                 i = n;
1649                 j = k = 0;
1650                 break;
1651
1652             case 2:
1653                 j = n%(1<<ScaleNb);
1654                 i = n/(1<<ScaleNb);
1655                 k = 0;
1656                 break;
1657
1658             case 3:
1659                 k = n%(1<<ScaleNb);
1660                 j = (n%(1<<(2*ScaleNb)))/(1<<
1661         ScaleNb);
1662                 i = n/(1<<(2*ScaleNb));
1663                 break;
1664         };
1665
1666         FileWrite(output, FORMAT, cell(i,j,k)->magField(1));
1667         FileWrite(output, FORMAT, cell(i,j,k)->magField(2));
1668         FileWrite(output, FORMAT, cell(i,j,k)->magField(3));
1669     }
1670 }
1671 }else{
1672     // MHD
1673     if (ScalarEqNb == 1)
1674         FileWrite(output, FORMAT, cell(i,j,k)->average(
1675     Dimension+3)/cell(i,j,k)->density());
1676     else
1677         FileWrite(output, FORMAT, cell(i,j,k)->density());
1678
1679     FileWrite(output, FORMAT, cell(i,j,k)->pressure()*

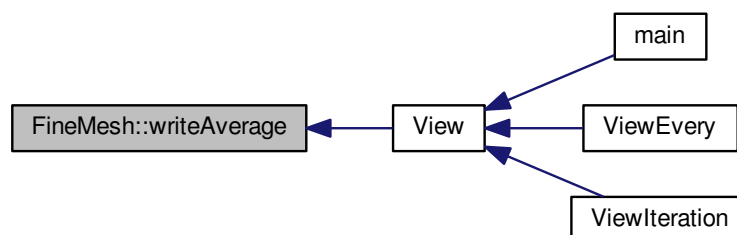
```

```

Gamma*Ma*Ma); // Dimensionless pressure
1680     FileWrite(output, FORMAT, cell(i,j,k)->temperature());
1681     FileWrite(output, FORMAT, cell(i,j,k)->energy());
1682     if (PostProcessing == 1) FileWrite(output,
FORMAT, 0.);
1683         for (int AxisNo = 1; AxisNo <= Dimension; AxisNo++)
1684             FileWrite(output, FORMAT, cell(i,j,k)->velocity(AxisNo));
1685     }
1686 }
1687
1688
1689     if (!DataIsBinary)
1690         fprintf(output, "\n");
1691
1692     if (PostProcessing == 1)
1693     {
1694         if (j==(1<<ScaleNb)-1)
1695             fprintf(output, "\n");
1696
1697         if (k==(1<<ScaleNb)-1)
1698             fprintf(output, "\n");
1699     }
1700 }
1701 }
1702 fclose(output);
1703
1704 // --- Eventually refine grid
1705
1706 if (PrintMoreScales == -1)
1707 {
1708     ScaleNb++;
1709     refine();
1710 }
1711 }
1712 else
1713 {
1714     cout << "FineMesh.cpp: In method `void FineMesh::writeAverage()':\n";
1715     cout << "FineMesh.cpp: cannot open file " << FileName << '\n';
1716     cout << "carmen: *** [FineMesh.o] Execution error\n";
1717     cout << "carmen: abort execution.\n";
1718     exit(1);
1719 }
1720 }

```

Here is the caller graph for this function:



### 5.2.3.17 void FineMesh::writeHeader ( const char \* *FileName* ) const

Write header for GNUplot, Data Explorer, TecPlot and VTK into file *FileName*.

#### Parameters

| <i>FileName</i> | Name of the file to write. |
|-----------------|----------------------------|
|-----------------|----------------------------|

## Returns

void

```

1097 {
1098     // --- Local variables ---
1099
1100     real dx=0., dy=0., dz=0.;           // deltas in x, y, and z
1101     FILE *output;                       // Pointer to output file
1102     int GridPoints;                     // Grid points
1103     char DependencyType[12];           // positions or connections
1104
1105
1106     // --- For the final data, use positions instead of connections ---
1107
1108     if (WriteAsPoints)
1109     {
1110         GridPoints = (1<<(ScaleNb+PrintMoreScales));
1111         sprintf(DependencyType,"positions");
1112     }
1113     else
1114     {
1115         GridPoints = (1<<(ScaleNb+PrintMoreScales))+1;
1116         sprintf(DependencyType,"connections");
1117     }
1118
1119     // --- Open file ---
1120
1121     if ((output = fopen(FileName,"w")) != NULL)
1122     {
1123         // --- Header ---
1124
1125
1126         dx = (XMax[1]-XMin[1])/((1<<(ScaleNb+PrintMoreScales))-1);
1127         dy = (XMax[2]-XMin[2])/((1<<(ScaleNb+PrintMoreScales))-1);
1128         dz = (XMax[3]-XMin[3])/((1<<(ScaleNb+PrintMoreScales))-1);
1129
1130         // GNUPLOT
1131
1132         switch(PostProcessing)
1133         {
1134             // GNUPLOT
1135             case 1:
1136                 fprintf(output,"#");
1137                 fprintf(output, TXTFORMAT, " x");
1138                 fprintf(output, TXTFORMAT, "Density");
1139                 fprintf(output, TXTFORMAT, "Pressure");
1140                 fprintf(output, TXTFORMAT, "Temperature");
1141                 fprintf(output, TXTFORMAT, "Energy");
1142                 fprintf(output, TXTFORMAT, "Velocity");
1143                 fprintf(output, "\n");
1144                 break;
1145
1146             // DATA EXPLORER
1147             case 2:
1148                 fprintf(output, "# Data Explorer file\n# generated by Carmen\n");
1149
1150                 switch(Dimension)
1151                 {
1152                     case 2:
1153                         dx = (XMax[1]-XMin[1])/(1<<(ScaleNb+
1154 PrintMoreScales));
1155                         dy = (XMax[2]-XMin[2])/(1<<(ScaleNb+
1156 PrintMoreScales));
1157                         fprintf(output, "grid = %d x %d\n", GridPoints, GridPoints);
1158                         fprintf(output, "positions = %f, %f, %f, %f\n#\n",XMin[1],dx,
1159 XMin[2],dy );
1160                         break;
1161
1162                     case 3:
1163                         dx = (XMax[1]-XMin[1])/(1<<(ScaleNb+
1164 PrintMoreScales));
1165                         dy = (XMax[2]-XMin[2])/(1<<(ScaleNb+
1166 PrintMoreScales));
1167                         dz = (XMax[3]-XMin[3])/(1<<(ScaleNb+
1168 PrintMoreScales));
1169                         fprintf(output, "grid = %d x %d x %d\n", GridPoints, GridPoints, GridPoints);
1170                         fprintf(output, "positions = %f, %f, %f, %f, %f, %f\n#\n",
1171 XMin[1],dx,XMin[2],dy,XMin[3],dz);
1172                         break;
1173                 }
1174             };
1175         }
1176     }
1177 }

```

```

1168         if (DataIsBinary)
1169             fprintf(output, "format = binary\n");
1170         else
1171             fprintf(output, "format = ascii\n");
1172
1173         fprintf(output, "interleaving = field\n");
1174         fprintf(output, "field = density, pressure, temperature, energy, velocity\n");
1175         fprintf(output, "structure = scalar, scalar, scalar, scalar, %d-vector\n",
Dimension);
1176         fprintf(output, "type = %s, %s, %s, %s, %s\n", REAL, REAL,
REAL, REAL, REAL);
1177         fprintf(output, "dependency = %s, %s, %s, %s, %s\n", DependencyType, DependencyType,
DependencyType, DependencyType, DependencyType);
1178
1179
1180         fprintf(output, "header = marker \"START_DATA\\n\" \n");
1181         fprintf(output, "end\n");
1182         fprintf(output, "START_DATA\n");
1183
1184         break;
1185
1186     // TECPLOT
1187     case 3:
1188         fprintf(output, "VARIABLES = \"x\"\n");
1189         if (Dimension > 1)
1190             fprintf(output, "\"y\"\n");
1191         if (Dimension > 2)
1192             fprintf(output, "\"z\"\n");
1193         fprintf(output, "\"RHO\\n\\P\\n\\T\\n\\E\\n\\U\"\n");
1194         if (Dimension > 1)
1195             fprintf(output, "\"V\"\n");
1196         if (Dimension > 2)
1197             fprintf(output, "\"W\"\n");
1198         fprintf(output, "ZONE T=\"Carmen %3.1f\"\n", CarmenVersion);
1199         fprintf(output, "I=%i, ", GridPoints-1);
1200         if (Dimension > 1)
1201             fprintf(output, "J=%i, ", GridPoints-1);
1202         if (Dimension > 2)
1203             fprintf(output, "K=%i, ", GridPoints-1);
1204         fprintf(output, "F=POINT\n");
1205         break;
1206
1207
1208     case 4:
1209         int N=(1<<ScaleNb);
1210         fprintf(output, "# vtk DataFile Version 2.8\nSolucao MHD\n");
1211         if(DataIsBinary)
1212             fprintf(output, "BINARY\n");
1213         else
1214             fprintf(output, "ASCII\n");
1215
1216         fprintf(output, "DATASET STRUCTURED_GRID\n");
1217         if (Dimension == 2)
1218         {
1219             fprintf(output, "DIMENSIONS %d %d %d \n", N,N,1);
1220             fprintf(output, "POINTS %d FLOAT\n", N*N);
1221             for (int i = 0; i < N; i++)
1222                 for (int j = 0; j < N; j++)
1223                     fprintf(output, "%f %f %f \n", XMin[1] + i*dx,
XMin[2] + j*dy, 0.0);
1224
1225             fprintf(output, "\n\nPOINT_DATA %d \n", N*N);
1226         }
1227         if (Dimension == 3)
1228         {
1229             fprintf(output, "DIMENSIONS %d %d %d \n", N,N,N);
1230             fprintf(output, "POINTS %d FLOAT\n", N*N*N);
1231             for (int i = 0; i < N; i++)
1232                 for (int j = 0; j < N; j++)
1233                     for (int k = 0; k < N; k++)
1234                         fprintf(output, "%f %f %f \n", XMin[1] + i*dx,
XMin[2] + j*dy, XMin[3] + k*dz);
1235
1236             fprintf(output, "\n\nPOINT_DATA %d \n", N*N*N);
1237         }
1238
1239         break;
1240     };
1241
1242     fclose(output);
1243     return;
1244 }
1245 else
1246 {
1247     cout << "FineMesh.cpp: In method `void FineMesh::writeHeader()':\n";
1248     cout << "FineMesh.cpp: cannot open file " << FileName << '\n';
1249 }

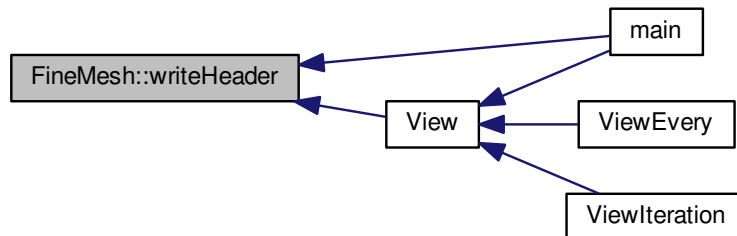
```

```

1250         cout << "carmen: *** [FineMesh.o] Execution error\n";
1251         cout << "carmen: abort execution.\n";
1252         exit(1);
1253     }
1254 }

```

Here is the caller graph for this function:



### 5.2.3.18 void FineMesh::writeTimeAverage ( const char \* *FileName* ) const

Write time-averages into file *FileName*.

#### Parameters

|                 |                            |
|-----------------|----------------------------|
| <i>FileName</i> | Name of the file to write. |
|-----------------|----------------------------|

#### Returns

void

```

1836 {
1837     // --- Local variables ---
1838
1839     int n=0, i=0, j=0, k=0;
1840
1841     real dx, dy, dz;           // deltas in x, y, and z
1842     real x=0, y=0, z=0;       // positions
1843     FILE *output;             // Pointer to output file
1844     int GridPoints = (1<<ScaleNb)+1; // Grid points
1845
1846     // --- Open file ---
1847
1848     if ((output = fopen(FileName,"w")) != NULL)
1849     {
1850         // --- Header ---
1851
1852         switch(PostProcessing)
1853         {
1854             // GNUPLOT
1855             case 1:
1856                 fprintf(output, "# x           Velocity           Stress\n");
1857                 break;
1858
1859             // DATA EXPLORER
1860             case 2:
1861                 fprintf(output, "# Data Explorer file\n# generated by Carmen\n");
1862
1863                 switch(Dimension)
1864                 {
1865                     case 2:
1866                         dx = (XMax[1]-XMin[1])/(1<<ScaleNb);
1867                         dy = (XMax[2]-XMin[2])/(1<<ScaleNb);
1868                         fprintf(output, "grid = %d x %d\n", GridPoints, GridPoints);
1869                         fprintf(output, "positions = %f, %f, %f, %f\n#\n", XMin[1], dx,
1870
1871                             XMin[2], dy );

```

```

1870         break;
1871
1872         case 3:
1873             dx = (XMax[1]-XMin[1])/(1<<ScaleNb);
1874             dy = (XMax[2]-XMin[2])/(1<<ScaleNb);
1875             dz = (XMax[3]-XMin[3])/(1<<ScaleNb);
1876             fprintf(output, "grid = %d x %d x %d\n", GridPoints, GridPoints, GridPoints);
1877             fprintf(output, "positions = %f, %f, %f, %f, %f, %f\n#\n",
XMin[1],dx,XMin[2],dy,XMin[3],dz);
1878         break;
1879     };
1880
1881     if (DataIsBinary)
1882         fprintf(output, "format = binary\n");
1883     else
1884         fprintf(output, "format = ascii\n");
1885
1886     fprintf(output, "interleaving = field\n");
1887
1888     fprintf(output, "field = U, V");
1889
1890     if (Dimension > 2)
1891         fprintf(output, ", W");
1892
1893     fprintf(output, ", U'U', U'V', V'V'");
1894
1895     if (Dimension > 2)
1896         fprintf(output, ", U'W', V'W', W'W'");
1897
1898     fprintf(output, "\n");
1899
1900     fprintf(output, "structure = scalar");
1901     for (n=1; n < (Dimension*(Dimension+3))/2 ; n++)
1902         fprintf(output, ", scalar");
1903     fprintf(output, "\n");
1904
1905     fprintf(output, "type = %s", REAL);
1906     for (n=1; n < (Dimension*(Dimension+3))/2 ; n++)
1907         fprintf(output, ", %s", REAL);
1908     fprintf(output, "\n");
1909
1910     fprintf(output, "dependency = connections");
1911     for (n=1; n < (Dimension*(Dimension+3))/2 ; n++)
1912         fprintf(output, ", connections");
1913     fprintf(output, "\n");
1914
1915     fprintf(output, "header = marker \"START_DATA\\n\" \n");
1916     fprintf(output, "end\n");
1917     fprintf(output, "START_DATA\n");
1918
1919     break;
1920
1921     // TECPLOT
1922     case 3:
1923
1924         // --- Write axes (x,y,z) ---
1925
1926         fprintf(output, "VARIABLES = \"x\"\n");
1927         if (Dimension > 1)
1928             fprintf(output, "\"y\"\n");
1929         if (Dimension > 2)
1930             fprintf(output, "\"z\"\n");
1931
1932     // --- Write averages U, V, W ---
1933
1934     fprintf(output, "\"U\"\n");
1935     if (Dimension > 1)
1936         fprintf(output, "\"V\"\n");
1937     if (Dimension > 2)
1938         fprintf(output, "\"W\"\n");
1939
1940     // --- Write stress tensor U'U', U'V', V'V', U'W', V'W', W'W' ---
1941
1942     fprintf(output, "\"U'U'\"\n");
1943     if (Dimension > 1)
1944     {
1945         fprintf(output, "\"U'V'\"\n");
1946         fprintf(output, "\"V'V'\"\n");
1947     }
1948     if (Dimension > 2)
1949     {
1950         fprintf(output, "\"U'W'\"\n");
1951         fprintf(output, "\"V'W'\"\n");
1952         fprintf(output, "\"W'W'\"\n");
1953     }
1954
1955     fprintf(output, "ZONE T=\"Carmen %3.1f\"\n", CarmenVersion);

```



```

1956         fprintf(output,"I=%i, ",(1<<ScaleNb));
1957         if (Dimension > 1)
1958             fprintf(output,"J=%i, ",(1<<ScaleNb));
1959         if (Dimension > 2)
1960             fprintf(output,"K=%i, ",(1<<ScaleNb));
1961         fprintf(output,"F=POINT\n");
1962         break;
1963     };
1964
1965     // --- Loop on all cells ---
1966
1967     for (n=0; n < (1<<(Dimension*ScaleNb)); n++)
1968     {
1969
1970         // -- Compute i, j, k --
1971
1972         // For Gnuplot and DX, loop order: for i... {for j... {for k...} }
1973
1974         if (PostProcessing != 3)
1975         {
1976             switch(Dimension)
1977             {
1978                 case 1:
1979                     i = n;
1980                     break;
1981
1982                 case 2:
1983                     j = n%(1<<ScaleNb);
1984                     i = n/(1<<ScaleNb);
1985                     break;
1986
1987                 case 3:
1988                     k = n%(1<<ScaleNb);
1989                     j = (n%(1<<(2*ScaleNb)))/(1<<ScaleNb);
1990                     i = n/(1<<(2*ScaleNb));
1991                     break;
1992             };
1993         }
1994         else
1995         {
1996             // For Tecplot, loop order: for k... {for j... {for i...} }
1997
1998             i = n%(1<<ScaleNb);
1999             if (Dimension > 1) j = (n%(1<<(2*ScaleNb)))/(1<<
ScaleNb);
2000             if (Dimension > 2) k = n/(1<<(2*ScaleNb));
2001         }
2002
2003         // Compute x,y,z
2004
2005         x = cell(i,j,k)->center(1);
2006         if (Dimension > 1)
2007             y = cell(i,j,k)->center(2);
2008         if (Dimension > 2)
2009             z = cell(i,j,k)->center(3);
2010
2011         // Write cell-center coordinates (only for Gnuplot and Tecplot)
2012
2013         if (PostProcessing != 2)
2014         {
2015             fprintf(output, FORMAT, x);
2016             if (Dimension > 1)
2017                 fprintf(output, FORMAT, y);
2018             if (Dimension > 2)
2019                 fprintf(output, FORMAT, z);
2020         }
2021
2022         for (int AxisNo = 1; AxisNo <= Dimension; AxisNo++)
2023             FileWrite(output, FORMAT,MyTimeAverageGrid->
velocity(i,j,k,AxisNo));
2024
2025         for (int AxisNo = 1; AxisNo <= (Dimension*(Dimension+1))/2; AxisNo++)
2026             FileWrite(output, FORMAT,MyTimeAverageGrid->
stress(i,j,k,AxisNo));
2027
2028         if (!DataIsBinary)
2029             fprintf(output,"\n");
2030
2031         if (PostProcessing == 1)
2032         {
2033             if (j==(1<<ScaleNb)-1)
2034                 fprintf(output,"\n");
2035
2036             if (k==(1<<ScaleNb)-1)
2037                 fprintf(output,"\n");
2038         }
2039     }

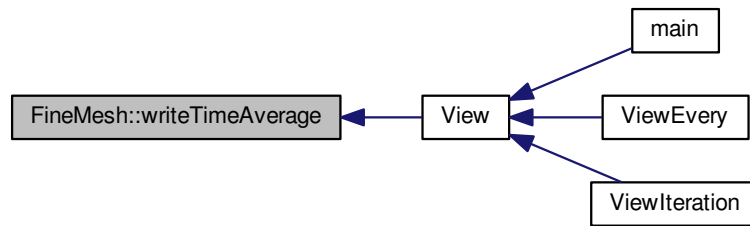
```

```

2040     fclose(output);
2041 }
2042 else
2043 {
2044     cout << "FineMesh.cpp: In method `void FineMesh::writeTimeAverage()':\n";
2045     cout << "FineMesh.cpp: cannot open file " << FileName << '\n';
2046     cout << "carmen: *** [FineMesh.o] Execution error\n";
2047     cout << "carmen: abort execution.\n";
2048     exit(1);
2049 }
2050 }

```

Here is the caller graph for this function:



## 5.2.4 Member Data Documentation

### 5.2.4.1 `Cell*` `FineMesh::MeshCell`

Array of cells

### 5.2.4.2 `Cell***` `FineMesh::Neighbour_iL`

Parallel variable

### 5.2.4.3 `Cell ***` `FineMesh::Neighbour_iU`

Parallel variable

### 5.2.4.4 `Cell ***` `FineMesh::Neighbour_jL`

Parallel variable

### 5.2.4.5 `Cell ***` `FineMesh::Neighbour_jU`

Parallel variable

### 5.2.4.6 `Cell ***` `FineMesh::Neighbour_kL`

Parallel variable

## 5.2.4.7 Cell \*\*\* FineMesh::Neighbour\_kU

Parallel variable

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

- [FineMesh.h](#)
- [FineMesh.cpp](#)

## 5.3 Matrix Class Reference

Standard class for every matrix in Carmen.

```
#include <Matrix.h>
```

### Public Member Functions

- [Matrix](#) ()  
*Constructor of matrix class. Generates a 1,1 matrix equal to zero.*
- [Matrix](#) (const int i, const int j)  
*Generates a matrix with i lines and j columns, each component being equal to zero. Example :*
- [Matrix](#) (const int i)  
*Generates a square matrix with i lines and i columns, each component being equal to zero. Example :*
- [Matrix](#) (const [Matrix](#) &M)  
*Generates a copy of the matrix M. Example :*
- [Matrix](#) (const [Vector](#) &V)  
*Generates a vector-matrix identical to V. Example :*
- [~Matrix](#) ()  
*Destructor of matrix class. Deallocate memory of the matrix.*
- void [setValue](#) (const int i, const int j, const [real](#) a)  
*Sets the component i, j to value a.*
- void [setZero](#) ()  
*Sets all the components to zero.*
- [real value](#) (const int i, const int j) const  
*Returns the value of the component i, j.*
- int [lines](#) () const  
*Returns the number of lines of the matrix.*
- int [columns](#) () const  
*Returns the number of columns of the matrix.*
- bool [operator==](#) (const [Matrix](#) &M) const  
*Compares the current matrix to a matrix M and returns true if they are equal.*
- void [operator=](#) (const [Matrix](#) &M)  
*Set the current matrix to the dimension and the value of M.*
- void [operator+=](#) (const [Matrix](#) &M)  
*Adds M to the current matrix.*
- [Matrix operator+](#) (const [Matrix](#) &M) const  
*Returns the addition of the current matrix and M.*
- void [operator-=](#) (const [Matrix](#) &M)  
*Subtracts M to the current matrix.*
- [Matrix operator-](#) (const [Matrix](#) &M) const  
*Returns the difference between the current matrix and M.*

- **Matrix operator-** ( ) const  
*Returns the opposite of the current matrix.*
- void **operator\*=** (const **real** a)  
*Multiplies the current matrix by a real a.*
- **Matrix operator\*** (const **real** a) const  
*Returns the product of the current matrix and a real a.*
- void **operator/=** (const **real** a)  
*Divides the current matrix by a real a.*
- **Matrix operator/** (const **real** a) const  
*Returns the division of the current matrix by a real a.*
- **Matrix operator\*** (const **Matrix** &M) const  
*Returns the product of the current matrix and a matrix M.*
- **Vector operator\*** (const **Vector** &V) const  
*Returns the product of the current matrix and a vector V.*
- void **setEigenMatrix** (const bool isLeft, const int AxisNo, const **Vector** V, const **real** c, const **real** h=0.)  
*Sets matrix as eigenmatrix.*

## Public Attributes

- int **Lines**
- int **Columns**
- **real** \* **U**

### 5.3.1 Detailed Description

Standard class for every matrix in Carmen.

It contains the following data:

the number of lines *Lines* ;

the number of columns *Columns* ;

the array of reals \**U*.

### 5.3.2 Constructor & Destructor Documentation

#### 5.3.2.1 Matrix::Matrix ( )

Constructor of matrix class. Generates a 1,1 matrix equal to zero.

Example :

```
#include "Matrix.h"
```

```
Matrix M;
```

```
31 {
32     Lines = Columns = 1;
33     U = new real;
34     *U = 0.;
35 }
```

#### 5.3.2.2 Matrix::Matrix ( const int i, const int j )

Generates a matrix with *i* lines and *j* columns, each component being equal to zero. Example :

```
#include "Matrix.h"
```

```
Matrix M(4, 5);
```

Parameters

|     |  |
|-----|--|
| $i$ |  |
| $j$ |  |

```

41 {
42     int n; // Counter
43
44     Lines = i;
45     Columns = j;
46
47     // If the size of the matrix is equal to zero, do not allocate memory
48     if (Lines==0 && Columns==0) return;
49
50     U = new real[Lines*Columns];
51
52     for (n = 1; n <= Lines*Columns; n++)
53         *(U+n-1) = 0.;
54 }

```

### 5.3.2.3 Matrix::Matrix ( const int $i$ )

Generates a square matrix with  $i$  lines and  $i$  columns, each component being equal to zero. Example :

```
#include "Matrix.h"
```

```
Matrix M(4);
```

Parameters

|     |  |
|-----|--|
| $i$ |  |
|-----|--|

```

60 {
61     int n; // Counter
62
63     Lines = i;
64     Columns = i;
65
66     U = new real[Lines*Columns];
67
68     for (n = 1; n <= Lines*Columns; n++)
69         *(U+n-1) = 0.;
70 }

```

### 5.3.2.4 Matrix::Matrix ( const Matrix & $M$ )

Generates a copy of the matrix  $M$ . Example :

```
#include "Matrix.h"
```

```
Matrix M(2,3);
```

```
Matrix P(M)
```

Parameters

|     |        |
|-----|--------|
| $M$ | Matrix |
|-----|--------|

```

76 {
77     int i,j;
78
79     Lines = M.lines();
80     Columns = M.columns();
81
82     U = new real[Lines*Columns];
83
84     for (i = 1; i <= Lines; i++)
85         for (j = 1; j <= Columns; j++)
86             setValue(i, j, M.value(i,j));
87 }

```

### 5.3.2.5 Matrix::Matrix ( const Vector & V )

Generates a vector-matrix identical to  $V$ . Example :

```
#include "Matrix.h"
```

```
Matrix V(3);
```

```
Matrix P(V);
```

Parameters

|     |  |
|-----|--|
| $V$ |  |
|-----|--|

```
93 {
94     int i;
95
96     Lines = V.dimension();
97     Columns = 1;
98
99     U = new real[Lines*Columns];
100
101     for (i = 1; i <= Lines; i++)
102         setValue(i, 1, V.value(i));
103 }
```

### 5.3.2.6 Matrix::~Matrix ( )

Destructor of matrix class. Deallocate memory of the matrix.

```
113 {
114     // If the size of the matrix is equal to zero, do not deallocate memory
115     if (Lines==0 && Columns==0) return;
116
117     delete[] U;
118 }
```

## 5.3.3 Member Function Documentation

### 5.3.3.1 int Matrix::columns ( ) const [inline]

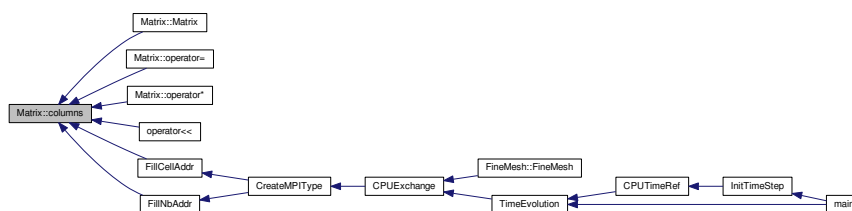
Returns the number of columns of the matrix.

Returns

int

```
496 {
497     return Columns;
498 }
```

Here is the caller graph for this function:



## 5.3.3.2 int Matrix::lines ( ) const [inline]

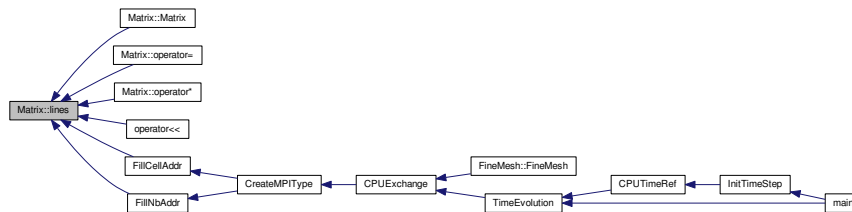
Returns the number of lines of the matrix.

## Returns

int

```
487 {
488     return Lines;
489 }
```

Here is the caller graph for this function:



## 5.3.3.3 Matrix Matrix::operator\* ( const real a ) const

Returns the product of the current matrix and a real *a*.

Example :

```
#include "Matrix.h"
Matrix M(2,2);
Matrix P;
real x = 2.;
...
P = M*x;
```

The operation  $P = x*M$  can also be done. See [operator\\*\(const real a, const Matrix& M\)](#).

Parameters

|   |            |
|---|------------|
| a | Real value |
|---|------------|

Returns

Matrix

```
281 {
282     int i,j;
283     Matrix result(Lines, Columns);
284
285     for (i = 1; i <= Lines; i++)
286         for (j = 1; j <= Columns; j++)
287             result.setValue(i, j, value(i,j)*a);
288
289     return result;
290 }
```

### 5.3.3.4 Matrix Matrix::operator\*( const Matrix & M ) const

Returns the product of the current matrix and a matrix  $M$ .

Example :

```
#include "Matrix.h"

Matrix M(2,3);
Matrix P(3,1);
Matrix Q;

...

Q = M*P;
```

Parameters

|     |                        |
|-----|------------------------|
| $M$ | <a href="#">Matrix</a> |
|-----|------------------------|

Returns

[Matrix](#)

```
332 {
333     int i, j, k;
334     Matrix result(lines(), M.columns());
335
336     for (i = 1; i <= lines(); i++)
337         for (j = 1; j <= M.columns(); j++)
338             for (k = 1; k <= columns(); k++)
339                 result.setValue(i, j, result.value(i, j) + value(i, k) * M.value(k, j));
340
341     return result;
342 }
```

### 5.3.3.5 Vector Matrix::operator\*( const Vector & V ) const

Returns the product of the current matrix and a vector  $V$ .

Example :

```
#include "Matrix.h"

Matrix M(2,3);
Vector V(3);
Vector P;

...

P = M*V;
```

Parameters

|     |                        |
|-----|------------------------|
| $V$ | <a href="#">Vector</a> |
|-----|------------------------|

Returns

[Vector](#)

```
351 {
352     int i, k;
353
354     Vector result(lines());
355
356     for (i = 1; i <= lines(); i++)
357         for (k = 1; k <= columns(); k++)
358             result.setValue(i, result.value(i) + value(i, k) * V.value(k));
```



```

359
360     return result;
361
362 }

```

### 5.3.3.6 void Matrix::operator\*=( const real a )

Multiplies the current matrix by a real *a*.

Example :

```

#include "Matrix.h"

Matrix M(2,2);

real x = 2.;

...

M *= x;

```

Parameters

|  |   |            |
|--|---|------------|
|  | a | Real value |
|--|---|------------|

Returns

void

```

269 {
270     int n;
271
272     for (n=1; n<=Lines*Columns; n++)
273         *(U+n-1) *= a;
274 }

```

### 5.3.3.7 Matrix Matrix::operator+( const Matrix & M ) const

Returns the addition of the current matrix and *M*.

Example :

```

#include "Matrix.h"

Matrix M(2,2);

Matrix P(2,2);

Matrix U;

...

U = M + P;

```

Parameters

|  |          |        |
|--|----------|--------|
|  | <i>M</i> | Matrix |
|--|----------|--------|

Returns

Matrix

```

202 {
203     int i, j;
204     Matrix result(Lines, Columns);
205
206     for (i = 1; i <= Lines; i++)
207         for (j = 1; j <= Columns; j++)
208             result.setValue(i, j, value(i,j) + M.value(i,j));
209
210     return result;
211 }

```

**5.3.3.8 void Matrix::operator+=( const Matrix & M )**

Adds *M* to the current matrix.

Example :

```
#include "Matrix.h"
Matrix M(2,2);
Matrix P(2,2);
...
P += M;
```

Parameters

|          |                        |
|----------|------------------------|
| <i>M</i> | <a href="#">Matrix</a> |
|----------|------------------------|

Returns

void

```
188 {
189     int i, j;
190
191     for (i = 1; i <= Lines; i++)
192     for (j = 1; j <= Columns; j++)
193         setValue(i, j, value(i, j) + M.value(i, j));
194 }
```

**5.3.3.9 Matrix Matrix::operator-( const Matrix & M ) const**

Returns the difference between the current matrix and *M*.

Example :

```
#include "Matrix.h"
Matrix M(2,2);
Matrix P(2,2);
Matrix U;
...
U = M - P;
```

Parameters

|          |                        |
|----------|------------------------|
| <i>M</i> | <a href="#">Matrix</a> |
|----------|------------------------|

Returns

[Matrix](#)

```
235 {
236     int i, j;
237     Matrix result(Lines, Columns);
238
239     for (i = 1; i <= Lines; i++)
240     for (j = 1; j <= Columns; j++)
241         result.setValue(i, j, value(i, j) - M.value(i, j));
242
243     return result;
244 }
```

**5.3.3.10 Matrix Matrix::operator-( ) const**

Returns the opposite of the current matrix.

Example :

```
#include "Matrix.h"
Matrix M(2,2);
Matrix P;
...
P = -M;
```

Returns

[Matrix](#)

```
250 {
251     int i, j;
252     Matrix result(Lines, Columns);
253
254     for (i = 1; i <= Lines; i++)
255         for (j = 1; j <= Columns; j++)
256             result.setValue(i, j, -value(i, j));
257
258     return result;
259 }
```

**5.3.3.11 void Matrix::operator-= ( const Matrix & M )**

Subtracts *M* to the current matrix.

Example :

```
#include "Matrix.h"
Matrix M(2,2);
Matrix P(2,2);
...
P -= M;
```

Parameters

|          |                        |
|----------|------------------------|
| <i>M</i> | <a href="#">Matrix</a> |
|----------|------------------------|

Returns

void

```
221 {
222     int i, j;
223
224     for (i = 1; i <= Lines; i++)
225         for (j = 1; j <= Columns; j++)
226             setValue(i, j, value(i, j) - M.value(i, j));
227 }
```

**5.3.3.12 Matrix Matrix::operator/( const real a ) const**

Returns the division of the current matrix by a real *a*.

Example :

```
#include "Matrix.h"
```

```
Matrix M(2,2);
```

```
Matrix P;
```

```
real x = 2.;
```

```
...
```

```
P = M / x;
```

Parameters

|   |            |
|---|------------|
| a | Real value |
|---|------------|

Returns

**Matrix**

```
313 {
314     int i,j;
315     Matrix result(Lines, Columns);
316
317     for (i = 1; i <= Lines; i++)
318         for (j = 1; j <= Columns; j++)
319             result.setValue(i, j, value(i,j)/a);
320
321     return result;
322 }
```

### 5.3.3.13 void Matrix::operator/= ( const real a )

Divides the current matrix by a real *a*.

Example :

```
#include "Matrix.h"
```

```
Matrix M(2,2);
```

```
real x = 2.;
```

```
...
```

```
M /= x;
```

Parameters

|   |            |
|---|------------|
| a | Real value |
|---|------------|

Returns

void

```
300 {
301     int n;
302
303     for (n=1; n<=Lines*Columns; n++)
304         *(U+n-1) /= a;
305 }
```

### 5.3.3.14 void Matrix::operator= ( const Matrix & M )

Set the current matrix to the dimension and the value of *M*.

Example :

```
#include "Matrix.h"
```

```
Matrix M(2,2);
```

```
Matrix P;
```

```
...
```

```
P = M;
```

Parameters

|          |                        |
|----------|------------------------|
| <i>M</i> | <a href="#">Matrix</a> |
|----------|------------------------|

Returns

```
void

162 {
163     int i, j;
164
165     if (M.lines() != Lines || M.columns() != Columns)
166     {
167         delete[] U;
168
169         Lines = M.lines();
170         Columns = M.columns();
171
172         U = new real[Lines*Columns];
173     }
174
175     for (i = 1; i <= Lines; i++)
176     for (j = 1; j <= Columns; j++)
177         setValue(i, j, M.value(i, j));
178 }
```

### 5.3.3.15 bool Matrix::operator==( const Matrix & M ) const

Compares the current matrix to a matrix *M* and returns true if they are equal.

Example :

```
#include "Matrix.h"

Matrix M(2,2);
Matrix P(2,2);

real x;

...

if (M == P)

x = M.value(1,1);
```

Parameters

|          |                        |
|----------|------------------------|
| <i>M</i> | <a href="#">Matrix</a> |
|----------|------------------------|

Returns

```
bool

144 {
145     int i, j;
146
147     for (i = 1; i <= Lines; i++)
148     for (j = 1; j <= Columns; j++)
149         if ( value(i,j) != M.value(i,j) ) return false;
150
151     return true;
152 }
```

### 5.3.3.16 void Matrix::setEigenMatrix ( const bool *isLeft*, const int *AxisNo*, const Vector *V*, const real *c*, const real *h=0.* )

Sets matrix as eigenmatrix.

## Parameters

|               |                                                    |
|---------------|----------------------------------------------------|
| <i>isLeft</i> | Boolean variable. True if location is on the left. |
| <i>AxisNo</i> | Axis of interest.                                  |
| <i>V</i>      | <a href="#">Vector</a>                             |
| <i>c</i>      | Real value                                         |
| <i>h</i>      | Real value                                         |

## Returns

void

```

371 {
372
373     // --- Local variables ---
374
375     int n; // Counter
376     real u = V.value(1);
377     real v = (Dimension > 1) ? V.value(2): 0.;
378     real w = (Dimension > 2) ? V.value(3): 0.;
379
380     // --- Allocate memory and set to zero ---
381
382     delete[] U;
383
384     Lines = Columns = QuantityNb;
385
386     U = new real[Lines*Columns];
387
388     for (n = 1; n <= Lines*Columns; n++)
389         *(U+n-1) = 0.;
390
391     // --- Fill values ---
392
393     switch (Dimension)
394     {
395
396         // --- 1D CASE -----
397
398         case 1:
399             if (isLeft)
400             {
401                 // --- Left eigenmatrix ---
402
403                 //
404                 //
405                 //
406                 //
407                 //
408                 //
409                 //
410                 //
411                 //
412                 //
413                 //
414
415                 //
416                 //
417                 //
418                 //
419                 //
420                 //
421                 //
422                 //
423                 //
424                 //
425                 //
426                 //
427                 //
428
429                 //
430                 //
431                 //
432                 //
433                 //
434                 //
435                 //
436                 //
437                 //
438                 //
439                 //
440                 //

```

```

441 //
442 //         setValue(1,3, 1./(c*c));
443 //         setValue(2,3, (u-c)/(c*c));
444 //         setValue(3,3, (h-u*c)/(c*c));
445
446
447 //         matrices de Vinokur
448
449         setValue(1,1, 1.);
450         setValue(2,1, u);
451         setValue(3,1, .5*(u*u));
452
453         setValue(1,2, 1.);
454         setValue(2,2, (u+c));
455         setValue(3,2, (h+ c*u));
456
457         setValue(1,3, 1.);
458         setValue(2,3, (u-c));
459         setValue(3,3, (h-u*c));
460
461     }
462     break;
463
464
465 // --- 2D CASE -----
466
467
468 case 2:
469     if (isLeft)
470     {
471         // --- Left eigenmatrix ---
472
473         switch (AxisNo)
474         {
475             case 1:
476 //         setValue(1,1, (c*c)*(1.-((Gamma-1.)*(u*u+v*v)/(2.*c*c)));
477 //         setValue(2,1, (c*c)*(-v));
478 //         setValue(3,1, (c*c)*(.5*( (Gamma-1.)*(u*u+v*v)/(2.*c*c) - u/c ));
479 //         setValue(4,1, (c*c)*(.5*( (Gamma-1.)*(u*u+v*v)/(2.*c*c) + u/c ));
480 //
481 //         setValue(1,2, (c*c)*((Gamma-1.)*u/(c*c));
482 //         setValue(2,2, 0.);
483 //         setValue(3,2, (c*c)*((c/(Gamma-1.))-u)*((Gamma-1.)/(2.*c*c)));
484 //         setValue(4,2, (c*c)*((-c/(Gamma-1.))-u)*((Gamma-1.)/(2.*c*c)));
485 //
486 //         setValue(1,3, (c*c)*((Gamma-1.)*v/(c*c));
487 //         setValue(2,3, (c*c)*1.);
488 //         setValue(3,3, (c*c)*(-(Gamma-1.)*v/(2.*c*c));
489 //         setValue(4,3, (c*c)*(-(Gamma-1.)*v/(2.*c*c));
490 //
491 //         setValue(1,4, (c*c)*(-(Gamma-1.)/(c*c));
492 //         setValue(2,4, 0.);
493 //         setValue(3,4, (c*c)*((Gamma-1.)/(2.*c*c));
494 //         setValue(4,4, (c*c)*((Gamma-1.)/(2.*c*c));
495
496
497 //         matrix vinokur
498
499         setValue(1,1, (1.-((Gamma-1.)*(u*u+v*v)/(2.*c*c)));
500         setValue(2,1, (-v));
501         setValue(3,1, (.5*( (Gamma-1.)*(u*u+v*v)/(2.*c*c) - u/c ));
502         setValue(4,1, (.5*( (Gamma-1.)*(u*u+v*v)/(2.*c*c) + u/c ));
503
504         setValue(1,2, ((Gamma-1.)*u/(c*c));
505         setValue(2,2, 0.);
506         setValue(3,2, ((c/(Gamma-1.))-u)*((Gamma-1.)/(2.*c*c)));
507         setValue(4,2, ((-c/(Gamma-1.))-u)*((Gamma-1.)/(2.*c*c)));
508
509         setValue(1,3, ((Gamma-1.)*v/(c*c));
510         setValue(2,3, 1.);
511         setValue(3,3, -(Gamma-1.)*v/(2.*c*c));
512         setValue(4,3, -(Gamma-1.)*v/(2.*c*c));
513
514         setValue(1,4, -(Gamma-1.)/(c*c));
515         setValue(2,4, 0.);
516         setValue(3,4, ((Gamma-1.)/(2.*c*c));
517         setValue(4,4, ((Gamma-1.)/(2.*c*c));
518
519
520         break;
521
522     case 2:
523 //         setValue(1,1, (c*c)*(1.-((Gamma-1.)*(u*u+v*v)/(2.*c*c)));
524 //         setValue(2,1, (c*c)*(-v));
525 //         setValue(3,1, (c*c)*(.5*( (Gamma-1.)*(u*u+v*v)/(2.*c*c) - u/c ));
526 //         setValue(4,1, (c*c)*(.5*( (Gamma-1.)*(u*u+v*v)/(2.*c*c) + u/c ));
527 //

```

```

528 //          setValue(1,2, (c*c)*((Gamma-1.)*u/(c*c)));
529 //          setValue(2,2, 0.);
530 //          setValue(3,2, (c*c)*((c/(Gamma-1.)-u)*((Gamma-1.)/(2.*c*c)));
531 //          setValue(4,2, (c*c)*((-c/(Gamma-1.)-u)*((Gamma-1.)/(2.*c*c)));
532 //
533 //          setValue(1,3, (c*c)*((Gamma-1.)*v/(c*c)));
534 //          setValue(2,3, (c*c)*1.);
535 //          setValue(3,3, (c*c)*(-(Gamma-1.)*v/(2.*c*c)));
536 //          setValue(4,3, (c*c)*(-(Gamma-1.)*v/(2.*c*c)));
537 //
538 //          setValue(1,4, (c*c)*(-(Gamma-1.)/(c*c)));
539 //          setValue(2,4, 0.);
540 //          setValue(3,4, (c*c)*((Gamma-1.)/(2.*c*c)));
541 //          setValue(4,4, (c*c)*((Gamma-1.)/(2.*c*c)));
542
543
544 //      matrix vinokur
545 //          setValue(1,1, (1.-((Gamma-1.)*(u*u+v*v)/(2.*c*c)));
546 //          setValue(2,1, (-v));
547 //          setValue(3,1, (.5*((Gamma-1.)*(u*u+v*v)/(2.*c*c) - u/c ));
548 //          setValue(4,1, (.5*((Gamma-1.)*(u*u+v*v)/(2.*c*c) + u/c ));
549
550 //          setValue(1,2, ((Gamma-1.)*u/(c*c)));
551 //          setValue(2,2, 0.);
552 //          setValue(3,2, ((c/(Gamma-1.)-u)*((Gamma-1.)/(2.*c*c)));
553 //          setValue(4,2, ((-c/(Gamma-1.)-u)*((Gamma-1.)/(2.*c*c)));
554
555 //          setValue(1,3, ((Gamma-1.)*v/(c*c)));
556 //          setValue(2,3, 1.);
557 //          setValue(3,3, (-(Gamma-1.)*v/(2.*c*c)));
558 //          setValue(4,3, (-(Gamma-1.)*v/(2.*c*c)));
559
560 //          setValue(1,4, (-(Gamma-1.)/(c*c)));
561 //          setValue(2,4, 0.);
562 //          setValue(3,4, ((Gamma-1.)/(2.*c*c)));
563 //          setValue(4,4, ((Gamma-1.)/(2.*c*c)));
564
565
566 //          break;
567 //      };
568 //  }
569 //  else
570 //  {
571 //      // --- Right eigenmatrix ---
572
573 //      switch(AxisNo)
574 //      {
575 //          case 1:
576 //              setValue(1,1, 1./(c*c));
577 //              setValue(2,1, u/(c*c));
578 //              setValue(3,1, v/(c*c));
579 //              setValue(4,1, .5*(u*u+v*v)/(c*c));
580 //
581 //              setValue(1,2, 0.);
582 //              setValue(2,2, 0.);
583 //              setValue(3,2, 1./(c*c));
584 //              setValue(4,2, v/(c*c));
585 //
586 //              setValue(1,3, 1./(c*c));
587 //              setValue(2,3, (u+c)/(c*c));
588 //              setValue(3,3, v/(c*c));
589 //              setValue(4,3, (h+ c*u)/(c*c));
590 //
591 //              setValue(1,4, 1./(c*c));
592 //              setValue(2,4, (u-c)/(c*c));
593 //              setValue(3,4, v/(c*c));
594 //              setValue(4,4, (h- c*u)/(c*c));
595
596 //      matrix vinokur
597 //          setValue(1,1, 1.);
598 //          setValue(2,1, u);
599 //          setValue(3,1, v);
600 //          setValue(4,1, .5*(u*u+v*v));
601
602 //          setValue(1,2, 0.);
603 //          setValue(2,2, 0.);
604 //          setValue(3,2, 1.);
605 //          setValue(4,2, v);
606
607 //          setValue(1,3, 1.);
608 //          setValue(2,3, (u+c));
609 //          setValue(3,3, v);
610 //          setValue(4,3, (h+ c*u));
611
612 //          setValue(1,4, 1.);
613 //          setValue(2,4, (u-c));
614

```



```

615         setValue(3,4, v);
616         setValue(4,4, (h- c*u));
617
618         break;
619
620     case 2:
621 //         setValue(1,1, 0.);
622 //         setValue(2,1, 0.);
623 //         setValue(3,1, 1./(c*c));
624 //         setValue(4,1, u/(c*c));
625 //
626 //         setValue(1,2, 1./(c*c));
627 //         setValue(2,2, u/(c*c));
628 //         setValue(3,2, v/(c*c));
629 //         setValue(4,2, .5*(u+u+v*v)/(c*c));
630 //
631 //         setValue(1,3, 1./(c*c));
632 //         setValue(2,3, u/(c*c));
633 //         setValue(3,3, (v+c)/(c*c));
634 //         setValue(4,3, (h+ c*v)/(c*c));
635 //
636 //         setValue(1,4, 1./(c*c));
637 //         setValue(2,4, u/(c*c));
638 //         setValue(3,4, (v-c)/(c*c));
639 //         setValue(4,4, (h- c*v)/(c*c));
640
641
642 //     matrix vinokur
643
644         setValue(1,1, 0.);
645         setValue(2,1, 0.);
646         setValue(3,1, 1.);
647         setValue(4,1, u);
648
649         setValue(1,2, 1.);
650         setValue(2,2, u);
651         setValue(3,2, v);
652         setValue(4,2, .5*(u+u+v*v));
653
654         setValue(1,3, 1.);
655         setValue(2,3, u);
656         setValue(3,3, (v+c));
657         setValue(4,3, (h+ c*v));
658
659         setValue(1,4, 1.);
660         setValue(2,4, u);
661         setValue(3,4, (v-c));
662         setValue(4,4, (h- c*v));
663
664         break;
665     };
666 }
667 break;
668
669 // --- 3D CASE -----
670
671 case 3:
672     if (isLeft)
673     {
674         // --- Left eigenmatrix ---
675
676         switch(AxisNo)
677         {
678 //             case 1:
679 //                 setValue(1,1, (c*c)*(1.- (Gamma-1.)*(u+u+v*v+w*w)/(2.*c*c)));
680 //                 setValue(2,1, (c*c)*(-v));
681 //                 setValue(3,1, (c*c)*(-w));
682 //                 setValue(4,1, (c*c)*(.5*( Gamma-1.)*(u+u+v*v+w*w)/(2.*c*c) - u/c ));
683 //                 setValue(5,1, (c*c)*(.5*( Gamma-1.)*(u+u+v*v+w*w)/(2.*c*c) + u/c ));
684 //
685 //                 setValue(1,2, (c*c)*((Gamma-1.)*u/(c*c));
686 //                 setValue(2,2, 0.);
687 //                 setValue(3,2, 0.);
688 //                 setValue(4,2, (c*c)*((c/(Gamma-1.)-u)*((Gamma-1.)/(2.*c*c)));
689 //                 setValue(5,2, (c*c)*((-c/(Gamma-1.)-u)*((Gamma-1.)/(2.*c*c)));
690 //
691 //                 setValue(1,3, (c*c)*((Gamma-1.)*v/(c*c));
692 //                 setValue(2,3, (c*c)*(1.));
693 //                 setValue(3,3, 0.);
694 //                 setValue(4,3, (c*c)*(-v*(Gamma-1.)/(2.*c*c));
695 //                 setValue(5,3, (c*c)*(-v*(Gamma-1.)/(2.*c*c));
696 //
697 //                 setValue(1,4, (c*c)*((Gamma-1.)*w/(c*c));
698 //                 setValue(2,4, 0.);
699 //                 setValue(3,4, (c*c)*(1.));
700 //                 setValue(4,4, (c*c)*(-w*(Gamma-1.)/(2.*c*c));
701 //                 setValue(5,4, (c*c)*(-w*(Gamma-1.)/(2.*c*c));

```

```

702 //
703 //         setValue(1,5, (c*c)*(-(Gamma-1.)/(c*c));
704 //         setValue(2,5, 0.);
705 //         setValue(3,5, 0.);
706 //         setValue(4,5, (c*c)*((Gamma-1.)/(2.*c*c));
707 //         setValue(5,5, (c*c)*((Gamma-1.)/(2.*c*c));
708 //         break;
709
710 //     matrix vinokur
711 //     case 1:
712 //         setValue(1,1, (1.-(Gamma-1.)*(u*u+v*v+w*w)/(2.*c*c));
713 //         setValue(2,1, (-v));
714 //         setValue(3,1, (-w));
715 //         setValue(4,1, (.5*(Gamma-1.)*(u*u+v*v+w*w)/(2.*c*c) - u/c));
716 //         setValue(5,1, (.5*(Gamma-1.)*(u*u+v*v+w*w)/(2.*c*c) + u/c));
717
718 //         setValue(1,2, (Gamma-1.)*u/(c*c));
719 //         setValue(2,2, 0.);
720 //         setValue(3,2, 0.);
721 //         setValue(4,2, ((c/(Gamma-1.)-u)*((Gamma-1.)/(2.*c*c)));
722 //         setValue(5,2, ((-c/(Gamma-1.)-u)*((Gamma-1.)/(2.*c*c)));
723
724 //         setValue(1,3, ((Gamma-1.)*v/(c*c));
725 //         setValue(2,3, (1.));
726 //         setValue(3,3, 0.);
727 //         setValue(4,3, -v*(Gamma-1.)/(2.*c*c));
728 //         setValue(5,3, -v*(Gamma-1.)/(2.*c*c));
729
730 //         setValue(1,4, (Gamma-1.)*w/(c*c));
731 //         setValue(2,4, 0.);
732 //         setValue(3,4, (1.));
733 //         setValue(4,4, -w*(Gamma-1.)/(2.*c*c));
734 //         setValue(5,4, -w*(Gamma-1.)/(2.*c*c));
735
736 //         setValue(1,5, (-(Gamma-1.)/(c*c));
737 //         setValue(2,5, 0.);
738 //         setValue(3,5, 0.);
739 //         setValue(4,5, (Gamma-1.)/(2.*c*c));
740 //         setValue(5,5, (Gamma-1.)/(2.*c*c));
741 //         break;
742
743
744
745
746
747
748
749
750 //     case 2:
751 //         setValue(1,1, (c*c)*(-u));
752 //         setValue(2,1, (c*c)*(1.-(Gamma-1.)*(u*u+v*v+w*w)/(2.*c*c));
753 //         setValue(3,1, (c*c)*(-w));
754 //         setValue(4,1, (c*c)*(.5*(Gamma-1.)*(u*u+v*v+w*w)/(2.*c*c) - v/c));
755 //         setValue(5,1, (c*c)*(.5*(Gamma-1.)*(u*u+v*v+w*w)/(2.*c*c) + v/c));
756 //
757 //         setValue(1,2, 0.);
758 //         setValue(2,2, (c*c)*((Gamma-1.)*u/(c*c));
759 //         setValue(3,2, 0.);
760 //         setValue(4,2, (c*c)*(-u*(Gamma-1.)/(2.*c*c));
761 //         setValue(5,2, (c*c)*(-u*(Gamma-1.)/(2.*c*c));
762 //
763 //         setValue(1,3, (c*c)*(1.));
764 //         setValue(2,3, (c*c)*((Gamma-1.)*v/(c*c));
765 //         setValue(3,3, 0.);
766 //         setValue(4,3, (c*c)*((c/(Gamma-1.)-v)*((Gamma-1.)/(2.*c*c)));
767 //         setValue(5,3, (c*c)*((-c/(Gamma-1.)-v)*((Gamma-1.)/(2.*c*c)));
768 //
769 //         setValue(1,4, 0.);
770 //         setValue(2,4, (c*c)*((Gamma-1.)*w/(c*c));
771 //         setValue(3,4, (c*c)*(1.));
772 //         setValue(4,4, (c*c)*(-w*(Gamma-1.)/(2.*c*c));
773 //         setValue(5,4, (c*c)*(-w*(Gamma-1.)/(2.*c*c));
774 //
775 //         setValue(1,5, 0.);
776 //         setValue(2,5, (c*c)*(-(Gamma-1.)/(c*c));
777 //         setValue(3,5, 0.);
778 //         setValue(4,5, (c*c)*((Gamma-1.)/(2.*c*c));
779 //         setValue(5,5, (c*c)*((Gamma-1.)/(2.*c*c));
780 //         break;
781
782 //     matrix vinokur
783 //     case 2:
784 //         setValue(1,1, -u);
785 //         setValue(2,1, (1.-(Gamma-1.)*(u*u+v*v+w*w)/(2.*c*c));
786 //         setValue(3,1, -w);
787 //         setValue(4,1, (.5*(Gamma-1.)*(u*u+v*v+w*w)/(2.*c*c) - v/c));
788 //         setValue(5,1, (.5*(Gamma-1.)*(u*u+v*v+w*w)/(2.*c*c) + v/c));

```

```

789
790     setValue(1,2, 0.);
791     setValue(2,2, (Gamma-1.)*u/(c*c));
792     setValue(3,2, 0.);
793     setValue(4,2, (-u*(Gamma-1.)/(2.*c*c)));
794     setValue(5,2, (-u*(Gamma-1.)/(2.*c*c)));
795
796     setValue(1,3, (c*c)*(1.));
797     setValue(2,3, (c*c)*((Gamma-1.)*v/(c*c)));
798     setValue(3,3, 0.);
799     setValue(4,3, (c*c)*((c/(Gamma-1.))-v)*((Gamma-1.)/(2.*c*c)));
800     setValue(5,3, (c*c)*((-c/(Gamma-1.))-v)*((
Gamma-1.)/(2.*c*c))););
801
802     setValue(1,4, 0.);
803     setValue(2,4, ((Gamma-1.)*w/(c*c)));
804     setValue(3,4, 1.);
805     setValue(4,4, -w*(Gamma-1.)/(2.*c*c));
806     setValue(5,4, -w*(Gamma-1.)/(2.*c*c));
807
808     setValue(1,5, 0.);
809     setValue(2,5, -(Gamma-1.)/(c*c));
810     setValue(3,5, 0.);
811     setValue(4,5, (Gamma-1.)/(2.*c*c));
812     setValue(5,5, (Gamma-1.)/(2.*c*c));
813     break;
814
815
816
817 //     case 3:
818 //     setValue(1,1, (c*c)*(-u));
819 //     setValue(2,1, (c*c)*(-v));
820 //     setValue(3,1, (c*c)*(1-((Gamma-1.)*(u*u+v*v+w*w)/(2.*c*c))));
821 //     setValue(4,1, (c*c)*(1.5*((Gamma-1.)*(u*u+v*v+w*w)/(2.*c*c) - w/c)));
822 //     setValue(5,1, (c*c)*(1.5*((Gamma-1.)*(u*u+v*v+w*w)/(2.*c*c) + w/c)));
823 //
824 //     setValue(1,2, (c*c)*(1.));
825 //     setValue(2,2, 0.);
826 //     setValue(3,2, (c*c)*((Gamma-1.)*u/(c*c)));
827 //     setValue(4,2, (c*c)*(-(Gamma-1.)*u/(2.*c*c)));
828 //     setValue(5,2, (c*c)*(-(Gamma-1.)*u/(2.*c*c)));
829 //
830 //     setValue(1,3, 0.);
831 //     setValue(2,3, (c*c)*(1.));
832 //     setValue(3,3, (c*c)*((Gamma-1.)*v/(c*c)));
833 //     setValue(4,3, (c*c)*(-(Gamma-1.)*v/(2.*c*c)));
834 //     setValue(5,3, (c*c)*(-(Gamma-1.)*v/(2.*c*c)));
835 //
836 //     setValue(1,4, 0.);
837 //     setValue(2,4, 0.);
838 //     setValue(3,4, (c*c)*((Gamma-1.)*w/(c*c)));
839 //     setValue(4,4, (c*c)*((c/(Gamma-1.))-w)*((Gamma-1.)/(2.*c*c)));
840 //     setValue(5,4, (c*c)*((-c/(Gamma-1.))-w)*((Gamma-1.)/(2.*c*c))););
841 //
842 //     setValue(1,5, 0.);
843 //     setValue(2,5, 0.);
844 //     setValue(3,5, (c*c)*(-(Gamma-1.)/(c*c)));
845 //     setValue(4,5, (c*c)*((Gamma-1.)/(2.*c*c)));
846 //     setValue(5,5, (c*c)*((Gamma-1.)/(2.*c*c)));
847 //     break;
848
849
850 //     matrix vinokur
851 //     case 3:
852 //     setValue(1,1, -u);
853 //     setValue(2,1, -v);
854 //     setValue(3,1, (1-((Gamma-1.)*(u*u+v*v+w*w)/(2.*c*c))));
855 //     setValue(4,1, (1.5*((Gamma-1.)*(u*u+v*v+w*w)/(2.*c*c) - w/c)));
856 //     setValue(5,1, (1.5*((Gamma-1.)*(u*u+v*v+w*w)/(2.*c*c) + w/c))););
857
858 //     setValue(1,2, 1.);
859 //     setValue(2,2, 0.);
860 //     setValue(3,2, (Gamma-1.)*u/(c*c));
861 //     setValue(4,2, -(Gamma-1.)*u/(2.*c*c));
862 //     setValue(5,2, -(Gamma-1.)*u/(2.*c*c));
863
864 //     setValue(1,3, 0.);
865 //     setValue(2,3, 1.);
866 //     setValue(3,3, (Gamma-1.)*v/(c*c));
867 //     setValue(4,3, -(Gamma-1.)*v/(2.*c*c));
868 //     setValue(5,3, -(Gamma-1.)*v/(2.*c*c));
869
870 //     setValue(1,4, 0.);
871 //     setValue(2,4, 0.);
872 //     setValue(3,4, (Gamma-1.)*w/(c*c));
873 //     setValue(4,4, ((c/(Gamma-1.))-w)*((Gamma-1.)/(2.*c*c))););
874 //     setValue(5,4, ((-c/(Gamma-1.))-w)*((Gamma-1.)/(2.*c*c))););

```

```

875
876         setValue(1,5, 0.);
877         setValue(2,5, 0.);
878         setValue(3,5, -(Gamma-1.)/(c*c));
879         setValue(4,5, (Gamma-1.)/(2.*c*c));
880         setValue(5,5, (Gamma-1.)/(2.*c*c));
881         break;
882
883
884
885     };
886 }
887 else
888 {
889     // --- Right eigenmatrix ---
890
891     switch(AxisNo)
892     {
893     //         case 1:
894     //             setValue(1,1, 1./(c*c));
895     //             setValue(2,1, u/(c*c));
896     //             setValue(3,1, v/(c*c));
897     //             setValue(4,1, w/(c*c));
898     //             setValue(5,1, .5*(u*u+v*v+w*w)/(c*c));
899     //
900     //             setValue(1,2, 0.);
901     //             setValue(2,2, 0.);
902     //             setValue(3,2, 1./(c*c));
903     //             setValue(4,2, 0.);
904     //             setValue(5,2, v/(c*c));
905     //
906     //             setValue(1,3, 0.);
907     //             setValue(2,3, 0.);
908     //             setValue(3,3, 0.);
909     //             setValue(4,3, 1./(c*c));
910     //             setValue(5,3, w/(c*c));
911     //
912     //             setValue(1,4, 1./(c*c));
913     //             setValue(2,4, (u+c)/(c*c));
914     //             setValue(3,4, v/(c*c));
915     //             setValue(4,4, w/(c*c));
916     //             setValue(5,4, (h+ c*u)/(c*c));
917     //
918     //             setValue(1,5, 1.);
919     //             setValue(2,5, (u-c)/(c*c));
920     //             setValue(3,5, v/(c*c));
921     //             setValue(4,5, w/(c*c));
922     //             setValue(5,5, (h- c*u)/(c*c));
923     //             break;
924
925     //         matrix vinokur
926     //             case 1:
927     //                 setValue(1,1, 1.);
928     //                 setValue(2,1, u);
929     //                 setValue(3,1, v);
930     //                 setValue(4,1, w);
931     //                 setValue(5,1, .5*(u*u+v*v+w*w));
932     //
933     //                 setValue(1,2, 0.);
934     //                 setValue(2,2, 0.);
935     //                 setValue(3,2, 1.);
936     //                 setValue(4,2, 0.);
937     //                 setValue(5,2, v);
938     //
939     //                 setValue(1,3, 0.);
940     //                 setValue(2,3, 0.);
941     //                 setValue(3,3, 0.);
942     //                 setValue(4,3, 1.);
943     //                 setValue(5,3, w);
944     //
945     //                 setValue(1,4, 1.);
946     //                 setValue(2,4, u+c);
947     //                 setValue(3,4, v);
948     //                 setValue(4,4, w);
949     //                 setValue(5,4, h+ c*u);
950     //
951     //                 setValue(1,5, 1.);
952     //                 setValue(2,5, u-c);
953     //                 setValue(3,5, v);
954     //                 setValue(4,5, w);
955     //                 setValue(5,5, h- c*u);
956     //                 break;
957
958
959     //         matrix vinokur
960     //             case 2:
961     //                 setValue(1,1, 0.);

```

```

962         setValue(2,1, 0.);
963         setValue(3,1, 1.);
964         setValue(4,1, 0.);
965         setValue(5,1, u);
966
967         setValue(1,2, 1.);
968         setValue(2,2, u);
969         setValue(3,2, v);
970         setValue(4,2, w);
971         setValue(5,2, .5*(u*u+v*v+w*w));
972
973         setValue(1,3, 0.);
974         setValue(2,3, 0.);
975         setValue(3,3, 0.);
976         setValue(4,3, 1.);
977         setValue(5,3, w);
978
979         setValue(1,4, 1.);
980         setValue(2,4, u);
981         setValue(3,4, v+c);
982         setValue(4,4, w);
983         setValue(5,4, h+ c*v);
984
985         setValue(1,5, 1.);
986         setValue(2,5, u);
987         setValue(3,5, v-c);
988         setValue(4,5, w);
989         setValue(5,5, h- c*v);
990         break;
991
992 //         matrix vinokur
993
994         case 3:
995             setValue(1,1, 0.);
996             setValue(2,1, 1.);
997             setValue(3,1, 0.);
998             setValue(4,1, 0.);
999             setValue(5,1, u);
1000
1001             setValue(1,2, 0.);
1002             setValue(2,2, 0.);
1003             setValue(3,2, 1.);
1004             setValue(4,2, 0.);
1005             setValue(5,2, v);
1006
1007             setValue(1,3, 1.);
1008             setValue(2,3, u);
1009             setValue(3,3, v);
1010             setValue(4,3, w);
1011             setValue(5,3, .5*(u*u+v*v+w*w));
1012
1013             setValue(1,4, 1.);
1014             setValue(2,4, u);
1015             setValue(3,4, v);
1016             setValue(4,4, w+c);
1017             setValue(5,4, h+ c*w);
1018
1019             setValue(1,5, 1.);
1020             setValue(2,5, u);
1021             setValue(3,5, v);
1022             setValue(4,5, w-c);
1023             setValue(5,5, h- c*w);
1024             break;
1025         };
1026     }
1027     break;
1028 };
1029
1030 }

```

### 5.3.3.17 void Matrix::setValue ( const int *i*, const int *j*, const real *a* ) [inline]

Sets the component *i*, *j* to value *a*.

Example :

```

#include "Matrix.h"
Matrix M(2,2);
real x = 3.;

```

```

real y = 1.;
M.setValue(1,1,x);
M.setValue(2,1,y);

```

Parameters

|     |          |
|-----|----------|
| $i$ | Position |
| $j$ | Position |
| $a$ | Value    |

Returns

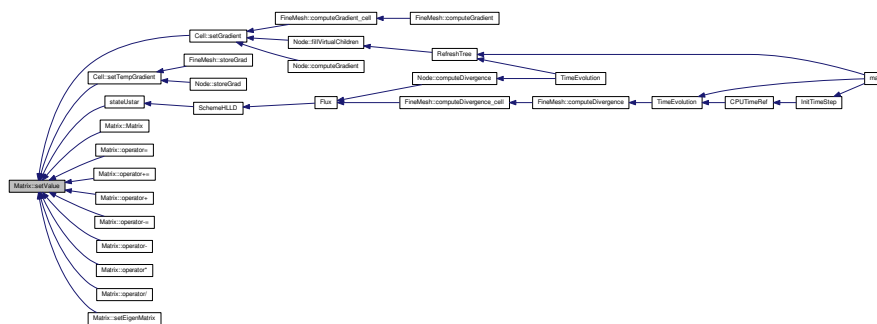
void

```

506 {
507     *( U + (i-1)*Columns + (j-1) ) = a;
508 }

```

Here is the caller graph for this function:



### 5.3.3.18 void Matrix::setZero ( )

Sets all the components to zero.

Returns

void

```

128 {
129     int n;
130
131     for (n = 1; n <= Lines*Columns; n++)
132         *(U+n-1) = 0.;
133 }

```

Here is the caller graph for this function:





- [Matrix.h](#)
- [Matrix.cpp](#)

## 5.4 Node Class Reference

An object [Node](#) is an element of a graded tree structure, used for multiresolution computations. Its contains the following informations:

```
#include <Node.h>
```

### Public Member Functions

- [Node](#) (const int l=0, const int i=0, const int j=0, const int k=0)  
*Constructor of [Node](#) class. Generates a new node at the position (l, i, j, k) in the tree structure. A new node is always a leaf. The array of pointers to the children is allocated, together with the informations on the corresponding cell: cell-center position and cell size.*
- [~Node](#) ()  
*Distructor of [Node](#) class. Removes the node from the tree structure. If the node is not a leaf, all the children are also removed.*
- int [cells](#) () const  
*Returns the number of cells in the tree.*
- int [leaves](#) () const  
*Returns the number of leaves in the tree.*
- int [adapt](#) ()  
*Computes the details in the leaves and its parent nodes and, in function of the threshold *Tolerance*, adapt the tree structure.*
- void [checkGradedTree](#) ()  
*Checks if the tree is graded. If not, an error is emitted. Only for debugging.*
- void [initValue](#) ()  
*Computes the initial value.*
- void [addLevel](#) ()  
*Adds levels when needed.*
- [Cell](#) \* [project](#) ()  
*Computes the cell-average values of all nodes that are not leaves by projection from the cell-averages values of the leaves. This procedure is required after a time evolution to refresh the internal nodes of the tree.*
- void [fillVirtualChildren](#) ()  
*Fills the cell-average values of every virtual leaf with values predicted from its parent and uncles. This procedure is required after a time evolution to refresh the virtual leaves of the tree.*
- void [store](#) ()  
*Stores cell-average values into temporary cell-average values.*
- void [storeGrad](#) ()  
*Stores gradient values into temporary gradient values.*
- void [computeDivergence](#) ()  
*Computes the divergence vector with the space discretization scheme.*
- void [RungeKutta](#) ()  
*Computes one Runge-Kutta step.*
- void [computeIntegral](#) ()  
*Computes integral values like e.g. flame velocity, global error, etc.*
- void [computeGradient](#) ()  
*Computes velocity gradient (only for Navier-Stokes).*
- void [computeCorrection](#) ()  
*Computes velocity gradient (only for Navier-Stokes).*



- void `checkStability ()`  
*Checks if the computation is numerically unstable, i.e. if one of the cell-averages is overflow. In case of numerical instability, the computation is stopped and a message appears.*
- void `writeTree (const char *FileName) const`  
*Writes tree structure into file FileName. Only for debugging.*
- void `writeAverage (const char *FileName)`  
*Writes cell-average values in multiresolution representation and the corresponding mesh into file FileName.*
- void `writeMesh (const char *FileName) const`  
*Writes mesh data for Gnuplot into file FileName.*
- void `writeHeader (const char *FileName) const`  
*Writes header for Data Explorer into file FileName.*
- void `writeFineGrid (const char *FileName, const int L=ScaleNb) const`  
*Writes cell-average values on a regular grid of level L into file FileName.*
- void `backup ()`  
*Backs up the tree structure and the cell-averages into a file `carmen.bak`. In further computations, the data can be recovered using `Restore()`.*
- void `restore ()`  
*Restores the tree structure and the cell-averages from the file `carmen.bak`. This file was created by the method `Backup()`.*
- void `restoreFineMesh ()`  
*Restores the tree structure and the cell-averages from the file `carmen.bak` in `FineMesh` format.*
- void `smooth ()`  
*Deletes the details in the highest level.*

### 5.4.1 Detailed Description

An object `Node` is an element of a graded tree structure, used for multiresolution computations. Its contains the following informations:

- A pointer to the root node `*Root` ;  
\* \* The corresponding cell `ThisCell` ;
- An array of pointers to the children nodes `**Child`. Each parent has  $2**Dimension$  children nodes ;
- The position of the node `Nl, Ni, Nj, Nk` into the tree structure (`Nl = level`) ;
- A `Flag` giving the kind of node : 0 = not a leaf, 1 = leaf, 2 = leaf with virtual children, 3 = virtual leaf.

A leaf is a node without children, a virtual leaf is an artificial leaf created only for the flux computations. No time evolution is made on virtual leaves.

### 5.4.2 Constructor & Destructor Documentation

#### 5.4.2.1 `Node::Node ( const int l = 0, const int i = 0, const int j = 0, const int k = 0 )`

Constructor of `Node` class. Generates a new node at the position  $(l, i, j, k)$  in the tree structure. A new node is always a leaf. The array of pointers to the children is allocated, together with the informations on the corresponding cell: cell-center position and cell size.

#### Parameters

---

|  |          |            |
|--|----------|------------|
|  | /        | Level      |
|  | <i>i</i> | Position x |
|  | <i>j</i> | Position y |
|  | <i>k</i> | Position z |

```

39 {
40     // Set Nl, Ni, Nj, Nk
41     Nl = 1;
42     Ni = i;
43     Nj = j;
44     Nk = k;
45
46     // --- If l = 0, then node is root ---
47
48     if (Nl == 0)
49     {
50         Root = this;
51         CellNb = 1;
52         LeafNb = 1;
53     }
54 // else if(CellNb<(power2(1<<ScaleNb))) CellNb++;
55 // else CellNb--;
56 // --- Increase the total number of cells ---
57
58     CellNb ++;
59
60     // --- Allocate array of pointers to children ---
61
62     Child = new Node* [ChildNb];
63
64     // --- A new node is a simple leaf ---
65
66     setSimpleLeaf();
67
68     // --- Set the coordinates and the size of the cell ---
69
70     // x-direction
71     ThisCell.setSize( 1, (XMax[1]-XMin[1])/(1<<Nl) );
72     ThisCell.setCenter( 1, XMin[1] + (Ni + .5)*ThisCell.size(1) );
73
74     // y-direction
75     if (Dimension > 1)
76     {
77         ThisCell.setSize( 2, (XMax[2]-XMin[2])/(1<<Nl) );
78         ThisCell.setCenter( 2, XMin[2] + (Nj + .5)*ThisCell.size(2) );
79     }
80
81     // z-direction
82     if (Dimension > 2)
83     {
84         ThisCell.setSize( 3, (XMax[3]-XMin[3])/(1<<Nl) );
85         ThisCell.setCenter( 3, XMin[3] + (Nk + .5)*ThisCell.size(3) );
86     }
87 }

```

#### 5.4.2.2 Node::~Node ( )

Destructor of [Node](#) class. Removes the node from the tree structure. If the node is not a leaf, all the children are also removed.

```

96 {
97     // --- Local variables -----
98
99     int n=0; // Counter on children
100
101     // --- Destructor procedure -----
102
103     // --- Decrease the total number of cells ---
104
105     CellNb --;
106
107     // --- If the node has children, delete them ---
108
109     if (hasChildren())
110         for (n = 0; n < ChildNb; n++) delete Child[n];
111
112     // --- Delete array of pointers to children ---
113
114     delete[] Child;
115 }

```

### 5.4.3 Member Function Documentation

#### 5.4.3.1 int Node::adapt ( )

Computes the details in the leaves and its parent nodes and, in function of the threshold *Tolerance*, adapt the tree structure.

#### Returns

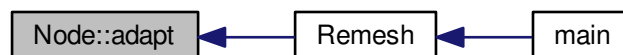
int

```

692 {
693     // --- Local variables ---
694
695     int     n;          // Counter on children
696     int     isDeletable; // Test if children are deletable (0 = all children are deletable)
697
698     // --- In case of time adaptivity, only remesh when a complete time evolution has been done ---
699
700     // --- Init ---
701
702     isDeletable = 0;
703
704     // --- Test to stop recursion ---
705
706     // If the node is not an internal node, this node can be deleted
707     if (!isInternalNode() || Nl >= ScaleNb) return 0;
708
709     // --- Recursion ---
710
711     // Test if children are deletable
712     for (n = 0; n < ChildNb; n++)
713         isDeletable += Child[n]->adapt();
714
715     // If all children are deletable, test if this node is also deletable
716
717     if (isDeletable == 0)
718     {
719         if (detailIsSmall())
720         {
721             if (!TimeAdaptivity || (TimeAdaptivity && isEndTimeCycle()))
722                 for (n = 0; n < ChildNb; n++) Child[n]->combine();
723
724             // Add value 0 to variable isDeletable of the parent
725             return 0;
726         }
727     }
728     else
729     {
730         if (!TimeAdaptivity || (TimeAdaptivity && isEndTimeCycle()))
731             for (n = 0; n < ChildNb; n++) Child[n]->split();
732
733         return 1;
734     }
735 }
736 // Add value 1 to variable Deletable of the parent
737 return 1;
738 }

```

Here is the caller graph for this function:



### 5.4.3.2 void Node::addLevel ( )

Adds levels when needed.

#### Returns

void

```

192 {
193     // --- Local variables -----
194
195     int n=0;    // Counter on children
196
197     // --- If level higher or equal to maximum scale number allowed, do not split
198
199     if ( Nl >= ScaleNb ) return;
200
201     // --- If node is not a leaf, recurse on children
202
203     if (isInternalNode())
204     {
205         for (n = 0 ; n < ChildNb; n++)
206             Child[n]->addLevel();
207     }
208     else
209     {
210         // If it is a virtual leaf, no splitting
211         if (isVirtualLeaf()) return;
212
213         // If it is on the wall, always split
214         if (UseBoundaryRegions && isOnBoundary())
215             split(true);
216
217         // Test on prediction error (always split on levels 0 and 1)
218         if (!detailIsSmall() || Nl <= 1)
219             split(true);
220     }
221 }
222 }

```

Here is the caller graph for this function:



### 5.4.3.3 void Node::backup ( )

Backs up the tree structure and the cell-averages into a file *carmen.bak*. In further computations, the data can be recovered using **Restore()**.

#### Returns

void

```

2717 {
2718     int n;                // Counter on children
2719     FILE* output;        // Output file
2720     int QuantityNo;      // Counter on quantities
2721
2722     // --- Init ---
2723
2724     if (Nl==0)
2725     {

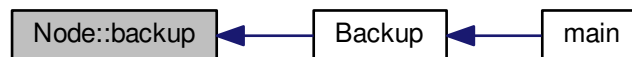
```

```

2726     output = fopen("carmen.bak","w");
2727
2728         // --- Write header ---
2729
2730     fprintf(output, "Backup at iteration %i, physical time %e\n",
IterationNo, ElapsedTime);
2731     }
2732     else
2733     output = fopen("carmen.bak","a");
2734
2735     // --- If node is not a leaf, recurse to children ---
2736
2737     if (isInternalNode())
2738     {
2739         fprintf(output,"N\n");
2740         fclose(output);
2741         for (n = 0; n < ChildNb; n++)
2742             Child[n]->backup();
2743     }
2744     else
2745     {
2746         for (QuantityNo=1; QuantityNo <= QuantityNb; QuantityNo++)
2747         {
2748             fprintf(output, FORMAT, ThisCell.average(QuantityNo));
2749             fprintf(output, "\n");
2750         }
2751         fclose(output);
2752     }
2753 }

```

Here is the caller graph for this function:



#### 5.4.3.4 int Node::cells ( ) const [inline]

Returns the number of cells in the tree.

##### Returns

int

```

685 {
686     return CellNb;
687 }

```

#### 5.4.3.5 void Node::checkGradedTree ( )

Checks if the tree is graded. If not, an error is emitted. Only for debugging.

##### Returns

void

```

2631 {
2632     // --- Local variables ---
2633
2634     int     n;                // Counter on children
2635     int     i, j, k;         // Counter in directions

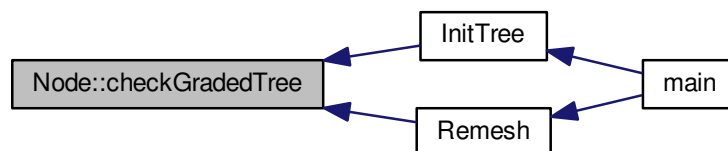
```

```

2636     int   ej, ek;           // 1 if this dimension is existing, 0 else.
2637
2638     // --- Init ---
2639     ej = (Dimension > 1)? 1:0;
2640     ek = (Dimension > 2)? 1:0;
2641
2642
2643     if (Nl == 0)
2644     {
2645         cout << "carmen: testing tree structure ... \n";
2646         for (n = 0; n < ChildNb; n++)
2647             Child[n]->checkGradedTree();
2648         cout << "carmen: tree structure OK. \n";
2649         return;
2650     }
2651
2652     // --- Test if neighbours are existing (eventually virtual) ---
2653
2654     for (i = -1; i <= 1; i+=1)
2655     for (j = -1*ej; j <= 1*ej; j+=1)
2656     for (k = -1*ek; k <= 1*ek; k+=1)
2657     {
2658         if (cell(Nl, Ni+i, Nj+j, Nk+k)==0)
2659         {
2660             cout << "carmen: Tree not graded' :\n";
2661             cout << "carmen: Node (" << Nl << ", " << Ni << ", " << Nj << ", " << Nk << ") \n";
2662             cout << "carmen: has missing neighbour (" << Nl << ", " << Ni+i << ", " << Nj+j << ", " << Nk+k <
2663 < ") \n";
2664             cout << "carmen: abort execution.\n";
2665             exit(1);
2666         }
2667     }
2668     // --- Recurse if it is a node ---
2669
2670     if (isInternalNode())
2671     {
2672         for (n = 0; n < ChildNb; n++)
2673             Child[n]->checkGradedTree();
2674     }
2675 }

```

Here is the caller graph for this function:



#### 5.4.3.6 void Node::checkStability ( )

Checks if the computation is numerically unstable, i.e. if one of the cell-averages is overflow. In case of numerical instability, the computation is stopped and a message appears.

#### Returns

void

```

2431 {
2432     // --- Local variables ---
2433
2434     int n,iaux;           // Counter on children
2435     real x=0., y=0., z=0.; // Real position
2436

```

```

2437     // --- Recursion ---
2438
2439     if (isInternalNode())
2440     {
2441         for (n = 0; n < ChildNb; n++)
2442             Child[n]->checkStability();
2443     }
2444     else
2445     {
2446         // --- Compute x, y, z ---
2447
2448         x = ThisCell.center(1);
2449         if (Dimension > 1) y = ThisCell.center(2);
2450         if (Dimension > 2) z = ThisCell.center(3);
2451
2452         if (ThisCell.isOverflow())
2453         {
2454             iaux=system("echo Unstable computation.>> carmen.prf");
2455             if (Cluster == 0) iaux=system("echo carmen: unstable computation. >> OUTPUT");
2456             cout << "carmen: instability detected at iteration no. "<<
IterationNo <<"\n";
2457             cout << "carmen: position ("<< x << ", "<<y<< ", "<<z<<")\n";
2458             cout << "carmen: abort execution.\n";
2459             exit(1);
2460         }
2461     }
2462 }

```

Here is the caller graph for this function:



#### 5.4.3.7 void Node::computeCorrection ( )

Computes velocity gradient (only for Navier-Stokes).

##### Returns

void

```

2189 {
2190     // --- Local variables ---
2191
2192     int n=0; // Counter on children
2193     real rho=0., psi=0.; // Variables density and psi
2194     int q=0, p=0; // Counter
2195     real Bx=0.; // Magnetic field
2196     real neweta=0.;
2197     real B1=0., B2=0., V=0., dx=0., DB=0., BR=0., BL=0., GB=0.;
2198     int ei=0, ej=0, ek=0;
2199     real udotB=0.;
2200
2201
2202     // --- Computation ---
2203
2204     if (requiresDivergenceComputation())
2205     {
2206         if (DivClean==1) // EGLM
2207         {
2208
2209             rho = ThisCell.density();
2210             psi = ThisCell.psi();
2211
2212             for (q=1; q <= 3; q++)
2213             {
2214                 Bx = ThisCell.magField(q);

```

```

2215         ThisCell.setAverage(q+1, ThisCell.average(q+1) -
TimeStep*Bx*Bdivergence/(ch*ch));
2216     }
2217
2218     ThisCell.setAverage(5, ThisCell.average(5) -
TimeStep*PsiGrad);
2219     ThisCell.setAverage(6, ThisCell.average(6)*exp(-(cr*
ch*TimeStep/SpaceStep));
2220
2221     }else if(DivClean==2)//GLM
2222     {
2223         psi = ThisCell.psi();
2224         ThisCell.setAverage(6, psi*exp(-(cr*ch*TimeStep/
SpaceStep));
2225     }else if(DivClean==3)
2226     {
2227         Bdivergence=0.;
2228         for (q=1; q <= Dimension; q++)
2229         {
2230             dx = ThisCell.size(q);
2231             dx *= 2.;
2232             B1=cell(Nl, Ni+ei, Nj+ej, Nk+ek)->magField(q);
2233             B2=cell(Nl, Ni-ei, Nj-ej, Nk-ek)->magField(q);
2234             Bdivergence += (B1-B2)/dx;
2235             udotB += ThisCell.velocity(q)*ThisCell.magField(q);
2236         }
2237
2238         ThisCell.setAverage(2, ThisCell.average(2) -
TimeStep*Bdivergence*ThisCell.magField(1));
2239         ThisCell.setAverage(3, ThisCell.average(3) -
TimeStep*Bdivergence*ThisCell.magField(2));
2240         ThisCell.setAverage(4, ThisCell.average(4) -
TimeStep*Bdivergence*ThisCell.magField(3));
2241         ThisCell.setAverage(5, ThisCell.average(5) -
TimeStep*Bdivergence*udotB);
2242         ThisCell.setAverage(7, ThisCell.average(7) -
TimeStep*Bdivergence*ThisCell.velocity(1));
2243         ThisCell.setAverage(8, ThisCell.average(8) -
TimeStep*Bdivergence*ThisCell.velocity(2));
2244         ThisCell.setAverage(9, ThisCell.average(9) -
TimeStep*Bdivergence*ThisCell.velocity(3));
2245
2246     }
2247 }
2248 }
2249
2250 // --- Recurse on children ---
2251
2252 if (isInternalNode())
2253     for (n = 0; n < ChildNb; n++){
2254         Child[n]->computeCorrection();
2255     }
2256 }

```

Here is the caller graph for this function:



#### 5.4.3.8 void Node::computeDivergence ( )

Computes the divergence vector with the space discretization scheme.



## Returns

void

## 2D resistive part of the model added to the Flux

```

1978 {
1979     // --- Local variables ---
1980
1981     int n=0; // Counter on children
1982     Vector FluxIn, FluxOut; // Ingoing and outgoing flux
1983     Vector InitAverage0; // Cell-average value of the initial condition
1984     Vector clean;
1985     real divCor=0.;
1986     // --- Computation ---
1987
1988     if (requiresDivergenceComputation())
1989     {
1990         // --- Compute source term -----
1991
1992         ThisCell.setDivergence(Source(ThisCell));
1993
1994         // --- Add flux in x-direction -----
1995
1996         // If the cell is a leaf with virtual children and its left cousin is a node, compute flux on upper
1997         level
1998         if (isLeafWithVirtualChildren() && node(Nl, Ni-1, Nj, Nk) != 0 && node(Nl, Ni-1, Nj, Nk)->
1999         isInternalNode() && FluxCorrection)
2000         {
2001             FluxIn = Flux( *childCell(-2,0,0), *childCell(-1,0,0), *childCell(0,0,0) , *childCell(1,0,
2002             0), 1 );
2003
2004             if (Dimension > 1)
2005                 FluxIn += Flux( *childCell(-2,1,0), *childCell(-1,1,0), *childCell(0,1,0), *childCell(1
2006             ,1,0), 1 );
2007
2008             if (Dimension > 2)
2009             {
2010                 FluxIn += Flux( *childCell(-2,0,1), *childCell(-1,0,1), *childCell(0,0,1), *childCell(1
2011             ,0,1), 1 );
2012                 FluxIn += Flux( *childCell(-2,1,1), *childCell(-1,1,1), *childCell(0,1,1), *childCell(1
2013             ,1,1), 1 );
2014             }
2015
2016             // Average flux
2017             FluxIn *= 1./(1<<(Dimension-1));
2018
2019             }
2020         else
2021             FluxIn = Flux( *cousinCell(-2,0,0), *cousinCell(-1,0,0), ThisCell, *cousinCell(1,0,0), 1 )
2022         ;
2023
2024         divCor = -auxvar;
2025         // If the cell is a leaf with virtual children and its right cousin is a node, compute flux on
2026         upper level
2027         if (isLeafWithVirtualChildren() && node(Nl, Ni+1, Nj, Nk) != 0 && node(Nl, Ni+1, Nj, Nk)->
2028         isInternalNode() && FluxCorrection)
2029         {
2030             FluxOut = Flux( *childCell(0,0,0), *childCell(1,0,0), *childCell(2,0,0), *childCell(3,0,0)
2031             , 1 );
2032
2033             if (Dimension > 1)
2034                 FluxOut += Flux( *childCell(0,1,0), *childCell(1,1,0), *childCell(2,1,0), *childCell(3,
2035             1,0), 1 );
2036
2037             if (Dimension > 2)
2038             {
2039                 FluxOut += Flux( *childCell(0,0,1), *childCell(1,0,1), *childCell(2,0,1), *childCell(3,
2040             0,1), 1 );
2041                 FluxOut += Flux( *childCell(0,1,1), *childCell(1,1,1), *childCell(2,1,1), *childCell(3,
2042             1,1), 1 );
2043             }
2044
2045             // Average flux
2046             FluxOut *= 1./(1<<(Dimension-1));
2047
2048             }
2049         else
2050             FluxOut = Flux( *cousinCell(-1,0,0), ThisCell, *cousinCell(1,0,0), *cousinCell(2,0,0), 1 );
2051
2052         divCor += auxvar;
2053         if(Resistivity){
2054             FluxIn = FluxIn - ResistiveTerms(ThisCell, *cousinCell(-1,0,0), *cousinCell(0,-

```

```

1,0), *cousinCell(0,0,-1), 1);
2045     FluxOut = FluxOut - ResistiveTerms(*cousinCell(1,0,0), ThisCell , *cousinCell(1,-
1,0), *cousinCell(1,0,-1), 1);
2046 }
2047
2048     // Add divergence in x-direction
2049     ThisCell.setDivergence( ThisCell.divergence() + (FluxIn - FluxOut)/(ThisCell
.size(1)));
2050
2051     // Variables \grad(psi) and \div(B) to evaluate GLM and EGLM divergence cleaning
2052     PsiGrad = ThisCell.average(7)*(FluxOut.value(7) - FluxIn
value(7) - divCor)/(ThisCell.size(1));
2053     Bdivergence = ((FluxOut.value(6) - FluxIn.value(6))/(ThisCell
.size(1)));
2054
2055     // --- Add flux in y-direction -----
2056
2057     if (Dimension > 1)
2058     {
2059         // If the cell is a leaf with virtual children and its front cousin is a node, compute flux on
upper level
2060
2061         if (isLeafWithVirtualChildren() && node(Nl, Ni, Nj-1, Nk) != 0 && node(Nl, Ni, Nj-1, Nk)->
isInternalNode() && FluxCorrection)
2062         {
2063             FluxIn = Flux( *childCell(0,-2,0), *childCell(0,-1,0), *childCell(0,0,0), *childCell(0
,1,0), 2 );
2064             FluxIn += Flux( *childCell(1,-2,0), *childCell(1,-1,0), *childCell(1,0,0) , *childCell(
1,1,0), 2 );
2065
2066             if (Dimension > 2)
2067             {
2068                 FluxIn += Flux( *childCell(0,-2,1), *childCell(0,-1,1), *childCell(0,0,1), *
childCell(0,1,1), 2 );
2069                 FluxIn += Flux( *childCell(1,-2,1), *childCell(1,-1,1), *childCell(1,0,1), *
childCell(1,1,1), 2 );
2070             }
2071
2072             // Average flux
2073             FluxIn *= 1./(1<<(Dimension-1));
2074
2075         }
2076         else
2077             FluxIn = Flux( *cousinCell(0,-2,0), *cousinCell(0,-1,0), ThisCell, *cousinCell(0,1,0),
2 );
2078
2079         divCor = -auxvar;
2080         // If the cell is a leaf with virtual children and its back cousin is a node, compute flux on
upper level
2081
2082         if (isLeafWithVirtualChildren() && node(Nl, Ni, Nj+1, Nk) != 0 && node(Nl, Ni, Nj+1, Nk)->
isInternalNode() && FluxCorrection)
2083         {
2084             FluxOut = Flux( *childCell(0,0,0), *childCell(0,1,0), *childCell(0,2,0), *childCell(0,
3,0), 2 );
2085             FluxOut += Flux( *childCell(1,0,0), *childCell(1,1,0), *childCell(1,2,0), *childCell(1,
3,0), 2 );
2086
2087             if (Dimension > 2)
2088             {
2089                 FluxOut += Flux( *childCell(0,0,1), *childCell(0,1,1), *childCell(0,2,1), *
childCell(0,3,1), 2 );
2090                 FluxOut += Flux( *childCell(1,0,1), *childCell(1,1,1), *childCell(1,2,1), *
childCell(1,3,1), 2 );
2091             }
2092
2093             // Average flux
2094             FluxOut *= 1./(1<<(Dimension-1));
2095
2096         }
2097         else
2098             FluxOut = Flux( *cousinCell(0,-1,0), ThisCell, *cousinCell(0,1,0), *cousinCell(0,2,0),
2 );
2099
2100         divCor += auxvar;
2101
2102         if(Resistivity){
2103             FluxIn = FluxIn - ResistiveTerms(ThisCell, *cousinCell(-1,0,0), *cousinCell
(0,-1,0), *cousinCell(0,0,-1), 2);
2104             FluxOut = FluxOut - ResistiveTerms(*cousinCell(0,1,0), *cousinCell(-1,1,0),
ThisCell , *cousinCell(0,1,-1), 2);
2105         }
2106
2107         // Add divergence in y-direction
2108         ThisCell.setDivergence( ThisCell.divergence() + (FluxIn - FluxOut)/(
ThisCell.size(2)));
2109

```

```

2110         // Variables \grad(psi) and \div(B) to evaluate GLM and EGLM divergence cleaning
2111         PsiGrad += ThisCell.average(8)*(FluxOut.value(8) - FluxIn.
value(8) - divCor)/(ThisCell.size(2));
2112         Bdivergence += ((FluxOut.value(6) - FluxIn.value(6))/(ThisCell.
size(2)));
2113
2114     }
2115 }
2116
2117     // --- Add flux in z-direction -----
2118
2119     if (Dimension > 2)
2120     {
2121     {
2122         // If the cell is a leaf with virtual children and its lower cousin is a node, compute flux on
upper level
2123
2124         if (isLeafWithVirtualChildren() && node(Nl, Ni, Nj, Nk-1) != 0 && node(Nl, Ni, Nj, Nk-1)->
isInternalNode() && FluxCorrection)
2125         {
2126             FluxIn = Flux( *childCell(0,0,-2), *childCell(0,0,-1), *childCell(0,0,0), *childCell(0
,0,1), 3 );
2127             FluxIn += Flux( *childCell(1,0,-2), *childCell(1,0,-1), *childCell(1,0,0), *childCell(1
,0,1), 3 );
2128             FluxIn += Flux( *childCell(0,1,-2), *childCell(0,1,-1), *childCell(0,1,0), *childCell(0
,1,1), 3 );
2129             FluxIn += Flux( *childCell(1,1,-2), *childCell(1,1,-1), *childCell(1,1,0), *childCell(1
,1,1), 3 );
2130
2131             // Average flux
2132             FluxIn *= 0.25;
2133         }
2134     }
2135     else
2136     {
2137         FluxIn = Flux( *cousinCell(0,0,-2), *cousinCell(0,0,-1), ThisCell, *cousinCell(0,0,1),
3 );
2138
2139         divCor = -auxvar;
2140
2141         // If the cell is a leaf with virtual children and its upper cousin is a node, compute flux on
upper level
2142
2143         if (isLeafWithVirtualChildren() && node(Nl, Ni, Nj, Nk+1) != 0 && node(Nl, Ni, Nj, Nk+1)->
isInternalNode() && FluxCorrection)
2144         {
2145             FluxOut = Flux( *childCell(0,0,0), *childCell(0,0,1), *childCell(0,0,2), *childCell(0
,0,3), 3 );
2146             FluxOut += Flux( *childCell(1,0,0), *childCell(1,0,1), *childCell(1,0,2), *childCell(1
,0,3), 3 );
2147             FluxOut += Flux( *childCell(0,1,0), *childCell(0,1,1), *childCell(0,1,2), *childCell(0
,1,3), 3 );
2148             FluxOut += Flux( *childCell(1,1,0), *childCell(1,1,1), *childCell(1,1,2), *childCell(1
,1,3), 3 );
2149
2150             // Average flux
2151             FluxOut *= 0.25;
2152         }
2153     }
2154     else
2155     {
2156         FluxOut = Flux( *cousinCell(0,0,-1), ThisCell, *cousinCell(0,0,1), *cousinCell(0,0,2),
3 );
2157
2158         divCor += auxvar;
2159
2160         if(Resistivity){
2161             FluxIn = FluxIn - ResistiveTerms(ThisCell, *cousinCell(-1,0,0), *cousinCell
(0,-1,0), *cousinCell(0,0,-1), 3);
2162             FluxOut = FluxOut - ResistiveTerms(*cousinCell(0,0,1), *cousinCell(-1,0,1), *
cousinCell(0,-1,1), ThisCell, 3);
2163         }
2164
2165         // Add divergence in z-direction
2166         ThisCell.setDivergence( ThisCell.divergence() + (FluxIn - FluxOut)/(
ThisCell.size(3)));
2167
2168         // Variables \grad(psi) and \div(B) to evaluate GLM and EGLM divergence cleaning
2169         PsiGrad += ThisCell.average(9)*(FluxOut.value(9) - FluxIn.
value(9) - divCor)/(ThisCell.size(3));
2170         Bdivergence += ((FluxOut.value(6) - FluxIn.value(6))/(ThisCell.
size(3)));
2171     }
2172 }
2173 // --- Recurse on children ---
2174
2175 if (isInternalNode())

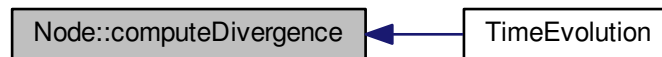
```

```

2176         for (n = 0; n < ChildNb; n++)
2177             Child[n]->computeDivergence();
2178     }

```

Here is the caller graph for this function:



#### 5.4.3.9 void Node::computeGradient ( )

Computes velocity gradient (only for Navier-Stokes).

##### Returns

void

```

2267 {
2268     // --- Local variables ---
2269
2270     int n=0; // Counter on children
2271     real rho1=0., rho2=0.; // Densities
2272     real rhoE1=0., rhoE2=0.; // Energies
2273     real V1=0., V2=0.; // Velocity
2274     real dx=0.; // Cell size
2275     real dxV=0.; // Correction of dx for the computation of GradV close to solid walls
2276     int p=0, q=0; // Counters
2277     int ei=0, ej=0, ek=0; // 1 if this direction is chosen, 0 elsewhere
2278
2279     // --- Computation ---
2280
2281     // if (requiresDivergenceComputation() || (CVS && isParentOfLeaf()))
2282
2283     if (requiresDivergenceComputation())
2284     {
2285         if (EquationType != 6)
2286         {
2287             cout << "Node.cpp: In method `void Node::computeGradient()':\n";
2288             cout << "Node.cpp: EquationType not equal to 6 \n";
2289             cout << "carmen: *** [Node.o] Execution error\n";
2290             cout << "carmen: abort execution.\n";
2291             exit(1);
2292         }
2293
2294         for (p=1;p <= Dimension; p++)
2295         {
2296             ei = (p==1)? 1:0;
2297             ej = (p==2)? 1:0;
2298             ek = (p==3)? 1:0;
2299
2300             dx = ThisCell.size(p);
2301             dx *= 2.;
2302
2303             // dxV = correction on dx for the computation of GradV close to solid walls
2304
2305             if (BoundaryRegion(cell(Nl, Ni+ei,Nj+ej,Nk+ek)->center()) > 3 ||
2306                 BoundaryRegion(cell(Nl, Ni-ei,Nj-ej,Nk-ek)->center()) > 3 )
2307                 dxV = 0.75*dx;
2308             else
2309                 dxV = dx;
2310
2311             rho1 = cell(Nl, Ni+ei,Nj+ej,Nk+ek)->density();
2312             rho2 = cell(Nl, Ni-ei,Nj-ej,Nk-ek)->density();
2313
2314             ThisCell.setGradient(p, 1, (rho1-rho2)/dx);
2315
2316             for (q=1; q <= Dimension; q++)

```

```

2317     {
2318         V1=cell(Nl, Ni+ei, Nj+ej, Nk+ek)->velocity(q);
2319         V2=cell(Nl, Ni-ei, Nj-ej, Nk-ek)->velocity(q);
2320         ThisCell.setGradient(p, q+1, (V1-V2)/dxV);
2321     }
2322
2323     rhoE1 = cell(Nl, Ni+ei, Nj+ej, Nk+ek)->energy();
2324     rhoE2 = cell(Nl, Ni-ei, Nj-ej, Nk-ek)->energy();
2325
2326     ThisCell.setGradient(p, Dimension+2, (rhoE1-rhoE2)/dx);
2327 }
2328 }
2329
2330 // --- Recurse on children ---
2331
2332 if (isInternalNode())
2333     for (n = 0; n < ChildNb; n++)
2334         Child[n]->computeGradient();
2335 }

```

#### 5.4.3.10 void Node::computeIntegral ( )

Computes integral values like e.g. flame velocity, global error, etc.

#### Returns

void

```

2471 {
2472     // --- Local variables ---
2473
2474     int QuantityNo; // Quantity number (0 to QuantityNb)
2475     int n; // Counter on children
2476     int AxisNo; // Counter on dimension
2477     real dx, dy=0., dz=0.; // Cell size
2478     Vector Center(Dimension); // local center of the flame ball
2479     real VelocityMax; // local maximum of the velocity
2480     real MaxSpeed;
2481     real MemoryCompression = 0.; // Memory compression
2482
2483     Vector GradDensity(Dimension); // gradient of density
2484     Vector GradPressure(Dimension); // gradient of pressure
2485     real divB=0;
2486     real modB=0.;
2487
2488     real B1=0., B2=0.; // Left and right magnetic field cells
2489
2490     int ei=0, ej=0, ek=0; // 1 if this direction is chosen, 0 elsewhere
2491
2492
2493     // --- Init ---
2494
2495     if (Nl == 0)
2496     {
2497         // Only if ExpectedCompression not equal to zero => variable tolerance
2498
2499         if (ExpectedCompression != 0.)
2500         {
2501             MemoryCompression = (1.*CellNb)/(1<<(ScaleNb*Dimension));
2502             Tolerance = Tolerance*(1.- (ExpectedCompression-
MemoryCompression));
2503             if (Tolerance > 1E+10)
2504             {
2505                 printf("carmen: ExpectedCompression unreachable\n");
2506                 printf("carmen: maximal compression is %5.2f %%", MemoryCompression*100.);
2507                 printf("carmen: abort execution.\n");
2508                 exit(1);
2509             }
2510         }
2511
2512         // Init integral values
2513         DIVB = 0.;
2514         //DIVBMax = 0.;
2515         FlameVelocity = 0.;
2516         GlobalMomentum = 0.;
2517         GlobalEnergy = 0.;
2518         GlobalEnstrophy = 0.;
2519         ExactMomentum = 0.;
2520         ExactEnergy = 0.;
2521
2522         GlobalReactionRate = 0.;

```

```

2523     AverageRadius      = 0.;
2524     ReactionRateMax    = 0.;
2525
2526     for (AxisNo=1; AxisNo <= Dimension; AxisNo++)
2527         Center.setValue(AxisNo,XCenter[AxisNo]);
2528
2529     ErrorMax           = 0.;
2530     ErrorMid           = 0.;
2531     ErrorL2            = 0.;
2532     ErrorNb           = 0;
2533
2534     RKFEError = 0.;
2535
2536     //Eigenvalue = 0.;
2537     QuantityMax.setZero();
2538     QuantityAverage.setZero();
2539
2540     IntVorticity=0.;
2541     IntDensity=0.;
2542     IntMomentum.setZero();
2543     BaroclinicEffect=0.;
2544
2545 }
2546
2547 // --- Recursion ---
2548
2549 if (isInternalNode())
2550 {
2551     for (n = 0; n < ChildNb; n++)
2552         Child[n]->computeIntegral();
2553 }
2554 else if (isLeaf())
2555 {
2556     // Whatever the equation, if ConstantTimeStep is false, compute RKFEError
2557
2558     if (!ConstantTimeStep && StepNb == 3)
2559     {
2560         for (QuantityNo = 1; QuantityNo <= QuantityNb; QuantityNo++)
2561         {
2562             if (Abs(ThisCell.average(QuantityNo)) >
2563                 RKFAccuracyFactor)
2564                 RKFEError = Max(RKFEError, Abs(1.-ThisCell.
2565                     lowAverage(QuantityNo)/ThisCell.average(QuantityNo)));
2566         }
2567
2568         dx = ThisCell.size(1);
2569         dy = (Dimension > 1) ? ThisCell.size(2) : 1.;
2570         dz = (Dimension > 2) ? ThisCell.size(3) : 1.;
2571
2572         // --- Compute the global momentum, global energy and global enstrophy ---
2573
2574         GlobalMomentum += ThisCell.average(2)*dx*dy*dz;
2575         GlobalEnergy    += .5*(ThisCell.magField()*ThisCell.
2576             magField() + ThisCell.density()*(ThisCell.velocity()*ThisCell.
2577             velocity())) + ThisCell.pressure()/(Gamma-1.0);
2578         GlobalEnergy    *= dx*dy*dz;
2579         Helicity        += (ThisCell.magField(2)*ThisCell.
2580             velocity(3) - ThisCell.magField(3)*ThisCell.velocity(2))*ThisCell.
2581             magField(1) +
2582             (ThisCell.magField(3)*ThisCell.velocity(1) - ThisCell.
2583             magField(1)*ThisCell.velocity(3))*ThisCell.magField(2) +
2584             (ThisCell.magField(1)*ThisCell.velocity(2) - ThisCell.
2585             magField(2)*ThisCell.velocity(1))*ThisCell.magField(3);
2586         Helicity        *= 2*dx*dy*dz;
2587
2588         // --- Compute maximum of the conservative quantities ---
2589
2590         for (QuantityNo=1; QuantityNo <=QuantityNb; QuantityNo++)
2591         {
2592             if ( QuantityMax.value(QuantityNo) < fabs(ThisCell.
2593                 average(QuantityNo)) )
2594                 QuantityMax.setValue(QuantityNo, fabs(ThisCell.
2595                     average(QuantityNo)) );
2596         }
2597
2598         // --- Compute the maximal eigenvalue ---
2599
2600         VelocityMax = 0.;
2601         MaxSpeed    = 0.;
2602         for (AxisNo=1; AxisNo <= Dimension; AxisNo++){
2603             VelocityMax = Max( VelocityMax, fabs(ThisCell.velocity(AxisNo)));
2604             MaxSpeed    = Max( MaxSpeed    , fabs(ThisCell.fastSpeed(AxisNo)));
2605         }
2606
2607         VelocityMax += MaxSpeed;
2608
2609     }

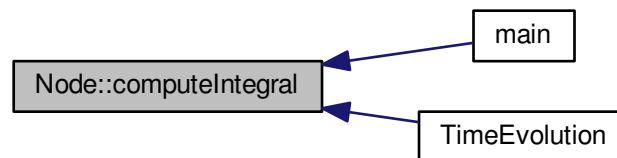
```

```

2600     EigenvalueMax = Max (EigenvalueMax, VelocityMax);
2601
2602
2603
2604     for (AxisNo = 1; AxisNo <= Dimension; AxisNo++)
2605     {
2606         ei = (AxisNo == 1)? 1:0;
2607         ej = (AxisNo == 2)? 1:0;
2608         ek = (AxisNo == 3)? 1:0;
2609
2610         dx = ThisCell.size(AxisNo);
2611         //dx *= 2.;
2612
2613         B1 = cell(Nl, Ni+ei, Nj+ej, Nk+ek)->magField(AxisNo);
2614         B2 = cell(Nl, Ni-ei, Nj-ej, Nk-ek)->magField(AxisNo);
2615         modB += (B1 + B2)/dx;
2616         divB += (B1-B2)/dx;
2617     }
2618     modB += 1.120e-13;
2619     DIVBMax = Max(DIVBMax, Abs(0.5*divB));
2620     DIVB = DIVBMax/modB;
2621 }
2622 }

```

Here is the caller graph for this function:



#### 5.4.3.11 void Node::fillVirtualChildren ( )

Fills the cell-average values of every virtual leaf with values predicted from its parent and uncles. This procedure is required after a time evolution to refresh the virtual leaves of the tree.

##### Returns

void

```

625 {
626     // --- Local variables ---
627
628     int n=0; // Counter on children
629
630     // --- Recursion ---
631
632     switch(Flag)
633     {
634         // If node is not a leaf or leaf with virtual children, recurse on children
635         case 0:
636             case 2:
637                 for (n = 0; n < ChildNb; n++)
638                     Child[n]->fillVirtualChildren();
639                 break;
640
641         // If node is a simple leaf, stop procedure
642         case 1:
643             return;
644             break;
645
646         // If node is a virtual leaf, compute value with prediction
647         case 3:

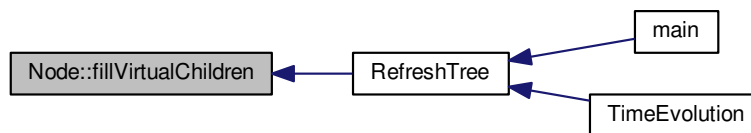
```

```

648         ThisCell.setAverage(predict ());
649         if (EquationType==6) ThisCell.setGradient (parentCell()->gradient ());
650         break;
651     };
652 };
653 }

```

Here is the caller graph for this function:



#### 5.4.3.12 void Node::initValue ( )

Computes the initial value.

##### Returns

void

```

125 {
126     int i,j,k; // Counters on directions
127
128     ThisCell.setAverageZero();
129
130     if (UseBoundaryRegions && isInsideBoundary())
131     {
132         ThisCell.setAverage(InitAverage( ThisCell.center(1),
133                                         (Dimension >1)? ThisCell.center(2):0.,
134                                         (Dimension > 2)? ThisCell.center(3):0. ) );
135     }
136     else
137     {
138         switch (Dimension)
139         {
140             case 1:
141                 for (i=0;i<=1;i++)
142                     ThisCell.setAverage( ThisCell.average()+.5*
InitAverage(
143                         ThisCell.center(1)+(i-0.5)*ThisCell.size(1)
144                     ) );
145                 break;
146             case 2:
147                 if(IcNb){
148                     for (i=0;i<=1;i++){
149                         for (j=0;j<=1;j++){
150                             ThisCell.setAverage( ThisCell.average()+.25*
InitAverage(
151                                 ThisCell.center(1)+(i-0.5)*ThisCell.size(1),
152                                 ThisCell.center(2)+(j-0.5)*ThisCell.size(2)));
153                             ThisCell.setRes(InitResistivity(ThisCell.
center(1), ThisCell.center(2)));
154                         }
155                     }
156                 }
157             }else{
158                 ThisCell.setAverage(InitAverage(ThisCell.
center(1), ThisCell.center(2)));
159                 ThisCell.setRes(InitResistivity(ThisCell.
center(1), ThisCell.center(2)));
160             }
161             break;
162             case 3:

```



```

165         if(IcNb){
166             for (i=0;i<=1;i++)
167                 for (j=0;j<=1;j++)
168                     for (k=0;k<=1;k++){
169                         ThisCell.setAverage( ThisCell.average()+.125*
170 InitAverage(
171                 ThisCell.center(1)+(i-0.5)*ThisCell.
172 size(1),
173                 ThisCell.center(2)+(j-0.5)*ThisCell.
174 size(2),
175                 ThisCell.center(3)+(k-0.5)*ThisCell.
176 size(3) );
177                 ThisCell.setRes(InitResistivity(ThisCell.
178 center(1), ThisCell.center(2),ThisCell.center(3)));
179             }
180         }else{
181             ThisCell.setAverage(InitAverage(ThisCell.
182 center(1), ThisCell.center(2),ThisCell.center(3)));
183             ThisCell.setRes(InitResistivity(ThisCell.
184 center(1), ThisCell.center(2),ThisCell.center(3)));
185             break;
186         }
187     };
188 }

```

Here is the caller graph for this function:



#### 5.4.3.13 int Node::leaves( ) const [inline]

Returns the number of leaves in the tree.

##### Returns

int

```

696 {
697     return LeafNb;
698 }

```

#### 5.4.3.14 Cell \* Node::project( )

Computes the cell-average values of all nodes that are not leaves by projection from the cell-averages values of the leaves. This procedure is required after a time evolution to refresh the internal nodes of the tree.

##### Returns

Cell\*

```

662 {
663     // --- Local variables ---
664
665     int n=0;    // Counter on children
666
667     // --- If cell is not a leaf, compute projection ie mean value of children ---
668

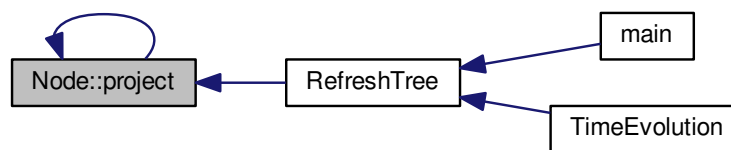
```

```

669     if (isInternalNode())
670     {
671         // Set value to zero
672         ThisCell.setAverageZero();
673
674         // Compute the mean value
675         for (n = 0; n < ChildNb; n++)
676             ThisCell.setAverage( ThisCell.average() + Child[n]->
project()->average() );
677
678         ThisCell.setAverage( ThisCell.average() / ChildNb );
679     }
680 }
681
682 return &ThisCell;
683 }

```

Here is the caller graph for this function:



#### 5.4.3.15 void Node::restore ( )

Restores the tree structure and the cell-averages from the file *carmen.bak*. This file was created by the method [Backup\(\)](#).

#### Returns

void

```

2762 {
2763
2764     int n=0;           // Counter on children
2765     int QuantityNo=0; // Counter on quantities
2766     char buf[256];    // Text buffer
2767     char* caux;
2768     // --- Init ---
2769
2770     if (Nl==0)
2771     {
2772         GlobalFile = fopen("carmen.bak","r");
2773         // fgets(buf, 256, GlobalFile);
2774     }
2775
2776     caux=fgets(buf,256,GlobalFile);
2777
2778     // If the first data is not a 'N', it means that the data has been created using FineMesh
2779
2780     if (buf[0] != 'N' && Nl==0)
2781     {
2782         fclose(GlobalFile);
2783         restoreFineMesh();
2784         return;
2785     }
2786
2787     // If end of file is reached, close file and return
2788     if (feof(GlobalFile))
2789     {
2790         fclose(GlobalFile);
2791         return;
2792     }
2793 }

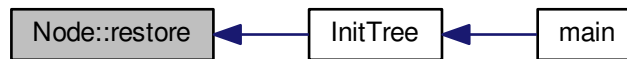
```

```

2794 // --- Recurse : if node is not a leaf, split it and restore children ---
2795
2796 if (buf[0]!='N')
2797 {
2798     split();
2799     for (n = 0; n < ChildNb; n++)
2800         Child[n]->restore();
2801 }
2802 else
2803 {
2804     ThisCell.setAverage(1,atof(buf));
2805     for (QuantityNo=2; QuantityNo <= QuantityNb; QuantityNo++)
2806     {
2807         caux=fgets(buf,256,GlobalFile);
2808         ThisCell.setAverage(QuantityNo, atof(buf));
2809     }
2810     return;
2811 }
2812 }

```

Here is the caller graph for this function:



#### 5.4.3.16 void Node::restoreFineMesh ( )

Restores the tree structure and the cell-averages from the file *carmen.bak* in [FineMesh](#) format.

##### Returns

void

```

2822 {
2823 // --- Local variables ---
2824
2825 int i=0,j=0,k=0; // Counters in the three directions
2826 int n=0,iaux; // Global counter
2827 int QuantityNo=0; // Counter on quantities
2828 FILE* input; // Input file
2829 real buf;
2830
2831 // --- Split the whole tree structure ---
2832
2833 splitAll();
2834
2835 // --- Get data from carmen.bak in the FineMesh format
2836
2837 // -- Open file --
2838
2839 input = fopen("carmen.bak","r");
2840
2841 // -- When there is no back-up file, return --
2842
2843 if (!input) return;
2844
2845 // -- Loop on fine-grid cells --
2846
2847 for (n = 0; n < 1<<(ScaleNb*Dimension); n++)
2848 {
2849 // -- Compute i, j, k --
2850
2851 i = n%(1<<ScaleNb);
2852 if (Dimension > 1) j = (n%(1<<(2*ScaleNb)))/(1<<ScaleNb);
2853 if (Dimension > 2) k = n/(1<<(2*ScaleNb));
2854
2855

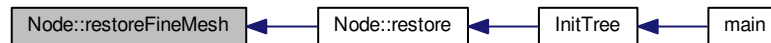
```

```

2856     for (QuantityNo=1; QuantityNo <= QuantityNb; QuantityNo++)
2857     {
2858        iaux=fscanf(input, BACKUP_FILE_FORMAT, &buf);
2859         cell(ScaleNb, i, j, k) -> setAverage(QuantityNo, buf);
2860     }
2861 }
2862
2863 fclose(input);
2864
2865 }

```

Here is the caller graph for this function:



#### 5.4.3.17 void Node::RungeKutta ( )

Computes one Runge-Kutta step.

##### Returns

void

```

2346 {
2347     // --- Local variables ---
2348
2349     int n=0;           // Counter on children
2350     real c1=0., c2=0., c3=0.; // Runge-Kutta coefficients
2351     real LocalTimeStep=TimeStep; // Local time step
2352
2353     Vector Q(QuantityNb), Qs(QuantityNb), D(QuantityNb); //
    Cell-average, temporary cell-average and divergence
2354
2355     // --- If node is in the tree, recurse on children -----
2356
2357     if (isInternalNode())
2358         for (n = 0; n < ChildNb; n++)
2359             Child[n]->RungeKutta();
2360
2361     // --- If the leaf is in the boundary, do not perform time evolution -----
2362
2363     if (UseBoundaryRegions)
2364     {
2365         if (isLeaf() && isInsideBoundary())
2366             return;
2367     }
2368
2369     // --- For leaves in the fluid region -----
2370
2371     if (requiresDivergenceComputation())
2372     {
2373         // --- Compute local time step in function of the scale ---
2374
2375         if (TimeAdaptivity)
2376         {
2377             // Compute local time step
2378             LocalTimeStep = TimeStep*(1<<(TimeAdaptivityFactor*(
    ScaleNb-N1)));
2379
2380             // At the end of the time cycle, Q <- Qlow (solution at end of cycle)
2381             if (isEndTimeCycle() && StepNo == 1 && N1 < ScaleNb)
2382                 ThisCell.setAverage( ThisCell.lowAverage());
2383         }
2384         // --- Define Runge-Kutta coefficients ---
2385
2386         switch(StepNo)
2387         {
2388             case 1:
2389                 c1 = 1.; c2 = 0.; c3 = 1.;

```

```

2390         break;
2391     case 2:
2392         if (StepNb == 2) {c1 = .5; c2 = .5; c3 = .5; }
2393         if (StepNb == 3) {c1 = .75; c2 = .25; c3 = .25; }
2394         break;
2395     case 3:
2396         c1 = 1./3.; c2 = 2.*c1; c3 = c2;
2397         break;
2398     };
2399
2400     // --- Runge-Kutta step ---
2401
2402     Q = ThisCell.average();
2403     Qs = ThisCell.tempAverage();
2404     D = ThisCell.divergence();
2405
2406     // Perform RK step only in fluid region
2407     ThisCell.setAverage( c1*Qs + c2*Q + (c3 * LocalTimeStep)*D );
2408
2409     // For the Runge-Kutta-Fehlberg 2(3) method, store second-stage with the RK2 coefficients
2410
2411     if (!ConstantTimeStep && (StepNo == 2) && (StepNb == 3)) // MODIFIED
10.12.05
2412         ThisCell.setLowAverage(0.5*(Qs + Q + LocalTimeStep*D));
2413
2414     // Time adaptivity :
2415     if (TimeAdaptivity)
2416     {
2417         if (isBeginTimeCycle() && StepNo == 1 && N1 < ScaleNb)
2418             storeTimeEvolution();
2419     }
2420 }
2421
2422 }

```

Here is the caller graph for this function:



#### 5.4.3.18 void Node::smooth ( )

Deletes the details in the highest level.

##### Returns

void

```

2875 {
2876     int n=0; // Child number
2877
2878     // --- Recurse on children ---
2879
2880     if (isInternalNode())
2881     {
2882         for (n = 0; n < ChildNb; n++)
2883             Child[n]->smooth();
2884     }
2885     else
2886         //if (N1 == ScaleNb)
2887         {
2888             if (parentCell()->average() == ThisCell.average())
2889                 return;
2890             else
2891                 ThisCell.setAverage(SmoothCoeff*predict() + (1.-
SmoothCoeff)*ThisCell.average());
2892         }

```

```

2893
2894     return;
2895 }

```

Here is the caller graph for this function:



#### 5.4.3.19 void Node::store ( )

Stores cell-average values into temporary cell-average values.

##### Returns

void

```

1914 {
1915     // --- Local variables ---
1916
1917     int n=0;    // Counter on children
1918
1919     // --- Store cell-average value Q into Qs ---
1920
1921     if ((EquationType==6) && SchemeNb > 5 ) || requiresTimeEvolution() ||
IterationNo == 1)
1922     {
1923         if (UseBoundaryRegions)
1924         {
1925             if (IterationNo == 1)
1926                 ThisCell.setOldAverage(ThisCell.average());
1927             else
1928                 ThisCell.setOldAverage(ThisCell.tempAverage());
1929         }
1930
1931         ThisCell.setTempAverage(ThisCell.average());
1932     }
1933
1934     // --- Recursion in nodes that have children (real or virtual) ---
1935
1936     if (hasChildren())
1937     {
1938         for (n = 0; n < ChildNb; n++)
1939             Child[n]->store();
1940     }
1941 }

```

Here is the caller graph for this function:



## 5.4.3.20 void Node::storeGrad ( )

Stores gradient values into temporary gradient values.

## Returns

void

```

1951 {
1952     // --- Local variables ---
1953
1954     int n=0;    // Counter on children
1955
1956     // --- Store gradient-average value Grad into Grads ---
1957
1958     if (((EquationType==6) && SchemeNb > 5) || requiresTimeEvolution() ||
IterationNo == 1)
1959         ThisCell.setTempGradient(ThisCell.gradient());
1960
1961     // --- Recursion in nodes that have children (real or virtual) ---
1962
1963     if (hasChildren())
1964     {
1965         for (n = 0; n < ChildNb; n++)
1966             Child[n]->storeGrad();
1967     }
1968 }
```

## 5.4.3.21 void Node::writeAverage ( const char \* FileName )

Writes cell-average values in multiresolution representation and the corresponding mesh into file *FileName*.

## Parameters

|                 |                   |
|-----------------|-------------------|
| <i>FileName</i> | Name of the file. |
|-----------------|-------------------|

## Returns

void

```

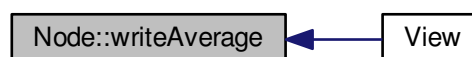
1090 {
1091     // --- Local variables ---
1092
1093     int n ;    // Counter on children
1094     FILE *output; // Pointer to output file
1095     Vector Qbuf(QuantityNb); // Buffer of the vector of conservative quantities
1096     real a=1., b=1.; // Weights for the synchronisation of conservative quantities
1097     int Nf=1;
1098
1099     // --- Open file ---
1100
1101     if ((output = fopen(FileName,"a")) != NULL)
1102     {
1103         // If node is not a leaf, recurse to children
1104         if (isInternalNode())
1105         {
1106             fclose(output);
1107             for (n = 0; n < ChildNb; n++)
1108                 Child[n]->writeAverage(FileName);
1109         }
1110         else
1111         {
1112             // In case of local time stepping, store value and synchronize it
1113             if ( TimeAdaptivity && Dimension == 1 && (N1 < (
ScaleNb-1)) )
1114             {
1115                 Qbuf = ThisCell.average();
1116
1117                 Nf = 1<<TimeAdaptivityFactor*(ScaleNb-N1);
1118                 a = 1.*(IterationNo%Nf - Nf/2);
1119                 b = 1.*(Nf - IterationNo%Nf);
1120
1121                 if ( (a+b) != 0. )
1122                 {
1123                     // Qbuf2 = ( b/(a+b) ) * ThisCell.average() + ( a/(a+b) ) * ThisCell.lowAverage();
1124                 }
```

```

1125             //             ThisCell.setAverage(Qbuf2);
1126
1127             //ThisCell.setAverage(( b/(a+b) ) * ThisCell.average()+ ( a/(a+b) ) *
ThisCell.lowAverage())
1128         }
1129     }
1130
1131     // write cell-average values and details from parent
1132
1133     fprintf(output, FORMAT, ThisCell.center(1));
1134     if (Dimension > 1) fprintf(output, FORMAT, ThisCell.
center(2));
1135     if (Dimension > 2) fprintf(output, FORMAT, ThisCell.
center(3));
1136
1137     if (Dimension > 1)
1138         fprintf(output, "%i", N1);
1139
1140     else{
1141         fprintf(output, FORMAT, ThisCell.density());
1142         fprintf(output, FORMAT, ThisCell.pressure());
1143         fprintf(output, FORMAT, ThisCell.psi());
1144         fprintf(output, FORMAT, ThisCell.energy());
1145         fprintf(output, FORMAT, ThisCell.velocity(1));
1146         fprintf(output, FORMAT, ThisCell.magField(1));
1147     }
1148
1149     fprintf(output, "\n");
1150
1151     // In case of local time stepping, return to the previous value
1152     if (TimeAdaptivity && Dimension == 1 && N1 <
ScaleNb-1)
1153         ThisCell.setAverage(Qbuf);
1154
1155     fclose(output);
1156 }
1157 }
1158 else
1159 {
1160     cout << "Node.cpp: In method 'void Node::writeAverage()':\n";
1161     cout << "Node.cpp: cannot open file " << FileName << '\n';
1162     cout << "carmen: *** [Node.o] Execution error\n";
1163     cout << "carmen: abort execution.\n";
1164     exit(1);
1165 }
1166 }

```

Here is the caller graph for this function:



#### 5.4.3.22 void Node::writeFineGrid ( const char \* *FileName*, const int *L* = ScaleNb ) const

Writes cell-average values on a regular grid of level *L* into file *FileName*.

##### Parameters

|                 |                  |
|-----------------|------------------|
| <i>FileName</i> | Name of the file |
| <i>L</i>        | Maximum level.   |



## Returns

void

```

1175 {
1176
1177     // --- Declarations -----
1178
1179     int l=0,i=0,j=0,k=0,n=0;    // counters
1180
1181     int ej = (Dimension > 1)? 1:0;
1182     int ek = (Dimension > 2)? 1:0;
1183
1184     real x=0., y=0., z=0., t=0.;    // Cell centers and time
1185     real dx=0., dy=0., dz=0.;    // Cell sizes
1186     int GridPoints;    // Grid points
1187     char DependencyType[12];    // positions or connections
1188
1189     PrintGrid FineGrid(L);
1190
1191     FILE *output;
1192
1193     // --- Execution -----
1194
1195     // --- Compute grid points and set dependency type ---
1196
1197     if (WriteAsPoints)
1198     {
1199         GridPoints = (1<<L);
1200         sprintf(DependencyType,"positions");
1201     }
1202     else
1203     {
1204         GridPoints = (1<<L)+1;
1205         sprintf(DependencyType,"connections");
1206     }
1207
1208     // --- Compute t, dx, dy, dz ---
1209
1210     t = ElapsedTime;
1211
1212     dx = (XMax[1]-XMin[1])/((1<<L)-1);
1213
1214     if (Dimension > 1)
1215         dy = (XMax[2]-XMin[2])/((1<<L)-1);
1216
1217     if (Dimension > 2)
1218         dz = (XMax[3]-XMin[3])/((1<<L)-1);
1219
1220
1221     // --- Compute result on fine mesh ---
1222
1223     for (l=0; l<=L; l++)
1224     {
1225         for (i = 0; i <= ((1<<l)-1); i++)
1226             for (j = 0; j <= ((1<<l)-1)*ej; j++)
1227                 for (k = 0; k <= ((1<<l)-1)*ek; k++)
1228                 {
1229                     if (node(l,i,j,k)==0)
1230                         FineGrid.predict(l,i,j,k);
1231                     else
1232                         FineGrid.setValue(i,j,k,cell(l,i,j,k)->average());
1233                 }
1234         FineGrid.refresh();
1235     }
1236
1237
1238     // --- Open file ---
1239
1240     if ((output = fopen(FileName,"w")) != NULL)
1241     {
1242         // --- Header ---
1243
1244         switch(PostProcessing)
1245         {
1246             // GNUPLOT
1247             case 1:
1248                 fprintf(output,"#");
1249                 fprintf(output, TXTFORMAT, " x");
1250                 fprintf(output, TXTFORMAT, "#Density");
1251                 fprintf(output, TXTFORMAT, "#Pressure");
1252                 fprintf(output, TXTFORMAT, "#Psi");
1253                 fprintf(output, TXTFORMAT, "#Energy");
1254                 fprintf(output, TXTFORMAT, "#Velocity");
1255                 fprintf(output, TXTFORMAT, "#MagField");
1256                 fprintf(output, TXTFORMAT, "#Div B");
1257                 fprintf(output, "\n");

```

```

1258         break;
1259
1260         // DATA EXPLOTER
1261     case 2:
1262         // Header for Data explorer
1263
1264         fprintf(output, "# Data Explorer file\n# generated by Carmen %3.1f\n",
1265 CarmenVersion);
1266
1267         switch(Dimension)
1268         {
1269             case 2:
1270                 fprintf(output, "#grid = %d x %d\n", GridPoints, GridPoints);
1271                 fprintf(output, "#positions = %f, %f, %f, %f\n#\n", XMin[1], dx,
1272 XMin[2], dy);
1273                 break;
1274             case 3:
1275                 fprintf(output, "#grid = %d x %d x %d\n", GridPoints, GridPoints, GridPoints);
1276                 fprintf(output, "#positions = %f, %f, %f, %f, %f, %f\n#\n",
1277 XMin[1], dx, XMin[2], dy, XMin[3], dz );
1278                 break;
1279         };
1280         if (DataIsBinary)
1281             fprintf(output, "#format = binary\n");
1282         else
1283             fprintf(output, "#format = ascii\n");
1284
1285         fprintf(output, "#interleaving = field\n");
1286
1287         // MHD
1288         fprintf(output, "#field = density, pressure, psi, energy, velocity, magField, Div B\n");
1289         fprintf(output, "#structure = scalar, scalar, scalar, scalar, 3-vector, 3-vector, scalar \n
1290 ");
1291         fprintf(output, "#type = %s, %s, %s, %s, %s, %s\n", REAL,
1292 REAL, REAL, REAL, REAL, REAL);
1293         fprintf(output, "#dependency = %s, %s, %s, %s, %s, %s\n", DependencyType, DependencyType,
1294 DependencyType, DependencyType, DependencyType, DependencyType);
1295
1296         fprintf(output, "#header = marker \"START_DATA\\\n\" \n");
1297         fprintf(output, "#end\n");
1298         fprintf(output, "#START_DATA\n");
1299
1300         break;
1301
1302         // TECPLOT
1303     case 3:
1304         fprintf(output, "VARIABLES = \"x\"\n");
1305         fprintf(output, "\"y\"\n");
1306         if (Dimension > 2)
1307             fprintf(output, "\"z\"\n");
1308
1309         fprintf(output, "\"RHO\"\n\"P\"\n\"PSI\"\n\"E\"\n\"U\"\n\"V\"\n\"W\"\n\"BX\"\n\"BY\"\n\"BZ\"
1310 \n");
1311
1312         fprintf(output, "ZONE T=\"Carmen %3.1f\"\n", CarmenVersion);
1313         fprintf(output, "I=%i, ", (1<<L));
1314         if (Dimension > 1)
1315             fprintf(output, "J=%i, ", (1<<L));
1316         if (Dimension > 2)
1317             fprintf(output, "K=%i, ", (1<<L));
1318         fprintf(output, "F=POINT\n");
1319         break;
1320
1321     case 4:
1322         int N=(1<<L);
1323         fprintf(output, "# vtk DataFile Version 2.8\nSolucao MHD\n");
1324         if(DataIsBinary)
1325             fprintf(output, "BINARY\n");
1326         else
1327             fprintf(output, "ASCII\n");
1328
1329         fprintf(output, "DATASET STRUCTURED_GRID\n");
1330         if (Dimension == 2)
1331         {
1332             fprintf(output, "DIMENSIONS %d %d %d \n", N,N,1);
1333             fprintf(output, "POINTS %d FLOAT\n", N*N);
1334             for (int i = 0; i < N; i++)
1335                 for (int j = 0; j < N; j++)
1336                     fprintf(output, "%f %f %f \n", XMin[1] + i*dx,
1337 XMin[2] + j*dy, 0.0);
1338
1339             fprintf(output, "\n\nPOINT_DATA %d \n", N*N);
1340         }
1341         if (Dimension == 3)
1342         {

```

```

1337         fprintf(output, "DIMENSIONS %d %d %d \n", N,N,N);
1338         fprintf(output, "POINTS %d FLOAT\n", N*N*N);
1339         for (int i = 0; i < N; i++)
1340             for (int j = 0; j < N; j++)
1341                 for (int k = 0; k < N; k++)
1342                     fprintf(output, "%f %f %f \n", XMin[1] + i*dx,
XMin[2] + j*dy, XMin[3] + k*dz);
1343
1344         fprintf(output, "\n\nPOINT_DATA %d \n", N*N*N);
1345     }
1346
1347     break;
1348
1349 };
1350
1351 // --- write values ---
1352
1353 for (n=0; n < (1<<(Dimension*L)); n++)
1354 {
1355     // -- Compute i, j, k --
1356
1357     // For Gnuplot and DX, loop order: for i... {for j... {for k...} }
1358     if (PostProcessing != 3)
1359     {
1360         switch(Dimension)
1361         {
1362             case 1:
1363                 i = n;
1364                 j = k = 0;
1365                 break;
1366             case 2:
1367                 j = n%(1<<L);
1368                 i = n/(1<<L);
1369                 k = 0;
1370                 break;
1371             case 3:
1372                 k = n%(1<<L);
1373                 j = (n%(1<<(2*L)))/(1<<L);
1374                 i = n/(1<<(2*L));
1375                 break;
1376         };
1377     }
1378     else
1379     {
1380         // For Tecplot, loop order: for k... {for j... {for i...} }
1381         i = n%(1<<L);
1382         if (Dimension > 1)
1383             j = (n%(1<<(2*L)))/(1<<L);
1384         else
1385             j = 0;
1386         if (Dimension > 2)
1387             k = n/(1<<(2*L));
1388         else
1389             k = 0;
1390     }
1391
1392     // Compute x, y, z
1393     if (PostProcessing == 1 || PostProcessing == 2)
1394     {
1395         x = XMin[1]+i*dx;
1396         if (Dimension > 1) y = XMin[2]+j*dy;
1397         if (Dimension > 2) z = XMin[3]+k*dz;
1398     }
1399     // For Tecplot, write coordinates
1400     if (PostProcessing == 3)
1401     {
1402         FileWrite(output, FORMAT, x);
1403         if (Dimension > 1) FileWrite(output,
FORMAT, y);
1404         if (Dimension > 2) FileWrite(output,
FORMAT, z);
1405     }
1406
1407     // MHD
1408     if (PostProcessing == 4)
1409     {
1410         /*
1411         fprintf(output, "\n\nSCALARS eta float\nLOOKUP_TABLE default\n");
1412         for (n=0; n < (1<<(Dimension*L)); n++){
1413             switch(Dimension)
1414             {
1415                 case 1:
1416                     i = n;
1417

```

```

1421             j = k = 0;
1422             break;
1423
1424         case 2:
1425             j = n%(1<<L);
1426             i = n/(1<<L);
1427             k = 0;
1428             break;
1429
1430         case 3:
1431             k = n%(1<<L);
1432             j = (n%(1<<(2*L)))/(1<<L);
1433             i = n/(1<<(2*L));
1434             break;
1435     };
1436     FileWrite(output, FORMAT, FineGrid.etaConst(i,j,k));
1437     fprintf(output, "\n");
1438 }
1439 */
1440 fprintf(output, "\n\nSCALARS Density float\nLOOKUP_TABLE default\n");
1441 for (n=0; n < (1<<(Dimension*L)); n++){
1442     switch(Dimension)
1443     {
1444         case 1:
1445             i = n;
1446             j = k = 0;
1447             break;
1448
1449         case 2:
1450             j = n%(1<<L);
1451             i = n/(1<<L);
1452             k = 0;
1453             break;
1454
1455         case 3:
1456             k = n%(1<<L);
1457             j = (n%(1<<(2*L)))/(1<<L);
1458             i = n/(1<<(2*L));
1459             break;
1460     };
1461     FileWrite(output, FORMAT, FineGrid.density(i,j,k));
1462     fprintf(output, "\n");
1463 }
1464
1465 fprintf(output, "\n\nSCALARS Pressure float\nLOOKUP_TABLE default\n");
1466 for (n=0; n < (1<<(Dimension*L)); n++){
1467     switch(Dimension)
1468     {
1469         case 1:
1470             i = n;
1471             j = k = 0;
1472             break;
1473
1474         case 2:
1475             j = n%(1<<L);
1476             i = n/(1<<L);
1477             k = 0;
1478             break;
1479
1480         case 3:
1481             k = n%(1<<L);
1482             j = (n%(1<<(2*L)))/(1<<L);
1483             i = n/(1<<(2*L));
1484             break;
1485     };
1486     FileWrite(output, FORMAT, FineGrid.pressure(i,j,k));
1487     fprintf(output, "\n");
1488 }
1489
1490 fprintf(output, "\n\nSCALARS Energy float\nLOOKUP_TABLE default\n");
1491 for (n=0; n < (1<<(Dimension*L)); n++){
1492     switch(Dimension)
1493     {
1494         case 1:
1495             i = n;
1496             j = k = 0;
1497             break;
1498
1499         case 2:
1500             j = n%(1<<L);
1501             i = n/(1<<L);
1502             k = 0;
1503             break;
1504
1505         case 3:
1506             k = n%(1<<L);
1507             j = (n%(1<<(2*L)))/(1<<L);

```

```

1508             i = n/(1<<(2*L));
1509             break;
1510         };
1511         FileWrite(output, FORMAT, FineGrid.energy(i,j,k));
1512         fprintf(output, "\n");
1513     }
1514
1515     fprintf(output, "\n\nSCALARS Vx float\nLOOKUP_TABLE default\n");
1516     for (n=0; n < (1<<(Dimension*L)); n++){
1517         switch(Dimension)
1518         {
1519             case 1:
1520                 i = n;
1521                 j = k = 0;
1522                 break;
1523
1524             case 2:
1525                 j = n%(1<<L);
1526                 i = n/(1<<L);
1527                 k = 0;
1528                 break;
1529
1530             case 3:
1531                 k = n%(1<<L);
1532                 j = (n%(1<<(2*L)))/(1<<L);
1533                 i = n/(1<<(2*L));
1534                 break;
1535         };
1536         FileWrite(output, FORMAT, FineGrid.velocity(i,j,k,1));
1537         fprintf(output, "\n");
1538     }
1539
1540     fprintf(output, "\n\nSCALARS Vy float\nLOOKUP_TABLE default\n");
1541     for (n=0; n < (1<<(Dimension*L)); n++){
1542         switch(Dimension)
1543         {
1544             case 1:
1545                 i = n;
1546                 j = k = 0;
1547                 break;
1548
1549             case 2:
1550                 j = n%(1<<L);
1551                 i = n/(1<<L);
1552                 k = 0;
1553                 break;
1554
1555             case 3:
1556                 k = n%(1<<L);
1557                 j = (n%(1<<(2*L)))/(1<<L);
1558                 i = n/(1<<(2*L));
1559                 break;
1560         };
1561         FileWrite(output, FORMAT, FineGrid.velocity(i,j,k,2));
1562         fprintf(output, "\n");
1563     }
1564
1565     fprintf(output, "\n\nSCALARS Vz float\nLOOKUP_TABLE default\n");
1566     for (n=0; n < (1<<(Dimension*L)); n++){
1567         switch(Dimension)
1568         {
1569             case 1:
1570                 i = n;
1571                 j = k = 0;
1572                 break;
1573
1574             case 2:
1575                 j = n%(1<<L);
1576                 i = n/(1<<L);
1577                 k = 0;
1578                 break;
1579
1580             case 3:
1581                 k = n%(1<<L);
1582                 j = (n%(1<<(2*L)))/(1<<L);
1583                 i = n/(1<<(2*L));
1584                 break;
1585         };
1586         FileWrite(output, FORMAT, FineGrid.velocity(i,j,k,3));
1587         fprintf(output, "\n");
1588     }
1589
1590     fprintf(output, "\n\nSCALARS Bx float\nLOOKUP_TABLE default\n");
1591     for (n=0; n < (1<<(Dimension*L)); n++){
1592         switch(Dimension)
1593         {
1594             case 1:

```

```

1595             i = n;
1596             j = k = 0;
1597             break;
1598
1599         case 2:
1600             j = n%(1<<L);
1601             i = n/(1<<L);
1602             k = 0;
1603             break;
1604
1605         case 3:
1606             k = n%(1<<L);
1607             j = (n%(1<<(2*L)))/(1<<L);
1608             i = n/(1<<(2*L));
1609             break;
1610     };
1611     FileWrite(output, FORMAT, FineGrid.magField(i,j,k,1));
1612     fprintf(output, "\n");
1613 }
1614
1615 fprintf(output, "\n\nSCALARS By float\nLOOKUP_TABLE default\n");
1616 for (n=0; n < (1<<(Dimension*L)); n++){
1617     switch(Dimension)
1618     {
1619         case 1:
1620             i = n;
1621             j = k = 0;
1622             break;
1623
1624         case 2:
1625             j = n%(1<<L);
1626             i = n/(1<<L);
1627             k = 0;
1628             break;
1629
1630         case 3:
1631             k = n%(1<<L);
1632             j = (n%(1<<(2*L)))/(1<<L);
1633             i = n/(1<<(2*L));
1634             break;
1635     };
1636     FileWrite(output, FORMAT, FineGrid.magField(i,j,k,2));
1637     fprintf(output, "\n");
1638 }
1639
1640 fprintf(output, "\n\nSCALARS Bz float\nLOOKUP_TABLE default\n");
1641 for (n=0; n < (1<<(Dimension*L)); n++){
1642     switch(Dimension)
1643     {
1644         case 1:
1645             i = n;
1646             j = k = 0;
1647             break;
1648
1649         case 2:
1650             j = n%(1<<L);
1651             i = n/(1<<L);
1652             k = 0;
1653             break;
1654
1655         case 3:
1656             k = n%(1<<L);
1657             j = (n%(1<<(2*L)))/(1<<L);
1658             i = n/(1<<(2*L));
1659             break;
1660     };
1661     FileWrite(output, FORMAT, FineGrid.magField(i,j,k,3));
1662     fprintf(output, "\n");
1663 }
1664
1665 fprintf(output, "\n\nSCALARS DivB float\nLOOKUP_TABLE default\n");
1666 for (n=0; n < (1<<(Dimension*L)); n++){
1667     switch(Dimension)
1668     {
1669         case 1:
1670             i = n;
1671             j = k = 0;
1672             break;
1673
1674         case 2:
1675             j = n%(1<<L);
1676             i = n/(1<<L);
1677             k = 0;
1678             break;
1679
1680         case 3:
1681             k = n%(1<<L);

```

```

1682             j = (n%(1<<(2*L)))/(1<<L);
1683             i = n/(1<<(2*L));
1684             break;
1685         };
1686         FileWrite(output, FORMAT, FineGrid.divergenceB(i,j,k));
1687         fprintf(output, "\n");
1688     }
1689     fprintf(output, "\n\nSCALARS DivB2 float\nLOOKUP_TABLE default\n");
1690     for (n=0; n < (1<<(Dimension*L)); n++){
1691         switch(Dimension)
1692         {
1693             case 1:
1694                 i = n;
1695                 j = k = 0;
1696                 break;
1697
1698             case 2:
1699                 j = n%(1<<L);
1700                 i = n/(1<<L);
1701                 k = 0;
1702                 break;
1703
1704             case 3:
1705                 k = n%(1<<L);
1706                 j = (n%(1<<(2*L)))/(1<<L);
1707                 i = n/(1<<(2*L));
1708                 break;
1709         };
1710         FileWrite(output, FORMAT, FineGrid.vorticity(i,j,k));
1711         fprintf(output, "\n");
1712     }
1713
1714     fprintf(output, "\n\nVECTORS Velocity float\n");
1715     for (n=0; n < (1<<(Dimension*L)); n++){
1716         switch(Dimension)
1717         {
1718             case 1:
1719                 i = n;
1720                 j = k = 0;
1721                 break;
1722
1723             case 2:
1724                 j = n%(1<<L);
1725                 i = n/(1<<L);
1726                 k = 0;
1727                 break;
1728
1729             case 3:
1730                 k = n%(1<<L);
1731                 j = (n%(1<<(2*L)))/(1<<L);
1732                 i = n/(1<<(2*L));
1733                 break;
1734         };
1735
1736         FileWrite(output, FORMAT, FineGrid.velocity(i,j,k,1));
1737         FileWrite(output, FORMAT, FineGrid.velocity(i,j,k,2));
1738         FileWrite(output, FORMAT, FineGrid.velocity(i,j,k,3));
1739     }
1740
1741     fprintf(output, "\n\nVECTORS MagField float\n");
1742     for (n=0; n < (1<<(Dimension*L)); n++){
1743         switch(Dimension)
1744         {
1745             case 1:
1746                 i = n;
1747                 j = k = 0;
1748                 break;
1749
1750             case 2:
1751                 j = n%(1<<L);
1752                 i = n/(1<<L);
1753                 k = 0;
1754                 break;
1755
1756             case 3:
1757                 k = n%(1<<L);
1758                 j = (n%(1<<(2*L)))/(1<<L);
1759                 i = n/(1<<(2*L));
1760                 break;
1761         };
1762
1763         FileWrite(output, FORMAT, FineGrid.magField(i,j,k,1));
1764         FileWrite(output, FORMAT, FineGrid.magField(i,j,k,2));
1765         FileWrite(output, FORMAT, FineGrid.magField(i,j,k,3));

```

```

1769     }
1770     }else{
1771         for (n=0; n < (1<<(Dimension*L)); n++)
1772         {
1773             FileWrite(output, FORMAT, FineGrid.density(i,j,k));
1774             FileWrite(output, FORMAT, FineGrid.pressure(i,j,k));
1775             FileWrite(output, FORMAT, FineGrid.psi(i,j,k));
1776             FileWrite(output, FORMAT, FineGrid.energy(i,j,k));
1777
1778             for (int AxisNo = 1; AxisNo <= 3; AxisNo++){
1779                 FileWrite(output, FORMAT, FineGrid.velocity(i,j,k,AxisNo));
1780                 FileWrite(output, FORMAT, FineGrid.magField(i,j,k,AxisNo));
1781             }
1782             FileWrite(output, FORMAT, FineGrid.divergenceB(i,j,k));
1783         }
1784     }
1785
1786     for (n=0; n < (1<<(Dimension*L)); n++){
1787         // For ASCII data, add a return at the end of the line
1788         if (!DataIsBinary)
1789             fprintf(output, "\n");
1790
1791         // For Gnuplot, add empty lines when j=jmax or k=kmax
1792         if (PostProcessing == 1)
1793         {
1794             if (j==(1<<ScaleNb)-1)
1795                 fprintf(output, "\n");
1796
1797             if (k==(1<<ScaleNb)-1)
1798                 fprintf(output, "\n");
1799         }
1800     }
1801     fclose(output);
1802 }
1803 }
1804 else
1805 {
1806     cout << "Node.cpp: In method 'void writeFineGrid(Node*, char*, int)':\n";
1807     cout << "Node.cpp: cannot open file " << FileName << '\n';
1808     cout << "carmen: *** [Node.o] Execution error\n";
1809     cout << "carmen: abort execution.\n";
1810     exit(1);
1811 }
1812 }/*

```

Here is the caller graph for this function:



#### 5.4.3.23 void Node::writeHeader ( const char \* *FileName* ) const

Writes header for Data Explorer into file *FileName*.

##### Parameters

|                 |                   |
|-----------------|-------------------|
| <i>FileName</i> | Name of the file. |
|-----------------|-------------------|

##### Returns

void

```

1029 {
1030     // --- Local variables ---
1031

```

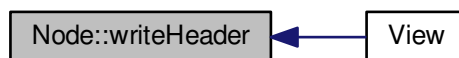


```

1032 FILE *output; // Pointer to output file
1033
1034 // --- Open file ---
1035
1036 if ((output = fopen(FileName,"w")) != NULL)
1037 {
1038 // --- Header ---
1039
1040 if (Dimension == 1)
1041 {
1042 // GNUPLOT
1043 fprintf(output, "#");
1044 fprintf(output, TXTFORMAT, " x");
1045 fprintf(output, TXTFORMAT, "Density");
1046 fprintf(output, TXTFORMAT, "Pressure");
1047 fprintf(output, TXTFORMAT, "Psi");
1048 fprintf(output, TXTFORMAT, "Energy");
1049 fprintf(output, TXTFORMAT, "Velocity");
1050 fprintf(output, TXTFORMAT, "MagField");
1051 fprintf(output, "\n");
1052 }
1053 else
1054 {
1055 fprintf(output, "# Data Explorer file\n# generated by Carmen %3.1f\n",
CarmenVersion);
1056 fprintf(output, "# points = %d\n", LeafNb);
1057 fprintf(output, "# format = ascii\n");
1058 fprintf(output, "# interleaving = field\n");
1059
1060 fprintf(output, "# field = locations, Q1\n");
1061 fprintf(output, "# structure = %d-vector, scalar\n", Dimension);
1062 fprintf(output, "# type = %s, %s\n", REAL, REAL);
1063 fprintf(output, "# dependency = positions, positions\n");
1064
1065 fprintf(output, "# header = marker \"START_DATA\\n\" \\n");
1066 fprintf(output, "# end\n");
1067 fprintf(output, "# START_DATA\n");
1068 }
1069
1070 fclose(output);
1071 return;
1072 }
1073 else
1074 {
1075 cout << "Node.cpp: In method `void Node::writeHeader()':\n";
1076 cout << "Node.cpp: cannot open file " << FileName << "\n";
1077 cout << "carmen: *** [Node.o] Execution error\n";
1078 cout << "carmen: abort execution.\n";
1079 exit(1);
1080 }
1081 }

```

Here is the caller graph for this function:



#### 5.4.3.24 void Node::writeMesh ( const char \* FileName ) const

Writes mesh data for Gnuplot into file *FileName*.

##### Parameters

| <i>FileName</i> | Name of the file. |
|-----------------|-------------------|
|-----------------|-------------------|

## Returns

void

```

1820 {
1821     // --- Local variables ---
1822
1823     int     n;           // Counter on children
1824     FILE    *output;    // Pointer to output file
1825
1826     // --- Open file ---
1827
1828     if ( (Nl == 0) ? (output = fopen(FileName,"w")) : (output = fopen(FileName,"a")) )
1829     {
1830         if (isInternalNode())
1831         {
1832             for (n = 0; n < ChildNb; n++)
1833                 Child[n]->writeMesh(FileName);
1834         }
1835         else
1836         {
1837             // x-direction
1838             fprintf(output, FORMAT, ThisCell.center(1)-.5*ThisCell.
size(1));
1839             if (Dimension >1) fprintf(output, FORMAT, ThisCell.
center(2)-.5*ThisCell.size(2));
1840             if (Dimension >2) fprintf(output, FORMAT, ThisCell.
center(3)-.5*ThisCell.size(3));
1841             fprintf(output, "%d", Nl);
1842             fprintf(output, "\n");
1843
1844             fprintf(output, FORMAT, ThisCell.center(1)+.5*ThisCell.
size(1));
1845             if (Dimension >1) fprintf(output, FORMAT, ThisCell.
center(2)-.5*ThisCell.size(2));
1846             if (Dimension >2) fprintf(output, FORMAT, ThisCell.
center(3)-.5*ThisCell.size(3));
1847             fprintf(output, "%d", Nl);
1848             fprintf(output, "\n\n");
1849
1850             // y-direction
1851             if (Dimension > 1)
1852             {
1853                 fprintf(output, FORMAT, ThisCell.center(1)-.5*ThisCell.
size(1));
1854                 fprintf(output, FORMAT, ThisCell.center(2)+.5*ThisCell.
size(2));
1855                 if (Dimension >2) fprintf(output, FORMAT, ThisCell.
center(3)-.5*ThisCell.size(3));
1856                 fprintf(output, "%d", Nl);
1857                 fprintf(output, "\n");
1858
1859                 fprintf(output, FORMAT, ThisCell.center(1)+.5*ThisCell.
size(1));
1860                 fprintf(output, FORMAT, ThisCell.center(2)+.5*ThisCell.
size(2));
1861                 if (Dimension >2) fprintf(output, FORMAT, ThisCell.
center(3)-.5*ThisCell.size(3));
1862                 fprintf(output, "%d", Nl);
1863                 fprintf(output, "\n\n");
1864             }
1865
1866             // z-direction
1867             if (Dimension > 2)
1868             {
1869                 fprintf(output, FORMAT, ThisCell.center(1)-.5*ThisCell.
size(1));
1870                 fprintf(output, FORMAT, ThisCell.center(2)-.5*ThisCell.
size(2));
1871                 fprintf(output, FORMAT, ThisCell.center(3)+.5*ThisCell.
size(3));
1872                 fprintf(output, "%d", Nl);
1873                 fprintf(output, "\n");
1874
1875                 fprintf(output, FORMAT, ThisCell.center(1)+.5*ThisCell.
size(1));
1876                 fprintf(output, FORMAT, ThisCell.center(2)-.5*ThisCell.
size(2));
1877                 fprintf(output, FORMAT, ThisCell.center(3)+.5*ThisCell.
size(3));
1878                 fprintf(output, "%d", Nl);
1879                 fprintf(output, "\n\n");

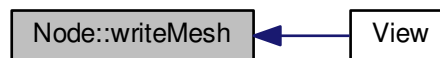
```

```

1880
1881         fprintf(output, FORMAT, ThisCell.center(1)-.5*ThisCell.
size(1));
1882         fprintf(output, FORMAT, ThisCell.center(2)+.5*ThisCell.
size(2));
1883         fprintf(output, FORMAT, ThisCell.center(3)+.5*ThisCell.
size(3));
1884         fprintf(output, "%d", N1);
1885         fprintf(output, "\n");
1886
1887         fprintf(output, FORMAT, ThisCell.center(1)+.5*ThisCell.
size(1));
1888         fprintf(output, FORMAT, ThisCell.center(2)+.5*ThisCell.
size(2));
1889         fprintf(output, FORMAT, ThisCell.center(3)+.5*ThisCell.
size(3));
1890         fprintf(output, "%d", N1);
1891         fprintf(output, "\n\n\n");
1892     }
1893     fprintf(output, "\n");
1894 }
1895 fclose(output);
1896 }
1897 else
1898 {
1899     cout << "Node.cpp: In method 'void Node::writeMesh()':\n";
1900     cout << "Node.cpp: cannot open file " << FileName << '\n';
1901     cout << "carmen: *** [Node.o] Execution error\n";
1902     cout << "carmen: abort execution.\n";
1903     exit(1);
1904 }
1905 }

```

Here is the caller graph for this function:



#### 5.4.3.25 void Node::writeTree ( const char \* FileName ) const

Writes tree structure into file *FileName*. Only for debugging.

##### Parameters

|                 |                   |
|-----------------|-------------------|
| <i>FileName</i> | Name of the file. |
|-----------------|-------------------|

##### Returns

void

```

956 {
957     // --- Local variables ---
958
959     int n, l;           // Counter
960
961     FILE *output;     // Pointer to output file
962
963     // --- Open file ---
964
965     if ( (N1 == 0) ? (output = fopen(FileName,"w")) : (output = fopen(FileName,"a")) )
966     {
967         for (l = 1; l <= N1; l++)
968             fprintf(output, "| " );
969
970         fprintf(output, "+ ");

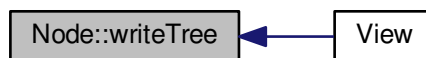
```

```

971     switch (Dimension)
972     {
973     case 1:
974         fprintf(output, "(%d, %d)", Nl, Ni );
975         break;
976
977     case 2:
978         fprintf(output, "(%d, %d, %d)", Nl, Ni, Nj);
979         break;
980
981     case 3:
982         fprintf(output, "(%d, %d, %d, %d)", Nl, Ni, Nj, Nk );
983         break;
984     };
985     switch(Flag)
986     {
987     case 0:
988         fprintf(output, "-- node --");
989         break;
990
991     case 1:
992         fprintf(output, "-- leaf --");
993         break;
994
995     case 2:
996         fprintf(output, "-- leaf with virtual children --");
997         break;
998
999     case 3:
1000         fprintf(output, "-- virtual leaf --");
1001         break;
1002     };
1003     fprintf(output, "\n");
1004     fclose(output);
1005     if (hasChildren())
1006     {
1007         for (n = 0; n < ChildNb; n++)
1008             Child[n]->writeTree(FileName);
1009     }
1010 }
1011 else
1012 {
1013     cout << "Node.cpp: In method 'void Node::writeText()':\n";
1014     cout << "Node.cpp: cannot open file " << FileName << '\n';
1015     cout << "carmen: *** [Node.o] Execution error\n";
1016     cout << "carmen: abort execution.\n";
1017     exit(1);
1018 }
1019 }
1020 }

```

Here is the caller graph for this function:



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

- [Node.h](#)
- [Node.cpp](#)

## 5.5 PrintGrid Class Reference

An object [PrintGrid](#) is a special regular grid created to write tree-structured data into an output file.

```
#include <PrintGrid.h>
```

## Public Member Functions

- [PrintGrid](#) (int L)
- [~PrintGrid](#) ()
- void [setValue](#) (const int i, const int j, const int k, const [Vector](#) &UserAverage)
 

*Sets the cell-average vector located at i, j, k to UserAverage.*
- [Vector value](#) (const int i, const int j, const int k) const
 

*Returns the cell-average vector located at i, j, k.*
- [real value](#) (const int i, const int j, const int k, const int QuantityNo) const
 

*Returns the quantity QuantityNo of the cell-average vector located at i, j, k.*
- [real cellValue](#) (const int i, const int j, const int k, const int QuantityNo) const
 

*Returns the quantity QuantityNo of the cell-average vector located at i, j, k, taking into account boundary conditions.*
- [real density](#) (const int i, const int j, const int k) const
 

*Returns the cell-average density located at i, j, k.*
- [real psi](#) (const int i, const int j, const int k) const
 

*Returns the cell-average density located at i, j, k.*
- [real pressure](#) (const int i, const int j, const int k) const
 

*Returns the cell-average pressure located at i, j, k.*
- [real temperature](#) (const int i, const int j, const int k) const
 

*Returns the cell-average temperature located at i, j, k.*
- [real concentration](#) (const int i, const int j, const int k) const
 

*Returns the cell-average concentration of the limiting reactant, located at i, j, k.*
- [real energy](#) (const int i, const int j, const int k) const
 

*Returns the cell-average energy per unit of volume located at i, j, k.*
- [real velocity](#) (const int i, const int j, const int k, const int AxisNo) const
 

*Returns the AxisNo-th component of the cell-average velocity located at i, j, k.*
- [Vector velocity](#) (const int i, const int j, const int k) const
 

*Returns the cell-average velocity located at i, j, k.*
- [real divergenceB](#) (const int i, const int j, const int k) const
 

*Returns the divergence of magnetic field B located at i, j, k.*
- [real vorticity](#) (const int i, const int j, const int k) const
 

*Returns 0 in 1D, the scalar vorticity in 2D, the norm of the cell-average vorticity in 3D, located at i, j, k. Does not work for MHD!*
- [real magField](#) (const int i, const int j, const int k, const int AxisNo) const
 

*Returns the AxisNo-th component of the cell-average velocity located at i, j, k.*
- [Vector magField](#) (const int i, const int j, const int k) const
 

*Returns the cell-average velocity located at i, j, k.*
- void [refresh](#) ()
 

*Stores the cell-average values of the current grid into temporary values, in order to compute cell-averages in the next finer grid.*
- void [predict](#) (const int l, const int i, const int j, const int k)
 

*Predicts the cell-average values of the current grid from the values stored in the temporary ones.*
- void [computePointValue](#) ()
 

*Transform cell-average values into point values.*

### 5.5.1 Detailed Description

An object `PrintGrid` is a special regular grid created to write tree-structured data into an output file.

It contains the following data:

- the scale number of the grid *LocalScaleNb* ;
- the current number of elements used in the grid *ElementNb* ;
- the array of cell-average values *\*Q* ;
- an array of temporary cell-average values *\*Qt*.

To write tree-structured data into a regular fine grid, one starts with the grid of level 0 and one stops at the level *L*. For a given grid of level *l* and a given element *i, j, k* of this grid, if the node of the tree corresponding to the element is a leaf, the value is replaced by the one of the node, else it is predicted from parents.

Such grid does not contain any cell information, in order to reduce memory requirements.

### 5.5.2 Constructor & Destructor Documentation

#### 5.5.2.1 `PrintGrid::PrintGrid ( int L )`

Constructor of `PrintGrid` class. Generates a grid of  $2^{*(Dimension*L)}$  elements.

```

31 {
32
33     int n=0; // Array size
34
35     localScaleNb=L;
36     elementNb = n = (1<<localScaleNb)+1;
37     if (Dimension > 1) elementNb *= n;
38     if (Dimension > 2) elementNb *= n;
39
40     Q = new Vector[elementNb];
41     Qt = new Vector[elementNb];
42
43     int i;
44     for (i=0;i<elementNb;i++)
45     {
46         Q[i].setDimension(QuantityNb);
47         Qt[i].setDimension(QuantityNb);
48     }
49
50
51 }
```

#### 5.5.2.2 `PrintGrid::~PrintGrid ( )`

Destructor of the `PrintGrid` class. Removes the grid.

```

60 {
61     delete[] Q;
62     delete[] Qt;
63 }
```

### 5.5.3 Member Function Documentation

#### 5.5.3.1 `real PrintGrid::cellValue ( const int i, const int j, const int k, const int QuantityNo ) const`

Returns the quantity *QuantityNo* of the cell-average vector located at *i, j, k*, taking into account boundary conditions.

## Parameters

|                   |                          |
|-------------------|--------------------------|
| <i>i</i>          | Position x               |
| <i>j</i>          | Position y               |
| <i>k</i>          | Position z               |
| <i>QuantityNo</i> | Number of MHD variables. |

## Returns

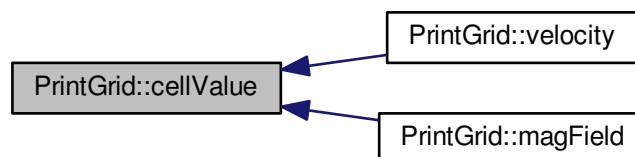
real

```

114 {
115     // --- Local variables ---
116
117     int n = (1<<localScaleNb)+1;           // n = 2^localScaleNb+1
118     int li=0, lj=0, lk=0;                // local i,j,k
119
120
121     if (CMin[1] == 2)
122         li = ((i+n)/n==1)? i : (2*n-i-1)%n; // Neumann
123     else
124         li = (i+n)%n;                      // Periodic
125
126     // -- in y --
127
128     if (Dimension > 1)
129     {
130         if (CMin[2] == 2)
131             lj = ((j+n)/n==1)? j : (2*n-j-1)%n; // Neumann
132         else
133             lj = (j+n)%n;                    // Periodic
134     }
135
136     // -- in z --
137
138     if (Dimension > 2)
139     {
140         if (CMin[3] == 2)
141             lk = ((k+n)/n==1)? k : (2*n-k-1)%n; // Neumann
142         else
143             lk = (k+n)%n;                    // Periodic
144     }
145
146     return (Q + li + n*(lj + n*lk))->value(QuantityNo);
147 }

```

Here is the caller graph for this function:



## 5.5.3.2 void PrintGrid::computePointValue ( )

Transform cell-average values into point values.

**Returns**

void

```

544 {
545     int i=0, j=0, k=0;
546     int l=localScaleNb;
547     int n=(1<<1);
548
549     switch(Dimension)
550     {
551         case 1:
552             for (i=0; i<=n; i++)
553                 setValue(i, j, k, .5*(tempValue(l, i-1, j, k)+tempValue(l, i, j, k)));
554             break;
555
556         case 2:
557             for (i=0; i<=n; i++)
558                 for (j=0; j<=n; j++)
559                     setValue(i, j, k, .25*(tempValue(l, i-1, j-1, k)+tempValue(l, i-1, j, k)+tempValue(l, i, j-1,
560 k)+tempValue(l, i, j, k)) );
561             break;
562
563         default:
564             for (i=0; i<=n; i++)
565                 for (j=0; j<=n; j++)
566                     for (k=0; k<=n; k++)
567                         setValue(i, j, k, .125*(tempValue(l, i-1, j-1, k-1)+tempValue(l, i-1, j, k-1)+tempValue(l, i,
568 j-1, k-1)+tempValue(l, i, j, k-1)
569                                     +tempValue(l, i-1, j-1, k)+tempValue(l, i-1, j, k)+tempValue(l, i,
570 j-1, k)+tempValue(l, i, j, k)) );
571             break;
572     };
573 }

```

**5.5.3.3 real PrintGrid::concentration ( const int *i*, const int *j*, const int *k* ) const**

Returns the cell-average concentration of the limiting reactant, located at *i, j, k*.

**Parameters**

|          |            |
|----------|------------|
| <i>i</i> | Position x |
| <i>j</i> | Position y |
| <i>k</i> | Position z |

**Returns**

real

```

220 {
221
222     if (EquationType >=3 && EquationType <=5)
223         return value(i, j, k, 2);
224
225     return 0.;
226 }

```

**5.5.3.4 real PrintGrid::density ( const int *i*, const int *j*, const int *k* ) const [inline]**

Returns the cell-average density located at *i, j, k*.

**Parameters**

|          |            |
|----------|------------|
| <i>i</i> | Position x |
| <i>j</i> | Position y |



|     |              |
|-----|--------------|
| $k$ | Position $z$ |
|-----|--------------|

**Returns**

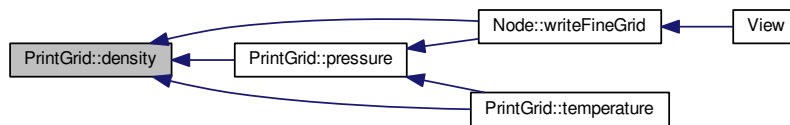
real

```

307 {
308     return value(i, j, k, l);
309 }

```

Here is the caller graph for this function:

**5.5.3.5 real PrintGrid::divergenceB ( const int  $i$ , const int  $j$ , const int  $k$  ) const**

Returns the divergence of magnetic field B located at  $i, j, k$ .

**Parameters**

|     |            |
|-----|------------|
| $i$ | Position x |
| $j$ | Position y |
| $k$ | Position z |

**Returns**

real

```

236 {
237     int L=localScaleNb;
238     int n = (l<<L);
239     real dx=0., dy=0., dz=0.;
240     real Div=0.;
241     real By1=0., By2=0., Bz1=0., Bz2=0.;
242     real Bx1=0., Bx2=0.;
243     real Bx =0., By=0.;
244     if (Dimension == 1){
245         dx = (XMax[1]-XMin[1])/n;
246
247         Bx1 = magField(BC(i-1,1,n), BC(j,2,n),BC(k,3,n),1);
248         Bx2 = magField(BC(i+1,1,n), BC(j,2,n),BC(k,3,n),1);
249
250         Div = (Bx2-Bx1)/(2.*dx);
251     }else if (Dimension == 2){
252
253         dx = (XMax[1]-XMin[1])/n;
254         dy = (XMax[2]-XMin[2])/n;
255         Bx1 = magField(BC(i-1,1,n), BC(j,2,n),BC(k,3,n),1);
256         Bx2 = magField(BC(i+1,1,n), BC(j,2,n),BC(k,3,n),1);
257         By1 = magField(BC(i,1,n), BC(j-1,2,n),BC(k,3,n),2);
258         By2 = magField(BC(i,1,n), BC(j+1,2,n),BC(k,3,n),2);
259         Bx = magField(BC(i,1,n), BC(j,2,n),BC(k,3,n),1);
260         By = magField(BC(i,1,n), BC(j,2,n),BC(k,3,n),2);
261
262         //if(Bx2!=Bx1 && By2!=By1)
263         Div = ((Bx2-Bx1)/(dx) + (By2-By1)/(dy))/((fabs(Bx1)+fabs(Bx2))/(dx) + (fabs(By1)+fabs(By2))/(dy
264 ) + 1.120e-13);
265         //else
266         // Div = ((Bx2-Bx1)/(2.*dx) + (By2-By1)/(2.*dy));

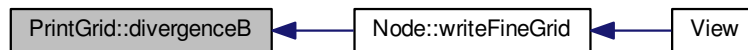
```

```

267     }else if (Dimension == 3){
268
269         dx = (XMax[1]-XMin[1])/n;
270         dy = (XMax[2]-XMin[2])/n;
271         dz = (XMax[3]-XMin[3])/n;
272
273         Bx1 = magField(BC(i-1,1,n), BC(j,2,n), BC(k,3,n),1);
274         Bx2 = magField(BC(i+1,1,n), BC(j,2,n), BC(k,3,n),1);
275         By1 = magField(BC(i,1,n), BC(j-1,2,n), BC(k,3,n),2);
276         By2 = magField(BC(i,1,n), BC(j+1,2,n), BC(k,3,n),2);
277         Bz1 = magField(BC(i,1,n), BC(j,2,n), BC(k-1,3,n),3);
278         Bz2 = magField(BC(i,1,n), BC(j,2,n), BC(k+1,3,n),3);
279
280         Div = (Bx2-Bx1)/(2.*dx) + (By2-By1)/(2.*dy) + (Bz2-Bz1)/(2.*dz);
281
282     }
283
284     return Div;
285 }

```

Here is the caller graph for this function:



#### 5.5.3.6 real PrintGrid::energy ( const int *i*, const int *j*, const int *k* ) const [inline]

Returns the cell-average energy per unit of volume located at *i*, *j*, *k*.

##### Parameters

|          |            |
|----------|------------|
| <i>i</i> | Position x |
| <i>j</i> | Position y |
| <i>k</i> | Position z |

##### Returns

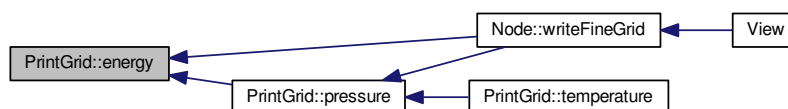
real

```

327 {
328     return value(i,j,k,5);
329 }

```

Here is the caller graph for this function:



#### 5.5.3.7 real PrintGrid::magField ( const int *i*, const int *j*, const int *k*, const int *AxisNo* ) const [inline]

Returns the *AxisNo*-th component of the cell-average velocity located at *i*, *j*, *k*.

## Parameters

|               |                  |
|---------------|------------------|
| <i>i</i>      | Position x       |
| <i>j</i>      | Position y       |
| <i>k</i>      | Position z       |
| <i>AxisNo</i> | Axis of interest |

## Returns

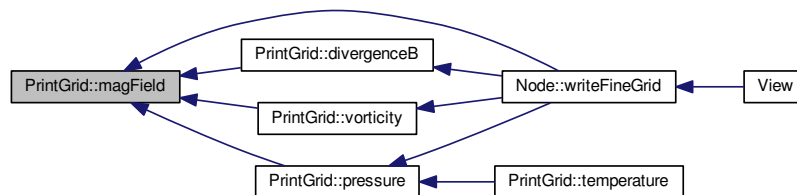
real

```

348 {
349     return cellValue(i, j, k, AxisNo+6);
350 }

```

Here is the caller graph for this function:

5.5.3.8 Vector PrintGrid::magField ( const int *i*, const int *j*, const int *k* ) const

Returns the cell-average velocity located at *i*, *j*, *k*.

## Parameters

|          |            |
|----------|------------|
| <i>i</i> | Position x |
| <i>j</i> | Position y |
| <i>k</i> | Position z |

## Returns

Vector

```

170 {
171     Vector V(3);
172
173     for (int AxisNo=1; AxisNo <= 3; AxisNo++)
174         V.setValue( AxisNo, cellValue(i, j, k, AxisNo+6));
175
176     return V;
177 }

```

5.5.3.9 void PrintGrid::predict ( const int *l*, const int *i*, const int *j*, const int *k* )

Predicts the cell-average values of the current grid from the values stored in the temporary ones.

## Parameters

|     |            |
|-----|------------|
| $i$ | Level      |
| $j$ | Position x |
| $k$ | Position y |
|     | Position z |

## Returns

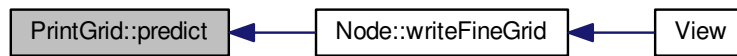
void

```

475 {
476     // --- Local variables ---
477
478     int pi=1, pj=1, pk=1; // Parity of i,j,k
479
480     Vector    Result(QuantityNb);
481
482     // --- Init result with the cell-average value of Qt ---
483
484     Result = tempValue(l-1, (i+4)/2-2, (j+4)/2-2, (k+4)/2-2);
485
486     // --- 1D case ---
487
488     pi = (i%2 == 0)?1:-1;
489     Result += (pi*-.125) * tempValue(l-1, (i+4)/2-2+1, (j+4)/2-2, (k+4)/2-2);
490     Result -= (pi*-.125) * tempValue(l-1, (i+4)/2-2-1, (j+4)/2-2, (k+4)/2-2);
491
492     // --- 2D case ---
493
494     if (Dimension > 1)
495     {
496         pj = (j%2 == 0)?1:-1;
497         Result += (pj*-.125) * tempValue(l-1, (i+4)/2-2, (j+4)/2-2+1, (k+4)/2-2);
498         Result -= (pj*-.125) * tempValue(l-1, (i+4)/2-2, (j+4)/2-2-1, (k+4)/2-2);
499
500         Result += (pi*pj*.015625) * tempValue(l-1, (i+4)/2-2+1, (j+4)/2-2+1, (k+4)/2-2);
501         Result -= (pi*pj*.015625) * tempValue(l-1, (i+4)/2-2+1, (j+4)/2-2-1, (k+4)/2-2);
502         Result -= (pi*pj*.015625) * tempValue(l-1, (i+4)/2-2-1, (j+4)/2-2+1, (k+4)/2-2);
503         Result += (pi*pj*.015625) * tempValue(l-1, (i+4)/2-2-1, (j+4)/2-2-1, (k+4)/2-2);
504     }
505
506     // --- 3D case ---
507
508     if (Dimension > 2)
509     {
510         pk = (k%2 == 0)?1:-1;
511         Result += (pk*-.125) * tempValue(l-1, (i+4)/2-2, (j+4)/2-2, (k+4)/2-2+1);
512         Result -= (pk*-.125) * tempValue(l-1, (i+4)/2-2, (j+4)/2-2, (k+4)/2-2-1);
513
514         Result += (pi*pk*.015625) * tempValue(l-1, (i+4)/2-2+1, (j+4)/2-2, (k+4)/2-2+1);
515         Result -= (pi*pk*.015625) * tempValue(l-1, (i+4)/2-2+1, (j+4)/2-2, (k+4)/2-2-1);
516         Result -= (pi*pk*.015625) * tempValue(l-1, (i+4)/2-2-1, (j+4)/2-2, (k+4)/2-2+1);
517         Result += (pi*pk*.015625) * tempValue(l-1, (i+4)/2-2-1, (j+4)/2-2, (k+4)/2-2-1);
518
519         Result += (pj*pk*.015625) * tempValue(l-1, (i+4)/2-2, (j+4)/2-2+1, (k+4)/2-2+1);
520         Result -= (pj*pk*.015625) * tempValue(l-1, (i+4)/2-2, (j+4)/2-2+1, (k+4)/2-2-1);
521         Result -= (pj*pk*.015625) * tempValue(l-1, (i+4)/2-2, (j+4)/2-2-1, (k+4)/2-2+1);
522         Result += (pj*pk*.015625) * tempValue(l-1, (i+4)/2-2, (j+4)/2-2-1, (k+4)/2-2-1);
523
524         Result += (pi*pj*pk*-.001953125) * tempValue(l-1, (i+4)/2-2+1, (j+4)/2-2+1, (k+4)/2-2+1);
525         Result -= (pi*pj*pk*-.001953125) * tempValue(l-1, (i+4)/2-2+1, (j+4)/2-2+1, (k+4)/2-2-1);
526         Result -= (pi*pj*pk*-.001953125) * tempValue(l-1, (i+4)/2-2+1, (j+4)/2-2-1, (k+4)/2-2+1);
527         Result += (pi*pj*pk*-.001953125) * tempValue(l-1, (i+4)/2-2+1, (j+4)/2-2-1, (k+4)/2-2-1);
528         Result -= (pi*pj*pk*-.001953125) * tempValue(l-1, (i+4)/2-2-1, (j+4)/2-2+1, (k+4)/2-2+1);
529         Result += (pi*pj*pk*-.001953125) * tempValue(l-1, (i+4)/2-2-1, (j+4)/2-2+1, (k+4)/2-2-1);
530         Result += (pi*pj*pk*-.001953125) * tempValue(l-1, (i+4)/2-2-1, (j+4)/2-2-1, (k+4)/2-2+1);
531         Result -= (pi*pj*pk*-.001953125) * tempValue(l-1, (i+4)/2-2-1, (j+4)/2-2-1, (k+4)/2-2-1);
532     }
533
534     setValue(i, j, k, Result);
535 }

```

Here is the caller graph for this function:



#### 5.5.3.10 real PrintGrid::pressure ( const int *i*, const int *j*, const int *k* ) const

Returns the cell-average pressure located at *i*, *j*, *k*.

##### Parameters

|          |            |
|----------|------------|
| <i>i</i> | Position x |
| <i>j</i> | Position y |
| <i>k</i> | Position z |

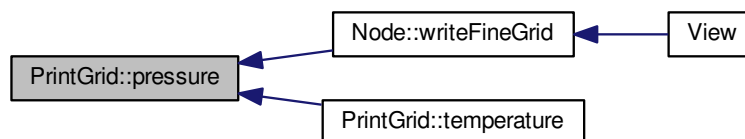
##### Returns

real

```

185 {
186   Vector V(3), B(3);
187   real rho, rhoE;
188
189   rho = density(i,j,k);
190   V   = velocity(i,j,k);
191   B   = magField(i,j,k);
192   rhoE = energy(i,j,k);
193
194   return (Gamma-1.)*(rhoE - .5*rho*(V*V) - .5*(B*B));
195 }
  
```

Here is the caller graph for this function:



#### 5.5.3.11 real PrintGrid::psi ( const int *i*, const int *j*, const int *k* ) const [inline]

Returns the cell-average density located at *i*, *j*, *k*.

## Parameters

|          |            |
|----------|------------|
| <i>i</i> | Position x |
| <i>j</i> | Position y |
| <i>k</i> | Position z |

## Returns

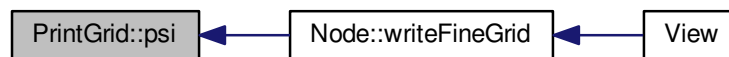
real

```

317 {
318     return value(i, j, k, 6);
319 }

```

Here is the caller graph for this function:



## 5.5.3.12 void PrintGrid::refresh ( )

Stores the cell-average values of the current grid into temporary values, in order to compute cell-averages in the next finer grid.

## Returns

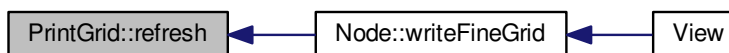
void

```

463 {
464     for (int n=0; n<elementNb; n++)
465         *(Qt+n) = *(Q+n);
466 }

```

Here is the caller graph for this function:

5.5.3.13 void PrintGrid::setValue ( const int *i*, const int *j*, const int *k*, const Vector & *UserAverage* )

Sets the cell-average vector located at *i*, *j*, *k* to *UserAverage*.

## Parameters

|                    |                    |
|--------------------|--------------------|
| <i>i</i>           | Position x         |
| <i>j</i>           | Position y         |
| <i>k</i>           | Position z         |
| <i>UserAverage</i> | Vector of averages |

## Returns

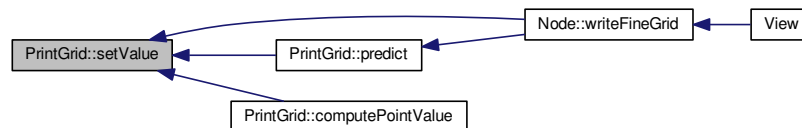
void

```

72 {
73     // --- Local variables ---
74
75     int n=(1<<localScaleNb)+1; // n = 2^localScaleNb+1
76
77     *(Q + i + n*(j + n*k)) = UserAverage;
78 }

```

Here is the caller graph for this function:

5.5.3.14 `real PrintGrid::temperature ( const int i, const int j, const int k ) const`

Returns the cell-average temperature located at *i, j, k*.

## Parameters

|          |            |
|----------|------------|
| <i>i</i> | Position x |
| <i>j</i> | Position y |
| <i>k</i> | Position z |

## Returns

real

```

203 {
204     real rho, p;
205
206     if (EquationType >=3 && EquationType <=5)
207         return value(i, j, k, 1);
208
209     rho = density(i, j, k);
210     p = pressure(i, j, k);
211
212     return Gamma*Ma*Ma*p/rho;
213 }

```

5.5.3.15 `Vector PrintGrid::value ( const int i, const int j, const int k ) const`

Returns the cell-average vector located at *i, j, k*.

## Parameters

|          |            |
|----------|------------|
| <i>i</i> | Position x |
| <i>j</i> | Position y |
| <i>k</i> | Position z |

## Returns

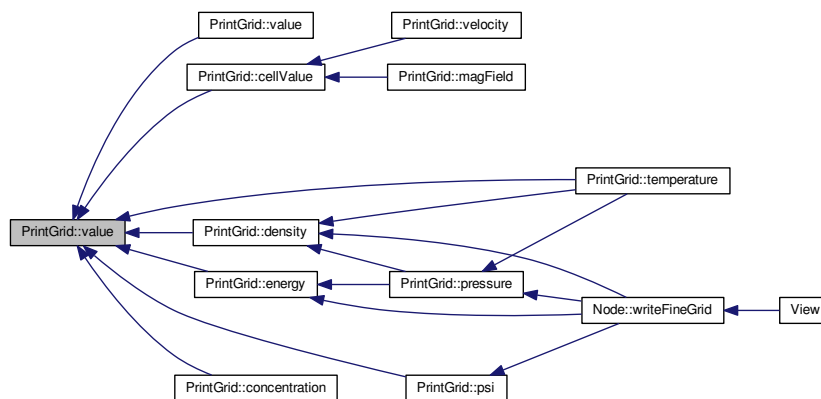
Vector

```

87 {
88     // --- Local variables ---
89
90     int n = (1<<localScaleNb)+1; // n = 2^localScaleNb
91
92     return *(Q + i + n*(j + n*k));
93 }

```

Here is the caller graph for this function:



### 5.5.3.16 real PrintGrid::value ( const int *i*, const int *j*, const int *k*, const int *QuantityNo* ) const

Returns the quantity *QuantityNo* of the cell-average vector located at *i*, *j*, *k*.

## Parameters

|                   |                          |
|-------------------|--------------------------|
| <i>i</i>          | Position x               |
| <i>j</i>          | Position y               |
| <i>k</i>          | Position z               |
| <i>QuantityNo</i> | Number of MHD variables. |

## Returns

real

```

100 {
101     // --- Local variables ---
102
103     int n = (1<<localScaleNb)+1; // n = 2^localScaleNb
104
105     return (Q + i + n*(j + n*k))->value(QuantityNo);
106 }

```



5.5.3.17 `real PrintGrid::velocity ( const int i, const int j, const int k, const int AxisNo ) const` `[inline]`

Returns the *AxisNo*-th component of the cell-average velocity located at *i*, *j*, *k*.

## Parameters

|               |                   |
|---------------|-------------------|
| <i>i</i>      | Position x        |
| <i>j</i>      | Position y        |
| <i>k</i>      | Position z        |
| <i>AxisNo</i> | Axis of interest. |

## Returns

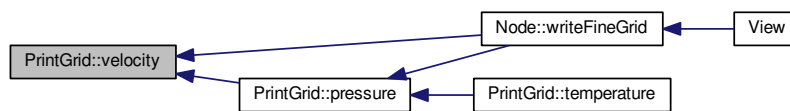
real

```

338 {
339     return cellValue(i, j, k, AxisNo+1)/cellValue(i, j, k, 1);
340 }

```

Here is the caller graph for this function:

5.5.3.18 Vector PrintGrid::velocity ( const int *i*, const int *j*, const int *k* ) const

Returns the cell-average velocity located at *i*, *j*, *k*.

## Parameters

|          |            |
|----------|------------|
| <i>i</i> | Position x |
| <i>j</i> | Position y |
| <i>k</i> | Position z |

## Returns

Vector

```

155 {
156     Vector V(3);
157
158     for (int AxisNo=1; AxisNo <= 3; AxisNo++)
159         V.setValue( AxisNo, cellValue(i, j, k, AxisNo+1)/cellValue(i, j, k, 1));
160
161     return V;
162 }

```

5.5.3.19 real PrintGrid::vorticity ( const int *i*, const int *j*, const int *k* ) const

Returns 0 in 1D, the scalar vorticity in 2D, the norm of the cell-average vorticity in 3D, located at *i*, *j*, *k*. Does not work for MHD!

## Parameters

|          |            |
|----------|------------|
| <i>i</i> | Position x |
| <i>j</i> | Position y |
| <i>k</i> | Position z |

## Returns

real

 $dx + 0.5 \cdot \text{abs}(\text{By}_1) / dy + 1.120e-13;$ 

```

295 {
296 /*
297   int L=localScaleNb;
298   real dx=0., dy=0., dz=0.;
299   real U=0.,V=0.,W=0.;
300   real Uy1=0., Uy2=0., Uz1=0., Uz2=0.;
301   real Vx1=0., Vx2=0., Vz1=0., Vz2=0.;
302   real Wx1=0., Wx2=0., Wy1=0., Wy2=0.;
303
304   int n = (1<<L);           // n = 2^localScaleNb
305
306   real result=0.;
307
308   if (Dimension == 1)
309     return 0.;
310
311   // Compute vorticity components
312
313   dx = (XMax[1]-XMin[1])/n;
314   dy = (XMax[2]-XMin[2])/n;
315
316   Vx1 = velocity(BC(i-1,1,n), BC(j, 2,n),BC(k,3,n),2);
317   Vx2 = velocity(BC(i+1,1,n), BC(j, 2,n),BC(k,3,n),2);
318   Uy1 = velocity(BC(i, 1,n), BC(j-1,2,n),BC(k,3,n),1);
319   Uy2 = velocity(BC(i, 1,n), BC(j+1,2,n),BC(k,3,n),1);
320
321   if (Dimension == 2)
322     W = (Vx2-Vx1)/(2.*dx) - (Uy2-Uy1)/(2.*dy);
323   else
324   {
325     dz = (XMax[3]-XMin[3])/(1<<L);
326
327     Uz1 = velocity(BC(i, 1,n), BC(j, 2,n),BC(k-1,3,n),1);
328     Uz1 = velocity(BC(i, 1,n), BC(j, 2,n),BC(k+1,3,n),1);
329
330     Vz1 = velocity(BC(i, 1,n), BC(j, 2,n),BC(k-1,3,n),2);
331     Vz1 = velocity(BC(i, 1,n), BC(j, 2,n),BC(k+1,3,n),2);
332
333     Wx1 = velocity(BC(i-1,1,n), BC(j, 2,n),BC(k, 3,n),3);
334     Wx2 = velocity(BC(i+1,1,n), BC(j, 2,n),BC(k, 3,n),3);
335
336     Wy1 = velocity(BC(i, 1,n), BC(j-1,2,n),BC(k, 3,n),3);
337     Wy2 = velocity(BC(i, 1,n), BC(j+1,2,n),BC(k, 3,n),3);
338
339     U = (Wy2-Wy1)/(2.*dy) - (Vz2-Vz1)/(2.*dz);
340     V = (Uz2-Uz1)/(2.*dz) - (Wx2-Wx1)/(2.*dx);
341     W = (Vx2-Vx1)/(2.*dx) - (Uy2-Uy1)/(2.*dy);
342
343   }
344
345   switch(Dimension)
346   {
347     case 2:
348       result = W;
349       break;
350
351     case 3:
352       result = sqrt(U*U+V*V+W*W);
353   };
354 */
355
356 // return result;
357   int L=localScaleNb;
358   int n = (1<<L);
359   real dx=0., dy=0., dz=0.;
360   real Div=0.;
361   real By1=0., By2=0., Bz1=0., Bz2=0.;
362   real Bx1=0., Bx2=0.;
363   real Bx =0., By=0.;
364   if (Dimension == 1){

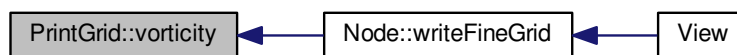
```

```

365     dx = (XMax[1]-XMin[1])/n;
366
367     Bx1 = magField(BC(i-1,1,n), BC(j,2,n),BC(k,3,n),1);
368     Bx2 = magField(BC(i+1,1,n), BC(j,2,n),BC(k,3,n),1);
369
370     Div = (Bx2-Bx1)/(2.*dx);
371
372 }else if (Dimension == 2){
373
374     dx = (XMax[1]-XMin[1])/n;
375     dy = (XMax[2]-XMin[2])/n;
376     Bx1 = magField(BC(i-1,1,n), BC(j,2,n),BC(k,3,n),1);
377     Bx2 = magField(BC(i+1,1,n), BC(j,2,n),BC(k,3,n),1);
378     By1 = magField(BC(i,1,n), BC(j-1,2,n),BC(k,3,n),2);
379     By2 = magField(BC(i,1,n), BC(j+1,2,n),BC(k,3,n),2);
380     Bx = magField(BC(i,1,n), BC(j,2,n),BC(k,3,n),1);
381     By = magField(BC(i,1,n), BC(j,2,n),BC(k,3,n),2);
382
383     //if(Bx2!=Bx1 && By2!=By1)
384     //Div = ((abs(Bx1)+abs(Bx2))/(2.*dx) + (abs(By1)+abs(By2))/(2.*dy) + 1.120e-13);
385     Div = 0.5*Abs(Bx1);
386     //else
387     //    Div = ((Bx2-Bx1)/(2.*dx) + (By2-By1)/(2.*dy));
388 }else if (Dimension == 3){
389
390     dx = (XMax[1]-XMin[1])/n;
391     dy = (XMax[2]-XMin[2])/n;
392     dz = (XMax[3]-XMin[3])/n;
393
394     Bx1 = magField(BC(i-1,1,n), BC(j,2,n),BC(k,3,n),1);
395     Bx2 = magField(BC(i+1,1,n), BC(j,2,n),BC(k,3,n),1);
396     By1 = magField(BC(i,1,n), BC(j-1,2,n),BC(k,3,n),2);
397     By2 = magField(BC(i,1,n), BC(j+1,2,n),BC(k,3,n),2);
398     Bz1 = magField(BC(i,1,n), BC(j,2,n),BC(k-1,3,n),3);
399     Bz2 = magField(BC(i,1,n), BC(j,2,n),BC(k+1,3,n),3);
400
401     Div = (Bx2-Bx1)/(2.*dx) + (By2-By1)/(2.*dy) + (Bz2-Bz1)/(2.*dz);
402
403 }
404
405 return Div;
406 }

```

Here is the caller graph for this function:



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

- [PrintGrid.h](#)
- [PrintGrid.cpp](#)

## 5.6 TimeAverageGrid Class Reference

Time Average Grid.

```
#include <TimeAverageGrid.h>
```

### Public Member Functions

- [TimeAverageGrid](#) (const int UserScaleNb, const int UserQuantityNb)  
*Constructor of [TimeAverageGrid](#) class. For a given variable number.*

- [TimeAverageGrid](#) (const int UserScaleNb)  
*Constructor of [TimeAverageGrid](#) class.*
- [~TimeAverageGrid](#) ()  
*Destructor of [TimeAverageGrid](#) clas.*
- void [updateValue](#) (const int ElementNo, const int QuantityNo, const [real](#) UserValue)  
*Update Values. For a given element.*
- void [updateValue](#) (const int i, const int j, const int k, const int QuantityNo, const [real](#) UserValue)  
*Update Values. At position i,j,k.*
- void [updateValue](#) (const int i, const int j, const int k, const [Vector](#) arg)  
*Update values.*
- void [updateSample](#) ()  
*Update number of samples.*
- [real value](#) (const int ElementNo, const int QuantityNo) const  
*Get value at the position ElementNo.*
- [real value](#) (const int i, const int j, const int k, const int QuantityNo) const  
*Get value at the position i,j,k.*
- [real density](#) (const int i, const int j, const int k) const  
*Get density at the position i,j,k.*
- [real velocity](#) (const int i, const int j, const int k, const int AxisNo) const  
*Get velocity at the position i,j,k.*
- [real stress](#) (const int i, const int j, const int k, const int No) const

### 5.6.1 Detailed Description

Time Average Grid.

### 5.6.2 Constructor & Destructor Documentation

#### 5.6.2.1 TimeAverageGrid::TimeAverageGrid ( const int *UserScaleNb*, const int *UserQuantityNb* )

Constructor of [TimeAverageGrid](#) class. For a given variable number.

##### Parameters

|                       |                 |
|-----------------------|-----------------|
| <i>UserScaleNb</i>    | Level           |
| <i>UserQuantityNb</i> | Variable number |

```

31 {
32
33     // Init SampleNb, LocalScaleNb, LocalQuantityNb, and ElementNb
34
35     SampleNb      = 0;
36     LocalScaleNb  = UserScaleNb;
37     LocalQuantityNb = UserQuantityNb;
38     ElementNb     = 1 << ( Dimension*LocalScaleNb );
39
40     // Allocate array of time-averages
41
42     // Q = new Vector[ElementNb](LocalQuantityNb); //!!!No ISO C++ compatible!!!
43     Q = new Vector[ElementNb];
44     int i;
45     for (i=0;i<ElementNb;i++) Q[i].setDimension(LocalQuantityNb);
46 }
```

#### 5.6.2.2 TimeAverageGrid::TimeAverageGrid ( const int *UserScaleNb* )

Constructor of [TimeAverageGrid](#) class.

## Parameters

| <i>UserScaleNb</i> | Level |
|--------------------|-------|
|--------------------|-------|

```

54 {
55
56 // Default quantity number (3 in 1D, 6 in 2D, 10 in 3D)
57
58     switch(Dimension)
59     {
60         case 1:
61             LocalQuantityNb = 3;
62             break;
63
64         case 2:
65             LocalQuantityNb = 6;
66             break;
67
68         case 3:
69             LocalQuantityNb = 10;
70             break;
71     };
72
73 // Init SampleNb, LocalScaleNb, LocalQuantityNb, and ElementNb
74
75     SampleNb           = 0;
76     LocalScaleNb      = UserScaleNb;
77     ElementNb         = 1 << ( Dimension*LocalScaleNb );
78
79 // Allocate array of time-averages
80
81 // Q = new Vector[ElementNb](LocalQuantityNb); !!!No ISO C++ comparable!!!
82 Q = new Vector[ElementNb];
83     int i;
84     for (i=0;i<ElementNb;i++) Q[i].setDimension(LocalQuantityNb);
85 }

```

## 5.6.2.3 TimeAverageGrid::~TimeAverageGrid ( )

Destructor of [TimeAverageGrid](#) clas.

```

95 {
96 // Deallocate array of time-averages
97
98     delete[] Q;
99 }

```

## 5.6.3 Member Function Documentation

5.6.3.1 real TimeAverageGrid::density ( const int *i*, const int *j*, const int *k* ) const [inline]

Get density at the position *i,j,k*.

## Parameters

|          |            |
|----------|------------|
| <i>i</i> | Position x |
| <i>j</i> | Position y |
| <i>k</i> | Position z |

## Returns

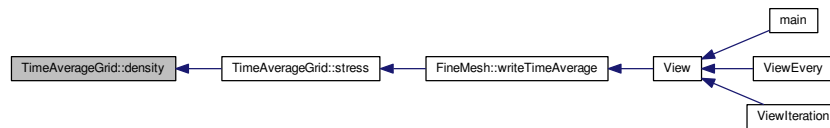
real

```

160 {
161     return value(i, j, k, l);
162 }

```

Here is the caller graph for this function:

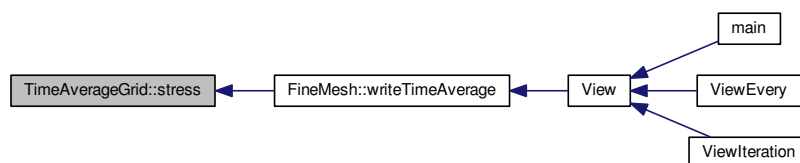


### 5.6.3.2 real TimeAverageGrid::stress ( const int *i*, const int *j*, const int *k*, const int *No* ) const

```

191 {
192     real rho = density(i,j,k);
193     real rhoV1=0., rhoV2=0., rhoV1V2=0.;
194
195     switch(No)
196     {
197     case 1:
198         rhoV1 = value(i,j,k,2); // rhoU
199         rhoV2 = value(i,j,k,2); // rhoU
200         rhoV1V2 = value(i,j,k,Dimension+2); // rhoUU
201         break;
202
203     case 2:
204         rhoV1 = value(i,j,k,2); // rhoU
205         rhoV2 = value(i,j,k,3); // rhoV
206         rhoV1V2 = value(i,j,k,Dimension+3); // rhoUV
207         break;
208
209     case 3:
210         rhoV1 = value(i,j,k,3); // rhoV
211         rhoV2 = value(i,j,k,3); // rhoV
212         rhoV1V2 = value(i,j,k,Dimension+4); // rhoVV
213         break;
214
215     case 4:
216         rhoV1 = value(i,j,k,2); // rhoU
217         rhoV2 = value(i,j,k,4); // rhoW
218         rhoV1V2 = value(i,j,k,Dimension+5); // rhoUW
219         break;
220
221     case 5:
222         rhoV1 = value(i,j,k,3); // rhoV
223         rhoV2 = value(i,j,k,4); // rhoW
224         rhoV1V2 = value(i,j,k,Dimension+6); // rhoVW
225         break;
226
227     case 6:
228         rhoV1 = value(i,j,k,4); // rhoW
229         rhoV2 = value(i,j,k,4); // rhoW
230         rhoV1V2 = value(i,j,k,Dimension+7); // rhoWW
231         break;
232     };
233
234     return ( ( rhoV1V2 - (rhoV1*rhoV2)/rho )/rho );
235
236 }
  
```

Here is the caller graph for this function:



### 5.6.3.3 void TimeAverageGrid::updateSample ( ) [inline]

Update number of samples.

#### Returns

void

```
172 {
173     SampleNb++;
174 }
```

Here is the caller graph for this function:



### 5.6.3.4 void TimeAverageGrid::updateValue ( const int *ElementNo*, const int *QuantityNo*, const real *UserValue* )

Update Values. For a given element.

#### Parameters

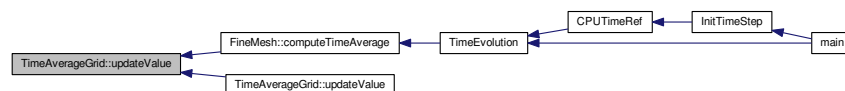
|                   |                 |
|-------------------|-----------------|
| <i>ElementNo</i>  | Element number  |
| <i>QuantityNo</i> | Variable number |
| <i>UserValue</i>  | Real value      |

#### Returns

void

```
108 {
109     real eps = 1./(SampleNb+1.);
110
111     (Q+ElementNo)->setValue(QuantityNo, eps*UserValue + (1-eps)*(Q+ElementNo)->
value(QuantityNo));
112 }
```

Here is the caller graph for this function:



### 5.6.3.5 void TimeAverageGrid::updateValue ( const int *i*, const int *j*, const int *k*, const int *QuantityNo*, const real *UserValue* )

Update Values. At position *i,j,k*.



## Parameters

|                   |                 |
|-------------------|-----------------|
| <i>i</i>          | Position x      |
| <i>j</i>          | Position y      |
| <i>k</i>          | Position z      |
| <i>QuantityNo</i> | Variable number |
| <i>UserValue</i>  | Real value      |

## Returns

void

```

122 {
123     // --- Local variables ---
124
125     int n = (1<<LocalScaleNb);
126     int ElementNo = i + n*(j + n*k);
127
128     updateValue(ElementNo, QuantityNo, UserValue);
129 }

```

5.6.3.6 void TimeAverageGrid::updateValue ( const int *i*, const int *j*, const int *k*, const Vector *arg* )

Update values.

## Parameters

|            |            |
|------------|------------|
| <i>i</i>   | Position x |
| <i>j</i>   | Position y |
| <i>k</i>   | Position z |
| <i>arg</i> | Vector     |

## Returns

void

```

138 {
139     real rho=0.;
140     real U=0., V=0., W=0.;
141
142     // Compute density and velocity
143
144     rho = arg.value(1);
145     U = arg.value(2)/rho;
146     if (Dimension > 1) V = arg.value(3)/rho;
147     if (Dimension > 2) W = arg.value(4)/rho;
148
149     // update values
150
151     updateValue(i, j, k, 1, rho);
152
153     switch(Dimension)
154     {
155     {
156         case 1:
157             updateValue(i, j, k, 2, rho*U);
158             updateValue(i, j, k, 3, rho*U*U);
159             break;
160
161         case 2:
162             updateValue(i, j, k, 2, rho*U);
163             updateValue(i, j, k, 3, rho*V);
164             updateValue(i, j, k, 4, rho*U*U);
165             updateValue(i, j, k, 5, rho*U*V);
166             updateValue(i, j, k, 6, rho*V*V);
167             break;
168
169         case 3:
170             updateValue(i, j, k, 2, rho*U);
171             updateValue(i, j, k, 3, rho*V);
172             updateValue(i, j, k, 4, rho*W);
173             updateValue(i, j, k, 5, rho*U*U);

```

```

174     updateValue(i, j, k, 6, rho*U*V);
175     updateValue(i, j, k, 7, rho*V*V);
176     updateValue(i, j, k, 8, rho*U*W);
177     updateValue(i, j, k, 9, rho*V*W);
178     updateValue(i, j, k, 10, rho*W*W);
179     break;
180 };
181
182 }

```

### 5.6.3.7 `real TimeAverageGrid::value ( const int ElementNo, const int QuantityNo ) const` `[inline]`

Get value at the position *ElementNo*.

#### Parameters

|                   |                 |
|-------------------|-----------------|
| <i>ElementNo</i>  | Element number  |
| <i>QuantityNo</i> | Variable number |

#### Returns

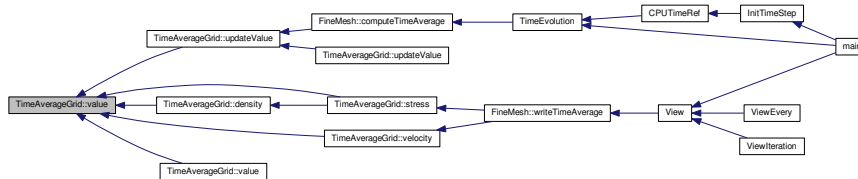
real

```

197 {
198     return (Q+ElementNo)->value(QuantityNo);
199 }

```

Here is the caller graph for this function:



### 5.6.3.8 `real TimeAverageGrid::value ( const int i, const int j, const int k, const int QuantityNo ) const` `[inline]`

Get value at the position *i,j,k*.

#### Parameters

|                   |                 |
|-------------------|-----------------|
| <i>i</i>          | Position x      |
| <i>j</i>          | Position y      |
| <i>k</i>          | Position z      |
| <i>QuantityNo</i> | Variable number |

#### Returns

real

```

209 {
210     return value(i + (1<<LocalScaleNb)*(j + (1<<LocalScaleNb)*k), QuantityNo);
211 }

```

### 5.6.3.9 `real TimeAverageGrid::velocity ( const int i, const int j, const int k, const int AxisNo ) const` `[inline]`

Get velocity at the position *i,j,k*.

## Parameters

|               |                  |
|---------------|------------------|
| <i>i</i>      | Position x       |
| <i>j</i>      | Position y       |
| <i>k</i>      | Position z       |
| <i>AxisNo</i> | Axis of interest |

## Returns

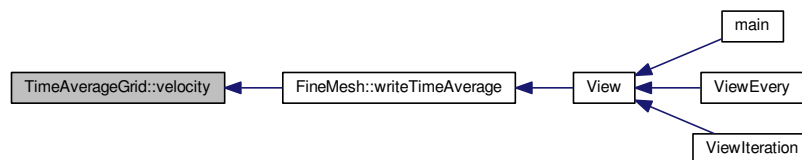
real

```

184 {
185     return value(i, j, k, AxisNo+1)/value(i, j, k, 1);
186 }

```

Here is the caller graph for this function:



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

- [TimeAverageGrid.h](#)
- [TimeAverageGrid.cpp](#)

## 5.7 Timer Class Reference

An object `Timer` gives information on the CPU time of long-time computations.

```
#include <Timer.h>
```

### Public Member Functions

- `Timer ()`  
*Constructor of `Timer` class. Initialize timer.*
- `void resetStart ()`  
*Resets time and start.*
- `void check ()`  
*Adds CPU time and real time to their buffers and resets. For long computations, it is recommended to do this operation at least once per iteration.*
- `void start ()`  
*Starts timer.*
- `double stop ()`  
*Stop timer and, if asked, returns CPU time from previous start in seconds.*
- `double CPUTime ()`  
*Returns CPU time from previous start in seconds.*
- `double realTime ()`

Returns real time from previous start in seconds.

- void `add` (double `cpuTime`, double `realTime`)  
Adds time to buffer (only when a computation is restarted).

### 5.7.1 Detailed Description

An object `Timer` gives information on the CPU time of long-time computations.

### 5.7.2 Constructor & Destructor Documentation

#### 5.7.2.1 `Timer::Timer ( )`

Constructor of `Timer` class. Initialize timer.

```
31 {
32     StartCPUtime = clock();
33     time(&StartRealTime);
34
35     sumCPUtime = 0.;
36     sumRealtime = 0.;
37     TimerOn = true;
38 }
```

### 5.7.3 Member Function Documentation

#### 5.7.3.1 `void Timer::add ( double cpuTime, double realTime )`

Adds time to buffer (only when a computation is restarted).

##### Parameters

|                 |           |
|-----------------|-----------|
| <i>cpuTime</i>  | CPU time  |
| <i>realTime</i> | Real time |

##### Returns

void

```
174 {
175     sumCPUtime=cpuTime;
176     sumRealtime=realTime;
177 }
```

#### 5.7.3.2 `void Timer::check ( )`

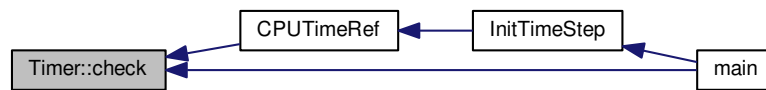
Adds CPU time and real time to their buffers and resets. For long computations, it is recommended to do this operation at least once per iteration.

##### Returns

void

```
104 {
105     // --- Local variables ---
106
107     clock_t EndCPUtime;           // end CPU time
108
109     // --- Execution ---
110
111     EndCPUtime = clock();
112     sumCPUtime += double((unsigned long int)EndCPUtime-StartCPUtime)/ (unsigned long int)CLOCKS_PER_SEC;
113     StartCPUtime = EndCPUtime;
114
115 }
```

Here is the caller graph for this function:



### 5.7.3.3 double Timer::CPUTime ( )

Returns CPU time from previous start in seconds.

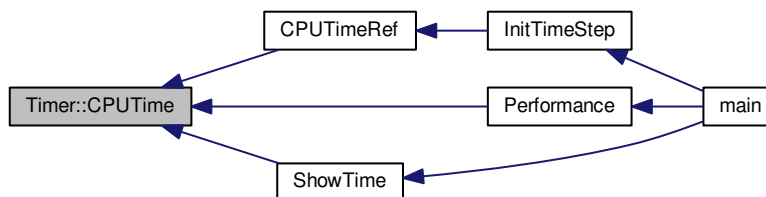
#### Returns

double

```

125 {
126     // --- Local variables ---
127     clock_t EndCPUTime;
128
129     // --- Execution ---
130     if (TimerOn)
131     {
132         // If timer is running, compute and return time in seconds
133         EndCPUTime = clock();
134         return sumCPUtime+double((unsigned long int)EndCPUTime-StartCPUTime)/ (unsigned long int)CLOCKS_PER_SEC
135     }
136     else
137         // else return time previously computed between last start and stop procedures
138         return sumCPUtime;
139 }
  
```

Here is the caller graph for this function:



### 5.7.3.4 double Timer::realTime ( )

Returns real time from previous start in seconds.

**Returns**

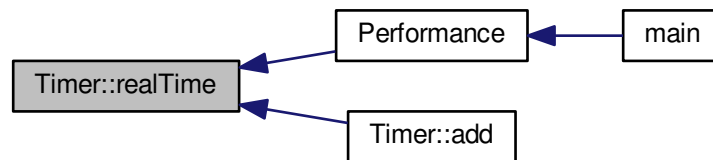
double

```

149 {
150     // --- Local variables ---
151
152     time_t EndRealTime;
153
154     // --- Execution ---
155
156     if (TimerOn)
157     {
158         // If timer is running, compute and return time in seconds
159         time(&EndRealTime);
160         return sumRealtime+difftime (EndRealTime, StartRealTime);
161     }
162     else
163         // else return time previously computed between last start and stop procedures
164         return sumRealtime;
165 }

```

Here is the caller graph for this function:

**5.7.3.5 void Timer::resetStart ( )**

Resets time and start.

**Returns**

void

```

47 {
48     StartCPUtime = clock();
49     time(&StartRealTime);
50
51     sumCPUtime = 0.;
52     sumRealtime = 0.;
53     TimerOn = true;
54 }

```

**5.7.3.6 void Timer::start ( )**

Starts timer.

## Returns

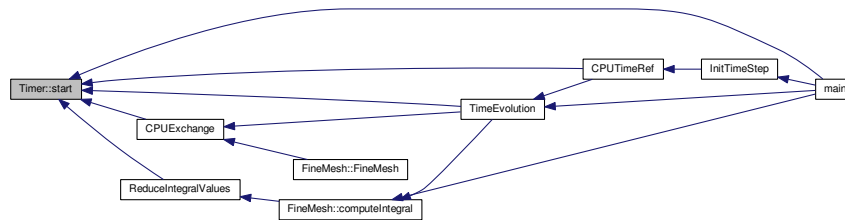
void

```

63 {
64   StartCPUTime = clock();
65   time(&StartRealTime);
66
67   TimerOn = true;
68   return;
69 }

```

Here is the caller graph for this function:



## 5.7.3.7 double Timer::stop ( )

Stop timer and, if asked, returns CPU time from previous start in seconds.

## Returns

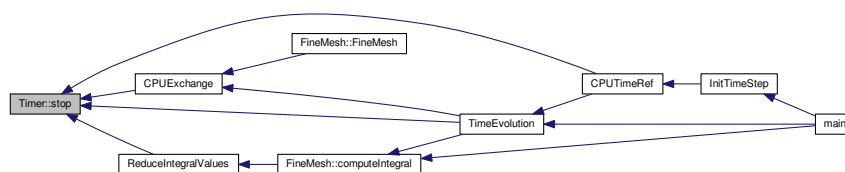
double

```

78 {
79   // --- Local variables ---
80
81   clock_t EndCPUTime;      // end CPU time
82   time_t EndRealTime;     // end real time
83
84
85   // --- Execution ---
86
87   EndCPUTime = clock();
88   time(&EndRealTime);
89
90   sumCPUTime += double((unsigned long int)EndCPUTime-StartCPUTime)/ (unsigned long int)CLOCKS_PER_SEC;
91   sumRealtime += difftime(EndRealTime, StartRealTime);
92
93   TimerOn = false;
94   return sumCPUTime;
95 }

```

Here is the caller graph for this function:



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

- [Timer.h](#)
- [Timer.cpp](#)

## 5.8 Vector Class Reference

Standard class for every vector in Carmen.

```
#include <Vector.h>
```

### Public Member Functions

- [Vector](#) ()  
*Generates a 1D vector equal to zero.*
- [Vector](#) (const int n)  
*Generates a vector of dimension n, each component being equal to zero.*
- [Vector](#) (const [real](#) x, const [real](#) y)  
*Generates the 2D vector (x,y).*
- [Vector](#) (const [real](#) x, const [real](#) y, const [real](#) z)  
*Generates the 3D vector (x,y,z).*
- [Vector](#) (const [Vector](#) &V)  
*.Generates a copy of the vector V.*
- [~Vector](#) ()  
*Destructor of [Vector](#) Class.*
- void [setValue](#) (const int n, const [real](#) a)  
*Sets the component n to value a.*
- void [setZero](#) ()  
*Sets all the components to zero.*
- void [setDimension](#) (const int n)  
*Sets the dimension of the vector to n and reset values to zero.*
- [real value](#) (const int n) const  
*Returns the value of the component n.*
- int [dimension](#) () const  
*Returns the dimension of the vector.*
- bool [operator==](#) (const [Vector](#) &V) const  
*Compares the current vector to a vector V and returns true if they are equal.*
- void [operator=](#) (const [Vector](#) &V)  
*Set the current vector to the dimension and the value of V.*
- void [operator+=](#) (const [Vector](#) &V)  
*Adds V to the current vector.*
- [Vector operator+](#) (const [Vector](#) &V) const  
*Returns the addition of the current vector and V.*
- void [operator-=](#) (const [Vector](#) &V)  
*Subtracts V to the current vector.*
- [Vector operator-](#) (const [Vector](#) &V) const  
*Returns the difference between the current vector and V.*
- [Vector operator-](#) () const  
*Returns the opposite of the current vector.*
- void [operator\\*=](#) (const [real](#) a)  
*Multiplies the current vector by a real a.*
- [Vector operator\\*](#) (const [real](#) a) const



- Returns the product of the current vector and a real  $a$ .*

  - void `operator/=` (const `real`  $a$ )

*Divides the current vector by a real  $a$ .*

  - `Vector operator/` (const `real`  $a$ ) const

*Returns the division of the current vector by a real  $a$ .*

  - `real operator*` (const `Vector` & $V$ ) const

*Returns the dot product of the current vector and  $V$ .*

  - `Vector operator|` (const `Vector` & $V$ ) const

*Returns the term-by-term product of the current vector and  $V$ .*

  - `Vector operator^` (const `Vector` & $V$ ) const

*Returns the vectorial product of the current vector and  $V$ .*

  - bool `isNaN` () const

*Returns true if one of the components of the current vector is not a number.*

## Public Attributes

- int `Columns`
- `real U` [9]

### 5.8.1 Detailed Description

Standard class for every vector in Carmen.

It contains the following data:

- the dimension of the vector *Columns* ;
- the array of reals *\*U*.

### 5.8.2 Constructor & Destructor Documentation

#### 5.8.2.1 `Vector::Vector ( )`

Generates a 1D vector equal to zero.

Example :

```
#include "Vector.h"
Vector V;
```

```
37 {
38     Columns = 1;
39     *U = 0.;
40 }
```

#### 5.8.2.2 `Vector::Vector ( const int $n$ )`

Generates a vector of dimension  $n$ , each component being equal to zero.

Example :

```
#include "Vector.h"
Vector V(4);
```

## Parameters

|     |  |
|-----|--|
| $n$ |  |
|-----|--|

```

46 {
47     Columns = n;
48
49     // If the size of the vector is 0, do not allocate memory
50     if (Columns == 0) return;
51
52     if (Columns==9) {
53         *U=0.;
54         *(U+1)=0.;
55         *(U+2)=0.;
56         *(U+3)=0.;
57         *(U+4)=0.;
58         *(U+5)=0.;
59         *(U+6)=0.;
60         *(U+7)=0.;
61         *(U+8)=0.;
62     }
63     else if (Columns==8) {
64         *U=0.;
65         *(U+1)=0.;
66         *(U+2)=0.;
67         *(U+3)=0.;
68         *(U+4)=0.;
69         *(U+5)=0.;
70         *(U+6)=0.;
71         *(U+7)=0.;
72     }
73     else if (Columns==7) {
74         *U=0.;
75         *(U+1)=0.;
76         *(U+2)=0.;
77         *(U+3)=0.;
78         *(U+4)=0.;
79         *(U+5)=0.;
80         *(U+6)=0.;
81     }
82     else if (Columns==6) {
83         *U=0.;
84         *(U+1)=0.;
85         *(U+2)=0.;
86         *(U+3)=0.;
87         *(U+4)=0.;
88         *(U+5)=0.;
89     }
90     else if (Columns==5) {
91         *U=0.;
92         *(U+1)=0.;
93         *(U+2)=0.;
94         *(U+3)=0.;
95         *(U+4)=0.;
96     }
97     else if (Columns==4) {
98         *U=0.;
99         *(U+1)=0.;
100        *(U+2)=0.;
101        *(U+3)=0.;
102    }
103    else if (Columns==3) {
104        *U=0.;
105        *(U+1)=0.;
106        *(U+2)=0.;
107    }
108    else if (Columns==2) {
109        *U=0.;
110        *(U+1)=0.;
111    }
112    else if (Columns==1) {
113        *U=0.;
114    }
115 }

```

### 5.8.2.3 Vector::Vector ( const real x, const real y )

Generates the 2D vector (x,y).

Example :

```
#include "Vector.h"
Vector V(0.,1.);
```

Parameters

|   |            |
|---|------------|
| x | Real value |
| y | Real value |

```
121 {
122     Columns = 2;
123     *U      = x;
124     *(U+1)  = y;
125 }
```

#### 5.8.2.4 Vector::Vector ( const real x, const real y, const real z )

Generates the 3D vector (x,y,z).

Example :

```
#include "Vector.h"
Vector V(0.,1.,0.);
```

Parameters

|   |            |
|---|------------|
| x | Real value |
| y | Real value |
| z | Real value |

```
131 {
132     Columns = 3;
133     *U      = x;
134     *(U+1)  = y;
135     *(U+2)  = z;
136 }
```

#### 5.8.2.5 Vector::Vector ( const Vector & V )

.Generates a copy of the vector V.

Example :

```
#include "Vector.h"
Vector V(0.,1.,0.);
Vector W(V);
```

Parameters

|   |        |
|---|--------|
| V | Vector |
|---|--------|

```
142 {
143     Columns = V.dimension();
144     if (Columns==9) {
145         *U = V.value(1);
146         *(U+1) = V.value(2);
147         *(U+2) = V.value(3);
148         *(U+3) = V.value(4);
149         *(U+4) = V.value(5);
150         *(U+5) = V.value(6);
151         *(U+6) = V.value(7);
152         *(U+7) = V.value(8);
153         *(U+8) = V.value(9);
154     }
155     else if (Columns==8) {
```

```

156     *U = V.value(1);
157     *(U+1) = V.value(2);
158     *(U+2) = V.value(3);
159     *(U+3) = V.value(4);
160     *(U+4) = V.value(5);
161     *(U+5) = V.value(6);
162     *(U+6) = V.value(7);
163     *(U+7) = V.value(8);
164 }
165 else if (Columns==7) {
166     *U = V.value(1);
167     *(U+1) = V.value(2);
168     *(U+2) = V.value(3);
169     *(U+3) = V.value(4);
170     *(U+4) = V.value(5);
171     *(U+5) = V.value(6);
172     *(U+6) = V.value(7);
173 }
174 else if (Columns==6) {
175     *U = V.value(1);
176     *(U+1) = V.value(2);
177     *(U+2) = V.value(3);
178     *(U+3) = V.value(4);
179     *(U+4) = V.value(5);
180     *(U+5) = V.value(6);
181 }
182 else if (Columns==5) {
183     *U = V.value(1);
184     *(U+1) = V.value(2);
185     *(U+2) = V.value(3);
186     *(U+3) = V.value(4);
187     *(U+4) = V.value(5);
188 }
189 else if (Columns==4) {
190     *U = V.value(1);
191     *(U+1) = V.value(2);
192     *(U+2) = V.value(3);
193     *(U+3) = V.value(4);
194 }
195 else if (Columns==3) {
196     *U = V.value(1);
197     *(U+1) = V.value(2);
198     *(U+2) = V.value(3);
199 }
200 else if (Columns==2) {
201     *U = V.value(1);
202     *(U+1) = V.value(2);
203 }
204 else if (Columns==1) {
205     *U = V.value(1);
206 }
207 }

```

### 5.8.2.6 Vector::~~Vector ( )

Destructor of [Vector](#) Class.

Deallocate memory of the vector.

```

217 {
218     // If the size of the vector is equal to zero, do not deallocate memory
219     if (Columns == 0) return;
220 }

```

## 5.8.3 Member Function Documentation

### 5.8.3.1 int Vector::dimension ( ) const [inline]

Returns the dimension of the vector.

Returns

int

```

536 {
537     return Columns;
538 }

```



## Parameters

|   |            |
|---|------------|
| a | Real value |
|---|------------|

## Returns

[Vector](#)

```

1032 {
1033     Vector result (Columns);
1034
1035     if (Columns==9) {
1036         result.setValue(1, value(1) *a);
1037         result.setValue(2, value(2) *a);
1038         result.setValue(3, value(3) *a);
1039         result.setValue(4, value(4) *a);
1040         result.setValue(5, value(5) *a);
1041         result.setValue(6, value(6) *a);
1042         result.setValue(7, value(7) *a);
1043         result.setValue(8, value(8) *a);
1044         result.setValue(9, value(9) *a);
1045     }
1046     else if (Columns==8) {
1047         result.setValue(1, value(1) *a);
1048         result.setValue(2, value(2) *a);
1049         result.setValue(3, value(3) *a);
1050         result.setValue(4, value(4) *a);
1051         result.setValue(5, value(5) *a);
1052         result.setValue(6, value(6) *a);
1053         result.setValue(7, value(7) *a);
1054         result.setValue(8, value(8) *a);
1055     }
1056     else if (Columns==7) {
1057         result.setValue(1, value(1) *a);
1058         result.setValue(2, value(2) *a);
1059         result.setValue(3, value(3) *a);
1060         result.setValue(4, value(4) *a);
1061         result.setValue(5, value(5) *a);
1062         result.setValue(6, value(6) *a);
1063         result.setValue(7, value(7) *a);
1064     }
1065     else if (Columns==6) {
1066         result.setValue(1, value(1) *a);
1067         result.setValue(2, value(2) *a);
1068         result.setValue(3, value(3) *a);
1069         result.setValue(4, value(4) *a);
1070         result.setValue(5, value(5) *a);
1071         result.setValue(6, value(6) *a);
1072     }
1073     else if (Columns==5) {
1074         result.setValue(1, value(1) *a);
1075         result.setValue(2, value(2) *a);
1076         result.setValue(3, value(3) *a);
1077         result.setValue(4, value(4) *a);
1078         result.setValue(5, value(5) *a);
1079     }
1080     else if (Columns==4) {
1081         result.setValue(1, value(1) *a);
1082         result.setValue(2, value(2) *a);
1083         result.setValue(3, value(3) *a);
1084         result.setValue(4, value(4) *a);
1085     }
1086     else if (Columns==3) {
1087         result.setValue(1, value(1) *a);
1088         result.setValue(2, value(2) *a);
1089         result.setValue(3, value(3) *a);
1090     }
1091     else if (Columns==2) {
1092         result.setValue(1, value(1) *a);
1093         result.setValue(2, value(2) *a);
1094     }
1095     else if (Columns==1) {
1096         result.setValue(1, value(1) *a);
1097     }
1098     return result;
1099 }

```

### 5.8.3.4 real Vector::operator\* ( const Vector & V ) const

Returns the dot product of the current vector and *V*.

Example :

```
#include "Vector.h"
Vector V(1.,0.,0.);
Vector W(1., 2., 1.);
real x;
...
x = V*W;
```

Parameters

|   |        |
|---|--------|
| V | Vector |
|---|--------|

Returns

real

```
1280 {
1281     real result = 0.;
1282
1283     #ifdef DEBUG
1284         if ( Columns != V.dimension())
1285             {
1286                 cout << "Vector.cpp: In method `real Vector::operator*(Vector&)':\n";
1287                 cout << "Vector.cpp: vectors have different dimensions\n";
1288                 cout << "carmen: *** [Vector.o] Execution error\n";
1289                 cout << "carmen: abort execution.\n";
1290                 exit(1);
1291             }
1292     #endif
1293
1294     if (Columns==9) {
1295         result += *U * V.value(1);
1296         result += *(U+1) * V.value(2);
1297         result += *(U+2) * V.value(3);
1298         result += *(U+3) * V.value(4);
1299         result += *(U+4) * V.value(5);
1300         result += *(U+5) * V.value(6);
1301         result += *(U+6) * V.value(7);
1302         result += *(U+7) * V.value(8);
1303         result += *(U+8) * V.value(9);
1304     }
1305     else if (Columns==8) {
1306         result += *U * V.value(1);
1307         result += *(U+1) * V.value(2);
1308         result += *(U+2) * V.value(3);
1309         result += *(U+3) * V.value(4);
1310         result += *(U+4) * V.value(5);
1311         result += *(U+5) * V.value(6);
1312         result += *(U+6) * V.value(7);
1313         result += *(U+7) * V.value(8);
1314     }
1315     else if (Columns==7) {
1316         result += *U * V.value(1);
1317         result += *(U+1) * V.value(2);
1318         result += *(U+2) * V.value(3);
1319         result += *(U+3) * V.value(4);
1320         result += *(U+4) * V.value(5);
1321         result += *(U+5) * V.value(6);
1322         result += *(U+6) * V.value(7);
1323     }
1324     else if (Columns==6) {
1325         result += *U * V.value(1);
1326         result += *(U+1) * V.value(2);
1327         result += *(U+2) * V.value(3);
1328         result += *(U+3) * V.value(4);
1329         result += *(U+4) * V.value(5);
1330         result += *(U+5) * V.value(6);
1331     }
1332     else if (Columns==5) {
1333         result += *U * V.value(1);
1334         result += *(U+1) * V.value(2);
1335         result += *(U+2) * V.value(3);
1336         result += *(U+3) * V.value(4);
1337         result += *(U+4) * V.value(5);
1338     }
1339     else if (Columns==4) {
1340         result += *U * V.value(1);
```

```

1341     result += *(U+1) * V.value(2);
1342     result += *(U+2) * V.value(3);
1343     result += *(U+3) * V.value(4);
1344 }
1345 else if (Columns==3) {
1346     result += *U * V.value(1);
1347     result += *(U+1) * V.value(2);
1348     result += *(U+2) * V.value(3);
1349 }
1350 else if (Columns==2) {
1351     result += *U * V.value(1);
1352     result += *(U+1) * V.value(2);
1353 }
1354 else if (Columns==1) {
1355     result += *U * V.value(1);
1356 }
1357
1358 return result;
1359
1360 }

```

### 5.8.3.5 void Vector::operator\*=( const real a )

Multiplies the current vector by a real  $a$ .

Example :

```

#include "Vector.h"
Vector V(1.,0.,0.);
real x = 2.;
...
V *= x;

```

Parameters

|   |            |
|---|------------|
| a | Real value |
|---|------------|

Returns

void

```

962 {
963     if (Columns==9) {
964         *U *= a;
965         *(U+1) *= a;
966         *(U+2) *= a;
967         *(U+3) *= a;
968         *(U+4) *= a;
969         *(U+5) *= a;
970         *(U+6) *= a;
971         *(U+7) *= a;
972         *(U+8) *= a;
973     }
974     else if (Columns==8) {
975         *U *= a;
976         *(U+1) *= a;
977         *(U+2) *= a;
978         *(U+3) *= a;
979         *(U+4) *= a;
980         *(U+5) *= a;
981         *(U+6) *= a;
982         *(U+7) *= a;
983     }
984     else if (Columns==7) {
985         *U *= a;
986         *(U+1) *= a;
987         *(U+2) *= a;
988         *(U+3) *= a;
989         *(U+4) *= a;
990         *(U+5) *= a;
991         *(U+6) *= a;
992     }
993     else if (Columns==6) {
994         *U *= a;

```



```

995     *(U+1) *= a;
996     *(U+2) *= a;
997     *(U+3) *= a;
998     *(U+4) *= a;
999     *(U+5) *= a;
1000 }
1001 else if (Columns==5) {
1002     *U *= a;
1003     *(U+1) *= a;
1004     *(U+2) *= a;
1005     *(U+3) *= a;
1006     *(U+4) *= a;
1007 }
1008 else if (Columns==4) {
1009     *U *= a;
1010     *(U+1) *= a;
1011     *(U+2) *= a;
1012     *(U+3) *= a;
1013 }
1014 else if (Columns==3) {
1015     *U *= a;
1016     *(U+1) *= a;
1017     *(U+2) *= a;
1018 }
1019 else if (Columns==2) {
1020     *U *= a;
1021     *(U+1) *= a;
1022 }
1023 else if (Columns==1) {
1024     *U *= a;
1025 }
1026 }

```

### 5.8.3.6 Vector Vector::operator+ ( const Vector & V ) const

Returns the addition of the current vector and *V*.

Example :

```

#include "Vector.h"
Vector V(1.,0.,0.);
Vector W(0.,-1.,2.);
Vector U;
...
U = V + W;

```

Parameters

|          |                        |
|----------|------------------------|
| <i>V</i> | <a href="#">Vector</a> |
|----------|------------------------|

Returns

[Vector](#)

```

625 {
626     Vector result(Columns);
627
628     #ifdef DEBUG
629     if ( Columns != V.dimension() )
630     {
631         cout << "Vector.cpp: In method 'Vector& Vector::operator+(Vector&)':\n";
632         cout << "Vector.cpp: vectors have different dimensions\n";
633         cout << "carmen: *** [Vector.o] Execution error\n";
634         cout << "carmen: abort execution.\n";
635         exit(1);
636     }
637     #endif
638
639     if (Columns==9) {
640         result.setValue(1, value(1) + V.value(1));
641         result.setValue(2, value(2) + V.value(2));
642         result.setValue(3, value(3) + V.value(3));
643         result.setValue(4, value(4) + V.value(4));

```

```

644     result.setValue(5, value(5) + V.value(5));
645     result.setValue(6, value(6) + V.value(6));
646     result.setValue(7, value(7) + V.value(7));
647     result.setValue(8, value(8) + V.value(8));
648     result.setValue(9, value(9) + V.value(9));
649 }
650 else if (Columns==8) {
651     result.setValue(1, value(1) + V.value(1));
652     result.setValue(2, value(2) + V.value(2));
653     result.setValue(3, value(3) + V.value(3));
654     result.setValue(4, value(4) + V.value(4));
655     result.setValue(5, value(5) + V.value(5));
656     result.setValue(6, value(6) + V.value(6));
657     result.setValue(7, value(7) + V.value(7));
658     result.setValue(8, value(8) + V.value(8));
659 }
660 else if (Columns==7) {
661     result.setValue(1, value(1) + V.value(1));
662     result.setValue(2, value(2) + V.value(2));
663     result.setValue(3, value(3) + V.value(3));
664     result.setValue(4, value(4) + V.value(4));
665     result.setValue(5, value(5) + V.value(5));
666     result.setValue(6, value(6) + V.value(6));
667     result.setValue(7, value(7) + V.value(7));
668 }
669 else if (Columns==6) {
670     result.setValue(1, value(1) + V.value(1));
671     result.setValue(2, value(2) + V.value(2));
672     result.setValue(3, value(3) + V.value(3));
673     result.setValue(4, value(4) + V.value(4));
674     result.setValue(5, value(5) + V.value(5));
675     result.setValue(6, value(6) + V.value(6));
676 }
677 else if (Columns==5) {
678     result.setValue(1, value(1) + V.value(1));
679     result.setValue(2, value(2) + V.value(2));
680     result.setValue(3, value(3) + V.value(3));
681     result.setValue(4, value(4) + V.value(4));
682     result.setValue(5, value(5) + V.value(5));
683 }
684 else if (Columns==4) {
685     result.setValue(1, value(1) + V.value(1));
686     result.setValue(2, value(2) + V.value(2));
687     result.setValue(3, value(3) + V.value(3));
688     result.setValue(4, value(4) + V.value(4));
689 }
690 else if (Columns==3) {
691     result.setValue(1, value(1) + V.value(1));
692     result.setValue(2, value(2) + V.value(2));
693     result.setValue(3, value(3) + V.value(3));
694 }
695 else if (Columns==2) {
696     result.setValue(1, value(1) + V.value(1));
697     result.setValue(2, value(2) + V.value(2));
698 }
699 else if (Columns==1) {
700     result.setValue(1, value(1) + V.value(1));
701 }
702 return result;
703
704 }

```

### 5.8.3.7 void Vector::operator+=( const Vector & V )

Adds *V* to the current vector.

Example :

```

#include "Vector.h"
Vector V(1.,0.,0.);
Vector W(0.,-1.,2.);
...
W += V;

```

## Parameters

|   |        |
|---|--------|
| V | Vector |
|---|--------|

## Returns

void

```

542 {
543 #ifdef DEBUG
544     if ( Columns != V.dimension())
545     {
546         cout << "Vector.cpp: In method `void Vector::operator+=(Vector&)':\n";
547         cout << "Vector.cpp: vectors have different dimensions\n";
548         cout << "carmen: *** [Vector.o] Execution error\n";
549         cout << "carmen: abort execution.\n";
550         exit(1);
551     }
552 #endif
553
554     if (Columns==9) {
555         *U += V.value(1);
556         *(U+1) += V.value(2);
557         *(U+2) += V.value(3);
558         *(U+3) += V.value(4);
559         *(U+4) += V.value(5);
560         *(U+5) += V.value(6);
561         *(U+6) += V.value(7);
562         *(U+7) += V.value(8);
563         *(U+8) += V.value(9);
564     }
565     else if (Columns==8) {
566         *U += V.value(1);
567         *(U+1) += V.value(2);
568         *(U+2) += V.value(3);
569         *(U+3) += V.value(4);
570         *(U+4) += V.value(5);
571         *(U+5) += V.value(6);
572         *(U+6) += V.value(7);
573         *(U+7) += V.value(8);
574     }
575     else if (Columns==7) {
576         *U += V.value(1);
577         *(U+1) += V.value(2);
578         *(U+2) += V.value(3);
579         *(U+3) += V.value(4);
580         *(U+4) += V.value(5);
581         *(U+5) += V.value(6);
582         *(U+6) += V.value(7);
583     }
584     else if (Columns==6) {
585         *U += V.value(1);
586         *(U+1) += V.value(2);
587         *(U+2) += V.value(3);
588         *(U+3) += V.value(4);
589         *(U+4) += V.value(5);
590         *(U+5) += V.value(6);
591     }
592     else if (Columns==5) {
593         *U += V.value(1);
594         *(U+1) += V.value(2);
595         *(U+2) += V.value(3);
596         *(U+3) += V.value(4);
597         *(U+4) += V.value(5);
598     }
599     else if (Columns==4) {
600         *U += V.value(1);
601         *(U+1) += V.value(2);
602         *(U+2) += V.value(3);
603         *(U+3) += V.value(4);
604     }
605     else if (Columns==3) {
606         *U += V.value(1);
607         *(U+1) += V.value(2);
608         *(U+2) += V.value(3);
609     }
610     else if (Columns==2) {
611         *U += V.value(1);
612         *(U+1) += V.value(2);
613     }
614     else if (Columns==1) {
615         *U += V.value(1);
616     }
617 }

```

### 5.8.3.8 Vector Vector::operator-( const Vector & V ) const

Returns the difference between the current vector and *V*.

Example :

```
#include "Vector.h"
Vector V(1.,0.,0.);
Vector W(0.,-1.,2.);
Vector U;
...
U = V - W;
```

Parameters

|   |        |
|---|--------|
| V | Vector |
|---|--------|

Returns

Vector

```
798 {
799     Vector result (Columns);
800
801 #ifdef DEBUG
802     if ( Columns != V.dimension() )
803     {
804         cout << "Vector.cpp: In method 'Vector& Vector::operator-(Vector&)':\n";
805         cout << "Vector.cpp: vectors have different dimensions\n";
806         cout << "carmen: *** [Vector.o] Execution error\n";
807         cout << "carmen: abort execution.\n";
808         exit(1);
809     }
810 #endif
811
812     if (Columns==9) {
813         result.setValue(1, value(1) - V.value(1));
814         result.setValue(2, value(2) - V.value(2));
815         result.setValue(3, value(3) - V.value(3));
816         result.setValue(4, value(4) - V.value(4));
817         result.setValue(5, value(5) - V.value(5));
818         result.setValue(6, value(6) - V.value(6));
819         result.setValue(7, value(7) - V.value(7));
820         result.setValue(8, value(8) - V.value(8));
821         result.setValue(9, value(9) - V.value(9));
822     }
823     else if (Columns==8) {
824         result.setValue(1, value(1) - V.value(1));
825         result.setValue(2, value(2) - V.value(2));
826         result.setValue(3, value(3) - V.value(3));
827         result.setValue(4, value(4) - V.value(4));
828         result.setValue(5, value(5) - V.value(5));
829         result.setValue(6, value(6) - V.value(6));
830         result.setValue(7, value(7) - V.value(7));
831         result.setValue(8, value(8) - V.value(8));
832     }
833     else if (Columns==7) {
834         result.setValue(1, value(1) - V.value(1));
835         result.setValue(2, value(2) - V.value(2));
836         result.setValue(3, value(3) - V.value(3));
837         result.setValue(4, value(4) - V.value(4));
838         result.setValue(5, value(5) - V.value(5));
839         result.setValue(6, value(6) - V.value(6));
840         result.setValue(7, value(7) - V.value(7));
841     }
842     else if (Columns==6) {
843         result.setValue(1, value(1) - V.value(1));
844         result.setValue(2, value(2) - V.value(2));
845         result.setValue(3, value(3) - V.value(3));
846         result.setValue(4, value(4) - V.value(4));
847         result.setValue(5, value(5) - V.value(5));
848         result.setValue(6, value(6) - V.value(6));
849     }
850     else if (Columns==5) {
851         result.setValue(1, value(1) - V.value(1));
852         result.setValue(2, value(2) - V.value(2));
853         result.setValue(3, value(3) - V.value(3));
```

```

854     result.setValue(4, value(4) - V.value(4));
855     result.setValue(5, value(5) - V.value(5));
856 }
857 else if (Columns==4) {
858     result.setValue(1, value(1) - V.value(1));
859     result.setValue(2, value(2) - V.value(2));
860     result.setValue(3, value(3) - V.value(3));
861     result.setValue(4, value(4) - V.value(4));
862 }
863 else if (Columns==3) {
864     result.setValue(1, value(1) - V.value(1));
865     result.setValue(2, value(2) - V.value(2));
866     result.setValue(3, value(3) - V.value(3));
867 }
868 else if (Columns==2) {
869     result.setValue(1, value(1) - V.value(1));
870     result.setValue(2, value(2) - V.value(2));
871 }
872 else if (Columns==1) {
873     result.setValue(1, value(1) - V.value(1));
874 }
875 return result;
876
877 }

```

### 5.8.3.9 Vector Vector::operator-( ) const

Returns the opposite of the current vector.

Example

```

#include "Vector.h"
Vector V(1.,0.,0.);
Vector W;
...
W = -V;

```

Returns

Vector

```

885 {
886     Vector result(Columns);
887
888     if (Columns==9) {
889         result.setValue(1,-*U);
890         result.setValue(2,-*(U+1));
891         result.setValue(3,-*(U+2));
892         result.setValue(4,-*(U+3));
893         result.setValue(5,-*(U+4));
894         result.setValue(6,-*(U+5));
895         result.setValue(7,-*(U+6));
896         result.setValue(8,-*(U+7));
897         result.setValue(9,-*(U+8));
898     }
899     else if (Columns==8) {
900         result.setValue(1,-*U);
901         result.setValue(2,-*(U+1));
902         result.setValue(3,-*(U+2));
903         result.setValue(4,-*(U+3));
904         result.setValue(5,-*(U+4));
905         result.setValue(6,-*(U+5));
906         result.setValue(7,-*(U+6));
907         result.setValue(8,-*(U+7));
908     }
909     else if (Columns==7) {
910         result.setValue(1,-*U);
911         result.setValue(2,-*(U+1));
912         result.setValue(3,-*(U+2));
913         result.setValue(4,-*(U+3));
914         result.setValue(5,-*(U+4));
915         result.setValue(6,-*(U+5));
916         result.setValue(7,-*(U+6));
917     }
918     else if (Columns==6) {
919         result.setValue(1,-*U);

```

```

920     result.setValue(2,-*(U+1));
921     result.setValue(3,-*(U+2));
922     result.setValue(4,-*(U+3));
923     result.setValue(5,-*(U+4));
924     result.setValue(6,-*(U+5));
925 }
926 else if (Columns==5) {
927     result.setValue(1,-*U);
928     result.setValue(2,-*(U+1));
929     result.setValue(3,-*(U+2));
930     result.setValue(4,-*(U+3));
931     result.setValue(5,-*(U+4));
932 }
933 else if (Columns==4) {
934     result.setValue(1,-*U);
935     result.setValue(2,-*(U+1));
936     result.setValue(3,-*(U+2));
937     result.setValue(4,-*(U+3));
938 }
939 else if (Columns==3) {
940     result.setValue(1,-*U);
941     result.setValue(2,-*(U+1));
942     result.setValue(3,-*(U+2));
943 }
944 else if (Columns==2) {
945     result.setValue(1,-*U);
946     result.setValue(2,-*(U+1));
947 }
948 else if (Columns==1) {
949     result.setValue(1,-*U);
950 }
951 return result;
952 }

```

### 5.8.3.10 void Vector::operator-= ( const Vector & V )

Subtracts  $V$  to the current vector.

Example :

```

#include "Vector.h"
Vector V(1.,0.,0.);
Vector W(0.,-1.,2.);
...
W -= V;

```

Parameters

|     |                        |
|-----|------------------------|
| $V$ | <a href="#">Vector</a> |
|-----|------------------------|

Returns

void

```

715 {
716 #ifndef DEBUG
717     if ( Columns != V.dimension() )
718     {
719         cout << "Vector.cpp: In method 'void Vector::operator-=(Vector&)':\n";
720         cout << "Vector.cpp: vectors have different dimensions\n";
721         cout << "carmen: *** [Vector.o] Execution error\n";
722         cout << "carmen: abort execution.\n";
723         exit(1);
724     }
725 #endif
726
727     if (Columns==9) {
728         *U -= V.value(1);
729         *(U+1) -= V.value(2);
730         *(U+2) -= V.value(3);
731         *(U+3) -= V.value(4);
732         *(U+4) -= V.value(5);
733         *(U+5) -= V.value(6);
734         *(U+6) -= V.value(7);

```

```

735     *(U+7) -= V.value(8);
736     *(U+8) -= V.value(9);
737 }
738 else if (Columns==8) {
739     *U -= V.value(1);
740     *(U+1) -= V.value(2);
741     *(U+2) -= V.value(3);
742     *(U+3) -= V.value(4);
743     *(U+4) -= V.value(5);
744     *(U+5) -= V.value(6);
745     *(U+6) -= V.value(7);
746     *(U+7) -= V.value(8);
747 }
748 else if (Columns==7) {
749     *U -= V.value(1);
750     *(U+1) -= V.value(2);
751     *(U+2) -= V.value(3);
752     *(U+3) -= V.value(4);
753     *(U+4) -= V.value(5);
754     *(U+5) -= V.value(6);
755     *(U+6) -= V.value(7);
756 }
757 else if (Columns==6) {
758     *U -= V.value(1);
759     *(U+1) -= V.value(2);
760     *(U+2) -= V.value(3);
761     *(U+3) -= V.value(4);
762     *(U+4) -= V.value(5);
763     *(U+5) -= V.value(6);
764 }
765 else if (Columns==5) {
766     *U -= V.value(1);
767     *(U+1) -= V.value(2);
768     *(U+2) -= V.value(3);
769     *(U+3) -= V.value(4);
770     *(U+4) -= V.value(5);
771 }
772 else if (Columns==4) {
773     *U -= V.value(1);
774     *(U+1) -= V.value(2);
775     *(U+2) -= V.value(3);
776     *(U+3) -= V.value(4);
777 }
778 else if (Columns==3) {
779     *U -= V.value(1);
780     *(U+1) -= V.value(2);
781     *(U+2) -= V.value(3);
782 }
783 else if (Columns==2) {
784     *U -= V.value(1);
785     *(U+1) -= V.value(2);
786 }
787 else if (Columns==1) {
788     *U -= V.value(1);
789 }
790 }

```

### 5.8.3.11 Vector Vector::operator/( const real a ) const

Returns the division of the current vector by a real  $a$ .

Example :

```

#include "Vector.h"
Vector V(1.,0.,0.);
Vector W;
real x = 2.;
...
W = V / x;

```

## Parameters

|   |            |
|---|------------|
| a | Real value |
|---|------------|

## Returns

## Vector

```

1192 {
1193     Vector result(Columns);
1194
1195 #ifdef DEBUG
1196     if ( a == 0.)
1197     {
1198         cout << "Vector.cpp: In method `void Vector::operator/(real)`: \n";
1199         cout << "Vector.cpp: division by zero \n";
1200         cout << "carmen: *** [Vector.o] Execution error \n";
1201         cout << "carmen: abort execution. \n";
1202         exit(1);
1203     }
1204 #endif
1205
1206     if (Columns==9) {
1207         result.setValue(1, value(1) /a);
1208         result.setValue(2, value(2) /a);
1209         result.setValue(3, value(3) /a);
1210         result.setValue(4, value(4) /a);
1211         result.setValue(5, value(5) /a);
1212         result.setValue(6, value(6) /a);
1213         result.setValue(7, value(7) /a);
1214         result.setValue(8, value(8) /a);
1215         result.setValue(9, value(9) /a);
1216     }
1217     else if (Columns==8) {
1218         result.setValue(1, value(1) /a);
1219         result.setValue(2, value(2) /a);
1220         result.setValue(3, value(3) /a);
1221         result.setValue(4, value(4) /a);
1222         result.setValue(5, value(5) /a);
1223         result.setValue(6, value(6) /a);
1224         result.setValue(7, value(7) /a);
1225         result.setValue(8, value(8) /a);
1226     }
1227     else if (Columns==7) {
1228         result.setValue(1, value(1) /a);
1229         result.setValue(2, value(2) /a);
1230         result.setValue(3, value(3) /a);
1231         result.setValue(4, value(4) /a);
1232         result.setValue(5, value(5) /a);
1233         result.setValue(6, value(6) /a);
1234         result.setValue(7, value(7) /a);
1235     }
1236     else if (Columns==6) {
1237         result.setValue(1, value(1) /a);
1238         result.setValue(2, value(2) /a);
1239         result.setValue(3, value(3) /a);
1240         result.setValue(4, value(4) /a);
1241         result.setValue(5, value(5) /a);
1242         result.setValue(6, value(6) /a);
1243     }
1244     else if (Columns==5) {
1245         result.setValue(1, value(1) /a);
1246         result.setValue(2, value(2) /a);
1247         result.setValue(3, value(3) /a);
1248         result.setValue(4, value(4) /a);
1249         result.setValue(5, value(5) /a);
1250     }
1251     else if (Columns==4) {
1252         result.setValue(1, value(1) /a);
1253         result.setValue(2, value(2) /a);
1254         result.setValue(3, value(3) /a);
1255         result.setValue(4, value(4) /a);
1256     }
1257     else if (Columns==3) {
1258         result.setValue(1, value(1) /a);
1259         result.setValue(2, value(2) /a);
1260         result.setValue(3, value(3) /a);
1261     }
1262     else if (Columns==2) {
1263         result.setValue(1, value(1) /a);
1264         result.setValue(2, value(2) /a);
1265     }
1266     else if (Columns==1) {
1267         result.setValue(1, value(1) /a);
1268     }

```



```

1269     return result;
1270 }

```

### 5.8.3.12 void Vector::operator/= ( const real a )

Divides the current vector by a real *a*.

Example :

```
#include "Vector.h"
```

```
Vector V(1.,0.,0.);
```

```
real x = 2.;
```

```
...
```

```
V /= x;
```

Parameters

|   |            |
|---|------------|
| a | Real value |
|---|------------|

Returns

void

```

1109 {
1110 #ifdef DEBUG
1111     if ( a == 0.)
1112     {
1113         cout << "Vector.cpp: In method `void Vector::operator/=(real)`: \n";
1114         cout << "Vector.cpp: division by zero \n";
1115         cout << "carmen: *** [Vector.o] Execution error \n";
1116         cout << "carmen: abort execution. \n";
1117         exit(1);
1118     }
1119 #endif
1120
1121     if (Columns==9) {
1122         *U /= a;
1123         *(U+1) /= a;
1124         *(U+2) /= a;
1125         *(U+3) /= a;
1126         *(U+4) /= a;
1127         *(U+5) /= a;
1128         *(U+6) /= a;
1129         *(U+7) /= a;
1130         *(U+8) /= a;
1131     }
1132     else if (Columns==8) {
1133         *U /= a;
1134         *(U+1) /= a;
1135         *(U+2) /= a;
1136         *(U+3) /= a;
1137         *(U+4) /= a;
1138         *(U+5) /= a;
1139         *(U+6) /= a;
1140         *(U+7) /= a;
1141     }
1142     else if (Columns==7) {
1143         *U /= a;
1144         *(U+1) /= a;
1145         *(U+2) /= a;
1146         *(U+3) /= a;
1147         *(U+4) /= a;
1148         *(U+5) /= a;
1149         *(U+6) /= a;
1150     }
1151     else if (Columns==6) {
1152         *U /= a;
1153         *(U+1) /= a;
1154         *(U+2) /= a;
1155         *(U+3) /= a;
1156         *(U+4) /= a;
1157         *(U+5) /= a;
1158     }
1159     else if (Columns==5) {

```

```

1160     *U /= a;
1161     *(U+1) /= a;
1162     *(U+2) /= a;
1163     *(U+3) /= a;
1164     *(U+4) /= a;
1165 }
1166 else if (Columns==4) {
1167     *U /= a;
1168     *(U+1) /= a;
1169     *(U+2) /= a;
1170     *(U+3) /= a;
1171 }
1172 else if (Columns==3) {
1173     *U /= a;
1174     *(U+1) /= a;
1175     *(U+2) /= a;
1176 }
1177 else if (Columns==2) {
1178     *U /= a;
1179     *(U+1) /= a;
1180 }
1181 else if (Columns==1) {
1182     *U /= a;
1183 }
1184 }

```

### 5.8.3.13 void Vector::operator=( const Vector & V )

Set the current vector to the dimension and the value of V.

Example :

```

#include "Vector.h"
Vector V(1.,0.,0.);
Vector W;
...
W = V;

```

Parameters

|   |        |
|---|--------|
| V | Vector |
|---|--------|

Returns

void

```

462 {
463     if (V.dimension() != Columns)
464     {
465         Columns = V.dimension();
466     }
467
468     if (Columns==9) {
469         *U = V.value(1);
470         *(U+1) = V.value(2);
471         *(U+2) = V.value(3);
472         *(U+3) = V.value(4);
473         *(U+4) = V.value(5);
474         *(U+5) = V.value(6);
475         *(U+6) = V.value(7);
476         *(U+7) = V.value(8);
477         *(U+8) = V.value(9);
478     }
479     else if (Columns==8) {
480         *U = V.value(1);
481         *(U+1) = V.value(2);
482         *(U+2) = V.value(3);
483         *(U+3) = V.value(4);
484         *(U+4) = V.value(5);
485         *(U+5) = V.value(6);
486         *(U+6) = V.value(7);
487         *(U+7) = V.value(8);
488     }
489     else if (Columns==7) {

```

```

490     *U = V.value(1);
491     *(U+1) = V.value(2);
492     *(U+2) = V.value(3);
493     *(U+3) = V.value(4);
494     *(U+4) = V.value(5);
495     *(U+5) = V.value(6);
496     *(U+6) = V.value(7);
497 }
498 else if (Columns==6) {
499     *U = V.value(1);
500     *(U+1) = V.value(2);
501     *(U+2) = V.value(3);
502     *(U+3) = V.value(4);
503     *(U+4) = V.value(5);
504     *(U+5) = V.value(6);
505 }
506 else if (Columns==5) {
507     *U = V.value(1);
508     *(U+1) = V.value(2);
509     *(U+2) = V.value(3);
510     *(U+3) = V.value(4);
511     *(U+4) = V.value(5);
512 }
513 else if (Columns==4) {
514     *U = V.value(1);
515     *(U+1) = V.value(2);
516     *(U+2) = V.value(3);
517     *(U+3) = V.value(4);
518 }
519 else if (Columns==3) {
520     *U = V.value(1);
521     *(U+1) = V.value(2);
522     *(U+2) = V.value(3);
523 }
524 else if (Columns==2) {
525     *U = V.value(1);
526     *(U+1) = V.value(2);
527 }
528 else if (Columns==1) {
529     *U = V.value(1);
530 }
531 }

```

### 5.8.3.14 bool Vector::operator==( const Vector & V ) const

Compares the current vector to a vector *V* and returns true if they are equal.

Example :

```

#include "Vector.h"

Vector V(2);
Vector W(2);

real x;

...

if (V == W)

x = V.value(1);

```

Parameters

|   |                        |
|---|------------------------|
| V | <a href="#">Vector</a> |
|---|------------------------|

Returns

bool

```

374 {
375 #ifdef DEBUG
376     if ( Columns != V.dimension() )
377     {
378         cout << "Vector.cpp: In method `bool Vector::operator==(Vector&)':\n";
379         cout << "Vector.cpp: vectors have different dimensions\n";
380         cout << "carmen: *** [Vector.o] Execution error\n";

```

```

381         cout << "carmen: abort execution.\n";
382         exit(1);
383     }
384 #endif
385
386     if (Columns==9) {
387         if ( *U != V.value(1) ) return false;
388         if ( *(U+1) != V.value(2) ) return false;
389         if ( *(U+2) != V.value(3) ) return false;
390         if ( *(U+3) != V.value(4) ) return false;
391         if ( *(U+4) != V.value(5) ) return false;
392         if ( *(U+5) != V.value(6) ) return false;
393         if ( *(U+6) != V.value(7) ) return false;
394         if ( *(U+7) != V.value(8) ) return false;
395         if ( *(U+8) != V.value(9) ) return false;
396     }
397     else if (Columns==8) {
398         if ( *U != V.value(1) ) return false;
399         if ( *(U+1) != V.value(2) ) return false;
400         if ( *(U+2) != V.value(3) ) return false;
401         if ( *(U+3) != V.value(4) ) return false;
402         if ( *(U+4) != V.value(5) ) return false;
403         if ( *(U+5) != V.value(6) ) return false;
404         if ( *(U+6) != V.value(7) ) return false;
405         if ( *(U+7) != V.value(8) ) return false;
406     }
407     else if (Columns==7) {
408         if ( *U != V.value(1) ) return false;
409         if ( *(U+1) != V.value(2) ) return false;
410         if ( *(U+2) != V.value(3) ) return false;
411         if ( *(U+3) != V.value(4) ) return false;
412         if ( *(U+4) != V.value(5) ) return false;
413         if ( *(U+5) != V.value(6) ) return false;
414         if ( *(U+6) != V.value(7) ) return false;
415     }
416     else if (Columns==6) {
417         if ( *U != V.value(1) ) return false;
418         if ( *(U+1) != V.value(2) ) return false;
419         if ( *(U+2) != V.value(3) ) return false;
420         if ( *(U+3) != V.value(4) ) return false;
421         if ( *(U+4) != V.value(5) ) return false;
422         if ( *(U+5) != V.value(6) ) return false;
423     }
424     else if (Columns==5) {
425         if ( *U != V.value(1) ) return false;
426         if ( *(U+1) != V.value(2) ) return false;
427         if ( *(U+2) != V.value(3) ) return false;
428         if ( *(U+3) != V.value(4) ) return false;
429         if ( *(U+4) != V.value(5) ) return false;
430     }
431     else if (Columns==4) {
432         if ( *U != V.value(1) ) return false;
433         if ( *(U+1) != V.value(2) ) return false;
434         if ( *(U+2) != V.value(3) ) return false;
435         if ( *(U+3) != V.value(4) ) return false;
436     }
437     else if (Columns==3) {
438         if ( *U != V.value(1) ) return false;
439         if ( *(U+1) != V.value(2) ) return false;
440         if ( *(U+2) != V.value(3) ) return false;
441     }
442     else if (Columns==2) {
443         if ( *U != V.value(1) ) return false;
444         if ( *(U+1) != V.value(2) ) return false;
445     }
446     else if (Columns==1) {
447         if ( *U != V.value(1) ) return false;
448     }
449     return true;
450 }
451 }

```

### 5.8.3.15 Vector Vector::operator^ ( const Vector & V ) const

Returns the vectorial product of the current vector and  $V$ .

## Parameters

|   |        |
|---|--------|
| V | Vector |
|---|--------|

## Returns

Vector

```

1402 {
1403     Vector result(3);
1404
1405     #ifdef DEBUG
1406         if ( Columns != V.dimension())
1407             {
1408                 cout << "Vector.cpp: In method `real Vector::operator^(Vector&)`: \n";
1409                 cout << "Vector.cpp: vectors have different dimensions \n";
1410                 cout << "carmen: *** [Vector.o] Execution error \n";
1411                 cout << "carmen: abort execution. \n";
1412                 exit(1);
1413             }
1414     #endif
1415
1416     if (Columns > 1)
1417         {
1418             result.setValue(3, value(1)*V.value(2)-value(2)*V.value(1));
1419             if (Columns > 2)
1420                 {
1421                     result.setValue(1, value(2)*V.value(3)-value(3)*V.
value(2));
1422                     result.setValue(2, value(3)*V.value(1)-value(1)*V.
value(3));
1423                 }
1424         }
1425
1426     return result;
1427
1428 }

```

## 5.8.3.16 Vector Vector::operator|( const Vector &amp; V ) const

Returns the term-by-term product of the current vector and V.

## Parameters

|   |        |
|---|--------|
| V | Vector |
|---|--------|

## Returns

Vector

```

1371 {
1372     int n;
1373     Vector result(Columns);
1374
1375     #ifdef DEBUG
1376         if ( Columns != V.dimension())
1377             {
1378                 cout << "Vector.cpp: In method `real Vector::operator|(Vector&)`: \n";
1379                 cout << "Vector.cpp: vectors have different dimensions \n";
1380                 cout << "carmen: *** [Vector.o] Execution error \n";
1381                 cout << "carmen: abort execution. \n";
1382                 exit(1);
1383             }
1384     #endif
1385
1386     for (n=1; n<=Columns; n++)
1387         result.setValue(n, value(n) * V.value(n));
1388
1389     return result;
1390
1391 }

```

### 5.8.3.17 void Vector::setDimension ( const int *n* )

Sets the dimension of the vector to *n* and reset values to zero.

Example :

```
#include "Vector.h"

Vector V;

...

V.setDimension(3);
```

Parameters

|          |           |
|----------|-----------|
| <i>n</i> | Dimension |
|----------|-----------|

Returns

void

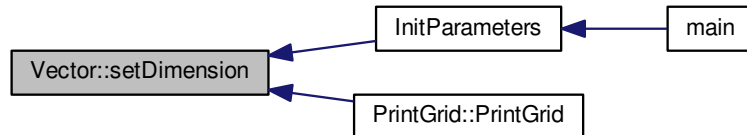
```
299 {
300     Columns = n;
301     if (Columns==9) {
302         *U=0.;
303         *(U+1)=0.;
304         *(U+2)=0.;
305         *(U+3)=0.;
306         *(U+4)=0.;
307         *(U+5)=0.;
308         *(U+6)=0.;
309         *(U+7)=0.;
310         *(U+8)=0.;
311     }
312     else if (Columns==8) {
313         *U=0.;
314         *(U+1)=0.;
315         *(U+2)=0.;
316         *(U+3)=0.;
317         *(U+4)=0.;
318         *(U+5)=0.;
319         *(U+6)=0.;
320         *(U+7)=0.;
321     }
322     else if (Columns==7) {
323         *U=0.;
324         *(U+1)=0.;
325         *(U+2)=0.;
326         *(U+3)=0.;
327         *(U+4)=0.;
328         *(U+5)=0.;
329         *(U+6)=0.;
330     }
331     else if (Columns==6) {
332         *U=0.;
333         *(U+1)=0.;
334         *(U+2)=0.;
335         *(U+3)=0.;
336         *(U+4)=0.;
337         *(U+5)=0.;
338     }
339     else if (Columns==5) {
340         *U=0.;
341         *(U+1)=0.;
342         *(U+2)=0.;
343         *(U+3)=0.;
344         *(U+4)=0.;
345     }
346     else if (Columns==4) {
347         *U=0.;
348         *(U+1)=0.;
349         *(U+2)=0.;
350         *(U+3)=0.;
351     }
352     else if (Columns==3) {
353         *U=0.;
354         *(U+1)=0.;
355         *(U+2)=0.;
356     }
357     else if (Columns==2) {
358         *U=0.;
```

```

359     *(U+1)=0.;
360   }
361   else if (Columns==1) {
362     *U=0.;
363   }
364 }

```

Here is the caller graph for this function:



#### 5.8.3.18 void Vector::setValue ( const int *n*, const real *a* ) [inline]

Sets the component *n* to value *a*.

Example :

```

#include "Vector.h"
Vector V(2);
real x = 3.;
real y = 1.;
V.setValue(1,x);
V.setValue(2,y);

```

Parameters

|          |                 |
|----------|-----------------|
| <i>n</i> | Variable number |
| <i>a</i> | Real value      |

Returns

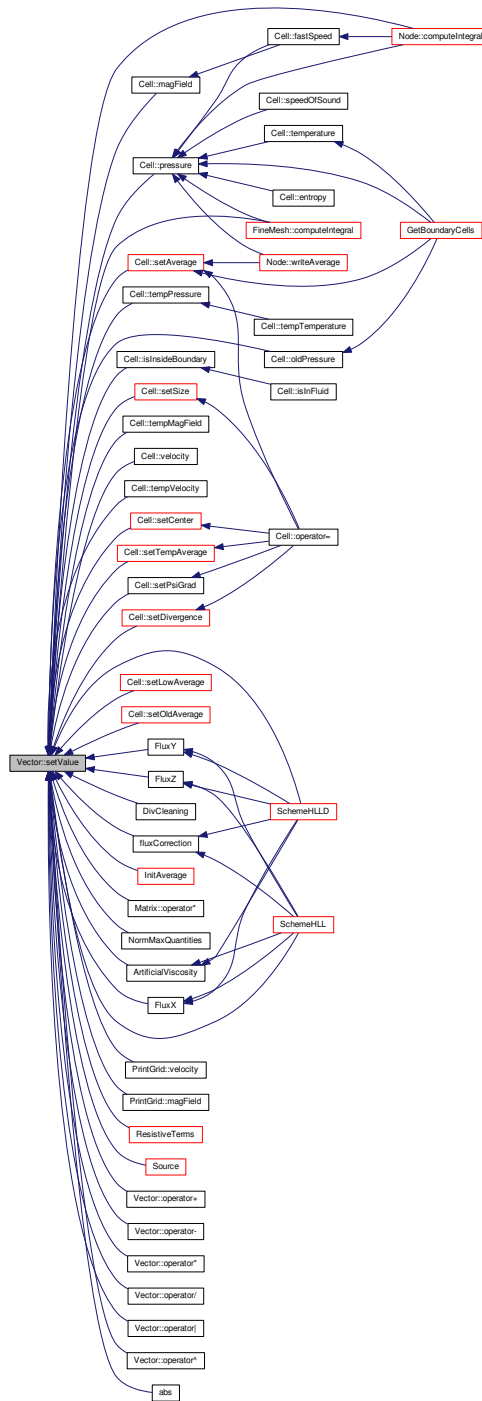
void

```

546 {
547
548 #ifdef DEBUG
549     if ( n <= 0 || n > Columns)
550     {
551         cout << "Vector.cpp: In method 'void Vector::setValue(int, real)':\n";
552         cout << "Vector.cpp: first argument out of range\n";
553         cout << "carmen: *** [Vector.o] Execution error\n";
554         cout << "carmen: abort execution.\n";
555         exit(1);
556     }
557 #endif
558
559     *(U+n-1) = a;
560 }

```

Here is the caller graph for this function:



### 5.8.3.19 void Vector::setZero ( )

Sets all the components to zero.



## Returns

void

```
229 {
230     if (Columns==9) {
231         *U=0.;
232         *(U+1)=0.;
233         *(U+2)=0.;
234         *(U+3)=0.;
235         *(U+4)=0.;
236         *(U+5)=0.;
237         *(U+6)=0.;
238         *(U+7)=0.;
239         *(U+8)=0.;
240     }
241     else if (Columns==8) {
242         *U=0.;
243         *(U+1)=0.;
244         *(U+2)=0.;
245         *(U+3)=0.;
246         *(U+4)=0.;
247         *(U+5)=0.;
248         *(U+6)=0.;
249         *(U+7)=0.;
250     }
251     else if (Columns==7) {
252         *U=0.;
253         *(U+1)=0.;
254         *(U+2)=0.;
255         *(U+3)=0.;
256         *(U+4)=0.;
257         *(U+5)=0.;
258         *(U+6)=0.;
259     }
260     else if (Columns==6) {
261         *U=0.;
262         *(U+1)=0.;
263         *(U+2)=0.;
264         *(U+3)=0.;
265         *(U+4)=0.;
266         *(U+5)=0.;
267     }
268     else if (Columns==5) {
269         *U=0.;
270         *(U+1)=0.;
271         *(U+2)=0.;
272         *(U+3)=0.;
273         *(U+4)=0.;
274     }
275     else if (Columns==4) {
276         *U=0.;
277         *(U+1)=0.;
278         *(U+2)=0.;
279         *(U+3)=0.;
280     }
281     else if (Columns==3) {
282         *U=0.;
283         *(U+1)=0.;
284         *(U+2)=0.;
285     }
286     else if (Columns==2) {
287         *U=0.;
288         *(U+1)=0.;
289     }
290     else if (Columns==1) {
291         *U=0.;
292     }
293 }
```



5.8.4.2 `real Vector::U[9]`

## Components

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

- [Vector.h](#)
- [Vector.cpp](#)



# Chapter 6

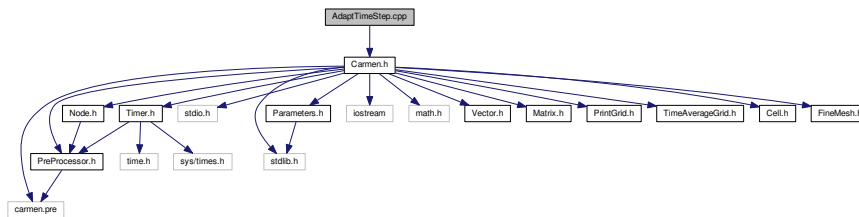
## File Documentation

### 6.1 AdaptTimeStep.cpp File Reference

This function computes the time step for the next iteration.

```
#include "Carmen.h"
```

Include dependency graph for AdaptTimeStep.cpp:



### Functions

- void [AdaptTimeStep](#) ()  
*Adapts time step when required.*

#### 6.1.1 Detailed Description

This function computes the time step for the next iteration.

#### 6.1.2 Function Documentation

##### 6.1.2.1 void AdaptTimeStep ( )

Adapts time step when required.

#### Returns

void

```
25 {  
26     int     RemainingIterations;  
27     real    RemainingTime;
```

```

28
29
30 // Security : do nothing if ConstantTimeStep is true
31
32 if (ConstantTimeStep)
33     return;
34
35 // Compute remaining time
36
37 RemainingTime = PhysicalTime-ElapsedTime;
38
39
40 // In this case, use time adaptivity based on CFL
41 if (Resistivity)
42     TimeStep = CFL*min(SpaceStep/Eigenvalue,
SpaceStep*SpaceStep/(4*eta));
43
44 else
45     TimeStep = CFL*SpaceStep/Eigenvalue;
46
47 // Recompute IterationNb
48
49 if (RemainingTime <= 0.)
50 {
51     IterationNb = IterationNo;
52 }
53 else if (RemainingTime < TimeStep)
54 {
55     TimeStep = RemainingTime;
56     IterationNb = IterationNo + 1;
57 }
58 else
59 {
60     RemainingIterations = (int)(RemainingTime/TimeStep);
61     IterationNb = IterationNo + RemainingIterations;
62 }
63
64 return;
65 }

```

Here is the caller graph for this function:

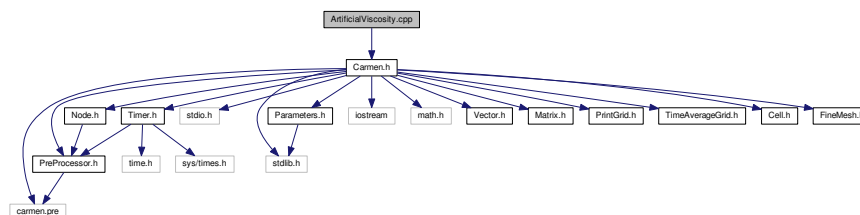


## 6.2 ArtificialViscosity.cpp File Reference

Computes the Laplacian terms for density, energy and momentum equations. It helps with the stability.

```
#include "Carmen.h"
```

Include dependency graph for ArtificialViscosity.cpp:



## Functions

- [Vector ArtificialViscosity](#) (const [Vector](#) &Cell1, const [Vector](#) &Cell2, [real](#) dx, int AxisNo)  
Returns the artificial diffusion source terms in the cell *UserCell*.

### 6.2.1 Detailed Description

Computes the Laplacian terms for density, energy and momentum equations. It helps with the stability.

#### Author

Anna Karina Fontes Gomes

#### Version

2.0

#### Date

Oct-2016

### 6.2.2 Function Documentation

#### 6.2.2.1 [Vector ArtificialViscosity](#) ( const [Vector](#) & *Cell1*, const [Vector](#) & *Cell2*, [real](#) *dx*, int *AxisNo* )

Returns the artificial diffusion source terms in the cell *UserCell*.

#### Parameters

|               |                  |
|---------------|------------------|
| <i>Cell1</i>  | Left cell value  |
| <i>Cell2</i>  | Right cell value |
| <i>AxisNo</i> | Axis of interest |

#### Returns

[Vector](#)

X - direction

Y - direction

Z - direction

```

12 {
13     // --- Local variables ---
14     Vector ML(3), MR(3);
15     Vector Result(QuantityNb);
16     real EL,ER,RL,RR;
17     real viscR, viscX, viscY, viscZ, viscE;
18
19
20     for(int i=1; i <= 3; i++){
21         ML.setValue(i, Cell1.value(i+1));
22         MR.setValue(i, Cell2.value(i+1));
23     }
24
25     RL = Cell1.value(1);
26     RR = Cell2.value(1);
27     EL = Cell1.value(5);
28     ER = Cell2.value(5);
29
30
31     if(AxisNo == 1){
32         viscR = (RR - RL)/dx;
33         viscE = (ER - EL)/dx;
34
35

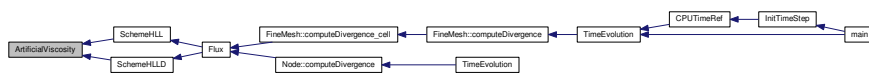
```

```

36     viscX = (MR.value(1) - ML.value(1))/dx;
37     viscY = (MR.value(2) - ML.value(2))/dx;
38     viscZ = (MR.value(3) - ML.value(3))/dx;
39
40
41     }else if(AxisNo == 2){
42
43         viscR = (RR - RL)/dx;
44         viscE = (ER - EL)/dx;
45
46         viscX = (MR.value(1) - ML.value(1))/dx;
47         viscY = (MR.value(2) - ML.value(2))/dx;
48         viscZ = (MR.value(3) - ML.value(3))/dx;
49
50     }else{
51
52         viscR = (RR - RL)/dx;
53         viscE = (ER - EL)/dx;
54
55         viscX = (MR.value(1) - ML.value(1))/dx;
56         viscY = (MR.value(2) - ML.value(2))/dx;
57         viscZ = (MR.value(3) - ML.value(3))/dx;
58
59     }
60
61
62     Result.setZero();
63
64     // These values will be added to the numerical flux
65     Result.setValue(1, chi*viscR);
66     Result.setValue(2, chi*viscX);
67     Result.setValue(3, chi*viscY);
68     Result.setValue(4, chi*viscZ);
69     Result.setValue(5, chi*viscE);
70
71     return Result;
72
73 }

```

Here is the caller graph for this function:

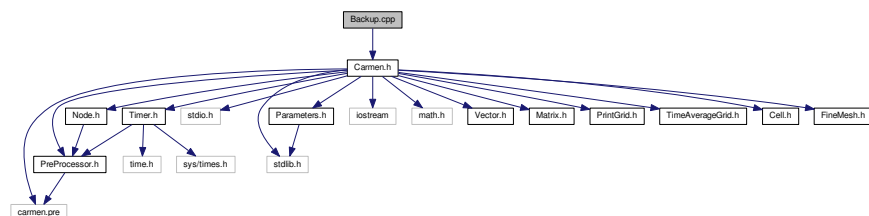


## 6.3 Backup.cpp File Reference

Backup the last simulation.

```
#include "Carmen.h"
```

Include dependency graph for Backup.cpp:



## Functions

- void [Backup](#) (Node \*Root)

*Stores the tree structure and data in order to restart a multiresolution computation.*

- void [Backup](#) (FineMesh \*Root)



Stores the data contained in a regular mesh *Root* in order to restart a finite volume computation.

### 6.3.1 Detailed Description

Backup the last simulation.

### 6.3.2 Function Documentation

#### 6.3.2.1 void Backup ( Node \* *Root* )

Stores the tree structure and data in order to restart a multiresolution computation.

- *Root* denotes the pointer to the first node of the tree structure.

#### Parameters

|             |      |
|-------------|------|
| <i>Root</i> | Root |
|-------------|------|

#### Returns

void

```
31 {
32     Root->backup ();
33 }
```

Here is the caller graph for this function:



#### 6.3.2.2 void Backup ( FineMesh \* *Root* )

Stores the data contained in a regular mesh *Root* in order to restart a finite volume computation.

#### Parameters

|             |      |
|-------------|------|
| <i>Root</i> | Root |
|-------------|------|

#### Returns

void

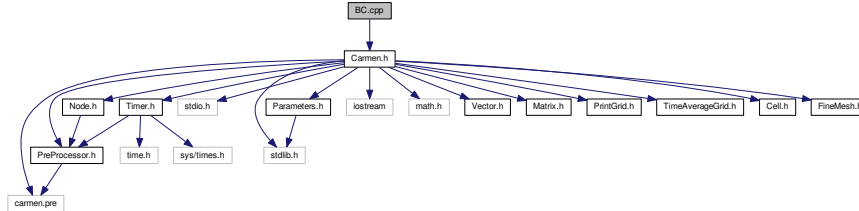
```
43 {
44     Root->backup ();
45 }
```

## 6.4 BC.cpp File Reference

This function returns the position of  $i$ , taking into account boundary conditions.

```
#include "Carmen.h"
```

Include dependency graph for BC.cpp:



### Functions

- `int BC (int i, int AxisNo, int N)`

Returns the position of  $i$  taking into account the boundary conditions in the direction  $AxisNo$ . The number of points in this direction is  $N$ .

Example: for  $AxisNo=1$  and for  $N=256$ ,  $i$  must be between 0 and 255. If  $i=-1$ , the function returns 255 for periodic boundary conditions and 0 for Neumann boundary conditions.

#### 6.4.1 Detailed Description

This function returns the position of  $i$ , taking into account boundary conditions.

#### 6.4.2 Function Documentation

##### 6.4.2.1 `int BC ( int i, int AxisNo, int N = (1 << ScaleNb) )`

Returns the position of  $i$  taking into account the boundary conditions in the direction  $AxisNo$ . The number of points in this direction is  $N$ .

Example: for  $AxisNo=1$  and for  $N=256$ ,  $i$  must be between 0 and 255. If  $i=-1$ , the function returns 255 for periodic boundary conditions and 0 for Neumann boundary conditions.

##### Parameters

|          |                                        |
|----------|----------------------------------------|
| $i$      | Position                               |
| $AxisNo$ | Axis of interest                       |
| $N$      | Defaults to $(1 \ll \text{ScaleNb})$ . |

##### Returns

int

```

38 {
39     int result=-999999;
40
41     if (AxisNo > Dimension)
42         return 0;
43
44 #if defined PARMPI
45     if (CMin[AxisNo] == 3) result=i;    //Periodic
46     else
47     {
48         if (i<0) if ((coords[0]==0 && AxisNo==1) || (coords[1]==0 && AxisNo==2) || (

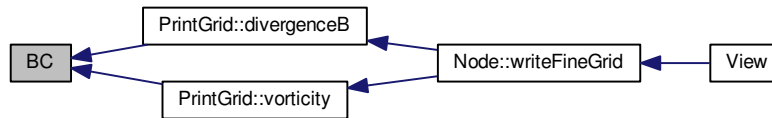
```

```

coords[2]==0 && AxisNo==3)) result=-i-1; //Neumann
49   if (i>=N)       if ((coords[0]==CartDims[0]-1 && AxisNo==1) || (
coords[1]==CartDims[1]-1 && AxisNo==2)
50   || (coords[2]==CartDims[2]-1 && AxisNo==3)) result=2
*N-i-1;
51   if (result==-999999) result=i; //Not the boundary, simple cell from another CPU
52   }
53
54
55 #else
56   if (CMin[AxisNo] == 3)
57     result = (i+N)%N; // Periodic
58   else if(CMin[AxisNo] == 2)
59     result = ((i+N)/N==1)? i : (2*N-i-1)%N; // Neumann
60
61 #endif
62
63   return result;
64 }

```

Here is the caller graph for this function:



## 6.5 BoundaryRegion.cpp File Reference

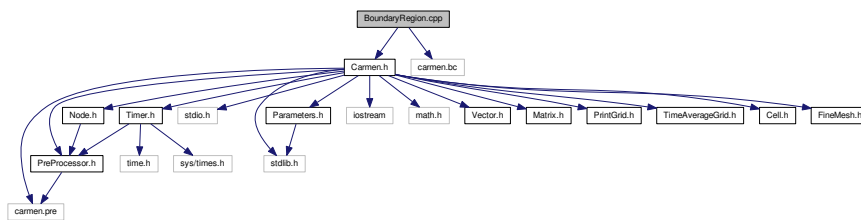
External boundary conditions ( if UseBoundaryRegions = true)

```

#include "Carmen.h"
#include "carmen.bc"

```

Include dependency graph for BoundaryRegion.cpp:



### Functions

- int [BoundaryRegion](#) (const [Vector](#) &X)
  - Returns the boundary region type at the position  $X=(x,y,z)$ .
  - The returned value correspond to: 0 = Fluid (not in the boundary) 1 = Inflow 2 = Outflow 3 = Solid with free-slip walls 4 = Solid with adiabatic walls 5 = Solid with isothermal walls.

### 6.5.1 Detailed Description

External boundary conditions ( if UseBoundaryRegions = true)

## 6.5.2 Function Documentation

### 6.5.2.1 int BoundaryRegion ( const Vector & X )

Returns the boundary region type at the position  $X=(x,y,z)$ .

The returned value correspond to: 0 = Fluid (not in the boundary) 1 = Inflow 2 = Outflow 3 = Solid with free-slip walls 4 = Solid with adiabatic walls 5 = Solid with isothermal walls.

#### Parameters

|   |        |
|---|--------|
| X | Vector |
|---|--------|

#### Returns

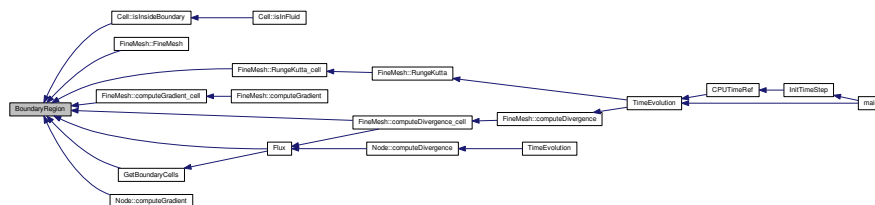
int

```

64 {
65     real x=0., y=0., z=0.;
66
67     int Fluid;
68     int Inflow;
69     int Outflow;
70     int FreeSlipSolid;
71     int AdiabaticSolid;
72     int IsothermalSolid;
73
74     int Region;
75
76     // Only in UseBoundaryRegions = true
77
78     if (!UseBoundaryRegions) return 0;
79
80     // --- Init values ---
81
82     Fluid      = 0;
83     Inflow     = 1;
84     Outflow    = 2;
85     FreeSlipSolid = 3;
86     AdiabaticSolid = 4;
87     IsothermalSolid = 5;
88
89     Region = Fluid;
90
91     x = X.value(1);
92     y = (Dimension > 1)? X.value(2):0.;
93     z = (Dimension > 2)? X.value(3):0.;
94
95     #include "carmen.bc"
96
97     return Region;
98 }

```

Here is the caller graph for this function:



## 6.6 Carmen.h File Reference

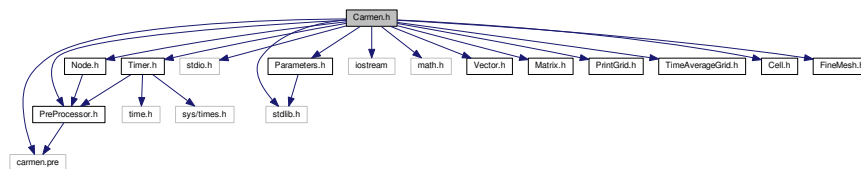
The .h that includes all functions headers.

```

#include "carmen.pre"
#include "PreProcessor.h"
#include <stdio.h>
#include <stdlib.h>
#include <iostream>
#include <math.h>
#include "Vector.h"
#include "Matrix.h"
#include "Timer.h"
#include "Parameters.h"
#include "PrintGrid.h"
#include "TimeAverageGrid.h"
#include "Cell.h"
#include "Node.h"
#include "FineMesh.h"

```

Include dependency graph for Carmen.h:



This graph shows which files directly or indirectly include this file:



## Macros

- #define **Max**(x, y) (((x) > (y)) ? (x):(y))
- #define **Max3**(x, y, z) (**Max**((x),**Max**((y),(z))))
- #define **Min**(x, y) (((x) < (y)) ? (x):(y))
- #define **Min3**(x, y, z) (**Min**((x),**Min**((y),(z))))
- #define **power2**(x) ((x)\*(x))
- #define **power3**(x) ((x)\*(x)\*(x))
- #define **Abs**(x) ( ((x) < 0)? -(x):(x) )

## Functions

- void **AdaptTimeStep** ()  
*Adapts time step when required.*
- int **BC** (int i, int AxisNo, int N=(1<< **ScaleNb**))  
*Returns the position of i taking into account the boundary conditions in the direction AxisNo. The number of points in this direction is N.  
Example: for AxisNo=1 and for N=256, i must be between 0 and 255. If i=-1, the function returns 255 for periodic boundary conditions and 0 for Neumann boundary conditions.*
- int **BoundaryRegion** (const **Vector** &X)  
*Returns the boundary region type at the position X=(x,y,z).  
The returned value correspond to: 0 = Fluid (not in the boundary) 1 = Inflow 2 = Outflow 3 = Solid with free-slip walls 4 = Solid with adiabatic walls 5 = Solid with isothermal walls.*
- void **Backup** (**Node** \*Root)

- Stores the tree structure and data in order to restart a multiresolution computation.*

  - void **Backup** (**FineMesh** \*Root)
- Stores the data contained in a regular mesh Root in order to restart a finite volume computation.*

  - double **CPUTimeRef** (int iterations, int scales)

*Returns the time required by a finite volume computation using iterations iterations and scales scales. It is use to estimate the CPU time compression.*
- **real ComputedTolerance** (const int ScaleNo)

*Returns the computed tolerance at the scale ScaleNo, either using Harten or Donoho thresholding (if CVS=true).*
- int **DigitNumber** (int arg)

*Returns the number of digits of the integer arg.*
- int **FileWrite** (FILE \*f, const char \*format, **real** arg)

*Writes in binary or ASCII mode the real number arg into the file f with the format format. The global parameter DataIsBinary determines this choice.*
- **Vector Flux** (**Cell** &Cell1, **Cell** &Cell2, **Cell** &Cell3, **Cell** &Cell4, int AxisNo)

*Returns the flux at the interface between Cell2 and Cell3. Here a 4-point space scheme is used. Cell2 and Cell3 are the first neighbours on the left and right sides. Cell1 and Cell4 are the second neighbours on the left and right sides.*
- void **GetBoundaryCells** (**Cell** &Cell1, **Cell** &Cell2, **Cell** &Cell3, **Cell** &Cell4, **Cell** &C1, **Cell** &C2, **Cell** &C3, **Cell** &C4, int AxisNo)

*Transform the 4 cells of the flux Cell1, Cell2, Cell3, Cell4 into C1, C2, C3, C4, to take into account boundary conditions (used in Flux.cpp).*
- **Vector InitAverage** (**real** x, **real** y=0., **real** z=0.)

*Returns the initial condition in (x, y, z) form the one defined in carmen.ini.*
- **real InitResistivity** (**real** x, **real** y=0., **real** z=0.)

*Returns the initial resistivity condition in (x, y, z) form the one defined in carmen.eta.*
- void **InitParameters** ()

*Inits parameters from file carmen.par. If a parameter is not mentioned in this file, the default value is used.*
- void **InitTimeStep** ()

*Inits time step and all the parameters which depend on it.*
- void **InitTree** (**Node** \*Root)

*Inits tree structure from initial condition, starting form the node Root. Only for multiresolution computations.*
- **Vector Limiter** (const **Vector** u, const **Vector** v)

*Returns the value of the slope limiter between the slopes u and v.*
- **real Limiter** (const **real** x)

*Returns the valur of slope limiter from a real value x.*
- **real MinAbs** (const **real** a, const **real** b)

*Returns the minimum in module of a and b.*
- **real NormMaxQuantities** (const **Vector** &V)

*Returns the Max-norm of the vector where every quantity is divided by its characteristic value.*
- void **Performance** (const char \*FileName)

*Computes the performance of the computation and, for cluster computations, write it into file FileName.*
- void **PrintIntegral** (const char \*FileName)

*Writes the integral values, like e.g flame velocity, global error, into file FileName.*
- void **RefreshTree** (**Node** \*Root)

*Refresh the tree structure, i.e. compute the cell-averages of the internal nodes by projection and those of the virtual leaves by prediction. The root node is Root. Only for multiresolution computations.*
- void **Remesh** (**Node** \*Root)

*Remesh the tree structure after a time evolution. The root node is Root. Only for multiresolution computations.*
- **Vector FluxX** (const **Vector** &Avg)

*Returns the physical flux of MHD equations in X direction.*
- **Vector FluxY** (const **Vector** &Avg)

*Returns the physical flux of MHD equations in Y direction.*
- **Vector FluxZ** (const **Vector** &Avg)

- Returns the physical flux of MHD equations in Z direction.*

  - **Vector SchemeHLL** (const **Cell** &Cell1, const **Cell** &Cell2, const **Cell** &Cell3, const **Cell** &Cell4, const int AxisNo)
 

*Returns the HLL numerical flux for MHD equations. The scheme uses four cells to estimate the flux at the interface. Cell2 and Cell3 are the first neighbours on the left and right sides. Cell1 and Cell4 are the second neighbours on the left and right sides.*
  - **Vector SchemeHLLD** (const **Cell** &Cell1, const **Cell** &Cell2, const **Cell** &Cell3, const **Cell** &Cell4, const int AxisNo)
 

*Returns the HLLD numerical flux for MHD equations. The scheme uses four cells to estimate the flux at the interface. Cell2 and Cell3 are the first neighbours on the left and right sides. Cell1 and Cell4 are the second neighbours on the left and right sides.*
  - **Matrix stateUstar** (const **Vector** &AvgL, const **Vector** &AvgR, const **real** prel, const **real** prer, **real** &slopeLeft, **real** &slopeRight, **real** &slopeM, **real** &slopeLeftStar, **real** &slopeRightStar, int AxisNo)
 

*Returns the intermediary states of HLLD numerical flux for MHD equations.*
  - void **fluxCorrection** (**Vector** &Flux, const **Vector** &AvgL, const **Vector** &AvgR, int AxisNo)
 

*This function apply the divergence-free correction to the numerical flux.*
  - void **ShowTime** (**Timer** arg)
 

*Writes on screen the estimation of total and remaining CPU times. These informations are stored in the timer arg.*
  - int **Sign** (const **real** a)
 

*Returns 1 if a is non-negative, -1 elsewhere.*
  - **Vector ArtificialViscosity** (const **Vector** &Cell1, const **Vector** &Cell2, **real** dx, int AxisNo)
 

*Returns the artificial diffusion source terms in the cell UserCell.*
  - **Vector ResistiveTerms** (**Cell** &Cell1, **Cell** &Cell2, **Cell** &Cell3, **Cell** &Cell4, int AxisNo)
 

*Returns the resistive source terms in the cell UserCell.*
  - **Vector Source** (**Cell** &UserCell)
 

*Returns the source term in the cell UserCell.*
  - **real Step** (**real** x)
 

*Returns a step (1 if  $x < 0$ , 0 if  $x > 0$ , 0.5 if  $x=0$ )*
  - void **TimeEvolution** (**FineMesh** \*Root)
 

*Computes a time evolution on the regular fine mesh Root. Only for finite volume computations.*
  - void **TimeEvolution** (**Node** \*Root)
 

*Computes a time evolution on the tree structure, the root node being Root. Only for multiresolution computations.*
  - void **View** (**FineMesh** \*Root, const char \*AverageFileName)
 

*Writes the current cell-averages of the fine mesh Root into file AverageFileName. Only for finite volume computations.*
  - void **View** (**Node** \*Root, const char \*TreeFileName, const char \*MeshFileName, const char \*AverageFileName)
 

*Writes the data of the tree structure into files TreeFileName (tree structure), MeshFileName (mesh) and AverageFileName (cell-averages). The root node is Root. Only for multiresolution computations.*
  - void **ViewEvery** (**FineMesh** \*Root, int arg)
 

*Same as previous for a fine mesh Root. Only for finite volume computations.*
  - void **ViewEvery** (**Node** \*Root, int arg)
 

*Writes into file the data of the tree structure at iteration arg. The output file names are AverageNNN.dat and MeshNN.dat, NNN being the iteration in an accurate format. The root node is Root. Only for multiresolution computations.*
  - void **ViewIteration** (**FineMesh** \*Root)
 

*Same as previous for a fine mesh Root. Only for finite volume computations.*
  - void **ViewIteration** (**Node** \*Root)
 

*Writes into file the data of the tree structure from physical time PrintTime1 to physical time PrintTime6. The output file names are Average\_N.dat and Mesh\_N.dat, N being between 1 and 6. The root node is Root. Only for multiresolution computations.*
  - void **CreateMPIType** (**FineMesh** \*Root)
  - void **CPUExchange** (**FineMesh** \*Root, int)
 

*Parallel function DOES NOT WORK!*
  - void **FreeMPIType** ()

- Parallel function DOES NOT WORK!*
- void [CreateMPITopology](#) ()
- Parallel function DOES NOT WORK!*
- void [CreateMPILinks](#) ()
- Parallel function DOES NOT WORK!*
- void [ReduceIntegralValues](#) ()
- Parallel function DOES NOT WORK!*

### 6.6.1 Detailed Description

The .h that includes all functions headers.

### 6.6.2 Macro Definition Documentation

#### 6.6.2.1 #define Abs( x ) ( ((x) < 0)? -(x):(x) )

Returns the absolute value of x.

#### 6.6.2.2 #define Max( x, y ) (((x) > (y)) ? (x):(y))

Returns the Maximum value between x and y.

#### 6.6.2.3 #define Max3( x, y, z ) (Max((x),Max((y),(z))))

Returns the Maximum value between x, y and z.

#### 6.6.2.4 #define Min( x, y ) (((x) < (y)) ? (x):(y))

Returns the minimum value between x and y.

#### 6.6.2.5 #define Min3( x, y, z ) (Min((x),Min((y),(z))))

Returns the minimum value between x, y and z.

#### 6.6.2.6 #define power2( x ) ((x)\*(x))

Returns the square of x.

#### 6.6.2.7 #define power3( x ) ((x)\*(x)\*(x))

Returns the cube of x.

### 6.6.3 Function Documentation

#### 6.6.3.1 void AdaptTimeStep ( )

Adapts time step when required.



## Returns

void

```

25 {
26     int     RemainingIterations;
27     real    RemainingTime;
28
29
30     // Security : do nothing if ConstantTimeStep is true
31
32     if (ConstantTimeStep)
33         return;
34
35     // Compute remaining time
36
37     RemainingTime = PhysicalTime-ElapsedTime;
38
39
40     // In this case, use time adaptivity based on CFL
41     if (Resistivity)
42         TimeStep = CFL*min(SpaceStep/Eigenvalue,
SpaceStep*SpaceStep/(4*eta));
43     else
44         TimeStep = CFL*SpaceStep/Eigenvalue;
45
46     // Recompute IterationNb
47
48     if (RemainingTime <= 0.)
49     {
50         IterationNb = IterationNo;
51     }
52     else if (RemainingTime < TimeStep)
53     {
54         TimeStep = RemainingTime;
55         IterationNb = IterationNo + 1;
56     }
57     else
58     {
59         RemainingIterations = (int)(RemainingTime/TimeStep);
60         IterationNb = IterationNo + RemainingIterations;
61     }
62
63     return;
64
65 }

```

Here is the caller graph for this function:



### 6.6.3.2 Vector ArtificialViscosity ( const Vector & Cell1, const Vector & Cell2, real dx, int AxisNo )

Returns the artificial diffusion source terms in the cell *UserCell*.

## Parameters

|              |                  |
|--------------|------------------|
| <i>Cell1</i> | Left cell value  |
| <i>Cell2</i> | Right cell value |

| AxisNo | Axis of interest |
|--------|------------------|
|--------|------------------|

## Returns

## Vector

X - direction

Y - direction

Z - direction

```

12 {
13     // --- Local variables ---
14     Vector ML(3), MR(3);
15     Vector Result(QuantityNb);
16     real EL,ER,RL,RR;
17     real viscR, viscX, viscY, viscZ, viscE;
18
19
20     for(int i=1; i <= 3; i++){
21         ML.setValue(i, Cell11.value(i+1));
22         MR.setValue(i, Cell12.value(i+1));
23     }
24
25     RL = Cell11.value(1);
26     RR = Cell12.value(1);
27     EL = Cell11.value(5);
28     ER = Cell12.value(5);
29
30
31     if(AxisNo == 1){
32         viscR = (RR - RL)/dx;
33         viscE = (ER - EL)/dx;
34
35         viscX = (MR.value(1) - ML.value(1))/dx;
36         viscY = (MR.value(2) - ML.value(2))/dx;
37         viscZ = (MR.value(3) - ML.value(3))/dx;
38
39
40     }else if(AxisNo == 2){
41         viscR = (RR - RL)/dx;
42         viscE = (ER - EL)/dx;
43
44         viscX = (MR.value(1) - ML.value(1))/dx;
45         viscY = (MR.value(2) - ML.value(2))/dx;
46         viscZ = (MR.value(3) - ML.value(3))/dx;
47
48     }else{
49         viscR = (RR - RL)/dx;
50         viscE = (ER - EL)/dx;
51
52         viscX = (MR.value(1) - ML.value(1))/dx;
53         viscY = (MR.value(2) - ML.value(2))/dx;
54         viscZ = (MR.value(3) - ML.value(3))/dx;
55
56     }
57
58     Result.setZero();
59
60     // These values will be added to the numerical flux
61     Result.setValue(1, chi*viscR);
62     Result.setValue(2, chi*viscX);
63     Result.setValue(3, chi*viscY);
64     Result.setValue(4, chi*viscZ);
65     Result.setValue(5, chi*viscE);
66
67     return Result;
68 }

```

Here is the caller graph for this function:



## 6.6.3.3 void Backup ( Node \* Root )

Stores the tree structure and data in order to restart a multiresolution computation.

- *Root* denotes the pointer to the first node of the tree structure.

## Parameters

|             |      |
|-------------|------|
| <i>Root</i> | Root |
|-------------|------|

## Returns

void

```
31 {
32     Root->backup ();
33 }
```

Here is the caller graph for this function:



## 6.6.3.4 void Backup ( FineMesh \* Root )

Stores the data contained in a regular mesh *Root* in order to restart a finite volume computation.

## Parameters

|             |      |
|-------------|------|
| <i>Root</i> | Root |
|-------------|------|

## Returns

void

```
43 {
44     Root->backup ();
45 }
```

6.6.3.5 int BC ( int *i*, int *AxisNo*, int *N* = (1 << **ScaleNb**) )

Returns the position of *i* taking into account the boundary conditions in the direction *AxisNo*. The number of points in this direction is *N*.

Example: for *AxisNo*=1 and for *N*=256, *i* must be between 0 and 255. If *i*=-1, the function returns 255 for periodic boundary conditions and 0 for Neumann boundary conditions.

## Parameters

|               |                                        |
|---------------|----------------------------------------|
| <i>i</i>      | Position                               |
| <i>AxisNo</i> | Axis of interest                       |
| <i>N</i>      | Defaults to $(1 \ll \text{ScaleNb})$ . |

## Returns

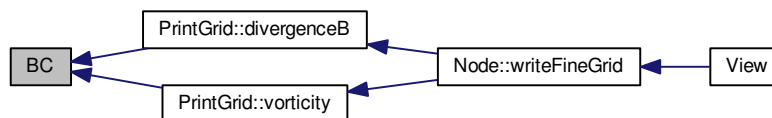
int

```

38 {
39   int result=-999999;
40
41   if (AxisNo > Dimension)
42     return 0;
43
44 #if defined PARMPI
45   if (CMin[AxisNo] == 3) result=i;    //Periodic
46   else
47   {
48     if (i<0) if ((coords[0]==0 && AxisNo==1) || (coords[1]==0 && AxisNo==2) || (
49 coords[2]==0 && AxisNo==3)) result=-i-1; //Neumann
50     if (i>=N) if ((coords[0]==CartDims[0]-1 && AxisNo==1) || (
51 coords[1]==CartDims[1]-1 && AxisNo==2)
52 || (coords[2]==CartDims[2]-1 && AxisNo==3)) result=2
53 *N-i-1;
54     if (result==-999999) result=i;    //Not the boundary, simple cell from another CPU
55   }
56 #else
57   if (CMin[AxisNo] == 3)
58     result = (i+N)%N;                // Periodic
59   else if(CMin[AxisNo] == 2)
60     result = ((i+N)/N==1)? i : (2*N-i-1)%N; // Neumann
61 #endif
62   return result;
63 }

```

Here is the caller graph for this function:



## 6.6.3.6 int BoundaryRegion ( const Vector &amp; X )

Returns the boundary region type at the position  $X=(x,y,z)$ .

The returned value correspond to: 0 = Fluid (not in the boundary) 1 = Inflow 2 = Outflow 3 = Solid with free-slip walls 4 = Solid with adiabatic walls 5 = Solid with isothermal walls.

## Parameters

|          |                        |
|----------|------------------------|
| <i>X</i> | <a href="#">Vector</a> |
|----------|------------------------|



### 6.6.3.8 void CPUExchange ( FineMesh \* Root, int )

Parallel function DOES NOT WORK!

#### Parameters

|             |           |
|-------------|-----------|
| <i>Root</i> | Fine mesh |
|-------------|-----------|

#### Returns

void

```

350                                     {
351     CommTimer.start();
352     #if defined PARMPI
353     int i,k;
354     int exNb=0;
355
356     WhatSend=WS;
357     CellElementsNb=0;
358
359     for (i=0;i<16;i++) {
360         k=1<<i;
361         if ((WS & k) != 0) CellElementsNb++;
362     }
363
364     static bool ft=true;
365     // if (ft==true) {
366     CreateMPIType(Root);
367     // CreateMPILinks();
368     // ft=false;
369     // }
370
371     // MPI_Startall(4*Dimension,req);
372
373
374     //Send
375     switch (MPISendType) {
376     case 0:
377         MPI_Ibsend(MPI_BOTTOM, 1, MPItypeSiL, rank_il, 100, comm_cart, &req[exNb++]);
378         MPI_Ibsend(MPI_BOTTOM, 1, MPItypeSiU, rank_iu, 200, comm_cart, &req[exNb++]);
379         break;
380
381     case 10:
382         MPI_Isend(MPI_BOTTOM, 1, MPItypeSiL, rank_il, 100, comm_cart, &req[exNb++]);
383         MPI_Isend(MPI_BOTTOM, 1, MPItypeSiU, rank_iu, 200, comm_cart, &req[exNb++]);
384         break;
385
386     case 20:
387         MPI_Issend(MPI_BOTTOM, 1, MPItypeSiL, rank_il, 100, comm_cart, &req[exNb++]);
388         MPI_Issend(MPI_BOTTOM, 1, MPItypeSiU, rank_iu, 200, comm_cart, &req[exNb++]);
389         break;
390     }
391
392     if (Dimension >= 2) {
393     switch (MPISendType) {
394     case 0:
395         MPI_Ibsend(MPI_BOTTOM, 1, MPItypeSjL, rank_jl, 300, comm_cart, &req[exNb++]);
396         MPI_Ibsend(MPI_BOTTOM, 1, MPItypeSjU, rank_ju, 400, comm_cart, &req[exNb++]);
397         break;
398
399     case 10:
400         MPI_Isend(MPI_BOTTOM, 1, MPItypeSjL, rank_jl, 300, comm_cart, &req[exNb++]);
401         MPI_Isend(MPI_BOTTOM, 1, MPItypeSjU, rank_ju, 400, comm_cart, &req[exNb++]);
402         break;
403
404     case 20:
405         MPI_Issend(MPI_BOTTOM, 1, MPItypeSjL, rank_jl, 300, comm_cart, &req[exNb++]);
406         MPI_Issend(MPI_BOTTOM, 1, MPItypeSjU, rank_ju, 400, comm_cart, &req[exNb++]);
407         break;
408     }
409     }
410
411     if (Dimension == 3) {
412     switch (MPISendType) {
413     case 0:
414         MPI_Ibsend(MPI_BOTTOM, 1, MPItypeSkL, rank_kl, 500, comm_cart, &req[exNb++]);
415         MPI_Ibsend(MPI_BOTTOM, 1, MPItypeSkU, rank_ku, 600, comm_cart, &req[exNb++]);
416         break;
417
418     case 10:
419         MPI_Isend(MPI_BOTTOM, 1, MPItypeSkL, rank_kl, 500, comm_cart, &req[exNb++]);
420         MPI_Isend(MPI_BOTTOM, 1, MPItypeSkU, rank_ku, 600, comm_cart, &req[exNb++]);

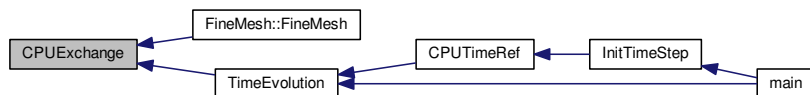
```

```

421         break;
422     }
423     case 20:
424         MPI_Issend(MPI_BOTTOM, 1, MPItypeSkL, rank_kl, 500, comm_cart, &req[exNb++]);
425         MPI_Issend(MPI_BOTTOM, 1, MPItypeSkU, rank_ku, 600, comm_cart, &req[exNb++]);
426         break;
427     }
428 }
429
430 //Recv
431
432 if (MPIRecvType==0) {
433     MPI_Recv(MPI_BOTTOM, 1, MPItypeRiL, rank_il, 200, comm_cart, &st[6]);
434     MPI_Recv(MPI_BOTTOM, 1, MPItypeRiU, rank_iu, 100, comm_cart, &st[7]);
435 } else
436 {
437     MPI_Irecv(MPI_BOTTOM, 1, MPItypeRiL, rank_il, 200, comm_cart, &req[exNb++]);
438     MPI_Irecv(MPI_BOTTOM, 1, MPItypeRiU, rank_iu, 100, comm_cart, &req[exNb++]);
439 }
440
441 if (Dimension >= 2) {
442     if (MPIRecvType==0) {
443         MPI_Recv(MPI_BOTTOM, 1, MPItypeRjL, rank_jl, 400, comm_cart, &st[8]);
444         MPI_Recv(MPI_BOTTOM, 1, MPItypeRjU, rank_ju, 300, comm_cart, &st[9]);
445     } else
446     {
447         MPI_Irecv(MPI_BOTTOM, 1, MPItypeRjL, rank_jl, 400, comm_cart, &req[exNb++]);
448         MPI_Irecv(MPI_BOTTOM, 1, MPItypeRjU, rank_ju, 300, comm_cart, &req[exNb++]);
449     }
450 }
451
452 if (Dimension == 3) {
453     if (MPIRecvType==0) {
454         MPI_Recv(MPI_BOTTOM, 1, MPItypeRkL, rank_kl, 600, comm_cart, &st[10]);
455         MPI_Recv(MPI_BOTTOM, 1, MPItypeRkU, rank_ku, 500, comm_cart, &st[11]);
456     } else
457     {
458         MPI_Irecv(MPI_BOTTOM, 1, MPItypeRkL, rank_kl, 600, comm_cart, &req[exNb++]);
459         MPI_Irecv(MPI_BOTTOM, 1, MPItypeRkU, rank_ku, 500, comm_cart, &req[exNb++]);
460     }
461 }
462
463 FreeMPIType();
464 #endif
465 CommTimer.stop();
466 }

```

Here is the caller graph for this function:



### 6.6.3.9 double CPUTimeRef ( int iterations, int scales )

Returns the time required by a finite volume computation using *iterations* iterations and *scales* scales. It is use to estimate the CPU time compression.

#### Parameters

|                   |                       |
|-------------------|-----------------------|
| <i>iterations</i> | Number of iterations. |
| <i>scales</i>     | Scales                |

#### Returns

double

```

24 {

```

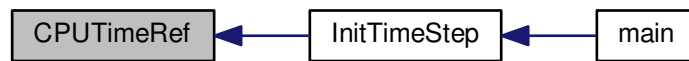
```

25 // --- Local variables -----
26
27 int OldIterationNb=0;
28 int OldScaleNb=0;
29 real OldTimeStep=0.;
30 bool ConstantTimeStepOld=ConstantTimeStep;
31
32 double result=0.;
33
34 Timer CPURef;
35 FineMesh* MeshRef;
36
37 // --- Execution -----
38
39 // Toggle on : Compute reference CPU time
40
41 ComputeCPUTimeRef = true;
42
43 // Toggle off : Constant time step
44
45 ConstantTimeStep = true;
46
47 // backup values of IterationNb and ScaleNb
48
49 OldIterationNb = IterationNb;
50 OldScaleNb = ScaleNb;
51 OldTimeStep = TimeStep;
52
53 // use reference values
54 IterationNb = iterations;
55 ScaleNb = scales;
56 TimeStep = 0.;
57
58 one_D=1; two_D=1;
59 if (Dimension >= 2) one_D=1<<ScaleNb;
60 if (Dimension == 3) two_D=1<<ScaleNb;
61
62 // init mesh
63 MeshRef = new FineMesh;
64
65 // Iterate on time
66
67 for (IterationNo = 1; IterationNo <= IterationNb;
IterationNo++)
68 {
69 // start timer
70 CPURef.start();
71
72 // Compute time evolution
73 TimeEvolution(MeshRef);
74
75 // check CPU Time
76 CPURef.check();
77
78 // stop timer
79 CPURef.stop();
80 }
81
82 // Compute CPUTimeRef
83 result = CPURef.CPUTime();
84 result *= 1./IterationNb;
85 result *= 1<<(Dimension*(OldScaleNb-ScaleNb));
86
87 // delete MeshRef
88 delete MeshRef;
89
90 // restore values of IterationNb and ScaleNb
91 IterationNb = OldIterationNb;
92 ScaleNb = OldScaleNb;
93 TimeStep = OldTimeStep;
94 IterationNo = 0;
95
96 one_D=1; two_D=1;
97 if (Dimension >= 2) one_D=1<<ScaleNb;
98 if (Dimension == 3) two_D=1<<ScaleNb;
99
100 // Toggle off : Compute reference CPU time
101
102 ComputeCPUTimeRef = false;
103
104 // Restore the value of ConstantTimeStep
105
106 ConstantTimeStep = ConstantTimeStepOld;
107
108 return result;
109 }

```



Here is the caller graph for this function:



### 6.6.3.10 void CreateMPLinks ( )

Parallel function DOES NOT WORK!

#### Returns

void

```

271     {
272     int exNb;
273     exNb=0;
274     #if defined PARMPI
275
276     //Send
277
278     switch (MPISendType) {
279     case 0:
280         MPI_Bsend_init(MPI_BOTTOM, 1, MPItypeSiL, rank_il, 100, comm_cart, &req[exNb++]);
281         MPI_Bsend_init(MPI_BOTTOM, 1, MPItypeSiU, rank_iu, 200, comm_cart, &req[exNb++]);
282         break;
283
284     case 10:
285         MPI_Send_init(MPI_BOTTOM, 1, MPItypeSiL, rank_il, 100, comm_cart, &req[exNb++]);
286         MPI_Send_init(MPI_BOTTOM, 1, MPItypeSiU, rank_iu, 200, comm_cart, &req[exNb++]);
287         break;
288
289     case 20:
290         MPI_Ssend_init(MPI_BOTTOM, 1, MPItypeSiL, rank_il, 100, comm_cart, &req[exNb++]);
291         MPI_Ssend_init(MPI_BOTTOM, 1, MPItypeSiU, rank_iu, 200, comm_cart, &req[exNb++]);
292         break;
293     }
294
295     if (Dimension >= 2) {
296         switch (MPISendType) {
297         case 0:
298             MPI_Bsend_init(MPI_BOTTOM, 1, MPItypeSjL, rank_jl, 300, comm_cart, &req[exNb++]);
299             MPI_Bsend_init(MPI_BOTTOM, 1, MPItypeSjU, rank_ju, 400, comm_cart, &req[exNb++]);
300             break;
301
302         case 10:
303             MPI_Send_init(MPI_BOTTOM, 1, MPItypeSjL, rank_jl, 300, comm_cart, &req[exNb++]);
304             MPI_Send_init(MPI_BOTTOM, 1, MPItypeSjU, rank_ju, 400, comm_cart, &req[exNb++]);
305             break;
306
307         case 20:
308             MPI_Ssend_init(MPI_BOTTOM, 1, MPItypeSjL, rank_jl, 300, comm_cart, &req[exNb++]);
309             MPI_Ssend_init(MPI_BOTTOM, 1, MPItypeSjU, rank_ju, 400, comm_cart, &req[exNb++]);
310             break;
311         }
312     }
313
314     if (Dimension == 3) {
315         switch (MPISendType) {
316         case 0:
317             MPI_Bsend_init(MPI_BOTTOM, 1, MPItypeSkL, rank_kl, 500, comm_cart, &req[exNb++]);
318             MPI_Bsend_init(MPI_BOTTOM, 1, MPItypeSkU, rank_ku, 600, comm_cart, &req[exNb++]);
319             break;
320
321         case 10:
322             MPI_Send_init(MPI_BOTTOM, 1, MPItypeSkL, rank_kl, 500, comm_cart, &req[exNb++]);
323             MPI_Send_init(MPI_BOTTOM, 1, MPItypeSkU, rank_ku, 600, comm_cart, &req[exNb++]);
324             break;
325

```

```

326         case 20:
327             MPI_Ssend_init(MPI_BOTTOM, 1, MPItypeSkL, rank_kl, 500, comm_cart, &req[exNb++]);
328             MPI_Ssend_init(MPI_BOTTOM, 1, MPItypeSkU, rank_ku, 600, comm_cart, &req[exNb++]);
329             break;
330         }
331     }
332
333 //Recv
334
335     MPI_Recv_init(MPI_BOTTOM, 1, MPItypeRiL, rank_il, 200, comm_cart, &req[exNb++]);
336     MPI_Recv_init(MPI_BOTTOM, 1, MPItypeRiU, rank_iu, 100, comm_cart, &req[exNb++]);
337
338     if (Dimension >= 2) {
339         MPI_Recv_init(MPI_BOTTOM, 1, MPItypeRjL, rank_jl, 400, comm_cart, &req[exNb++]);
340         MPI_Recv_init(MPI_BOTTOM, 1, MPItypeRjU, rank_ju, 300, comm_cart, &req[exNb++]);
341     }
342
343     if (Dimension == 3) {
344         MPI_Recv_init(MPI_BOTTOM, 1, MPItypeRkL, rank_kl, 600, comm_cart, &req[exNb++]);
345         MPI_Recv_init(MPI_BOTTOM, 1, MPItypeRkU, rank_ku, 500, comm_cart, &req[exNb++]);
346     }
347 #endif
348 }

```

### 6.6.3.11 void CreateMPITopology ( )

Parallel function DOES NOT WORK!

#### Returns

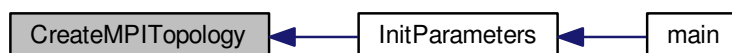
void

```

22     {
23 #if defined PARMPI
24     int src;
25     int periods[]={1,1,1};
26     CartDims[0]=CartDims[1]=CartDims[2]=0;
27
28     MPI_Dims_create(size,Dimension,CartDims);
29     MPI_Cart_create(MPI_COMM_WORLD,Dimension,CartDims,periods,1,&comm_cart);
30     MPI_Comm_rank(comm_cart, &rank);
31     MPI_Cart_coords(comm_cart,rank,Dimension,coords);
32
33     MPI_Cart_shift(comm_cart, 0, -1, &src, &rank_il);
34     MPI_Cart_shift(comm_cart, 0, 1, &src, &rank_iu);
35
36     if (Dimension >= 2) {
37         MPI_Cart_shift(comm_cart, 1, -1, &src, &rank_jl);
38         MPI_Cart_shift(comm_cart, 1, 1, &src, &rank_ju);
39     }
40
41     if (Dimension == 3) {
42         MPI_Cart_shift(comm_cart, 2, -1, &src, &rank_kl);
43         MPI_Cart_shift(comm_cart, 2, 1, &src, &rank_ku);
44     }
45 #endif
46 }

```

Here is the caller graph for this function:



## 6.6.3.12 void CreateMPIType ( FineMesh \* Root )

```

121                                     {
122   #if defined PARMPI
123     int i,j,k;
124     int n,d,l;
125
126     Cell *MeshCell;
127     MeshCell=Root->MeshCell;
128
129     n=0;
130     for (l=0;l<NeighbourNb;l++)
131         for (j=0;j<one_D;j++)
132             for (k=0;k<two_D;k++)   FillNbAddr (Root->Neighbour_iL,l,j,k,n);
133     MPI_Type_hindexed (CellElementsNb*NeighbourNb*one_D*two_D,blocklen,disp,
134                       MPI_Type,&MPItypeRiL);
135     MPI_Type_commit (&MPItypeRiL);
136
137     n=0;
138     for (l=0;l<NeighbourNb;l++)
139         for (j=0;j<one_D;j++)
140             for (k=0;k<two_D;k++)   FillNbAddr (Root->Neighbour_iU,l,j,k,n);
141     MPI_Type_hindexed (CellElementsNb*NeighbourNb*one_D*two_D,blocklen,disp,
142                       MPI_Type,&MPItypeRiU);
143     MPI_Type_commit (&MPItypeRiU);
144
145     n=0;
146     for (l=0;l<NeighbourNb;l++)
147         for (j=0;j<one_D;j++)
148             for (k=0;k<two_D;k++) {
149                 i=l;
150                 d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
151                 FillCellAddr (MeshCell,d,n);
152             }
153     MPI_Type_hindexed (CellElementsNb*NeighbourNb*one_D*two_D,blocklen,disp,
154                       MPI_Type,&MPItypeSiL);
155     MPI_Type_commit (&MPItypeSiL);
156
157     n=0;
158     for (l=0;l<NeighbourNb;l++)
159         for (j=0;j<one_D;j++)
160             for (k=0;k<two_D;k++) {
161                 i=(1<<ScaleNb)-NeighbourNb+1;
162                 d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
163                 FillCellAddr (MeshCell,d,n);
164             }
165     MPI_Type_hindexed (CellElementsNb*NeighbourNb*one_D*two_D,blocklen,disp,
166                       MPI_Type,&MPItypeSiU);
167     MPI_Type_commit (&MPItypeSiU);
168
169     if (Dimension >= 2) {
170         n=0;
171         for (l=0;l<NeighbourNb;l++)
172             for (i=0;i<one_D;i++)
173                 for (k=0;k<two_D;k++)   FillNbAddr (Root->
174               Neighbour_jL,l,i,k,n);
175         MPI_Type_hindexed (CellElementsNb*NeighbourNb*one_D*two_D,blocklen,disp,
176                           MPI_Type,&MPItypeRjL);
177         MPI_Type_commit (&MPItypeRjL);
178
179         n=0;
180         for (l=0;l<NeighbourNb;l++)
181             for (i=0;i<one_D;i++)
182                 for (k=0;k<two_D;k++)   FillNbAddr (Root->
183               Neighbour_jU,l,i,k,n);
184         MPI_Type_hindexed (CellElementsNb*NeighbourNb*one_D*two_D,blocklen,disp,
185                           MPI_Type,&MPItypeRjU);
186         MPI_Type_commit (&MPItypeRjU);
187
188         n=0;
189         for (l=0;l<NeighbourNb;l++)
190             for (i=0;i<one_D;i++)
191                 for (k=0;k<two_D;k++) {
192                     j=1;
193                     d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
194                     FillCellAddr (MeshCell,d,n);
195                 }
196         MPI_Type_hindexed (CellElementsNb*NeighbourNb*one_D*two_D,blocklen,disp,
197                           MPI_Type,&MPItypeSjL);
198         MPI_Type_commit (&MPItypeSjL);
199
200         n=0;
201         for (l=0;l<NeighbourNb;l++)
202             for (i=0;i<one_D;i++)
203                 for (k=0;k<two_D;k++) {
204                     j=(1<<ScaleNb)-NeighbourNb+1;
205                     d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);

```

```

197             FillCellAddr (MeshCell, d, n);
198         }
199         MPI_Type_hindexed(CellElementsNb*NeighbourNb*one_D*two_D, blocklen, disp,
MPI_Type, &MPITypeSjU);
200         MPI_Type_commit (&MPITypeSjU);
201     }
202
203
204     if (Dimension == 3) {
205         n=0;
206         for (l=0; l<NeighbourNb; l++)
207             for (i=0; i<one_D; i++)
208                 for (j=0; j<two_D; j++) FillNbAddr (Root->
Neighbour_kL, l, i, j, n);
209         MPI_Type_hindexed(CellElementsNb*NeighbourNb*one_D*two_D, blocklen, disp,
MPI_Type, &MPITypeRkL);
210         MPI_Type_commit (&MPITypeRkL);
211
212         n=0;
213         for (l=0; l<NeighbourNb; l++)
214             for (i=0; i<one_D; i++)
215                 for (j=0; j<two_D; j++) FillNbAddr (Root->
Neighbour_kU, l, i, j, n);
216         MPI_Type_hindexed(CellElementsNb*NeighbourNb*one_D*two_D, blocklen, disp,
MPI_Type, &MPITypeRkU);
217         MPI_Type_commit (&MPITypeRkU);
218
219         n=0;
220         for (l=0; l<NeighbourNb; l++)
221             for (i=0; i<one_D; i++)
222                 for (j=0; j<two_D; j++) {
223                     k=1;
224                     d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
225                     FillCellAddr (MeshCell, d, n);
226                 }
227         MPI_Type_hindexed(CellElementsNb*NeighbourNb*one_D*two_D, blocklen, disp,
MPI_Type, &MPITypeSkL);
228         MPI_Type_commit (&MPITypeSkL);
229
230         n=0;
231         for (l=0; l<NeighbourNb; l++)
232             for (i=0; i<one_D; i++)
233                 for (j=0; j<two_D; j++) {
234                     k=(1<<ScaleNb)-NeighbourNb+1;
235                     d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
236                     FillCellAddr (MeshCell, d, n);
237                 }
238         MPI_Type_hindexed(CellElementsNb*NeighbourNb*one_D*two_D, blocklen, disp,
MPI_Type, &MPITypeSkU);
239         MPI_Type_commit (&MPITypeSkU);
240     }
241
242 #endif
243 }

```

Here is the caller graph for this function:



### 6.6.3.13 int DigitNumber ( int arg )

Returns the number of digits of the integer *arg*.

#### Parameters

|            |          |
|------------|----------|
| <i>arg</i> | Argument |
|------------|----------|

**Returns**

int

```

23 {
24     int result;
25     int i;
26
27     result = 0;
28     i = arg;
29
30     while(i != 0)
31     {
32         i/=10;
33         result++;
34     }
35
36     return result;
37 }

```

Here is the caller graph for this function:

**6.6.3.14 int FileWrite ( FILE \* *f*, const char \* *format*, real *arg* )**

Writes in binary or ASCII mode the real number *arg* into the file *f* with the format *format*. The global parameter *DataIsBinary* determines this choice.

**Parameters**

|               |           |
|---------------|-----------|
| <i>f</i>      | File name |
| <i>format</i> | Format    |
| <i>arg</i>    | Argument  |

**Returns**

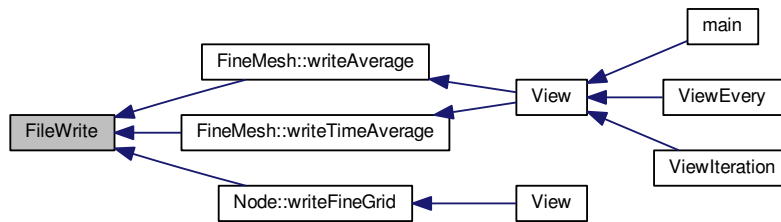
int

```

23 {
24     int result;
25     real x;
26
27     x = arg;
28
29     if (DataIsBinary)
30         result = fwrite(&x, sizeof(real), 1, f);
31     else
32         result = fprintf(f, format, x);
33
34
35     return result;
36 }

```

Here is the caller graph for this function:



### 6.6.3.15 Vector Flux ( Cell & Cell1, Cell & Cell2, Cell & Cell3, Cell & Cell4, int AxisNo )

Returns the flux at the interface between *Cell2* and *Cell3*. Here a 4-point space scheme is used. *Cell2* and *Cell3* are the first neighbours on the left and right sides. *Cell1* and *Cell4* are the second neighbours on the left and right sides.

#### Parameters

|               |                                    |
|---------------|------------------------------------|
| <i>Cell1</i>  | second neighbour on the left side  |
| <i>Cell2</i>  | first neighbour on the left side   |
| <i>Cell3</i>  | first neighbour on the right side  |
| <i>Cell4</i>  | second neighbour on the right side |
| <i>AxisNo</i> | Axis of interest                   |

#### Returns

##### Vector

```

23 {
24     // --- Local variables ---
25
26     Vector Result(QuantityNb);
27
28     Cell C1, C2, C3, C4;
29
30     int BoundaryCell1 = BoundaryRegion(Cell1.center());
31     int BoundaryCell2 = BoundaryRegion(Cell2.center());
32     int BoundaryCell3 = BoundaryRegion(Cell3.center());
33     int BoundaryCell4 = BoundaryRegion(Cell4.center());
34
35     bool UseBoundaryCells = (UseBoundaryRegions && (BoundaryCell1!=0 || BoundaryCell2!=0
|| BoundaryCell3!=0 || BoundaryCell4!=0));
36
37     // --- Take into account boundary conditions ---
38
39     if (UseBoundaryCells)
40         GetBoundaryCells(Cell1, Cell2, Cell3, Cell4, C1, C2, C3, C4, AxisNo);
41
42     switch(SchemeNb)
43     {
44     case 1:
45     default:
46         if (UseBoundaryCells)
47             Result = SchemeHLL(C1, C2, C3, C4, AxisNo);
48         else
49             Result = SchemeHLL(Cell1, Cell2, Cell3, Cell4, AxisNo);
50         break;
51
52     case 2:
53         if (UseBoundaryCells)
54             Result = SchemeHLLD(C1, C2, C3, C4, AxisNo);
55         else
56             Result = SchemeHLLD(Cell1, Cell2, Cell3, Cell4, AxisNo);
57         break;
  
```

```

58     break;
59   }
60 }
61
62
63 return Result;
64 }

```

Here is the caller graph for this function:



### 6.6.3.16 void fluxCorrection ( Vector & Flux, const Vector & AvgL, const Vector & AvgR, int AxisNo )

This function apply the divergence-free correction to the numerical flux.

#### Parameters

|               |                       |
|---------------|-----------------------|
| <i>Flux</i>   | Numerical flux vector |
| <i>AvgL</i>   | Left average vector   |
| <i>AvgR</i>   | Right average vector  |
| <i>AxisNo</i> | Axis of interest      |

#### Returns

void

```

10 {
11     auxvar = Flux.value(AxisNo+6);
12     Flux.setValue(AxisNo+6, Flux.value(AxisNo+6) + (AvgL.value(6) +
13     value(6) - AvgL.value(6))
14     - ch*.5*(AvgR.value(AxisNo+6) - AvgL.
15     value(AxisNo+6)));
16     Flux.setValue(6, ch*ch*(AvgL.value(AxisNo+6) + .5*(AvgR.
17     value(AxisNo+6) - AvgL.value(AxisNo+6))
18     - .5*(AvgR.value(6) - AvgL.value(6))/
19     ch));

```

Here is the caller graph for this function:



### 6.6.3.17 Vector FluxX ( const Vector & Avg )

Returns the physical flux of MHD equations in X direction.

## Parameters

|     |                 |
|-----|-----------------|
| Avg | Average vector. |
|-----|-----------------|

## Returns

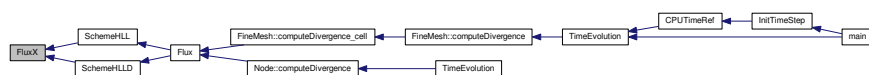
## Vector

```

9 {
10   real rho;
11   real vx, vy, vz;
12   real pre, e;
13   real Bx, By, Bz;
14   real Bx2, By2, Bz2, B2;
15   real vx2, vy2, vz2, v2;
16   real half = 0.5;
17   Vector F(QuantityNb);
18
19   //Variables
20   rho = Avg.value(1);
21   vx = Avg.value(2)/rho;
22   vy = Avg.value(3)/rho;
23   vz = Avg.value(4)/rho;
24   e = Avg.value(5);
25   Bx = Avg.value(7);
26   By = Avg.value(8);
27   Bz = Avg.value(9);
28
29   Bx2 = Bx*Bx;
30   By2 = By*By;
31   Bz2 = Bz*Bz;
32   B2 = half*(Bz2+Bx2+By2);
33
34   vx2 = vx*vx;
35   vy2 = vy*vy;
36   vz2 = vz*vz;
37   v2 = half*(vz2+vx2+vy2);
38
39   //pressure
40   pre = (Gamma -1.)*(e - rho*v2 - B2);
41
42   //Physical flux - x-direction
43   F.setValue(1, rho*vx);
44   F.setValue(2, rho*vx2 + pre + half*(Bz2+By2-Bx2));
45   F.setValue(3, rho*vx*vy - Bx*By);
46   F.setValue(4, rho*vx*vz - Bx*Bz);
47   F.setValue(5, (e + pre + B2)*vx - Bx*(vx*Bx + vy*By + vz*Bz) );
48   F.setValue(6, 0.0);
49   F.setValue(7, 0.0);
50   F.setValue(8, vx*By - vy*Bx);
51   F.setValue(9, vx*Bz - vz*Bx);
52
53   return F;
54 }
55 }

```

Here is the caller graph for this function:



### 6.6.3.18 Vector FluxY ( const Vector & Avg )

Returns the physical flux of MHD equations in Y direction.



Parameters

|     |                 |
|-----|-----------------|
| Avg | Average vector. |
|-----|-----------------|

Returns

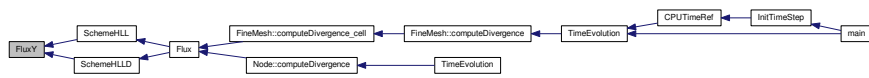
Vector

```

59 {
60   real rho;
61   real vx, vy, vz;
62   real pre, e;
63   real Bx, By, Bz;
64   real Bx2, By2, Bz2, B2;
65   real vx2, vy2, vz2, v2;
66   real half = 0.5;
67
68   Vector G(QuantityNb);
69
70   //Variables
71   rho = Avg.value(1);
72   vx = Avg.value(2)/rho;
73   vy = Avg.value(3)/rho;
74   vz = Avg.value(4)/rho;
75   e = Avg.value(5);
76   Bx = Avg.value(7);
77   By = Avg.value(8);
78   Bz = Avg.value(9);
79
80   Bx2 = Bx*Bx;
81   By2 = By*By;
82   Bz2 = Bz*Bz;
83   B2 = half*(Bz2+Bx2+By2);
84
85   vx2 = vx*vx;
86   vy2 = vy*vy;
87   vz2 = vz*vz;
88   v2 = half*(vz2+vx2+vy2);
89
90   //pressure
91   pre = (Gamma -1.)*(e - rho*v2 - B2);
92
93   //Physical flux - y-direction
94   G.setValue(1,rho*vy);
95   G.setValue(2,rho*vx*vy - Bx*By);
96   G.setValue(3,rho*vy2 + pre + half*(Bx2+Bz2-By2));
97   G.setValue(4,rho*vy*vz - By*Bz);
98   G.setValue(5,(e + pre + B2)*vy - By*(vx*Bx + vy*By + vz*Bz));
99   G.setValue(6,0.0);
100  G.setValue(7,vy*Bx - vx*By);
101  G.setValue(8,0.0);
102  G.setValue(9,vy*Bz - vz*By);
103  return G;
104 }

```

Here is the caller graph for this function:



6.6.3.19 Vector FluxZ ( const Vector & Avg )

Returns the physical flux of MHD equations in Z direction.

Parameters

|     |                 |
|-----|-----------------|
| Avg | Average vector. |
|-----|-----------------|

## Returns

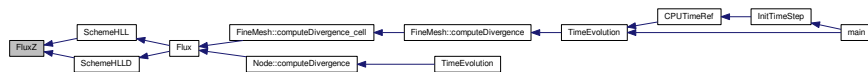
### Vector

```

107 {
108   real rho;
109   real vx, vy, vz;
110   real pre, e;
111   real Bx, By, Bz;
112   real Bx2, By2, Bz2, B2;
113   real vx2, vy2, vz2, v2;
114   real half = 0.5;
115
116   Vector H(QuantityNb);
117
118   //Variables
119   rho = Avg.value(1);
120   vx = Avg.value(2)/rho;
121   vy = Avg.value(3)/rho;
122   vz = Avg.value(4)/rho;
123   e = Avg.value(5);
124   Bx = Avg.value(7);
125   By = Avg.value(8);
126   Bz = Avg.value(9);
127
128   Bx2 = Bx*Bx;
129   By2 = By*By;
130   Bz2 = Bz*Bz;
131   B2 = half*(Bz2+Bx2+By2);
132
133   vx2 = vx*vx;
134   vy2 = vy*vy;
135   vz2 = vz*vz;
136   v2 = half*(vz2+vx2+vy2);
137
138   //pressure
139   pre = (Gamma -1.)*(e - rho*v2 - B2);
140
141   //Physical flux - y-direction
142   H.setValue(1, rho+vz);
143   H.setValue(2, rho+vz*vx - Bz*Bx);
144   H.setValue(3, rho+vz*vy - Bz*By);
145   H.setValue(4, rho+vz2 + pre + half*(Bx2+By2-Bz2));
146   H.setValue(5, (e + pre + B2)*vz - Bz*(vx*Bx + vy*By + vz*Bz));
147   H.setValue(6, 0.0);
148   H.setValue(7, vz*Bx - vx*Bz);
149   H.setValue(8, vz*By - vy*Bz);
150   H.setValue(9, 0.0);
151
152   return H;
153 }

```

Here is the caller graph for this function:



### 6.6.3.20 void FreeMPIType ( )

Parallel function DOES NOT WORK!

## Returns

void

```

247   {
248   #if defined PARMPI
249     MPI_Type_free (&MPITypeSiL);

```

```

250 MPI_Type_free (&MPItypeSiU);
251 MPI_Type_free (&MPItypeRiL);
252 MPI_Type_free (&MPItypeRiU);
253
254 if (Dimension >= 2) {
255     MPI_Type_free (&MPItypeSjL);
256     MPI_Type_free (&MPItypeSjU);
257     MPI_Type_free (&MPItypeRjL);
258     MPI_Type_free (&MPItypeRjU);
259 }
260
261 if (Dimension == 3) {
262     MPI_Type_free (&MPItypeSkL);
263     MPI_Type_free (&MPItypeSkU);
264     MPI_Type_free (&MPItypeRkL);
265     MPI_Type_free (&MPItypeRkU);
266 }
267 #endif
268 }
    
```

Here is the caller graph for this function:



**6.6.3.21 void GetBoundaryCells ( Cell & Cell1, Cell & Cell2, Cell & Cell3, Cell & Cell4, Cell & C1, Cell & C2, Cell & C3, Cell & C4, int AxisNo )**

Transform the 4 cells of the flux *Cell1*, *Cell2*, *Cell3*, *Cell4* into *C1*, *C2*, *C3*, *C4*, to take into account boundary conditions (used in [Flux.cpp](#)).

**Parameters**

|               |                                    |
|---------------|------------------------------------|
| <i>Cell1</i>  | second neighbour on the left side  |
| <i>Cell2</i>  | first neighbour on the left side   |
| <i>Cell3</i>  | first neighbour on the right side  |
| <i>Cell4</i>  | second neighbour on the right side |
| <i>C1</i>     | Auxiliar cell1                     |
| <i>C2</i>     | Auxiliar cell2                     |
| <i>C3</i>     | Auxiliar cell3                     |
| <i>C4</i>     | Auxiliar cell4                     |
| <i>AxisNo</i> | ...                                |

**Returns**

void

```

26 {
27     // --- Local variables ---
28
29     int InCell1, InCell2, InCell3, InCell4; // Boundary conditions in cells 1, 2, 3, 4
30     real P1, P2, P3, P4; // Pressures in cells 1, 2, 3, 4
31     real T1, T2, T3, T4; // Temperatures in cells 1, 2, 3, 4
32     real rho1, rho2, rho3, rho4; // Densities in cells 1, 2, 3, 4
33     Vector V1(Dimension), V2(Dimension), V3(Dimension), V4(
Dimension); // Velocities in cells 1, 2, 3, 4
34     real e1, e2, e3, e4; // Energies in cell 1, 2, 3, 4
35     real Y1=0., Y2=0., Y3=0., Y4=0.; // Partial mass in cell 1, 2, 3, 4
36
37     int i; // Counter
38
39     // --- Init C1, C2, C3, C4 ---
40
41     C1 = Cell1;
    
```

```

42     C2 = Cell2;
43     C3 = Cell3;
44     C4 = Cell4;
45
46     // --- Depending on the boundary region type, transform C1, C2, C3, C4 ---
47
48     InCell1 = BoundaryRegion(Cell1.center());
49     InCell2 = BoundaryRegion(Cell2.center());
50     InCell3 = BoundaryRegion(Cell3.center());
51     InCell4 = BoundaryRegion(Cell4.center());
52
53     // --- Cell2 IN THE BOUNDARY, Cell3 IN THE FLUID -----
54
55     if (InCell2 != 0 && InCell3 == 0)
56     {
57         switch(InCell2)
58         {
59             // INFLOW
60             case 1:
61                 // Dirichlet on temperature
62                 T2 = Cell2.temperature();
63                 T1 = Cell1.temperature();
64
65                 // Extrapolate pressure
66                 P2 = Cell3.oldPressure();
67                 P1 = P2;
68                 // P1 = 2*P2 - Cell3.pressure();
69
70                 // Compute density
71                 rho2 = Gamma*Ma*Ma*P2/T2;
72                 rho1 = Gamma*Ma*Ma*P1/T1;
73
74                 // Dirichlet on momentum
75                 V2 = (Cell2.density()/rho2)*Cell2.velocity();
76                 V1 = (Cell1.density()/rho1)*Cell1.velocity();
77
78                 // Dirichlet on partial mass
79                 if (ScalarEqNb == 1)
80                 {
81                     Y2 = Cell2.average(Dimension+3)/Cell2.
density();
82                     Y1 = Cell1.average(Dimension+3)/Cell1.
density();
83                 }
84
85                 // Compute energies
86                 e2 = P2/((Gamma-1.)*rho2) + 0.5*N2(V2);
87                 e1 = P1/((Gamma-1.)*rho1) + 0.5*N2(V1);
88
89                 // Correct C1 and C2
90                 C2.setAverage(1, rho2);
91                 C1.setAverage(1, rho1);
92                 for (i=1; i<=Dimension; i++)
93                 {
94                     C2.setAverage(i+1, rho2*V2.value(i));
95                     C1.setAverage(i+1, rho1*V1.value(i));
96                 }
97                 C2.setAverage(Dimension+2, rho2*e2);
98                 C1.setAverage(Dimension+2, rho1*e1);
99
100                if (ScalarEqNb == 1)
101                {
102                    C2.setAverage(Dimension+3, rho2*Y2);
103                    C1.setAverage(Dimension+3, rho1*Y1);
104                }
105                break;
106
107                // OUTFLOW : use the old value of the neighbour
108                case 2:
109                    C2.setAverage(Cell3.oldAverage());
110                    C1.setAverage(Cell3.oldAverage());
111
112                    // Also change the values in the boundary
113                    Cell2.setAverage(C2.average());
114                    Cell1.setAverage(C1.average());
115                    break;
116
117                // FREE-SLIP WALL : Neuman on all quantities
118                case 3:
119
120                    C2 = Cell3;
121                    C1 = Cell4;
122                    break;
123
124                // ADIABATIC WALL
125                case 4:
126

```

```

127         // Dirichlet on velocity
128         V2 = Cell12.velocity();
129         V1 = Cell11.velocity();
130
131         // Neuman on temperature
132         T2 = Cell13.temperature();
133         T1 = Cell14.temperature();
134
135         // Neuman on pressure
136         P2 = Cell13.pressure();
137         P1 = Cell14.pressure();
138
139         // Extrapolate pressure
140         //P2 = 2*Cell13.pressure()-Cell14.pressure();
141         //P1 = P2;
142
143         // Compute densities
144         rho2 = Gamma*Ma*Ma*P2/T2;
145         rho1 = Gamma*Ma*Ma*P1/T1;
146
147         // Compute energies
148         e2 = P2/((Gamma-1.)*rho2) + 0.5*N2(V2);
149         e1 = P1/((Gamma-1.)*rho1) + 0.5*N2(V1);
150
151         // Correct C1 and C2
152         C2.setAverage(1, rho2);
153         C1.setAverage(1, rho1);
154         for (i=1; i<=Dimension; i++)
155         {
156             C2.setAverage(i+1, rho2*V2.value(i));
157             C1.setAverage(i+1, rho1*V1.value(i));
158         }
159         C2.setAverage(Dimension+2, rho2*e2);
160         C1.setAverage(Dimension+2, rho1*e1);
161
162         // Neuman on partial mass
163         if (ScalarEqNb == 1)
164         {
165             C2.setAverage(Dimension+3, Cell13.average(
Dimension+3));
166             C1.setAverage(Dimension+3, Cell14.average(
Dimension+3));
167         }
168         break;
169
170     // ISOTHERMAL WALL
171     case 5:
172
173         // Dirichlet on velocity
174         V2 = Cell12.velocity();
175         V1 = Cell11.velocity();
176
177         // Dirichlet on temperature
178         T2 = Cell12.temperature();
179         T1 = Cell11.temperature();
180
181         // Neuman on pressure
182         P2 = Cell13.pressure();
183         P1 = Cell14.pressure();
184
185         // Extrapolate pressure
186         //P2 = 2*Cell13.pressure()-Cell14.pressure();
187         //P1 = P2;
188
189         // Compute densities
190         rho2 = Gamma*Ma*Ma*P2/T2;
191         rho1 = Gamma*Ma*Ma*P1/T1;
192
193         // Compute energies
194         e2 = P2/((Gamma-1.)*rho2) + 0.5*N2(V2);
195         e1 = P1/((Gamma-1.)*rho1) + 0.5*N2(V1);
196
197         // Correct C1 and C2
198         C2.setAverage(1, rho2);
199         C1.setAverage(1, rho1);
200         for (i=1; i<=Dimension; i++)
201         {
202             C2.setAverage(i+1, rho2*V2.value(i));
203             C1.setAverage(i+1, rho1*V1.value(i));
204         }
205         C2.setAverage(Dimension+2, rho2*e2);
206         C1.setAverage(Dimension+2, rho1*e1);
207
208         // Neuman on partial mass
209         if (ScalarEqNb == 1)
210         {
211

```

```

212         C2.setAverage(Dimension+3, Cell3.average(
Dimension+3));
213         C1.setAverage(Dimension+3, Cell4.average(
Dimension+3));
214     }
215
216         break;
217     };
218     return;
219 }
220
221 // --- Cell1 IN THE BOUNDARY, Cell2 IN THE FLUID -----
222
223     if (InCell1 != 0 && InCell2 == 0)
224     {
225         switch(InCell1)
226         {
227             // INFLOW
228             case 1:
229                 // Dirichlet on temperature
230                 T1 = Cell1.temperature();
231
232                 // Extrapolate pressure from old value
233                 P1 = Cell2.oldPressure();
234
235                 // Compute density
236                 rho1 = Gamma*Ma*Ma*P1/T1;
237
238                 // Dirichlet on momentum
239                 V1 = (Cell1.density()/rho1)*Cell1.velocity();
240
241                 // Dirichlet on partial mass
242                 if (ScalarEqNb == 1)
243                     Y1 = Cell1.average(Dimension+3)/Cell1.
density();
244
245                 // Compute energies
246                 e1 = P1/((Gamma-1.)*rho1) + 0.5*N2(V1);
247
248                 // Correct C1
249                 C1.setAverage(1, rho1);
250                 for (i=1; i<=Dimension; i++)
251                     C1.setAverage(i+1, rho1*V1.value(i));
252                 C1.setAverage(Dimension+2, rho1*e1);
253
254                 if (ScalarEqNb == 1)
255                     C1.setAverage(Dimension+3, rho1*Y1);
256                 break;
257
258             // OUTFLOW : Get old value of the neighbour
259             case 2:
260
261                 C1.setAverage(Cell2.oldAverage());
262                 break;
263
264             // FREE-SLIP WALL : Neuman on all quantities
265             case 3:
266
267                 C1 = Cell2;
268                 break;
269
270             // ADIABATIC WALL
271             case 4:
272
273                 // Dirichlet on velocity
274                 V1 = Cell1.velocity();
275
276                 // Neuman on temperature
277                 T1 = Cell2.temperature();
278
279                 // Neuman on pressure
280                 P1 = Cell2.pressure();
281
282                 // Extrapolate pressure
283                 //P1 = 2*Cell2.pressure()-Cell3.pressure();
284
285                 // Compute density
286                 rho1 = Gamma*Ma*Ma*P1/T1;
287
288                 // Compute energy
289                 e1 = P1/((Gamma-1.)*rho1) + 0.5*N2(V1);
290
291                 // Correct C1
292                 C1.setAverage(1, rho1);
293                 for (i=1; i<=Dimension; i++)
294                     C1.setAverage(i+1, rho1*V1.value(i));
295                 C1.setAverage(Dimension+2, rho1*e1);

```

```

296
297     // Neuman on partial mass
298     if (ScalarEqNb == 1)
299         C1.setAverage(Dimension+3, Cell2.average(
Dimension+3));
300
301         break;
302
303     // ISOTHERMAL WALL
304     case 5:
305
306         // Dirichlet on velocity
307         V1 = Cell1.velocity();
308
309         // Dirichlet on temperature
310         T1 = Cell1.temperature();
311
312         // Neuman on pressure
313         P1 = Cell2.pressure();
314
315         // Extrapolate pressure
316         //P1 = 2*Cell2.pressure()-Cell3.pressure();
317
318         // Compute density
319         rho1 = Gamma*Ma*Ma*P1/T1;
320
321         // Compute energies
322         e1 = P1/((Gamma-1.)*rho1) + 0.5*N2(V1);
323
324         // Correct C1
325         C1.setAverage(1, rho1);
326         for (i=1; i<=Dimension; i++)
327             C1.setAverage(i+1, rho1*V1.value(i));
328         C1.setAverage(Dimension+2, rho1*e1);
329
330         // Neuman on partial mass
331         if (ScalarEqNb == 1)
332             C1.setAverage(Dimension+3, Cell2.average(
Dimension+3));
333
334         break;
335     };
336     return;
337 }
338 // --- Cell13 IN THE BOUNDARY, Cell12 IN THE FLUID -----
339
340 if (InCell13 !=0 && InCell12 == 0)
341 {
342     switch(InCell13)
343     {
344         // INFLOW
345         case 1:
346             // Dirichlet on temperature
347             T3 = Cell13.temperature();
348             T4 = Cell14.temperature();
349
350             // Extrapolate pressure from old value
351             P3 = Cell2.oldPressure();
352             P4 = P3;
353             //P4 = 2*P3 - Cell2.pressure();
354
355             // Compute densities
356             rho3 = Gamma*Ma*Ma*P3/T3;
357             rho4 = Gamma*Ma*Ma*P4/T4;
358
359             // Dirichlet on momentum
360             V3 = (Cell13.density()/rho3)*Cell13.velocity();
361             V4 = (Cell14.density()/rho4)*Cell14.velocity();
362
363             // Dirichlet on partial mass
364             if (ScalarEqNb == 1)
365             {
366                 Y3 = Cell13.average(Dimension+3)/Cell13.
density();
367                 Y4 = Cell14.average(Dimension+3)/Cell14.
density();
368             }
369
370             // Compute energies
371             e3 = P3/((Gamma-1.)*rho3) + 0.5*N2(V3);
372             e4 = P4/((Gamma-1.)*rho4) + 0.5*N2(V4);
373
374             // Correct C1 and C2
375             C3.setAverage(1, rho3);
376             C4.setAverage(1, rho4);
377             for (i=1; i<=Dimension; i++)
378             {

```

```

379         C3.setAverage(i+1, rho3*V3.value(i));
380         C4.setAverage(i+1, rho4*V4.value(i));
381     }
382     C3.setAverage(Dimension+2, rho3*e3);
383     C4.setAverage(Dimension+2, rho4*e4);
384
385     if (ScalarEqNb == 1)
386     {
387         C3.setAverage(Dimension+3, rho3*Y3);
388         C4.setAverage(Dimension+3, rho4*Y4);
389     }
390     break;
391
392     // OUTFLOW
393     case 2:
394
395         C3.setAverage(Cell2.oldAverage());
396         C4.setAverage(Cell2.oldAverage());
397         //C4.setAverage(2*C3.average()-Cell2.average());
398
399         // Also change the values in the boundary
400         Cell3.setAverage(C3.average());
401         Cell4.setAverage(C4.average());
402         break;
403
404     // FREE-SLIP WALL : Neuman on all quantities
405     case 3:
406
407         C3 = Cell2;
408         C4 = Cell1;
409         break;
410
411     // ADIABATIC WALL
412     case 4:
413
414         // Dirichlet on velocity
415         V3 = Cell3.velocity();
416         V4 = Cell4.velocity();
417
418         // Neuman on temperature
419         T3 = Cell2.temperature();
420         T4 = Cell1.temperature();
421
422         // Neuman on pressure
423         P3 = Cell2.pressure();
424         P4 = Cell1.pressure();
425
426         // Extrapolate pressure
427         //P3 = 2*Cell2.pressure()-Cell1.pressure();
428         //P4 = P3;
429
430         // Compute densities
431         rho3 = Gamma*Ma*Ma*P3/T3;
432         rho4 = Gamma*Ma*Ma*P4/T4;
433
434         // Compute energies
435         e3 = P3/((Gamma-1.)*rho3) + 0.5*N2(V3);
436         e4 = P4/((Gamma-1.)*rho4) + 0.5*N2(V4);
437
438         // Correct C3 and C4
439         C3.setAverage(1, rho3);
440         C4.setAverage(1, rho4);
441         for (i=1; i<=Dimension; i++)
442         {
443             C3.setAverage(i+1, rho3*V3.value(i));
444             C4.setAverage(i+1, rho4*V4.value(i));
445         }
446         C3.setAverage(Dimension+2, rho3*e3);
447         C4.setAverage(Dimension+2, rho4*e4);
448
449         // Neuman on partial mass
450         if (ScalarEqNb == 1)
451         {
452             C3.setAverage(Dimension+3, Cell2.average(
Dimension+3));
453             C4.setAverage(Dimension+3, Cell1.average(
Dimension+3));
454         }
455         break;
456
457     // ISOTHERMAL WALL
458     case 5:
459
460         // Dirichlet on velocity
461         V3 = Cell3.velocity();
462         V4 = Cell4.velocity();
463

```



```

464
465 // Dirichlet on temperature
466 T3 = Cell3.temperature();
467 T4 = Cell4.temperature();
468
469 // Neuman on pressure
470 P3 = Cell2.pressure();
471 P4 = Cell1.pressure();
472
473 // Extrapolate pressure
474 //P3 = 2*Cell2.pressure()-Cell1.pressure();
475 //P4 = P3;
476
477 // Compute densities
478 rho3 = Gamma*Ma*Ma*P3/T3;
479 rho4 = Gamma*Ma*Ma*P4/T4;
480
481 // Compute energies
482 e3 = P3/((Gamma-1.)*rho3) + 0.5*N2(V3);
483 e4 = P4/((Gamma-1.)*rho4) + 0.5*N2(V4);
484
485 // Correct C3 and C4
486 C3.setAverage(1, rho3);
487 C4.setAverage(1, rho4);
488 for (i=1; i<=Dimension; i++)
489 {
490     C3.setAverage(i+1, rho3*V3.value(i));
491     C4.setAverage(i+1, rho4*V4.value(i));
492 }
493 C3.setAverage(Dimension+2, rho3*e3);
494 C4.setAverage(Dimension+2, rho4*e4);
495
496 // Neuman on partial mass
497 if (ScalarEqNb == 1)
498 {
499     C3.setAverage(Dimension+3, Cell2.average(
500 Dimension+3));
501     C4.setAverage(Dimension+3, Cell1.average(
502 Dimension+3));
503 }
504 break;
505 };
506 return;
507 }
508 // --- Cell4 IN THE BOUNDARY, Cell3 IN THE FLUID -----
509
510 if (InCell4 != 0 && InCell3 == 0)
511 {
512     switch(InCell4)
513     {
514         // INFLOW
515         case 1:
516             // Dirichlet on temperature
517             T4 = Cell4.temperature();
518
519             // Extrapolate pressure from old value
520             P4 = Cell3.oldPressure();
521
522             // Compute density
523             rho4 = Gamma*Ma*Ma*P4/T4;
524
525             // Dirichlet on momentum
526             V4 = (Cell4.density()/rho4)*Cell4.velocity();
527
528             // Dirichlet on partial mass
529             if (ScalarEqNb == 1)
530                 Y4 = Cell4.average(Dimension+3)/Cell4.
531 density();
532
533             // Compute energies
534             e4 = P4/((Gamma-1.)*rho4) + 0.5*N2(V4);
535
536             // Correct C4
537             C4.setAverage(1, rho4);
538             for (i=1; i<=Dimension; i++)
539                 C4.setAverage(i+1, rho4*V4.value(i));
540             C4.setAverage(Dimension+2, rho4*e4);
541
542             if (ScalarEqNb == 1)
543                 C4.setAverage(Dimension+3, rho4*Y4);
544             break;
545
546             // OUTFLOW : Use old cell-average values of the neighbour
547             case 2:

```

```

548         C4.setAverage(Cell13.oldAverage());
549         break;
550
551     // FREE-SLIP WALL : Neuman on all quantities
552     case 3:
553
554         C4 = Cell13;
555         break;
556
557     // ADIABATIC WALL
558     case 4:
559
560         // Dirichlet on velocity
561         V4 = Cell14.velocity();
562
563         // Neuman on temperature
564         T4 = Cell13.temperature();
565
566         // Neuman on pressure
567         P4 = Cell13.pressure();
568
569         // Extrapolate pressure
570         //P4 = 2*Cell13.pressure()-Cell12.pressure();
571
572         // Compute density
573         rho4 = Gamma*Ma*Ma*P4/T4;
574
575         // Compute energy
576         e4 = P4/((Gamma-1.)*rho4) + 0.5*N2(V4);
577
578         // Correct C4
579         C4.setAverage(1, rho4);
580         for (i=1; i<=Dimension; i++)
581             C4.setAverage(i+1, rho4*V4.value(i));
582         C4.setAverage(Dimension+2, rho4*e4);
583
584         // Neuman on partial mass
585         if (ScalarEqNb == 1)
586             C4.setAverage(Dimension+3, Cell13.average(
Dimension+3));
587
588         break;
589
590     // ISOTHERMAL WALL
591     case 5:
592
593         // Dirichlet on velocity
594         V4 = Cell14.velocity();
595
596         // Dirichlet on temperature
597         T4 = Cell14.temperature();
598
599         // Neuman on pressure
600         P4 = Cell13.pressure();
601
602         // Extrapolate pressure
603         //P4 = 2*Cell13.pressure()-Cell12.pressure();
604
605         // Compute density
606         rho4 = Gamma*Ma*Ma*P4/T4;
607
608         // Compute energies
609         e4 = P4/((Gamma-1.)*rho4) + 0.5*N2(V4);
610
611         // Correct C4
612         C4.setAverage(1, rho4);
613         for (i=1; i<=Dimension; i++)
614             C4.setAverage(i+1, rho4*V4.value(i));
615         C4.setAverage(Dimension+2, rho4*e4);
616
617         // Neuman on partial mass
618         if (ScalarEqNb == 1)
619             C4.setAverage(Dimension+3, Cell13.average(
Dimension+3));
620
621         break;
622     };
623     return;
624 }
625
626 }

```

Here is the caller graph for this function:



### 6.6.3.22 Vector InitAverage ( real x, real y = 0., real z = 0. )

Returns the initial condition in  $(x, y, z)$  form the one defined in *carmen.ini*.

#### Parameters

|   |                             |
|---|-----------------------------|
| x | Position x                  |
| y | Position y. Defaults to 0.. |
| z | Position z. Defaults to 0.. |

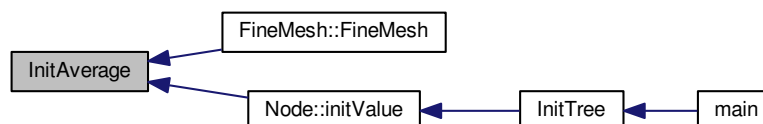
#### Returns

Vector

```

23 {
24     // --- Local variables ---
25
26     Vector Result(QuantityNb);
27     real *Q;
28     Q = new real [QuantityNb+1];
29     int n;
30
31     // --- Init Q ---
32
33     for (n = 1; n <= QuantityNb; n++)
34         Q[n]=0.;
35
36     // --- Use definition of initial Q contained in file 'initial' ---
37
38     #include "carmen.ini"
39
40     // --- Fill vector Result and return it ---
41
42     for (n = 1; n <= QuantityNb; n++)
43         Result.setValue(n, Q[n]);
44
45     delete[] Q;
46
47     return Result;
48 }
  
```

Here is the caller graph for this function:



### 6.6.3.23 void InitParameters ( )

Initns parameters from file *carmen.par*. If a parameter is not mentioned in this file, the default value is used.

## Returns

void

— Compute ch -----

```

284 {
285     // --- Local variables -----
286
287     int    i;          // Counter
288
289     // --- Set global variables from file "carmen.par" -----
290
291     #include "carmen.par"
292
293     // --- Adapt IterationNbRef to the dimension -----
294
295     IterationNbRef=(int) (exp((4.-Dimension)*log(10.)));
296
297     // --- Compute the number of children of a given parent cell ---
298
299     ChildNb = (1<<Dimension);
300
301 #if defined PARMPI
302
303     AllTaskScaleNb=ScaleNb;
304     for (i=0;i<4;i++)
305     {
306         AllXMax[i]=XMax[i];
307         AllXMin[i]=XMin[i];
308     }
309
310     //some combinations give deadlock...
311     MPISendType = 10;    //0 - Ibsend; 10 - Isend; 20 - Issend;
312     MPIRecvType = 1;    //0 - Recv; 1 - Irecv;
313
314     CPUScales=0;
315     int tmp=size;
316     while ((tmp=(tmp>>1))>0) CPUScales++;
317     ScaleNb=CPUScales/Dimension;
318
319     one_D=1; two_D=1;
320     if (Dimension >= 2) one_D=1<<ScaleNb;
321     if (Dimension == 3) two_D=1<<ScaleNb;
322
323 //#if defined PARMPI
324
325     NeighbourNb=2;
326     MaxCellElementsNb=6;
327
328     // -- Create memory arrays that are needs for the MPI Type creation ---
329     disp = new MPI_Aint [NeighbourNb*MaxCellElementsNb*
one_D*two_D];
330     blocklen = new int [NeighbourNb*MaxCellElementsNb*
one_D*two_D];
331
332     // --- Allocate additional memory for MPI buffer send---
333     Cell tc;
334     int CellElNb,bufsize;
335     CellElNb=tc.size().dimension()+tc.center().dimension()+tc.average().dimension()+tc.
tempAverage().dimension()+tc.divergence().dimension();
336
337     if (EquationType==6)
338         CellElNb += tc.gradient().lines()*tc.gradient().columns();
339
340     bufsize=(CellElNb*one_D*two_D*NeighbourNb+MPI_BSEND_OVERHEAD)*2*
Dimension+1024;
341     MPIbuffer=new real [bufsize];
342     MPI_Buffer_attach(MPIbuffer,bufsize*sizeof(real));
343
344 #else
345     NeighbourNb=0;
346 #endif
347
348 #if defined PARMPI
349     CreateMPITopology();
350
351     // --- Compute domain coordinates for the processors ---
352     XMin[1] = AllXMin[1] + coords[0]*(AllXMax[1]-AllXMin[1])/
CartDims[0];
353     XMax[1] = AllXMin[1] + (coords[0]+1)*(AllXMax[1]-
AllXMin[1])/CartDims[0];
354
355     if (Dimension >= 2)
356     {

```

```

357     XMin[2] = AllXMin[2] + coords[1]*(AllXMax[2]-
AllXMin[2])/CartDims[1];
358     XMax[2] = AllXMin[2] + (coords[1]+1)*(AllXMax[2]-
AllXMin[2])/CartDims[1];
359 }
360
361 if (Dimension == 3)
362 {
363     XMin[3] = AllXMin[3] + coords[2]*(AllXMax[3]-
AllXMin[3])/CartDims[2];
364     XMax[3] = AllXMin[3] + (coords[2]+1)*(AllXMax[3]-
AllXMin[3])/CartDims[2];
365 }
366
367 // --- Set the backup file name for the current processor
368 sprintf(BackupName,"%d_%d_%d_%s",coords[0],coords[1],
coords[2],"carmen.bak");
369 #else
370 sprintf(BackupName,"%s","carmen.bak");
371 #endif
372
373
374 // --- Use CVS only if Dimension > 1 -----
375
376 if (Dimension == 1)
377     CVS = false;
378
379 // --- TimeAveraging always false if not Navier-Stokes -----
380
381 if (EquationType != 6)
382     TimeAveraging = false;
383
384 // --- If there is no file "carmen.bak", set Recovery=false -----
385
386 if (!fopen(BackupName,"r"))
387     Recovery = false;
388
389 // --- If PrintMoreScales != 0 or 1 with Multiresolution = false, print error and stop ---
390
391 if (!Multiresolution && (!(PrintMoreScales == 0 ||
PrintMoreScales == -1)) )
392 {
393     cout << "Parameters.cpp: In method 'void InitParameters()':\n";
394     cout << "Parameters.cpp: value of PrintMoreScales incompatible with FV computations\n";
395     cout << "Parameters.cpp: must be 0 or -1\n";
396     cout << "carmen: *** [Parameters.o] Execution error\n";
397     cout << "carmen: abort execution.\n";
398     exit(1);
399 }
400
401 // --- Compute global volume -----
402
403 GlobalVolume = fabs(XMax[1]-XMin[1]);
404
405 if (Dimension > 1)
406     GlobalVolume *= fabs(XMax[2]-XMin[2]);
407
408 if (Dimension > 2)
409     GlobalVolume *= fabs(XMax[3]-XMin[3]);
410
411 // --- Compute PostProcessing and DataIsBinary -----
412
413 // In 1D, use Gnuplot instead of Data Explorer
414
415 if (Dimension == 1 && PostProcessing == 2)
416     PostProcessing = 1;
417
418 // In 2D-3D, use Data Explorer instead of Gnuplot
419
420 if (Dimension != 1 && PostProcessing == 1)
421     PostProcessing = 2;
422
423 // --- Compute number of conservative quantities -----
424
425 QuantityNb = 9;
426
427 // --- Set the dimension of QuantityMax to QuantityNb -----
428
429 QuantityMax.setDimension(QuantityNb);
430
431 // --- Set the dimension of QuantityAverage to 4 (pressure, vorticity, entropy, volume)
432
433 QuantityAverage.setDimension(4);
434
435 // --- Set the dimension of IntMomentum to dimension -----
436
437 IntMomentum.setDimension(Dimension);

```

```

438
439 // --- Compute minimal space step -----
440
441 SpaceStep = fabs(XMax[1]-XMin[1]);
442
443 for (i = 2; i <= Dimension; i++)
444     SpaceStep = Min(SpaceStep, fabs(XMax[i]-XMin[i]));
445
446 SpaceStep /= (1<<ScaleNb);
447
448 // --- Compute time step from CFL if TimeStep = 0 -----
449
450 if (TimeStep == 0.)
451 {
452     if (fabs(Eigenvalue)>0.0e-20)
453         TimeStep = CFL*SpaceStep/Eigenvalue;
454     else
455         TimeStep = 0.0001;
456 }
457 else
458     ConstantTimeStep = true;
459
460 ch = CFL*SpaceStep/TimeStep;
461
462
463 }

```

Here is the caller graph for this function:



#### 6.6.3.24 real InitResistivity ( real x, real y=0., real z=0. )

Returns the initial resistivity condition in (x, y, z) form the one defined in *carmen.eta*.

##### Parameters

|   |                             |
|---|-----------------------------|
| x | Position x                  |
| y | Position y. Defaults to 0.. |
| z | Position z. Defaults to 0.. |

##### Returns

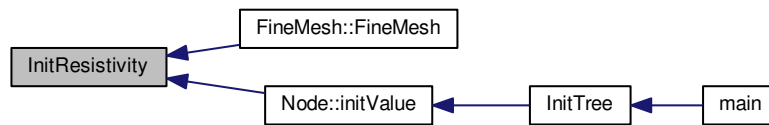
double

```

51 {
52     // --- Local variables ---
53
54     real Result=0.;
55     real Res = 0.;
56
57     #include "carmen.eta"
58
59     Result = Res;
60
61     return Result;
62 }

```

Here is the caller graph for this function:



### 6.6.3.25 void InitTimeStep ( )

Init time step and all the parameters which depend on it.

#### Returns

void

```

23 {
24
25     // --- Init TimeStep -----
26
27     if (TimeStep == 0)
28     {
29         if (Resistivity) TimeStep = CFL*SpaceStep/(
Eigenvalue + eta);
30         else TimeStep = CFL*SpaceStep/Eigenvalue;
31     }
32
33     // --- Compute number of iterations -----
34
35     if (PhysicalTime != 0. && IterationNb == 0)
36         IterationNb = (int)(ceil(PhysicalTime/TimeStep));
37
38     // --- Compute Refresh -----
39
40     if (Refresh == 0)
41         Refresh = (int)(ceil(IterationNb/(RefreshNb*1.)));
42
43     // --- Compute PrintEvery -----
44
45     if ((PrintEvery == 0)&&(ImageNb != 0))
46         PrintEvery = (int)(ceil(IterationNb/(ImageNb*1.)));
47
48     // --- Compute iterations for print -----
49
50     if (PrintTime1 != 0.)
51         PrintIt1 = (int)(ceil(PrintTime1/TimeStep));
52
53     if (PrintTime2 != 0.)
54         PrintIt2 = (int)(ceil(PrintTime2/TimeStep));
55
56     if (PrintTime3 != 0.)
57         PrintIt3 = (int)(ceil(PrintTime3/TimeStep));
58
59     if (PrintTime4 != 0.)
60         PrintIt4 = (int)(ceil(PrintTime4/TimeStep));
61
62     if (PrintTime5 != 0.)
63         PrintIt5 = (int)(ceil(PrintTime5/TimeStep));
64
65     if (PrintTime6 != 0.)
66         PrintIt6 = (int)(ceil(PrintTime6/TimeStep));
67
68     // --- Compute FV reference time -----
69
70     if (Multiresolution)
71         FVTimeRef = CPUTimeRef(IterationNbRef,
ScaleNbRef);
72
73 }
  
```

Here is the caller graph for this function:



### 6.6.3.26 void InitTree ( Node \* Root )

Inits tree structure from initial condition, starting from the node *Root*. Only for multiresolution computations.

#### Parameters

|             |      |
|-------------|------|
| <i>Root</i> | Root |
|-------------|------|

#### Returns

void

```

23 {
24     // --- Local variables ---
25
26     int l; // Counter on levels
27
28     // --- Init cell-average value in root and split it ---
29
30     if (Recovery && UseBackup)
31         Root->restore();
32     else
33     {
34         Root->initValue();
35
36         // --- Add and init nodes in different levels, when necessary ---
37
38         for (l=1; l <= ScaleNb; l++)
39             Root->addLevel();
40     }
41
42     // -- Check if tree is graded ---
43
44     if (debug) Root->checkGradedTree();
45
46 }
47
48 }
  
```

Here is the caller graph for this function:





### 6.6.3.27 Vector Limiter ( const Vector $u$ , const Vector $v$ )

Returns the value of the slope limiter between the slopes  $u$  and  $v$ .

## Parameters

|     |        |
|-----|--------|
| $u$ | Vector |
| $v$ | Vector |

## Returns

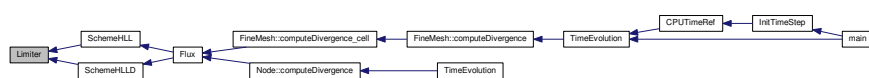
Vector

```

57 {
58     // Min Mod limiter
59
60     int LimiterNo = 3;
61
62     Vector Result(u.dimension());
63     int i;
64     real x, y; // slopes
65
66     for (i=1; i<=u.dimension(); i++)
67     {
68         x = u.value(i);
69         y = v.value(i);
70
71         switch(LimiterNo)
72         {
73             // MIN-MOD
74             case 1:
75                 if (x == y)
76                     Result.setValue(i, 0.);
77                 else
78                     Result.setValue(i, Min(1., fabs(x)/fabs(x-y)));
79                 break;
80
81             // VAN LEER
82             case 3:
83             default:
84                 if ((fabs(x) + fabs(y)) == 0.)
85                     Result.setValue(i, 0.);
86                 else
87                     Result.setValue(i, fabs(x) / (fabs(x)+fabs(y)));
88                 break;
89         };
90     }
91
92     return Result;
93 }

```

Here is the caller graph for this function:



## 6.6.3.28 real Limiter ( const real x )

Returns the value of slope limiter from a real value x.

## Parameters

|     |     |
|-----|-----|
| $x$ | ... |
|-----|-----|

## Returns

double

```

23 {
24     real Result = 0.;
25
26     switch (LimiterNo)

```

```

27 {
28     case 1: // Min-Mod
29         Result = Max(0., Min(1., r));
30         break;
31
32     case 2: // Van Albada
33         Result = (r<=0.)? 0.: (r*r+r)/(r*r+1.);
34         break;
35
36     case 3: // Van Leer
37         Result = (r<=0.) ? 0.: (r+Abs(r))/(1.+Abs(r));
38         break;
39
40     case 4: // Superbee
41         Result = (r<=0.) ? 0.: Max(0., Max(Min(2.*r, 1.), Min(r, 2.)));
42         break;
43     case 5: // Monotonized Central
44         Result = max(0.0, min(min(2*r, 0.5*(1+r)), 2.0));
45         break;
46
47 };
48
49     return Result;
50 }

```

### 6.6.3.29 real MinAbs ( const real a, const real b )

Returns the minimum in module of  $a$  and  $b$ .

#### Parameters

|     |            |
|-----|------------|
| $a$ | Real value |
| $b$ | Real value |

#### Returns

double

```

23 {
24     return (fabs(a) <= fabs(b)) ? a:b;
25 }

```

### 6.6.3.30 real NormMaxQuantities ( const Vector & V )

Returns the Max-norm of the vector where every quantity is divided by its characteristic value.

#### Parameters

|     |        |
|-----|--------|
| $V$ | Vector |
|-----|--------|

#### Returns

double

```

26 {
27     Vector W(QuantityNb);
28     int AxisNo=1;
29     real MomentumMax=0.;
30     real MagMax=0.;
31
32
33
34     W.setZero();
35
36 /*
37     // Density
38     W.setValue(1, V.value(1)/QuantityMax.value(1));
39
40     // Momentum
41     W.setValue(2, V.value(2)/QuantityMax.value(2));
42     W.setValue(3, V.value(3)/QuantityMax.value(3));

```

```

43     W.setValue(4 , V.value(4)/QuantityMax.value(4) );
44
45     // Energy
46     W.setValue(5 , V.value(5)/QuantityMax.value(5) );
47
48     // psi
49     // W.setValue(6, V.value(6)/QuantityMax.value(6));
50
51     // Magnetic Field
52     W.setValue(7 , V.value(7)/QuantityMax.value(7) );
53     W.setValue(8 , V.value(8)/QuantityMax.value(8) );
54     W.setValue(9 , V.value(9)/QuantityMax.value(9) );
55 */
56
57 // --- Compute Linf norm --
58
59 W.setValue(1, (V.value(1))/QuantityMax.value(1));
60 W.setValue(5, (V.value(5))/QuantityMax.value(5));
61 //W.setValue(6, (V.value(6))/QuantityMax.value(6));
62
63
64 for (AxisNo = 1; AxisNo <= Dimension; AxisNo++)
65 {
66     MomentumMax = Max( MomentumMax, QuantityMax.value(AxisNo+1) );
67     MagMax      = Max( MagMax,      QuantityMax.value(AxisNo+6) );
68     W.setValue(2, W.value(2) + V.value(AxisNo+1)*V.value(AxisNo+1));
69     W.setValue(7, W.value(7) + V.value(AxisNo+6)*V.value(AxisNo+6));
70 }
71
72 if(Dimension==2){
73     W.setValue(4, (V.value(4))/QuantityMax.value(4));
74     W.setValue(9, (V.value(9))/QuantityMax.value(9));
75 }
76
77 W.setValue(2, sqrt(W.value(2))/MomentumMax);
78 W.setValue(7, sqrt(W.value(7))/MagMax );
79
80 if(IterationNo==0) return NMax(V);
81 return NMax(W);
82 }

```

### 6.6.3.31 void Performance ( const char \* FileName )

Computes the performance of the computation and, for cluster computations, write it into file *FileName*.

#### Parameters

| <i>FileName</i> | Name of the file. |
|-----------------|-------------------|
|-----------------|-------------------|

#### Returns

void

```

23 {
24     // --- Local variables ---
25
26     bool EndComputation;           // True if end computation
27     int FineCellNb;               // Number of cells on fine grid
28     int CellPVirt;
29     FILE *output;                 // Pointer to output file
30
31     double realtimefull;          //full real time
32     double ftime;                 // real time
33     double ctime;                 // CPU time
34     unsigned int ttime, rtime;    // total and remaining real time (in seconds)
35     unsigned int tctime=0, rctime=0; // total and remaining CPU time (in seconds)
36     unsigned int rest;
37     int day, hour, min, sec;
38
39     // --- Init EndComputation
40
41     EndComputation = (IterationNo > IterationNb);
42
43     // --- Compute FineCellNb ---
44
45     FineCellNb = 1<<(ScaleNb*Dimension);
46     CellPVirt = 1<<(ScaleNb*(Dimension-1));
47     CellPVirt = CellPVirt*2*Dimension;
48     // --- Write in file ---
49

```

```

50 /*
51   char CPUFileName[255];
52   #if defined PARMPI
53     sprintf(CPUFileName, "%d_%d_%d_%s", coords[0], coords[1], coords[2], FileName);
54   // strcpy(CPUFileName, FileName);
55   #else
56     strcpy(CPUFileName, FileName);
57   #endif
58 */
59
60   if ((output = fopen(FileName, "w"))
61   {
62
63     realtimefull=ftime = CPUTime.realTime();
64     ctime = CPUTime.CPUTime();
65
66     if (!EndComputation)
67     {
68       ttime = (unsigned int)((ftime*IterationNb)/IterationNo);
69       rtime = (unsigned int)((ftime*(IterationNb-IterationNo))/
IterationNo);
70       tctime = (unsigned int)((ctime*IterationNb)/IterationNo);
71       rctime = (unsigned int)((ctime*(IterationNb-IterationNo))/
IterationNo);
72     }
73
74     fprintf(output, "Dimension           : %12i\n", Dimension);
75
76     if (EndComputation)
77       fprintf(output, "Iterations           : %12i\n", IterationNb);
78     else
79     {
80       fprintf(output, "Iterations (total)   : %12i\n", IterationNb);
81       fprintf(output, "Iterations (elapsed) : %12i\n", IterationNo);
82       fprintf(output, "In progress         : %13.6f %%\n", 100.*
IterationNo/(1.*IterationNb));
83     }
84
85     fprintf(output, "Scales (max)        : %12i\n", ScaleNb);
86     fprintf(output, "Cells (max)         : %12i\n", (1<<(ScaleNb*Dimension)));
87
88     if (Multiresolution)
89       fprintf(output, "Solver              :           MR\n");
90     else
91       fprintf(output, "Solver              :           FV\n");
92
93     //fprintf(output, "Time integration    :           explicit\n");
94     fprintf(output, "Time accuracy order : %12i\n", StepNb);
95     fprintf(output, "Time step           : %13.6e s\n", TimeStep);
96     fprintf(output, "Threshold parameter : %13.6e \n", Tolerance);
97     fprintf(output, "Threshold norm      : %12i\n", ThresholdNorm);
98     fprintf(output, "CFL                 : %13.6e \n", CFL);
99
100    if(Resistivity)
101      fprintf(output, "Eta                 : %13.6e \n", eta);
102    if(Diffusivity)
103      fprintf(output, "Chi                 : %13.6e \n", chi);
104
105    if (EndComputation)
106    {
107      fprintf(output, "Physical time       : %13.6e s\n", ElapsedTime); //TimeStep *
IterationNb);
108      fprintf(output, "CPU time (s)        : %13.6e s\n", ctime);
109
110      if (Multiresolution)
111        fprintf(output, "CPU time / it. x pt : %13.6e s\n", ctime/
TotalLeafNb);
112      else
113        fprintf(output, "CPU time / it. x pt : %13.6e s\n", ctime/((1<<(
ScaleNb*Dimension))*IterationNb));
114
115      if (Multiresolution)
116      {
117
118        fprintf(output, "Leaf compression   : %13.6f %% \n", (100.*
TotalLeafNb)/(1.0*IterationNb*FineCellNb));
119        //fprintf(output, "Memory compression : %13.6f %% \n",
(100.*TotalCellNb)/(1.0*IterationNb*(FineCellNb)));
120        fprintf(output, "Memory compression : %13.6f %% \n", (100.*
TotalCellNb)/(1.0*IterationNb*(FineCellNb + CellPVirt)));
121        fprintf(output, "CPU compression    : %13.6f %% \n", (100.*ctime)/(IterationNb*
FVTimeRef));
122      }
123      else
124      {
125        fprintf(output, "Leaf compression   : %13.6f %% \n", 100.);
126        fprintf(output, "Memory compression : %13.6f %% \n", 100.);

```

```

127         fprintf(output, "CPU compression      :%13.6f %% \n", 100.);
128     }
129 }
130     else
131     {
132         fprintf(output, "Total physical time  :%13.6e s\n", TimeStep * IterationNb);
133         fprintf(output, "Elapsed physical time :%13.6e s\n", TimeStep *
IterationNb);
134         if (Multiresolution)
135         {
136             fprintf(output, "Leaf compression      :%13.6f %% \n", (100.*
TotalLeafNb)/(1.0*IterationNb*FineCellNb));
137             fprintf(output, "Memory compression   :%13.6f %% \n", (100.*
TotalCellNb)/(1.0*IterationNo*FineCellNb));
138             fprintf(output, "CPU compression      :%13.6f %% \n", (100.*ctime)/(
IterationNo*FVTimeRef));
139         }
140         else
141         {
142             fprintf(output, "Leaf compression      :%13.6f %% \n", 100.);
143             fprintf(output, "Memory compression   :%13.6f %% \n", 100.);
144             fprintf(output, "CPU compression      :%13.6f %% \n", 100.);
145         }
146     }
147
148     fprintf(output, "\n");
149
150     if (EndComputation)
151     {
152         // --- Print final time -----
153
154         rest = (unsigned int)(ctime);
155         day = rest/86400;
156         rest %= 86400;
157         hour = rest/3600;
158         rest %= 3600;
159         min = rest/60;
160         rest %= 60;
161         sec = rest;
162         rest = (unsigned int)(ctime);
163
164         if (rest >= 86400)
165             fprintf(output, "CPU time : %5d day %2d h %2d min %2d s\n", day, hour, min, sec);
166
167         if ((rest < 86400)&&(rest >= 3600))
168             fprintf(output, "CPU time : %2d h %2d min %2d s\n", hour, min, sec);
169
170         if ((rest < 3600)&&(rest >= 60))
171             fprintf(output, "CPU time : %2d min %2d s\n", min, sec);
172
173         if (rest < 60)
174             fprintf(output, "CPU time : %2d s\n", sec);
175     }
176     else
177     {
178         // --- Print total time -----
179
180         rest = tctime;
181         day = rest/86400;
182         rest %= 86400;
183         hour = rest/3600;
184         rest %= 3600;
185         min = rest/60;
186         rest %= 60;
187         sec = rest;
188
189         if (tctime >= 86400)
190             fprintf(output, "Total CPU time (estimation) : %5d day %2d h %2d min %2d s\n", day,
hour, min, sec);
191
192         if ((tctime < 86400)&&(tctime >= 3600))
193             fprintf(output, "Total CPU time (estimation) : %2d h %2d min %2d s\n", hour, min, sec);
194
195         if ((tctime < 3600)&&(tctime >= 60))
196             fprintf(output, "Total CPU time (estimation) : %2d min %2d s\n", min, sec);
197
198         if (tctime < 60)
199             fprintf(output, "Total CPU time (estimation) : %2d s\n", sec);
200
201         // --- Print remaining time -----
202
203         rest = rctime;
204         day = rest/86400;
205         rest %= 86400;
206         hour = rest/3600;
207         rest %= 3600;
208         min = rest/60;

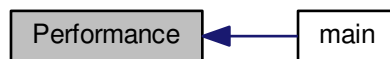
```

```

209         rest %= 60;
210         sec = rest;
211
212         if (rctime >= 86400)
213             fprintf(output, "Remaining CPU time (estimation) : %5d day %2d h %2d min %2d s\n", day,
hour, min, sec);
214
215         if ((rctime < 86400)&&(rctime >= 3600))
216             fprintf(output, "Remaining CPU time (estimation) : %2d h %2d min %2d s\n", hour, min, sec);
217
218         if ((rctime < 3600)&&(rctime >= 60))
219             fprintf(output, "Remaining CPU time (estimation) : %2d min %2d s\n", min, sec);
220
221         if (rctime < 60)
222             fprintf(output, "Remaining CPU time (estimation) : %2d s\n", sec);
223
224     }
225
226
227 #if defined PARMPI
228     fprintf(output, "\n");
229     fprintf(output, "Real time (time() function) :%lf\n", realtimefull);
230     fprintf(output, "clock() function :%lf\n", ctime);
231     fprintf(output, "\nCommunications real timer: %lf\n", CommTimer.
realTime());
232     fprintf(output, "Communications clock():%lf\n", CommTimer.
CPUTime());
233 #endif
234
235     fprintf(output, "\n");
236     fclose (output);
237 }
238 else
239 {
240     cout << "Performance.cpp: In method `void Performance(Node*, char*)':\n";
241     cout << "Performance.cpp: cannot open file " << FileName << '\n';
242     cout << "carmen: *** [Performance.o] Execution error\n";
243     cout << "carmen: abort execution.\n";
244     exit(1);
245 }
246 }

```

Here is the caller graph for this function:



### 6.6.3.32 void PrintIntegral ( const char \* *FileName* )

Writes the integral values, like e.g flame velocity, global error, into file *FileName*.

#### Parameters

|                 |                  |
|-----------------|------------------|
| <i>FileName</i> | Name of the file |
|-----------------|------------------|

#### Returns

void

```

31 {
32     // --- Local variables ---
33
34     real    t;           // time
35     FILE    *output;    // output file
36     int    i;           // counter

```

```

37     real Volume=1.; // Total volume
38
39     // --- Open file ---
40
41     if ( (IterationNo == 0) ? (output = fopen(FileName,"w")) : (output = fopen(FileName,"a")) )
42     {
43         // HEADER
44
45         if (IterationNo == 0)
46         {
47
48             fprintf(output, "#");
49             fprintf(output, TXTFORMAT, " Time");
50             fprintf(output, TXTFORMAT, " CFL");
51             fprintf(output, TXTFORMAT, " Energy");
52             fprintf(output, TXTFORMAT, " Div B Max");
53             fprintf(output, TXTFORMAT, " ch");
54             fprintf(output, TXTFORMAT, " Helicity");
55             fprintf(output, TXTFORMAT, " DivB Max norm");
56 /*
57             if (Multiresolution)
58             {
59                 fprintf(output, TXTFORMAT, "Memory comp.");
60                 fprintf(output, TXTFORMAT, "CPU comp.");
61                 if (ExpectedCompression != 0. || CVS)
62                     fprintf(output, TXTFORMAT, "Tolerance");
63             // if (CVS)
64             //     fprintf(output, TXTFORMAT, "Av. Pressure");
65             }
66 */
67 */
68             if (!ConstantTimeStep)
69             {
70                 if (StepNb == 3) fprintf(output, TXTFORMAT, "RKF Error");
71                 fprintf(output, TXTFORMAT, "Next time step");
72                 fprintf(output, "%13s ", "IterationNo");
73                 fprintf(output, "%13s ", "IterationNb");
74             }
75
76             fprintf(output, "\n");
77         }
78
79         if (ConstantTimeStep)
80             t=IterationNo*TimeStep;
81         else
82             t = ElapsedTime;
83
84         fprintf(output, FORMAT, t);
85
86         // --- Compute total volume ---
87
88         for (i=1; i<= Dimension; i++)
89             Volume *= fabs(XMax[i]-XMin[i]);
90
91         // Print CFL
92         fprintf(output, FORMAT, Eigenvalue*TimeStep/SpaceStep);
93
94         // Print momentum and energy
95         //fprintf(output, FORMAT, GlobalMomentum);
96         fprintf(output, FORMAT, GlobalEnergy);
97         fprintf(output, FORMAT, DIVBMax);
98         fprintf(output, FORMAT, ch);
99         fprintf(output, FORMAT, Helicity);
100        fprintf(output, FORMAT, DIVB);
101
102
103
104 /*
105         if (Multiresolution)
106         {
107             fprintf(output, FORMAT, (1.*CellNb)/(1<<(ScaleNb*Dimension)));
108             fprintf(output, FORMAT, CPUTime.CPUTime()/(IterationNo*FVTimeRef));
109
110             if (ExpectedCompression != 0.)
111                 fprintf(output, FORMAT, Tolerance);
112
113             // if (CVS)
114             //{
115             //     fprintf(output, FORMAT, ComputedTolerance(ScaleNb));
116             //     fprintf(output, FORMAT, QuantityAverage.value(1));
117             //}
118         }
119 */
120         if (!ConstantTimeStep)
121         {
122             if (StepNb == 3) fprintf(output, FORMAT, RKFError);
123             fprintf(output, FORMAT, TimeStep);

```



```

124         fprintf(output, "%13i ", IterationNo);
125         fprintf(output, "%13i ", IterationNb);
126     }
127
128     fprintf(output, "\n");
129     fclose(output);
130 }
131 else
132 {
133     cout << "PrintIntegral.cpp: In method 'void PrintIntegral(Node*, char*)':\n";
134     cout << "PrintIntegral.cpp: cannot open file " << FileName << "\n";
135     cout << "carmen: *** [PrintIntegral.o] Execution error\n";
136     cout << "carmen: abort execution.\n";
137     exit(1);
138 }
139 }

```

Here is the caller graph for this function:



### 6.6.3.33 void ReduceIntegralValues ( )

Parallel function DOES NOT WORK!

#### Returns

void

```

469     {
470     real rb; //Recieve Buffer
471     rb=0.0;
472     CommTimer.start();
473     #if defined PARMPI
474     MPI_Reduce (&ErrorMax, &rb, 1, MPI_Type, MPI_MAX, 0, MPI_COMM_WORLD);
475     ErrorMax=rb;
476
477     MPI_Reduce (&ErrorMid, &rb, 1, MPI_Type, MPI_SUM, 0, MPI_COMM_WORLD);
478     ErrorMid=rb/size;
479
480     MPI_Reduce (&ErrorL2, &rb, 1, MPI_Type, MPI_SUM, 0, MPI_COMM_WORLD);
481     ErrorL2=rb/size;
482
483     MPI_Reduce (&ErrorNb, &rb, 1, MPI_Type, MPI_SUM, 0, MPI_COMM_WORLD);
484     ErrorNb=rb;
485
486     MPI_Allreduce (&FlameVelocity, &rb, 1, MPI_Type, MPI_SUM, MPI_COMM_WORLD);
487     FlameVelocity=rb;
488
489     MPI_Allreduce (&GlobalMomentum, &rb, 1, MPI_Type, MPI_SUM, MPI_COMM_WORLD);
490     GlobalMomentum=rb;
491
492     MPI_Allreduce (&GlobalEnergy, &rb, 1, MPI_Type, MPI_SUM, MPI_COMM_WORLD);
493     GlobalEnergy=rb;
494
495     MPI_Reduce (&ExactMomentum, &rb, 1, MPI_Type, MPI_SUM, 0, MPI_COMM_WORLD);
496     ExactMomentum=rb;
497
498     MPI_Reduce (&ExactEnergy, &rb, 1, MPI_Type, MPI_SUM, 0, MPI_COMM_WORLD);
499     ExactEnergy=rb;
500
501     MPI_Allreduce (&GlobalReactionRate, &rb, 1, MPI_Type, MPI_SUM, MPI_COMM_WORLD);
502     GlobalReactionRate=rb;
503
504     MPI_Allreduce (&EigenvalueMax, &rb, 1, MPI_Type, MPI_MAX, MPI_COMM_WORLD);
505     EigenvalueMax=rb;

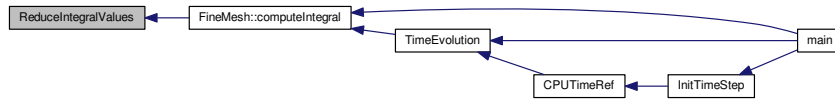
```

```

506
507 #endif
508   CommTimer.stop();
509 }

```

Here is the caller graph for this function:



#### 6.6.3.34 void RefreshTree ( Node \* Root )

Refresh the tree structure, i.e. compute the cell-averages of the internal nodes by projection and those of the virtual leaves by prediction. The root node is *Root*. Only for multiresolution computations.

Parameters

|             |      |
|-------------|------|
| <i>Root</i> | Root |
|-------------|------|

Returns

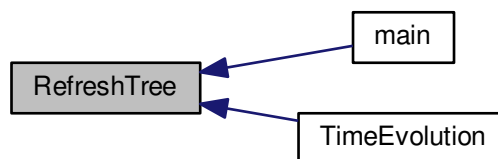
void

```

23 {
24   // --- Project : compute cell-average values in all nodes ---
25   Root->project();
26
27   // --- Fill virtual children with predicted values ---
28   Root->fillVirtualChildren();
29 }

```

Here is the caller graph for this function:



#### 6.6.3.35 void Remesh ( Node \* Root )

Remesh the tree structure after a time evolution. The root node is *Root*. Only for multiresolution computations.

## Parameters

|             |      |
|-------------|------|
| <i>Root</i> | Root |
|-------------|------|

## Returns

void

```

23 {
24     // --- Refresh tree structure ---
25     // RefreshTree(Root);
26
27     // --- Check if tree is graded ---
28     if (debug) Root->checkGradedTree();
29
30     // --- Adapt : depending on details, refine or combine ---
31     Root->adapt();
32
33     // --- Check if tree is graded ---
34     if (debug) Root->checkGradedTree();
35 }

```

Here is the caller graph for this function:



### 6.6.3.36 Vector ResistiveTerms ( Cell & Cell1, Cell & Cell2, Cell & Cell3, Cell & Cell4, int AxisNo )

Returns the resistive source terms in the cell *UserCell*.

## Parameters

|               |                  |
|---------------|------------------|
| <i>Cell1</i>  | Cell 1           |
| <i>Cell2</i>  | Cell 2           |
| <i>Cell3</i>  | Cell 3           |
| <i>Cell4</i>  | Cell 4           |
| <i>AxisNo</i> | Axis of interest |

## Returns

Vector

X - direction

2D

Y - direction

2D

Z - direction

3D

```

12 {
13     // --- Local variables ---
14     Vector B(3), Bi(3), Bj(3), Bk(3);

```

```

15 Vector Result(QuantityNb);
16 Vector Bavg(3);
17 real Jx = 0., Jy = 0., Jz = 0.;
18 real dx, dy, dz;
19 real ResX= 0., ResY= 0., ResZ= 0., ResE= 0.;
20 real eta0=0., etai=0., etaj=0., etak=0., etaR=0.;
21
22 dx = Cell2.size(1);
23 dy = Cell2.size(2);
24 dz = Cell2.size(3);
25
26 eta0 = Cell1.Res;
27 etai = Cell2.Res;
28 etaj = Cell3.Res;
29 etak = Cell4.Res;
30
31 for(int i=1; i <= 3; i++){
32     B.setValue(i, Cell1.average(i+6));
33     Bi.setValue(i, Cell2.average(i+6));
34     Bj.setValue(i, Cell3.average(i+6));
35     Bk.setValue(i, Cell4.average(i+6));
36 }
37
38
39 if(AxisNo == 1){
40     etaR = (eta0 + etai)/2.;
41
42     Bavg.setValue(2, 0.5*(B.value(2) + Bi.value(2)));
43     Bavg.setValue(3, 0.5*(B.value(3) + Bi.value(3)));
44
45     Jy = -(B.value(3) - Bi.value(3))/dx;
46     Jz = (B.value(2) - Bi.value(2))/dx;
47
48     Jz = Jz - (B.value(1) - Bj.value(1))/dy;
49
50     if(Dimension==3){
51         Jy = Jy + (B.value(1) - Bk.value(1))/dz;
52     }
53
54     ResE = etaR*(Bavg.value(2)*Jz - Bavg.value(3)*Jy);
55     ResX = 0.;
56     ResY = etaR*Jz;
57     ResZ = -etaR*Jy;
58
59 }else if(AxisNo == 2){
60     etaR = (eta0 + etaj)/2.;
61
62     Bavg.setValue(1, 0.5*(B.value(1) + Bj.value(1)));
63     Bavg.setValue(3, 0.5*(B.value(3) + Bj.value(3)));
64
65     Jx = (B.value(3) - Bj.value(3))/dy;
66     Jz = -(B.value(1) - Bj.value(1))/dy;
67
68     Jz = Jz + (B.value(2) - Bi.value(2))/dx;
69
70     if(Dimension==3){
71         Jx = Jx + (B.value(2) - Bk.value(2))/dz;
72     }
73
74     ResE = etaR*(Bavg.value(3)*Jx - Bavg.value(1)*Jz);
75     ResX = -etaR*Jz;
76     ResY = 0.;
77     ResZ = etaR*Jx;
78
79 }else{
80     etaR = (eta0 + etak)/2.;
81
82     Bavg.setValue(1, 0.5*(B.value(1) + Bk.value(1)));
83     Bavg.setValue(2, 0.5*(B.value(2) + Bk.value(2)));
84
85     Jx = -(B.value(2) - Bk.value(2))/dz;
86     Jy = (B.value(1) - Bk.value(1))/dz;
87
88     Jx = Jx + (B.value(3) - Bj.value(3))/dy;
89     Jy = Jy - (B.value(3) - Bi.value(3))/dx;
90
91     ResE = etaR*(Bavg.value(1)*Jy - Bavg.value(2)*Jx);
92     ResX = etaR*Jy;
93     ResY = -etaR*Jx;
94     ResZ = 0.;
95 }
96
97 Result.setZero();
98
99 // These values will be added to the numerical flux

```

```

108     Result.setValue(5, ResE);
109     Result.setValue(7, ResX);
110     Result.setValue(8, ResY);
111     Result.setValue(9, ResZ);
112
113     return Result;
114 }
115 }

```

Here is the caller graph for this function:



### 6.6.3.37 Vector SchemeHLL ( const Cell & Cell1, const Cell & Cell2, const Cell & Cell3, const Cell & Cell4, const int AxisNo )

Returns the HLL numerical flux for MHD equations. The scheme uses four cells to estimate the flux at the interface. *Cell2* and *Cell3* are the first neighbours on the left and right sides. *Cell1* and *Cell4* are the second neighbours on the left and right sides.

#### Parameters

|               |                                    |
|---------------|------------------------------------|
| <i>Cell1</i>  | second neighbour on the left side  |
| <i>Cell2</i>  | first neighbour on the left side   |
| <i>Cell3</i>  | first neighbour on the right side  |
| <i>Cell4</i>  | second neighbour on the right side |
| <i>AxisNo</i> | Axis of interest.                  |

#### Returns

##### Vector

```

12 {
13
14     // General variables
15
16     Vector    LeftAverage(QuantityNb); //
17     Vector    RightAverage(QuantityNb); // Conservative quantities
18     Vector    Result(QuantityNb);      // MHD numerical flux
19     int aux=0;
20     // Variables for the HLL scheme
21     Vector    FL(QuantityNb), FR(QuantityNb); //Left and right physical fluxes
22     Vector    VL(3), VR(3); // Left and right velocities
23     Vector    BL(3), BR(3); // Left and right velocities
24     real rhoL=0., rhoR=0.; // Left and right densities
25     real eL=0., eR=0.; // Left and right energies
26     real preL=0., preR=0.;
27     real bkL=0., bkR=0.;
28     real aL=0., aR=0.;
29     real bL=0., bR=0.;
30     real cfL=0., cfR=0.;
31     real SL=0., SR=0.;
32     real dx=0.;
33     dx = Cell2.size(AxisNo);
34     real r, Limit, LeftSlope = 0., RightSlope = 0.; // Left and right slopes
35     int i;
36
37     // --- Limiter function -----
38
39     for (i=1; i<=QuantityNb; i++)
40     {
41         // --- Compute left cell-average value ---
42
43         if (Cell2.average(i) != Cell1.average(i))
44         {
45             RightSlope = Cell3.average(i)-Cell2.average(i);

```

```

46     LeftSlope = Cell2.average(i)-Cell1.average(i);
47     r        = RightSlope/LeftSlope;
48     Limit    = Limiter(r);
49     LeftAverage.setValue(i, Cell2.average(i) + 0.5*Limit*LeftSlope);
50     aux = 1;
51 }
52 else
53     LeftAverage.setValue(i, Cell2.average(i));
54
55 // --- Compute right cell-average value ---
56
57 if (Cell3.average(i) != Cell2.average(i))
58 {
59     RightSlope = Cell4.average(i)-Cell3.average(i);
60     LeftSlope = Cell3.average(i)-Cell2.average(i);
61     r        = RightSlope/LeftSlope;
62     Limit    = Limiter(r);
63     RightAverage.setValue(i, Cell3.average(i) - 0.5*Limit*LeftSlope);
64     aux = 1;
65 }
66 else
67     RightAverage.setValue(i, Cell3.average(i));
68 }
69
70
71 // --- HLL scheme -----
72
73 // --- Conservative variables ---
74
75 // Left and right densities
76 rhoL = LeftAverage.value(1);
77 rhoR = RightAverage.value(1);
78
79 // Left and right momentum and magnetic field
80 for (int i=1;i<=3;i++)
81 {
82     VL.setValue(i, LeftAverage.value(i+1));
83     VR.setValue(i, RightAverage.value(i+1));
84     BL.setValue(i, LeftAverage.value(i+6));
85     BR.setValue(i, RightAverage.value(i+6));
86 }
87
88 // Left and right energies
89 eL = LeftAverage.value(5);
90 eR = RightAverage.value(5);
91
92 // Left and right pressures
93 preL = (Gamma - 1.)*(eL - 0.5*(VL*VL)/rhoL - 0.5*(BL*BL));
94 preR = (Gamma - 1.)*(eR - 0.5*(VR*VR)/rhoR - 0.5*(BR*BR));
95
96 // --- Magnetoacoustic waves calculations --
97
98 bkL = power2(BL.value(AxisNo))/rhoL;
99 bkR = power2(BR.value(AxisNo))/rhoR;
100
101 aL = Gamma*preL/rhoL;
102 aR = Gamma*preR/rhoR;
103
104 bL = (BL*BL)/rhoL;
105 bR = (BR*BR)/rhoR;
106
107 // Left and Right fast speeds
108 cfL = sqrt(0.5*(aL + bL + sqrt(power2(aL + bL) - 4.0*aL*bkL)));
109 cfR = sqrt(0.5*(aR + bR + sqrt(power2(aR + bR) - 4.0*aR*bkR)));
110
111 // Left and right slopes
112 SL = Min(Min(VL.value(AxisNo)/rhoL - cfL, VR.value(AxisNo)/rhoR - cfR),0.0);
113 SR = Max(Max(VL.value(AxisNo)/rhoL + cfL, VR.value(AxisNo)/rhoR + cfR),0.0);
114
115 // --- Physical flux ---
116 if(AxisNo ==1){
117     EigenvalueX = Max(Max(Abs(SL),Abs(SR)),
EigenvalueX);
118     FL = FluxX(LeftAverage);
119     FR = FluxX(RightAverage);
120 }else if(AxisNo ==2){
121     EigenvalueY = Max(Max(Abs(SL),Abs(SR)),
EigenvalueY);
122     FL = FluxY(LeftAverage);
123     FR = FluxY(RightAverage);
124 }else{
125     EigenvalueZ = Max(Max(Abs(SL),Abs(SR)),
EigenvalueZ);
126     FL = FluxZ(LeftAverage);
127     FR = FluxZ(RightAverage);
128 }
129

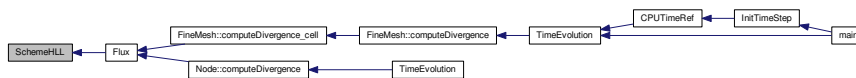
```

```

130
131 // --- HLL Riemann Solver ---
132
133 for(int i=1;i<=QuantityNb;i++)
134 {
135     Result.setValue(i, (SR*FL.value(i) - SL*FR.value(i) + SR*SL*(RightAverage.value(i) - LeftAverage.
value(i)))/(SR-SL));
136 }
137
138 // Parabolic-Hyperbolic divergence Cleaning (Dedner, 2002)
139 fluxCorrection(Result, LeftAverage, RightAverage, AxisNo);
140
141 // Artificial diffusion terms
142 if(Diffusivity && aux==1) Result = Result - ArtificialViscosity(
LeftAverage,RightAverage,dx,AxisNo);
143
144 return Result;
145
146 }

```

Here is the caller graph for this function:



### 6.6.3.38 Vector SchemeHLLD ( const Cell & Cell1, const Cell & Cell2, const Cell & Cell3, const Cell & Cell4, const int AxisNo )

Returns the HLLD numerical flux for MHD equations. The scheme uses four cells to estimate the flux at the interface. *Cell2* and *Cell3* are the first neighbours on the left and right sides. *Cell1* and *Cell4* are the second neighbours on the left and right sides.

#### Parameters

|               |                                    |
|---------------|------------------------------------|
| <i>Cell1</i>  | second neighbour on the left side  |
| <i>Cell2</i>  | first neighbour on the left side   |
| <i>Cell3</i>  | first neighbour on the right side  |
| <i>Cell4</i>  | second neighbour on the right side |
| <i>AxisNo</i> | Axis of interest.                  |

#### Returns

##### Vector

```

13 {
14
15 // General variables
16
17 Vector LeftAverage(QuantityNb); //
18 Vector RightAverage(QuantityNb); // Conservative quantities
19 Vector Result(QuantityNb); // MHD numerical flux
20 int aux=0;
21
22 // Variables for the HLL scheme
23 Vector FL(QuantityNb), FR(QuantityNb); //Left and right physical fluxes
24 Vector VL(3), VR(3); // Left and right velocities
25 Vector BL(3), BR(3); // Left and right velocities
26 real rhoL=0., rhoR=0.; // Left and right densities
27 real eL=0., eR=0.; // Left and right energies
28 real preL=0., preR=0.;
29 real bkL=0., bkR=0.;
30 real aL=0., aR=0.;
31 real bL=0., bR=0.;
32 real cfL=0., cfR=0.;
33 real SL=0., SR=0.;
34 real SLS=0., SRS=0.;
35 real SM=0.;

```

```

36 Matrix U(QuantityNb,4);
37 Matrix F(QuantityNb,2);
38 real dx=0.;
39 dx = Cell2.size(AxisNo);
40 real r, Limit, LeftSlope = 0., RightSlope = 0.; // Left and right slopes
41 int i;
42
43 // --- Limiter function -----
44
45 for (i=1; i<=QuantityNb; i++)
46 {
47     // --- Compute left cell-average value ---
48
49     if (Cell2.average(i) != Cell1.average(i))
50     {
51         RightSlope = Cell3.average(i)-Cell2.average(i);
52         LeftSlope = Cell2.average(i)-Cell1.average(i);
53         r = RightSlope/LeftSlope;
54         Limit = Limiter(r);
55         LeftAverage.setValue(i, Cell2.average(i) + 0.5*Limit*LeftSlope);
56         aux = 1;
57     }
58     else
59         LeftAverage.setValue(i, Cell2.average(i));
60
61     // --- Compute right cell-average value ---
62
63     if (Cell3.average(i) != Cell2.average(i))
64     {
65         RightSlope = Cell4.average(i)-Cell3.average(i);
66         LeftSlope = Cell3.average(i)-Cell2.average(i);
67         r = RightSlope/LeftSlope;
68         Limit = Limiter(r);
69         RightAverage.setValue(i, Cell3.average(i) - 0.5*Limit*LeftSlope);
70         aux = 1;
71     }
72     else
73         RightAverage.setValue(i, Cell3.average(i));
74 }
75
76 // --- HLLD scheme -----
77
78 // --- Conservative variables ---
79
80 // Left and right densities
81 rhoL = LeftAverage.value(1);
82 rhoR = RightAverage.value(1);
83
84 // Left and right momentum and magnetic field
85 for (int i=1;i<=3;i++)
86 {
87     VL.setValue(i, LeftAverage.value(i+1));
88     VR.setValue(i, RightAverage.value(i+1));
89     BL.setValue(i, LeftAverage.value(i+6));
90     BR.setValue(i, RightAverage.value(i+6));
91 }
92
93 // Left and right energies
94 eL = LeftAverage.value(5);
95 eR = RightAverage.value(5);
96
97 // Left and right pressures
98 preL = (Gamma -1.)*(eL - 0.5*(VL*VL)/rhoL - 0.5*(BL*BL));
99 preR = (Gamma -1.)*(eR - 0.5*(VR*VR)/rhoR - 0.5*(BR*BR));
100
101 // --- Magnetoacoustic waves computation --
102 bkL = power2(BL.value(AxisNo))/rhoL;
103 bkR = power2(BR.value(AxisNo))/rhoR;
104
105 aL = Gamma*preL/rhoL;
106 aR = Gamma*preR/rhoR;
107
108 bL = (BL*BL)/rhoL;
109 bR = (BR*BR)/rhoR;
110
111 // Left and Right fast speeds
112 cfL = sqrt(0.5*(aL + bL + sqrt(power2(aL + bL) - 4.0*aL*bL)));
113 cfR = sqrt(0.5*(aR + bR + sqrt(power2(aR + bR) - 4.0*aR*bR)));
114
115 // Left and Right slopes
116 SL = Min((VL.value(AxisNo))/rhoL, (VR.value(AxisNo))/rhoR) - Max(cfL,cfR);
117 SR = Max((VL.value(AxisNo))/rhoL, (VR.value(AxisNo))/rhoR) + Max(cfL,cfR);
118
119 // --- Physical flux ---
120 if(AxisNo ==1){
121     EigenvalueX = Max(Max(Abs(SL), Abs(SR)),
EigenvalueX);

```

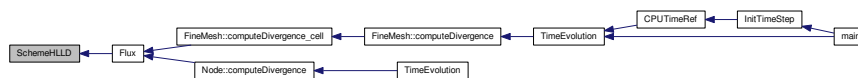


```

122     FL = FluxX(LeftAverage);
123     FR = FluxX(RightAverage);
124 }else if (AxisNo ==2){
125     EigenvalueY = Max(Max(Abs(SL),Abs(SR)),
EigenvalueY);
126     FL = FluxY(LeftAverage);
127     FR = FluxY(RightAverage);
128 }else{
129     EigenvalueZ = Max(Max(Abs(SL),Abs(SR)),
EigenvalueZ);
130     FL = FluxZ(LeftAverage);
131     FR = FluxZ(RightAverage);
132 }
133
134 // Intermediary states U* and U**
135 U = stateUstar(LeftAverage, RightAverage, preL, preR, SL, SR, SM, SLS, SRS,AxisNo);
136
137 // --- HLLD Riemann Solver ---
138
139 for(int i=1;i<=QuantityNb;i++)
140 {
141     //Flux Function - Equation 66
142     //F_L
143     if(SL>=0.)
144         Result.setValue(i, FL.value(i));
145     //F-star left // FL=FLstar
146     else if(SLS>=0. && SL<0.)
147         Result.setValue(i, FL.value(i) + SL*(U.value(i,1) - LeftAverage.value(i)));
148     //F-star-star left
149     else if(SM>=0. && SLS<0.)
150         Result.setValue(i, FL.value(i) + SLS*U.value(i,3) - (SLS - SL)*U.value(i,1) - SL*
LeftAverage.value(i));
151     //F-star-star right
152     else if(SRS>=0. && SM<0.)
153         Result.setValue(i, FR.value(i) + SRS*U.value(i,4) - (SRS - SR)*U.value(i,2) - SR*
RightAverage.value(i));
154     //F-star right
155     else if(SR>=0. && SRS<0.)
156         Result.setValue(i, FR.value(i) + SR*(U.value(i,2) - RightAverage.value(i)));
157     //F_R
158     else
159         Result.setValue(i, FR.value(i));
160 }
161 // Parabolic-Hyperbolic divergence Cleaning (Dedner, 2002)
162 //fluxCorrection(Result, Cell2.average(), Cell3.average(), AxisNo);
163 fluxCorrection(Result, LeftAverage, RightAverage, AxisNo);
164
165 // Artificial diffusion terms
166 if(Diffusivity && aux==1) Result = Result - ArtificialViscosity(
LeftAverage,RightAverage,dx,AxisNo);
167
168 return Result;
169
170 }

```

Here is the caller graph for this function:



### 6.6.3.39 void ShowTime ( Timer arg )

Writes on screen the estimation of total and remaining CPU times. These informations are stored in the timer *arg*.

#### Parameters

| <i>arg</i> | Argument |
|------------|----------|
|------------|----------|

## Returns

void

```

24 {
25 // double ftime;                // real time
26 double ctime;                 // CPU time
27 // unsigned int   ttime, rtime; // total and remaining real time (in seconds)
28 unsigned int   tctime, rctime; // total and remaining CPU time (in seconds)
29
30 int day, hour, min, sec;
31 unsigned int rest;
32
33 // --- Write total and remaining estimated time -----
34
35 // ftime = arg.GetRealTime();
36 ctime = arg.CPUTime();
37 // ttime = (unsigned int)((ftime*IterationNb)/IterationNo);
38 // rtime = (unsigned int)((ftime*(IterationNb-IterationNo))/IterationNo);
39 tctime = (unsigned int)((ctime*IterationNb)/IterationNo);
40 rctime = (unsigned int)((ctime*(IterationNb-IterationNo))/
IterationNo);
41
42 // --- Show total time -----
43
44 rest = tctime;
45 day = rest/86400;
46 rest %= 86400;
47 hour = rest/3600;
48 rest %= 3600;
49 min = rest/60;
50 rest %= 60;
51 sec = rest;
52
53 printf("\033[1A\033[1A");
54
55 if (tctime >= 86400)
56     printf("Total CPU time      (estimation) : %5d day %2d h %2d min %2d s\n", day, hour, min, sec);
57
58 if ((tctime < 86400)&&(tctime >= 3600))
59     printf("Total CPU time      (estimation) : %2d h %2d min %2d s          \n", hour, min, sec);
60
61 if ((tctime < 3600)&&(tctime >= 60))
62     printf("Total CPU time      (estimation) : %2d min %2d s          \n", min, sec);
63
64 if (tctime < 60)
65     printf("Total CPU time      (estimation) : %2d s          \n", sec);
66
67 // --- Show remaining time -----
68
69 rest = rctime;
70 day = rest/86400;
71 rest %= 86400;
72 hour = rest/3600;
73 rest %= 3600;
74 min = rest/60;
75 rest %= 60;
76 sec = rest;
77
78 if (rctime >= 86400)
79     printf("Remaining CPU time (estimation) : %5d day %2d h %2d min %2d s\n", day, hour, min, sec);
80
81 if ((rctime < 86400)&&(rctime >= 3600))
82     printf("Remaining CPU time (estimation) : %2d h %2d min %2d s          \n", hour, min, sec);
83
84 if ((rctime < 3600)&&(rctime >= 60))
85     printf("Remaining CPU time (estimation) : %2d min %2d s          \n", min, sec);
86
87 if (rctime < 60)
88     printf("Remaining CPU time (estimation) : %2d s          \n", sec);
89
90 }

```

Here is the caller graph for this function:



#### 6.6.3.40 int Sign ( const real a )

Returns 1 if *a* is non-negative, -1 elsewhere.

##### Parameters

|          |            |
|----------|------------|
| <i>a</i> | Real value |
|----------|------------|

##### Returns

int

```

23 {
24   if (a >= 0)
25     return 1;
26   else
27     return -1;
28 }
  
```

#### 6.6.3.41 Vector Source ( Cell & UserCell )

Returns the source term in the cell *UserCell*.

##### Parameters

|                 |            |
|-----------------|------------|
| <i>UserCell</i> | Cell value |
|-----------------|------------|

##### Returns

Vector

##### Gravity vector

```

24 {
25   // --- Local variables ---
26
27   Vector Force(Dimension);
28   Vector Result(QuantityNb);
29   Result.setZero();
30
31   Vector V(3);
32   real Gx=0., Gy=0., Gz=0., rho=0.;
33   for(int i=1;i<=3;i++)
34     V.setValue(i,UserCell.average(i+1));
35   rho = UserCell.density();
36   Gz = 0.2;
37   Result.setValue(2,rho*Gx);
38   Result.setValue(3,rho*Gy);
39   Result.setValue(4,rho*Gz);
40   Result.setValue(5,rho*(Gx*V.value(1) + Gy*V.value(2) + Gz*V.value(3)));
41
42   Result.setZero();
43   return Result;
44 }
  
```

Here is the caller graph for this function:



### 6.6.3.42 Matrix stateUstar ( const Vector & AvgL, const Vector & AvgR, const real prel, const real prer, real & slopeLeft, real & slopeRight, real & slopeM, real & slopeLeftStar, real & slopeRightStar, int AxisNo )

Returns the intermediary states of HLLD numerical flux for MHD equations.

#### Parameters

|                       |                                        |
|-----------------------|----------------------------------------|
| <i>AvgL</i>           | Left average vector                    |
| <i>AvgR</i>           | Right average vector                   |
| <i>prel</i>           | Left pressure                          |
| <i>prer</i>           | Right pressure                         |
| <i>slopeLeft</i>      | Left slope                             |
| <i>slopeRight</i>     | Right slope                            |
| <i>slopeM</i>         | Slope value computed for HLLD          |
| <i>slopeLeftStar</i>  | Left star slope (intermediary states)  |
| <i>slopeRightStar</i> | Right star slope (intermediary states) |
| <i>AxisNo</i>         | Axis of interest                       |

#### Returns

[Matrix](#)

variables U-star

variables U-star-star

```

13 {
14   real rhol=0., rhor=0.;
15   real psil=0., psir=0.;
16   real vxl=0., vxr=0., vyl=0., vyr=0., vzl=0., vzr=0.;
17   real Bxr=0., Bxl=0., Byl=0., Byr=0., Bzl=0., Bzr=0.;
18   real vl=0., vr=0., mf=0.;
19   real Bl=0., Br=0.;
20   real pTl=0., pTr=0.;
21   real vBl=0., vBr=0.;
22   real el=0., er=0.;
23   real rholss=0., rhorss=0.;
24   real vxlss=0., vxrss=0., vylss=0., vyrss=0., vzlss=0., vzrss=0.;
25   real Bxlls=0., Bxrss=0., Bylls=0., Byrss=0., Bzlls=0., Bzrss=0.;
26   real pTss=0.;
27   real vBlss=0., vBrss=0.;
28   real Elss=0., Erss=0.;
29   real rholss=0., rhorss=0.;
30   real vxlss=0., vxrss=0., vylss=0., vyrss=0., vzlss=0., vzrss=0.;
31   real Bxlls=0., Bxrss=0., Bylls=0., Byrss=0., Bzlls=0., Bzrss=0.;
32   real vBlss=0., vBrss=0.;
33   real Elss=0., Erss=0.;
34   int Bsign=0;
35   real half=0.5;
36   real epsilon2=1e-16;
37   real auxl=0., auxr=0., Sqrtrhols=0.;
38
39   Matrix U(QuantityNb,4);
40
41   //Variables
42   rhol = AvgL.value(1);
43   vxl  = AvgL.value(2)/rhol;
44   vyl  = AvgL.value(3)/rhol;
45   vzl  = AvgL.value(4)/rhol;
46   el   = AvgL.value(5);

```

```

47     psil = AvgL.value(6);
48     Bxl = AvgL.value(7);
49     Byl = AvgL.value(8);
50     Bzl = AvgL.value(9);
51
52     rhor = AvgR.value(1);
53     vxr = AvgR.value(2)/rhor;
54     vyr = AvgR.value(3)/rhor;
55     vzr = AvgR.value(4)/rhor;
56     er = AvgR.value(5);
57     psir = AvgR.value(6);
58     Bxr = AvgR.value(7);
59     Byr = AvgR.value(8);
60     Bzr = AvgR.value(9);
61
62     //v_x and v_y
63     vl = AvgL.value(AxisNo+1)/rhol;
64     vr = AvgR.value(AxisNo+1)/rhor;
65
66     // Average B value
67     mf = (AvgL.value(AxisNo+6)+AvgR.value(AxisNo+6))/(double (2.0));
68
69     if(AxisNo==1)
70     {
71         //|B|
72         Bl = sqrt( mf*mf + Byl*Byl + Bzl*Bzl );
73         Br = sqrt( mf*mf + Byr*Byr + Bzr*Bzr );
74         //Inner product v.B
75         vBl = vxl*mf + vyl*Byl + vzl*Bzl;
76         vBr = vxr*mf + vyr*Byr + vzr*Bzr;
77     }else if(AxisNo==2){
78         //|B|
79         Bl = sqrt( Bxl*Bxl + mf*mf + Bzl*Bzl );
80         Br = sqrt( Bxr*Bxr + mf*mf + Bzr*Bzr );
81         //Inner product v.B
82         vBl = vxl*Bxl + vyl*mf + vzl*Bzl;
83         vBr = vxr*Bxr + vyr*mf + vzr*Bzr;
84     }else{
85         //|B|
86         Bl = sqrt( Bxl*Bxl + Byl*Byl + mf*mf );
87         Br = sqrt( Bxr*Bxr + Byr*Byr + mf*mf );
88         //Inner product v.B
89         vBl = vxl*Bxl + vyl*Byl + vzl*mf;
90         vBr = vxr*Bxr + vyr*Byr + vzr*mf;
91     }
92
93     //Total Pressure
94     pTl = prel + half*Bl*Bl;
95     pTr = prer + half*Br*Br;
96
97     // S_M - Equation 38
98     slopeM = ((slopeRight - vr)*rhor*vr - (slopeLeft - vl)*rhol*vl - pTr +pTl)/
99             ((slopeRight - vr)*rhor - (slopeLeft - vl)*rhol);
100
101     //Sign function
102     if(mf > 0) Bsign = 1;
103     else Bsign = -1;
104
105
106     //density - Equation 43
107     rhols = rhol*(slopeLeft-vl)/(slopeLeft-slopeM);
108     rhors = rhor*(slopeRight-vr)/(slopeRight-slopeM);
109
110     if(AxisNo==1){
111         //velocities
112         //Equation 39
113         vxls = slopeM;
114         vxrs = vxls;
115         //B_x
116         Bxls = mf;
117         Bxrs = Bxls;
118
119         auxl = (rhol*(slopeLeft - vl)*(slopeLeft - slopeM)-mf*mf);
120         auxr = (rhor*(slopeRight - vr)*(slopeRight - slopeM)-mf*mf);
121
122         if( fabs(auxl)<epsilon2 ||
123            ( ( fabs(slopeM -vl) ) < epsilon2 ) &&
124            ( ( fabs(Byl) + fabs(Bzl) ) < epsilon2 ) &&
125            ( ( mf*mf ) > (Gamma*prel) ) ) ) {
126             vyls = vyl;
127             vzls = vzl;
128             Byls = Byl;
129             Bzls = Bzl;
130         }else{
131             //Equation 44
132             vyls = vyl - mf*Byl*(slopeM-vl)/auxl;
133             //Equation 46
134

```

```

135         vzls = vzl - mf*Bzl*(slopeM-vl)/auxl;
136         //Equation 45
137         Byls = Byl*(rhol*(slopeLeft-vl)*(slopeLeft-vl) - mf*mf)/auxl;
138         //Equation 47
139         Bzls = Bzl*(rhol*(slopeLeft-vl)*(slopeLeft-vl) - mf*mf)/auxl;
140     }
141     if( fabs(auxr)<epsilon2 ||
142         ( ( ( fabs(slopeM -vr) ) < epsilon2 ) &&
143           ( ( fabs(Byr) + fabs(Bzr) ) < epsilon2 ) &&
144           ( ( mf*mf ) > (Gamma*prer) ) ) ) {
145         vyrs = vyr;
146         vzrs = vzr;
147         Byrs = Byr;
148         Bzrs = Bzr;
149     }else{
150         //Equation 44
151         vyrs = vyr - mf*Byr*(slopeM-vr)/auxr;
152         //Equation 46
153         vzrs = vzr - mf*Bzr*(slopeM-vr)/auxr;
154         //Equation 45;
155         Byrs = Byr*(rhor*(slopeRight-vr)*(slopeRight-vr) - mf*mf)/auxr;
156         //Equation 47
157         Bzrs = Bzr*(rhor*(slopeRight-vr)*(slopeRight-vr) - mf*mf)/auxr;
158     }
159 }else if(AxisNo==2){
160     //velocities
161     //Equation 39
162     vyls = slopeM;
163     vyrs = vyls;
164     //B_y
165     Byls = mf;
166     Byrs = Byls;
167
168     auxl= (rhol*(slopeLeft - vl)*(slopeLeft - slopeM)-mf*mf);
169     auxr= (rhor*(slopeRight - vr)*(slopeRight - slopeM)-mf*mf);
170
171     if( fabs(auxl)<epsilon2 ||
172         ( ( ( fabs(slopeM -vl) ) < epsilon2 ) &&
173           ( ( fabs(Bxl) + fabs(Bzl) ) < epsilon2 ) &&
174           ( ( mf*mf ) > (Gamma*prel) ) ) ) {
175         vxls = vxl;
176         vzls = vzl;
177         Bxls = Bxl;
178         Bzls = Bzl;
179     }else{
180         //Equation 44
181         vxls = vxl - mf*Bxl*(slopeM-vl)/auxl;
182         //Equation 46
183         vzls = vzl - mf*Bzl*(slopeM-vl)/auxl;
184         //Equation 45
185         Bxls = Bxl*(rhol*(slopeLeft-vl)*(slopeLeft-vl) - mf*mf)/auxl;
186         //Equation 47
187         Bzls = Bzl*(rhol*(slopeLeft-vl)*(slopeLeft-vl) - mf*mf)/auxl;
188     }
189
190     if( fabs(auxr)<epsilon2 ||
191         ( ( ( fabs(slopeM -vr) ) < epsilon2 ) &&
192           ( ( fabs(Bxr) + fabs(Bzr) ) < epsilon2 ) &&
193           ( ( mf*mf ) > (Gamma*prer) ) ) ) {
194         vxrs = vxr;
195         vzrs = vzr;
196         Bxrs = Bxr;
197         Bzrs = Bzr;
198     }else{
199         //Equation 44
200         vxrs = vxr - mf*Bxr*(slopeM-vr)/auxr;
201         //Equation 46
202         vzrs = vzr - mf*Bzr*(slopeM-vr)/auxr;
203         //Equation 45
204         Bxrs = Bxr*(rhor*(slopeRight-vr)*(slopeRight-vr) - mf*mf)/auxr;
205         //Equation 47
206         Bzrs = Bzr*(rhor*(slopeRight-vr)*(slopeRight-vr) - mf*mf)/auxr;
207     }
208 }else{
209     //velocities
210     //Equation 39
211     vzls = slopeM;
212     vzrs = vzls;
213     //B_z
214     Bzls = mf;
215     Bzrs = Bzls;
216
217     auxl= (rhol*(slopeLeft - vl)*(slopeLeft - slopeM)-mf*mf);
218     auxr= (rhor*(slopeRight - vr)*(slopeRight - slopeM)-mf*mf);
219
220     if( fabs(auxl)<epsilon2 ||

```

```

222         ( ( ( fabs(slopeM -v1) ) < epsilon2 )  &&
223         ( ( fabs(Bx1) + fabs(By1) ) < epsilon2 )  &&
224         ( ( mf*mf ) > (Gamma*prel) ) ) ) {
225     vxls = vx1;
226     vyls = vyl;
227     Bxls = Bx1;
228     Byls = By1;
229 }else{
230     //Equation 44
231     vxls = vx1 - mf*Bx1*(slopeM-v1)/aux1;
232     //Equation 46
233     vyls = vyl - mf*By1*(slopeM-v1)/aux1;
234     //Equation 45
235     Bxls = Bx1*(rhol*(slopeLeft-v1)*(slopeLeft-v1) - mf*mf)/aux1;
236     //Equation 47
237     Byls = By1*(rhol*(slopeLeft-v1)*(slopeLeft-v1) - mf*mf)/aux1;
238 }
239
240     if( fabs(auxr)<epsilon2 ||
241         ( ( ( fabs(slopeM -vr) ) < epsilon2 )  &&
242         ( ( fabs(Bxr) + fabs(Byr) ) < epsilon2 )  &&
243         ( ( mf*mf ) > (Gamma*prer) ) ) ) ) {
244     vxrs = vxr;
245     vyrs = vyr;
246     Bxrs = Bxr;
247     Byrs = Byr;
248 }else{
249     //Equation 44
250     vxrs = vxr - mf*Bxr*(slopeM-vr)/auxr;
251     //Equation 46
252     vyrs = vyr - mf*Byr*(slopeM-vr)/auxr;
253     //Equation 45
254     Bxrs = Bxr*(rhor*(slopeRight-vr)*(slopeRight-vr) - mf*mf)/auxr;
255     //Equation 47
256     Byrs = Byr*(rhor*(slopeRight-vr)*(slopeRight-vr) - mf*mf)/auxr;
257 }
258 }
259
260 //total pressure - Equation 41
261 pTs = pTl + rhol*(slopeLeft - v1)*(slopeM - v1);
262
263 //inner product vs*Bs
264 vBls = vxls*Bxls + vyls*Byls + vzls*Bzls;
265 vBrs = vxrs*Bxrs + vyrs*Byrs + vzrs*Bzrs;
266
267 //energy - Equation 48
268 Els = ((slopeLeft -v1)*el - pTl*v1+pTs*slopeM+mf*(vBl - vBls))/(slopeLeft - slopeM);
269 Ers = ((slopeRight-vr)*er - pTr*vr+pTs*slopeM+mf*(vBr - vBrs))/(slopeRight - slopeM);
270
271 //U-star variables
272 //Left
273 U.setValue(1,1,rhols);
274 U.setValue(2,1,rhols*vxls);
275 U.setValue(3,1,rhols*vyls);
276 U.setValue(4,1,rhols*vzls);
277 U.setValue(5,1,Els);
278 U.setValue(6,1,psil);
279 U.setValue(7,1,Bxls);
280 U.setValue(8,1,Byls);
281 U.setValue(9,1,Bzls);
282 //Right
283 U.setValue(1,2,rhors);
284 U.setValue(2,2,rhors*vxrs);
285 U.setValue(3,2,rhors*vyrs);
286 U.setValue(4,2,rhors*vzrs);
287 U.setValue(5,2,Ers);
288 U.setValue(6,2,psir);
289 U.setValue(7,2,Bxrs);
290 U.setValue(8,2,Byrs);
291 U.setValue(9,2,Bzrs);
292
293
294
295
296
297 Sqrtrhols = (sqrt(rhols)+sqrt(rhors));
298
299 if((mf*mf/min((Bl*Bl),(Br*Br))) < epsilon2){
300 for(int k=1;k<=QuantityNb;k++){
301     U.setValue(k,3, U.value(k,1));
302     U.setValue(k,4, U.value(k,2));
303 }
304 }else{
305     //density - Equation 49
306     rholss = rhols;
307     rhorss = rhors;
308
309     if(AxisNo==1){

```

```

310         //velocities
311         //Equation 39
312         vxlss = slopeM;
313         vxrss = slopeM;
314         //Equation 59
315         vylss = (sqrt(rhols)*vyls + sqrt(rhors)*vyrs + (Byrs - Byls)*Bsign)/Sqrtrhols;
316         vyrrs = vylss;
317         //Equation 60
318         vzlss = (sqrt(rhols)*vzls + sqrt(rhors)*vzrs + (Bzrs - Bzls)*Bsign)/Sqrtrhols;
319         vzrrs = vzlss;
320
321         //Magnetic Field components
322         //B_x
323         Bxlss = mf;
324         Bxrss = mf;
325         //Equation 61
326         Bylss = (sqrt(rhols)*Byrs + sqrt(rhors)*Byls + sqrt(rhols*rhors)*(vyrs - vyls)*Bsign)/
Sqrtrhols;
327         Byrrs = Bylss;
328         //Equation 62
329         Bzlss = (sqrt(rhols)*Bzrs + sqrt(rhors)*Bzls + sqrt(rhols*rhors)*(vzrs - vzls)*Bsign)/
Sqrtrhols;
330         Bzrrs = Bzlss;
331     }
332     else if(AxisNo==2){
333         //velocities
334         //Equation 59
335         vxlss = (sqrt(rhols)*vxls + sqrt(rhors)*vxrs + (Bxrs - Bxls)*Bsign)/Sqrtrhols;
336         vxrrs = vxlss;
337         //Equation 39
338         vylss = slopeM;
339         vyrrs = slopeM;
340         //Equation 60
341         vzlss = (sqrt(rhols)*vzls + sqrt(rhors)*vzrs + (Bzrs - Bzls)*Bsign)/Sqrtrhols;
342         vzrrs = vzlss;
343
344         //Magnetic Field components
345         //Equation 61
346         Bxlss = (sqrt(rhols)*Bxrs + sqrt(rhors)*Bxls + sqrt(rhols*rhors)*(vxrs - vxls)*Bsign)/
Sqrtrhols;
347         Bxrss = Bxlss;
348         //B_y
349         Bylss = mf;
350         Byrrs = mf;
351         //Equation 62
352         Bzlss = (sqrt(rhols)*Bzrs + sqrt(rhors)*Bzls + sqrt(rhols*rhors)*(vzrs - vzls)*Bsign)/
Sqrtrhols;
353         Bzrrs = Bzlss;
354
355     }else{
356         //velocities
357         //Equation 59
358         vxlss = (sqrt(rhols)*vxls + sqrt(rhors)*vxrs + (Bxrs - Bxls)*Bsign)/Sqrtrhols;
359         vxrrs = vxlss;
360         //Equation 60
361         vylss = (sqrt(rhols)*vyls + sqrt(rhors)*vyrs + (Byrs - Byls)*Bsign)/Sqrtrhols;
362         vyrrs = vylss;
363         //Equation 39
364         vzlss = slopeM;
365         vzrrs = slopeM;
366
367         //Magnetic Field components
368         //Equation 61
369         Bxlss = (sqrt(rhols)*Bxrs + sqrt(rhors)*Bxls + sqrt(rhols*rhors)*(vxrs - vxls)*Bsign)/
Sqrtrhols;
370         Bxrss = Bxlss;
371         //Equation 62
372         Bylss = (sqrt(rhols)*Byrs + sqrt(rhors)*Byls + sqrt(rhols*rhors)*(vyrs - vyls)*Bsign)/
Sqrtrhols;
373         Byrrs = Bylss;
374         //B_y
375         Bzlss = mf;
376         Bzrrs = mf;
377
378     }
379
380
381     //inner product vss*Bss
382     vBlss = vxlss*Bxlss + vylss*Bylss + vzlss*Bzlss;
383     vBrss = vxrss*Bxrss + vyrrs*Byrrs + vzrrs*Bzrrs;
384
385     //Energy - Equation 63
386     Elss = Els - sqrt(rhols)*(vBls - vBlss)*Bsign;
387     Erss = Ers + sqrt(rhors)*(vBrS - vBrss)*Bsign;
388
389
390

```

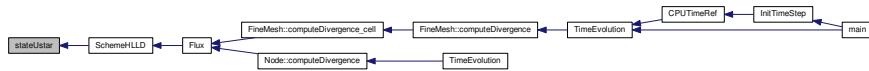


```

391
392         //U-star-star variables
393         //Left
394         U.setValue(1,3,rholss);
395         U.setValue(2,3,rholss*vxlss);
396         U.setValue(3,3,rholss*vylss);
397         U.setValue(4,3,rholss*vzls);
398         U.setValue(5,3,Elss);
399         U.setValue(6,3,psil);
400         U.setValue(7,3,Bxlss);
401         U.setValue(8,3,Bylss);
402         U.setValue(9,3,Bzls);
403         //Right
404         U.setValue(1,4,rhorss);
405         U.setValue(2,4,rhorss*vxrss);
406         U.setValue(3,4,rhorss*vyrss);
407         U.setValue(4,4,rhorss*vzrss);
408         U.setValue(5,4,Erss);
409         U.setValue(6,4,psir);
410         U.setValue(7,4,Bxrss);
411         U.setValue(8,4,Byrss);
412         U.setValue(9,4,Bzrss);
413     }
414     //Equation 61 - S-star left and S-star right
415     slopeLeftStar = slopeM - Abs(mf)/sqrt(rhols);
416     slopeRightStar = slopeM + Abs(mf)/sqrt(rhors);
417
418     return U;
419 }

```

Here is the caller graph for this function:



### 6.6.3.43 real Step ( real x )

Returns a step (1 if x <0, 0 if x >0, 0.5 if x=0)

#### Parameters

|   |            |
|---|------------|
| x | Real value |
|---|------------|

#### Returns

double

```

25 {
26     if (x < 0.)
27         return 1.;
28     else if (x > 0.)
29         return 0.;
30     else
31         return .5;
32 }

```

### 6.6.3.44 void TimeEvolution ( FineMesh \* Root )

Computes a time evolution on the regular fine mesh *Root*. Only for finite volume computations.

## Parameters

|             |           |
|-------------|-----------|
| <i>Root</i> | Fine mesh |
|-------------|-----------|

## Returns

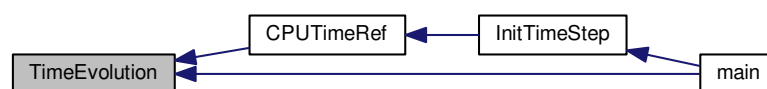
void

```

78 {
79
80     // --- Store cell-average values into temporary ---
81     Root->store();
82
83     for (StepNo = 1; StepNo <= StepNb; StepNo++)
84     {
85         // --- Compute divergence for neighbour cells ---
86         //The same conception with background computations, see upper...
87         Root->computeDivergence(1);
88         // --- Runge-Kutta step for neighbour cells ---
89         Root->RungeKutta(1);
90         // --- Divergence cleaning source term
91         Root->computeCorrection(1);
92         // --- Start inter-CPU exchanges ---
93         CPUExchange(Root, SendQ);
94         // --- Compute divergence for internal cells ---
95         Root->computeDivergence(0);
96         // --- Runge-Kutta step for internal cells ---
97         Root->RungeKutta(0);
98         // --- Divergence cleaning source term
99         Root->computeCorrection(0);
100
101     #if defined PARMPI
102         CommTimer.start(); //Communication Timer Start
103         //Waiting while inter-CPU exchanges are finished
104         if (MPIRecvType == 1) //for nonblocking receive...
105             MPI_Waitall(4*Dimension, req, st);
106         CommTimer.stop();
107     #endif
108     }
109
110     // --- Check stability ---
111     Root->checkStability();
112
113     // --- Compute integral values ---
114     Root->computeIntegral();
115
116     // --- Compute elapsed time and adapt time step ---
117
118     if (!ComputeCPUTimeRef)
119     {
120         Eigenvalue = Max(EigenvalueX, Max(
121             EigenvalueY, EigenvalueZ));
122         ElapsedTime += TimeStep;
123         if (!ConstantTimeStep) AdaptTimeStep();
124         // --- Compute divergence-free correction constant
125         //ch = CFL*SpaceStep/TimeStep;
126         ch = Max(CFL*SpaceStep/TimeStep, Eigenvalue);
127     }
128
129     // --- Compute time-average values ---
130
131     if (TimeAveraging)
132         Root->computeTimeAverage();
133
134 }

```

Here is the caller graph for this function:



## 6.6.3.45 void TimeEvolution ( Node \* Root )

Computes a time evolution on the tree structure, the root node being *Root*. Only for multiresolution computations.

## Parameters

|             |           |
|-------------|-----------|
| <i>Root</i> | Root node |
|-------------|-----------|

## Returns

void

```

20 {
21
22     // --- Smooth data ---
23
24     if ( SmoothCoeff != 0.)
25         Root->smooth();
26
27     // --- Store cell-average values of leaves ---
28     Root->store();
29
30     // --- Refresh tree structure ---
31     RefreshTree(Root);
32
33     for (StepNo = 1; StepNo <= StepNb; StepNo++)
34     {
35         // --- Compute divergence ---
36         Root->computeDivergence();
37         // --- Runge-Kutta step ---
38         Root->RungeKutta();
39         // --- Divergence cleaning source-terms
40         Root->computeCorrection();
41
42     }
43
44     // --- Refresh tree structure ---
45     RefreshTree(Root);
46
47
48
49     // --- Check stability ---
50     Root->checkStability();
51
52     // --- Compute integral values ---
53     Root->computeIntegral();
54
55     // --- Compute total number of cells and leaves ---
56
57     TotalCellNb += CellNb;
58     TotalLeafNb += LeafNb;
59     //cout<<"eigen= "<<Eigenvalue<<endl;
60     // --- Compute elapsed time and adapt time step ---
61     Eigenvalue = Max(EigenvalueX,Max(EigenvalueY,
EigenvalueZ));
62     ElapsedTime += TimeStep;
63     if (!ConstantTimeStep) AdaptTimeStep();
64
65     // --- Compute divergence-free correction constant
66     //ch = CFL*SpaceStep/TimeStep;
67     ch = Max(CFL*SpaceStep/TimeStep, Eigenvalue);
68
69 }
```

## 6.6.3.46 void View ( FineMesh \* Root, const char \* AverageFileName )

Writes the current cell-averages of the fine mesh *Root* into file *AverageFileName*. Only for finite volume computations.

## Parameters

|                        |           |
|------------------------|-----------|
| <i>Root</i>            | Fine mesh |
| <i>AverageFileName</i> | File name |

### Returns

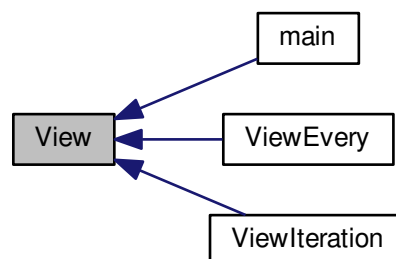
void

```

88 {
89     char buf[256];
90     intiaux;
91
92
93     char CPUFileName[255];
94     #if defined PARMPI
95         sprintf(CPUFileName, "%d_%d_%d_%s", coords[0], coords[1], coords[2], AverageFileName);
96     #else
97         strcpy(CPUFileName, AverageFileName);
98     #endif
99
100     // write header for graphic visualization
101     Root->writeHeader(CPUFileName);
102
103     // write cell-average values for graphic visualization
104     Root->writeAverage(CPUFileName);
105
106     // Compress data
107     if (Dimension != 1)
108     {
109         if (ZipData)
110         {
111             sprintf(buf, "gzip %s", CPUFileName);
112            iaux=system(buf);
113         }
114     }
115
116     // --- Write time-average values into file ---
117
118     if (TimeAveraging)
119         Root->writeTimeAverage("TimeAverage.dat");
120
121 }

```

Here is the caller graph for this function:



**6.6.3.47** void View ( Node \* Root, const char \* TreeFileName, const char \* MeshFileName, const char \* AverageFileName )

Writes the data of the tree structure into files *TreeFileName* (tree structure), *MeshFileName* (mesh) and *AverageFileName* (cell-averages). The root node is *Root*. Only for multiresolution computations.

## Parameters

|                        |                   |
|------------------------|-------------------|
| <i>Root</i>            | Root node         |
| <i>TreeFileName</i>    | Tree file name    |
| <i>MeshFileName</i>    | Mesh file name    |
| <i>AverageFileName</i> | Average file name |

## Returns

void

```

36 {
37     char buf[256];
38     intiaux;
39
40     // write tree (debugging only)
41     if (debug) Root->writeTree(TreeFileName);
42
43 // Root->computeCorrection();
44
45     // write mesh for graphic visualisation
46     if (Dimension != 1)
47     {
48         Root->writeHeader(MeshFileName);
49         Root->writeAverage(MeshFileName);
50
51         // Compress data (if parameter ZipData is true)
52         if (ZipData)
53         {
54             sprintf(buf, "gzip %s", MeshFileName);
55            iaux=system(buf);
56         }
57     }
58     else
59         Root->writeMesh(MeshFileName);
60
61
62     // write cell-averages in multiresolution representation (1D) or on fine grid (2-3D)
63     if (Dimension != 1)
64     {
65         Root->writeFineGrid(AverageFileName, ScaleNb+
PrintMoreScales);
66
67         // Compress data
68         if (ZipData)
69         {
70             sprintf(buf, "gzip %s", AverageFileName);
71            iaux=system(buf);
72         }
73     }
74     else
75     {
76         Root->writeHeader(AverageFileName);
77         Root->writeAverage(AverageFileName);
78     }
79 }

```

## 6.6.3.48 void ViewEvery ( FineMesh \* Root, int arg )

Same as previous for a fine mesh *Root*. Only for finite volume computations.

## Parameters

|             |           |
|-------------|-----------|
| <i>Root</i> | Fine mesh |
| <i>arg</i>  | argument  |

## Returns

void

```

54 {
55     char AverageName[256]; // File name for AverageNNN.dat

```

```

56     char AverageFormat[256];    // File format for AverageNNN.dat
57
58     sprintf(AverageFormat, "Average%s0%i.vtk", "%", DigitNumber(
IterationNb));
59     sprintf(AverageName, AverageFormat, arg);
60
61     View(Root, AverageName);
62 }

```

Here is the caller graph for this function:



#### 6.6.3.49 void ViewEvery ( Node \* Root, int arg )

Writes into file the data of the tree structure at iteration *arg*. The output file names are *AverageNNN.dat* and *MeshNNN.dat*, *NNN* being the iteration in an accurate format. The root node is *Root*. Only for multiresolution computations.

##### Parameters

|             |           |
|-------------|-----------|
| <i>Root</i> | Root node |
| <i>arg</i>  | Argument  |

##### Returns

void

```

33 {
34     char AverageName[256];    // File name for AverageNNN.dat
35     char MeshName[256];    // File name for MeshNNN.dat
36     char AverageFormat[256];    // File format for AverageNNN.dat
37     char MeshFormat[256];    // File format for MeshNNN.dat
38
39     sprintf(AverageFormat, "Average%s0%i.vtk", "%", DigitNumber(
IterationNb));
40     sprintf(AverageName, AverageFormat, arg);
41     sprintf(MeshFormat, "Mesh%s0%i.dat", "%", DigitNumber(
IterationNb));
42     sprintf(MeshName, MeshFormat, arg);
43
44     View(Root, "Tree.dat", MeshName, AverageName);
45 }

```

#### 6.6.3.50 void ViewIteration ( FineMesh \* Root )

Same as previous for a fine mesh *Root*. Only for finite volume computations.

##### Parameters

|             |           |
|-------------|-----------|
| <i>Root</i> | Fine mesh |
|-------------|-----------|

## Returns

void

```

61 {
62     if (IterationNo == PrintIt1)
63         View(Root, "Average_1.vtk");
64
65     if (IterationNo == PrintIt2)
66         View(Root, "Average_2.vtk");
67
68     if (IterationNo == PrintIt3)
69         View(Root, "Average_3.vtk");
70
71     if (IterationNo == PrintIt4)
72         View(Root, "Average_4.vtk");
73
74     if (IterationNo == PrintIt5)
75         View(Root, "Average_5.vtk");
76
77     if (IterationNo == PrintIt6)
78         View(Root, "Average_6.vtk");
79 }

```

Here is the caller graph for this function:



## 6.6.351 void ViewIteration ( Node \* Root )

Writes into file the data of the tree structure from physical time *PrintTime1* to physical time *PrintTime6*. The output file names are *Average\_N.dat* and *Mesh\_N.dat*, *N* being between 1 and 6. The root node is *Root*. Only for multiresolution computations.

## Parameters

|             |           |
|-------------|-----------|
| <i>Root</i> | Root node |
|-------------|-----------|

## Returns

void

```

34 {
35     if (IterationNo == PrintIt1)
36         View(Root, "Tree.dat", "Mesh_1.dat", "Average_1.vtk");
37
38     if (IterationNo == PrintIt2)
39         View(Root, "Tree.dat", "Mesh_2.dat", "Average_2.vtk");
40
41     if (IterationNo == PrintIt3)
42         View(Root, "Tree.dat", "Mesh_3.dat", "Average_3.vtk");
43
44     if (IterationNo == PrintIt4)
45         View(Root, "Tree.dat", "Mesh_4.dat", "Average_4.vtk");
46
47     if (IterationNo == PrintIt5)
48         View(Root, "Tree.dat", "Mesh_5.dat", "Average_5.vtk");
49
50     if (IterationNo == PrintIt6)
51         View(Root, "Tree.dat", "Mesh_6.dat", "Average_6.vtk");
52 }

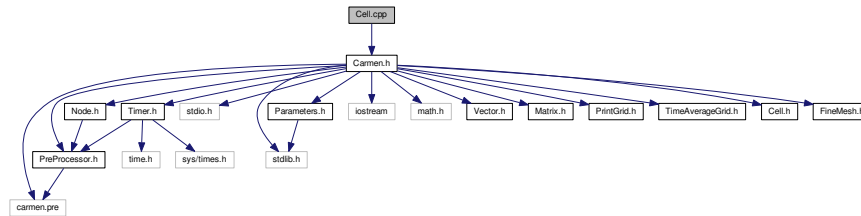
```

## 6.7 Cell.cpp File Reference

Constructor and destructor of the cell class. Also computes the cell-averages of the MHD variables.

```
#include "Carmen.h"
```

Include dependency graph for Cell.cpp:



### 6.7.1 Detailed Description

Constructor and destructor of the cell class. Also computes the cell-averages of the MHD variables.

## 6.8 Cell.h File Reference

This graph shows which files directly or indirectly include this file:



### Classes

- class [Cell](#)

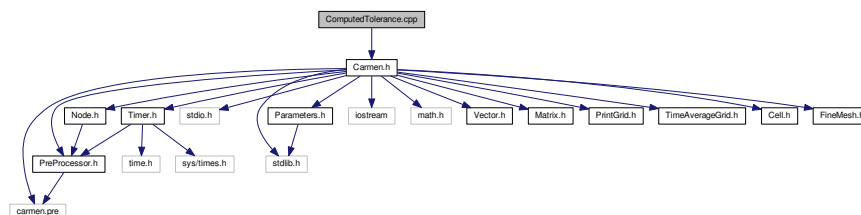
An object [Cell](#) contains all the informations of a cell for both multiresolution and finite volume computations.

## 6.9 ComputedTolerance.cpp File Reference

Adapt threshold parameter or use it fixed.

```
#include "Carmen.h"
```

Include dependency graph for ComputedTolerance.cpp:





Functions

- [real ComputedTolerance](#) (const int ScaleNo)  
Returns the computed tolerance at the scale ScaleNo, either using Harten or Donoho thresholding (if CVS=true).

6.9.1 Detailed Description

Adapt trheshold parameter or use it fixed.

6.9.2 Function Documentation

6.9.2.1 real ComputedTolerance ( const int ScaleNo )

Returns the computed tolerance at the scale ScaleNo, either using Harten or Donoho thresholding (if CVS=true).

Parameters

|         |                    |
|---------|--------------------|
| ScaleNo | Level of interest. |
|---------|--------------------|

Returns

double

```

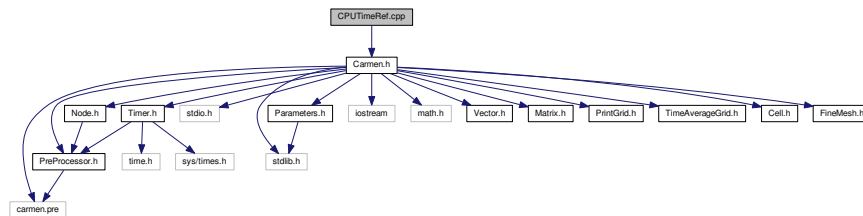
23 {
24 //if ThresholdNorm==0 const Tolerance, else L1 Harten norm
25
26     if(ThresholdNorm)
27         return((Tolerance/GlobalVolume)*exp(Dimension*(ScaleNo-
ScaleNb+1)*log(2.)));
28     else
29         return(Tolerance);
30
31 }
```

6.10 CPUTimeRef.cpp File Reference

Compute the reference CPU time with the finite volume solver. The output is the CPU time for 1 iteration.

```
#include "Carmen.h"
```

Include dependency graph for CPUTimeRef.cpp:



Functions

- double [CPUTimeRef](#) (int iterations, int scales)  
Returns the time required by a finite volume computation using iterations iterations and scales scales. It is use to estimate the CPU time compression.

## 6.10.1 Detailed Description

Compute the reference CPU time with the finite volume solver. The output is the CPU time for 1 iteration.

## 6.10.2 Function Documentation

### 6.10.2.1 `double CPUTimeRef ( int iterations, int scales )`

Returns the time required by a finite volume computation using *iterations* iterations and *scales* scales. It is use to estimate the CPU time compression.

#### Parameters

|                   |                       |
|-------------------|-----------------------|
| <i>iterations</i> | Number of iterations. |
| <i>scales</i>     | Scales                |

#### Returns

double

```

24 {
25     // --- Local variables -----
26
27     int OldIterationNb=0;
28     int OldScaleNb=0;
29     real OldTimeStep=0.;
30     bool ConstantTimeStepOld=ConstantTimeStep;
31
32     double result=0.;
33
34     Timer CPURef;
35     FineMesh* MeshRef;
36
37     // --- Execution -----
38
39     // Toggle on : Compute reference CPU time
40
41     ComputeCPUTimeRef = true;
42
43     // Toggle off : Constant time step
44
45     ConstantTimeStep = true;
46
47     // backup values of IterationNb and ScaleNb
48
49     OldIterationNb = IterationNb;
50     OldScaleNb = ScaleNb;
51     OldTimeStep = TimeStep;
52
53     // use reference values
54     IterationNb = iterations;
55     ScaleNb = scales;
56     TimeStep = 0.;
57
58     one_D=1; two_D=1;
59     if (Dimension >= 2) one_D=1<<ScaleNb;
60     if (Dimension == 3) two_D=1<<ScaleNb;
61
62     // init mesh
63     MeshRef = new FineMesh;
64
65     // Iterate on time
66
67     for (IterationNo = 1; IterationNo <= IterationNb;
IterationNo++)
68     {
69         // start timer
70         CPURef.start();
71
72         // Compute time evolution
73         TimeEvolution(MeshRef);
74
75         // check CPU Time
76         CPURef.check();
77
78         // stop timer
79         CPURef.stop();

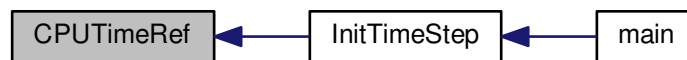
```

```

80     }
81
82     // Compute CPUTimeRef
83     result = CPURef.CPUTime();
84     result *= 1./IterationNb;
85     result *= 1<<(Dimension*(OldScaleNb-ScaleNb));
86
87     // delete MeshRef
88     delete MeshRef;
89
90     // restore values of IterationNb and ScaleNb
91     IterationNb = OldIterationNb;
92     ScaleNb = OldScaleNb;
93     TimeStep = OldTimeStep;
94     IterationNo = 0;
95
96     one_D=1; two_D=1;
97     if (Dimension >= 2) one_D=1<<ScaleNb;
98     if (Dimension == 3) two_D=1<<ScaleNb;
99
100    // Toggle off : Compute reference CPU time
101
102    ComputeCPUTimeRef = false;
103
104    // Restore the value of ConstantTimeStep
105
106    ConstantTimeStep = ConstantTimeStepOld;
107
108    return result;
109 }

```

Here is the caller graph for this function:

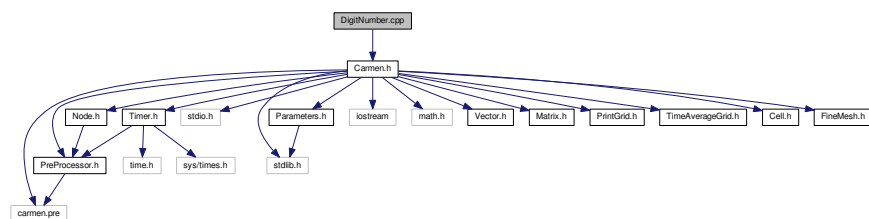


## 6.11 DigitNumber.cpp File Reference

This function returns the number of digits of an integer.

```
#include "Carmen.h"
```

Include dependency graph for DigitNumber.cpp:



### Functions

- int [DigitNumber](#) (int arg)

Returns the number of digits of the integer arg.

### 6.11.1 Detailed Description

This function returns the number of digits of an integer.

### 6.11.2 Function Documentation

#### 6.11.2.1 int DigitNumber ( int *arg* )

Returns the number of digits of the integer *arg*.

#### Parameters

|            |          |
|------------|----------|
| <i>arg</i> | Argument |
|------------|----------|

#### Returns

int

```

23 {
24     int result;
25     int i;
26
27     result = 0;
28     i = arg;
29
30     while(i != 0)
31     {
32         i/=10;
33         result++;
34     }
35
36     return result;
37 }
```

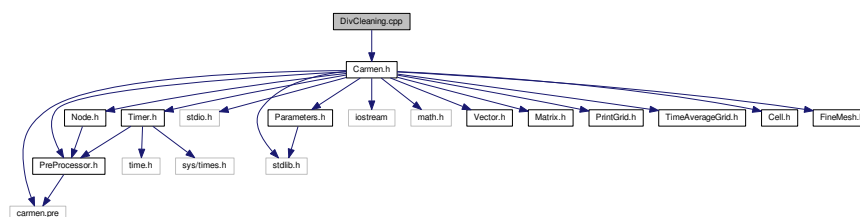
Here is the caller graph for this function:



## 6.12 DivCleaning.cpp File Reference

```
#include "Carmen.h"
```

Include dependency graph for DivCleaning.cpp:



## Functions

- [Vector DivCleaning](#) (const [Cell](#) &Cell0, const [Cell](#) &Cell1, const [Cell](#) &Cell2, int AxisNo)

### 6.12.1 Function Documentation

#### 6.12.1.1 Vector DivCleaning ( const Cell & Cell0, const Cell & Cell1, const Cell & Cell2, int AxisNo )

```

11
12
13     Vector LeftAvg(QuantityNb);
14     Vector RightAvg(QuantityNb);
15     Vector Avg(QuantityNb);
16     Vector p8wave(QuantityNb);
17     real Bdiv=0;
18     real udotB=0;
19     real psiGrad=0;
20     real dx=0;
21
22     LeftAvg = Cell1.average();
23     RightAvg = Cell2.average();
24     Avg = Cell0.average();
25
26     dx = Cell0.size(AxisNo);
27     dx = 2*dx;
28     Bdiv = (RightAvg.value(AxisNo+6) - LeftAvg.value(AxisNo+6))/dx;
29     p8wave.setZero();
30
31     if(DivClean==1){
32         udotB = Avg.value(2)*Avg.value(7) + Avg.value(3)*Avg.value(8) + Avg.value(4)*Avg.value(9);
33
34         p8wave.setValue(2, -Bdiv*Avg.value(7));
35         p8wave.setValue(3, -Bdiv*Avg.value(8));
36         p8wave.setValue(4, -Bdiv*Avg.value(9));
37         p8wave.setValue(7, -Bdiv*Avg.value(2)/Avg.value(1));
38         p8wave.setValue(8, -Bdiv*Avg.value(3)/Avg.value(1));
39         p8wave.setValue(9, -Bdiv*Avg.value(4)/Avg.value(1));
40         p8wave.setValue(5, -Bdiv*udotB/Avg.value(1));
41
42         return p8wave;
43     }
44     }else if(DivClean==2){
45
46         psiGrad = (RightAvg.value(6) - LeftAvg.value(6))/dx;
47
48         p8wave.setValue(2, -Bdiv*Avg.value(7));
49         p8wave.setValue(3, -Bdiv*Avg.value(8));
50         p8wave.setValue(4, -Bdiv*Avg.value(9));
51         p8wave.setValue(5, -Avg.value(AxisNo+6)*psiGrad);
52         p8wave.setValue(6, Avg.value(6)*exp(-(cr*ch*TimeStep/
SpaceStep)));
53
54         return p8wave;
55     }else if(DivClean==3){
56         if(AxisNo==3)
57             p8wave.setValue(6, Avg.value(6)*exp(-(cr*ch*TimeStep/
SpaceStep)));
58         return p8wave;
59     }else
60         return p8wave;
61
62
63 }

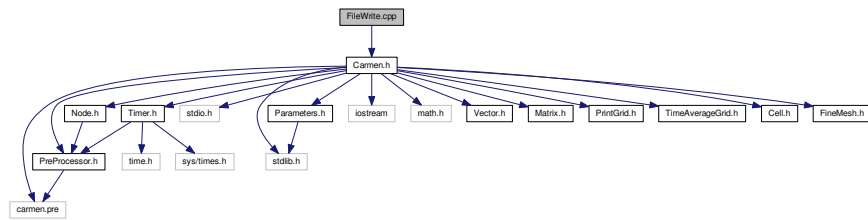
```

## 6.13 FileWrite.cpp File Reference

Writes in binary or ASCII mode the real number *arg* into the file *f*. The global parameter *DatalsBinary* determines this choice.

```
#include "Carmen.h"
```

Include dependency graph for FileWrite.cpp:



## Functions

- int [FileWrite](#) (FILE \*f, const char \*format, real arg)

Writes in binary or ASCII mode the real number *arg* into the file *f* with the format *format*. The global parameter *DataIsBinary* determines this choice.

### 6.13.1 Detailed Description

Writes in binary or ASCII mode the real number *arg* into the file *f*. The global parameter *DataIsBinary* determines this choice.

### 6.13.2 Function Documentation

#### 6.13.2.1 int FileWrite ( FILE \* f, const char \* format, real arg )

Writes in binary or ASCII mode the real number *arg* into the file *f* with the format *format*. The global parameter *DataIsBinary* determines this choice.

#### Parameters

|               |           |
|---------------|-----------|
| <i>f</i>      | File name |
| <i>format</i> | Format    |
| <i>arg</i>    | Argument  |

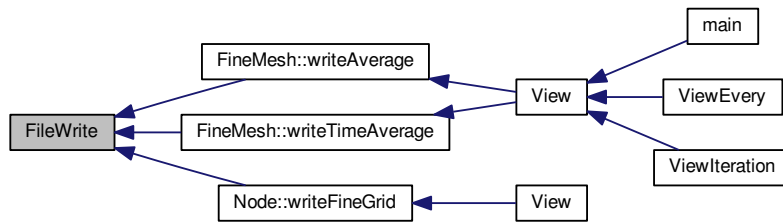
#### Returns

int

```

23 {
24     int result;
25     real x;
26
27     x = arg;
28
29     if (DataIsBinary)
30         result = fwrite(&x, sizeof(real), 1, f);
31     else
32         result = fprintf(f, format, x);
33
34
35     return result;
36 }
  
```

Here is the caller graph for this function:

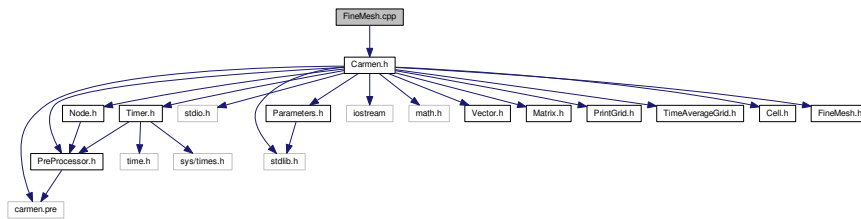


## 6.14 FineMesh.cpp File Reference

Fine mesh simulation functions.

```
#include "Carmen.h"
```

Include dependency graph for FineMesh.cpp:



### 6.14.1 Detailed Description

Fine mesh simulation functions.

Version

2.0

Date

November-2016

## 6.15 FineMesh.h File Reference

This graph shows which files directly or indirectly include this file:



## Classes

- class [FineMesh](#)

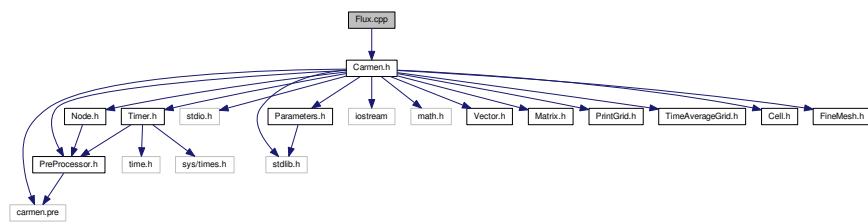
An object [FineMesh](#) is a regular fine mesh, used for finite volume computations. It is not used for multiresolution computations.

## 6.16 Flux.cpp File Reference

Computes the numerical fluxes HLL and HLLD.

```
#include "Carmen.h"
```

Include dependency graph for Flux.cpp:



## Functions

- [Vector Flux](#) ([Cell](#) &[Cell1](#), [Cell](#) &[Cell2](#), [Cell](#) &[Cell3](#), [Cell](#) &[Cell4](#), int [AxisNo](#))

Returns the flux at the interface between *Cell2* and *Cell3*. Here a 4-point space scheme is used. *Cell2* and *Cell3* are the first neighbours on the left and right sides. *Cell1* and *Cell4* are the second neighbours on the left and right sides.

### 6.16.1 Detailed Description

Computes the numerical fluxes HLL and HLLD.

### 6.16.2 Function Documentation

#### 6.16.2.1 Vector Flux ( [Cell](#) & [Cell1](#), [Cell](#) & [Cell2](#), [Cell](#) & [Cell3](#), [Cell](#) & [Cell4](#), int [AxisNo](#) )

Returns the flux at the interface between *Cell2* and *Cell3*. Here a 4-point space scheme is used. *Cell2* and *Cell3* are the first neighbours on the left and right sides. *Cell1* and *Cell4* are the second neighbours on the left and right sides.

#### Parameters

|               |                                    |
|---------------|------------------------------------|
| <i>Cell1</i>  | second neighbour on the left side  |
| <i>Cell2</i>  | first neighbour on the left side   |
| <i>Cell3</i>  | first neighbour on the right side  |
| <i>Cell4</i>  | second neighbour on the right side |
| <i>AxisNo</i> | Axis of interest                   |



## Returns

Vector

```

23 {
24     // --- Local variables ---
25
26     Vector Result(QuantityNb);
27
28     Cell C1, C2, C3, C4;
29
30     int BoundaryCell1 = BoundaryRegion(Cell1.center());
31     int BoundaryCell2 = BoundaryRegion(Cell2.center());
32     int BoundaryCell3 = BoundaryRegion(Cell3.center());
33     int BoundaryCell4 = BoundaryRegion(Cell4.center());
34
35     bool UseBoundaryCells = (UseBoundaryRegions && (BoundaryCell1!=0 || BoundaryCell2!=0
|| BoundaryCell3!=0 || BoundaryCell4!=0));
36
37     // --- Take into account boundary conditions ---
38
39     if (UseBoundaryCells)
40         GetBoundaryCells(Cell1, Cell2, Cell3, Cell4, C1, C2, C3, C4, AxisNo);
41
42     switch(SchemeNb)
43     {
44     case 1:
45     default:
46         if (UseBoundaryCells)
47             Result = SchemeHLL(C1, C2, C3, C4, AxisNo);
48         else
49             Result = SchemeHLL(Cell1, Cell2, Cell3, Cell4, AxisNo);
50         break;
51
52     case 2:
53         if (UseBoundaryCells)
54             Result = SchemeHLLD(C1, C2, C3, C4, AxisNo);
55         else
56             Result = SchemeHLLD(Cell1, Cell2, Cell3, Cell4, AxisNo);
57         break;
58     break;
59
60     }
61
62
63     return Result;
64 }

```

Here is the caller graph for this function:

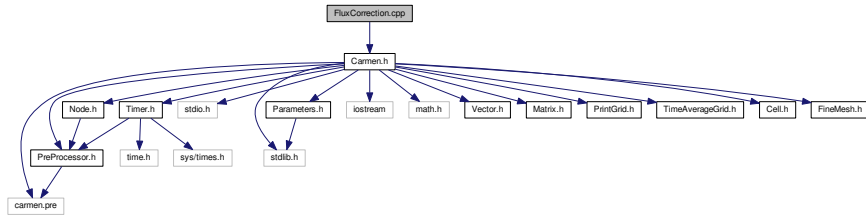


## 6.17 FluxCorrection.cpp File Reference

Computes the mixed correction in the numerical fluxes (Dedner, 2002)

```
#include "Carmen.h"
```

Include dependency graph for FluxCorrection.cpp:



## Functions

- void `fluxCorrection` (`Vector` &`Flux`, const `Vector` &`AvgL`, const `Vector` &`AvgR`, int `AxisNo`)  
*This function apply the divergence-free correction to the numerical flux.*

### 6.17.1 Detailed Description

Computes the mixed correction in the numerical fluxes (Dedner, 2002)

#### Author

Anna Karina Fontes Gomes

#### Version

2.0

### 6.17.2 Function Documentation

#### 6.17.2.1 void fluxCorrection ( `Vector` & `Flux`, const `Vector` & `AvgL`, const `Vector` & `AvgR`, int `AxisNo` )

This function apply the divergence-free correction to the numerical flux.

#### Parameters

|                     |                       |
|---------------------|-----------------------|
| <code>Flux</code>   | Numerical flux vector |
| <code>AvgL</code>   | Left average vector   |
| <code>AvgR</code>   | Right average vector  |
| <code>AxisNo</code> | Axis of interest      |

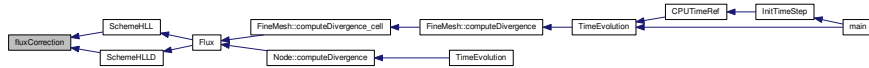
#### Returns

void

```

10 {
11     auxvar = Flux.value(AxisNo+6);
12
13     Flux.setValue(AxisNo+6, Flux.value(AxisNo+6) + (AvgL.value(6) + .5*(AvgR.
14     value(6) - AvgL.value(6))
15     - ch*.5*(AvgR.value(AxisNo+6) - AvgL.
16     value(AxisNo+6)));
17     Flux.setValue(6, ch*ch*(AvgL.value(AxisNo+6) + .5*(AvgR.
18     value(AxisNo+6) - AvgL.value(AxisNo+6))
19     - .5*(AvgR.value(6) - AvgL.value(6))/
20     ch);
21 }
  
```

Here is the caller graph for this function:

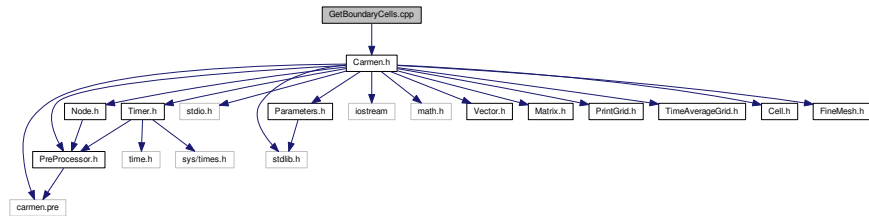


## 6.18 GetBoundaryCells.cpp File Reference

Computes the cells C1, C2, C3, C4 in function of the cells Cell1, Cell2, Cell3, Cell4 taking into account boundary conditions.

```
#include "Carmen.h"
```

Include dependency graph for GetBoundaryCells.cpp:



### Functions

- void [GetBoundaryCells](#) (Cell &Cell1, Cell &Cell2, Cell &Cell3, Cell &Cell4, Cell &C1, Cell &C2, Cell &C3, Cell &C4, const int AxisNo)
  - Transform the 4 cells of the flux Cell1, Cell2, Cell3, Cell4 into C1, C2, C3, C4, to take into account boundary conditions (used in [Flux.cpp](#)).

#### 6.18.1 Detailed Description

Computes the cells C1, C2, C3, C4 in function of the cells Cell1, Cell2, Cell3, Cell4 taking into account boundary conditions.

#### 6.18.2 Function Documentation

6.18.2.1 void [GetBoundaryCells](#) ( Cell & Cell1, Cell & Cell2, Cell & Cell3, Cell & Cell4, Cell & C1, Cell & C2, Cell & C3, Cell & C4, int AxisNo )

Transform the 4 cells of the flux Cell1, Cell2, Cell3, Cell4 into C1, C2, C3, C4, to take into account boundary conditions (used in [Flux.cpp](#)).

#### Parameters

|       |                                   |
|-------|-----------------------------------|
| Cell1 | second neighbour on the left side |
| Cell2 | first neighbour on the left side  |

|               |                                    |
|---------------|------------------------------------|
| <i>Cell3</i>  | first neighbour on the right side  |
| <i>Cell4</i>  | second neighbour on the right side |
| <i>C1</i>     | Auxiliar cell1                     |
| <i>C2</i>     | Auxiliar cell2                     |
| <i>C3</i>     | Auxiliar cell3                     |
| <i>C4</i>     | Auxiliar cell4                     |
| <i>AxisNo</i> | ...                                |

## Returns

void

```

26 {
27 // --- Local variables ---
28
29 int InCell1, InCell2, InCell3, InCell4; // Boundary conditions in cells 1, 2, 3, 4
30 real P1, P2, P3, P4; // Pressures in cells 1, 2, 3, 4
31 real T1, T2, T3, T4; // Temperatures in cells 1, 2, 3, 4
32 real rho1, rho2, rho3, rho4; // Densities in cells 1, 2, 3, 4
33 Vector V1(Dimension), V2(Dimension), V3(Dimension), V4(
Dimension); // Velocities in cells 1, 2, 3, 4
34 real e1, e2, e3, e4; // Energies in cell 1, 2, 3, 4
35 real Y1=0., Y2=0., Y3=0., Y4=0.; // Partial mass in cell 1, 2, 3, 4
36
37 int i; // Counter
38
39 // --- Init C1, C2, C3, C4 ---
40
41 C1 = Cell1;
42 C2 = Cell2;
43 C3 = Cell3;
44 C4 = Cell4;
45
46 // --- Depending on the boundary region type, transform C1, C2, C3, C4 ---
47
48 InCell1 = BoundaryRegion(Cell1.center());
49 InCell2 = BoundaryRegion(Cell2.center());
50 InCell3 = BoundaryRegion(Cell3.center());
51 InCell4 = BoundaryRegion(Cell4.center());
52
53 // --- Cell2 IN THE BOUNDARY, Cell3 IN THE FLUID -----
54
55 if (InCell2 !=0 && InCell3 == 0)
56 {
57 switch(InCell2)
58 {
59 // INFLOW
60 case 1:
61 // Dirichlet on temperature
62 T2 = Cell2.temperature();
63 T1 = Cell1.temperature();
64
65 // Extrapolate pressure
66 P2 = Cell3.oldPressure();
67 P1 = P2;
68 // P1 = 2*P2 - Cell3.pressure();
69
70 // Compute density
71 rho2 = Gamma*Ma*Ma*P2/T2;
72 rho1 = Gamma*Ma*Ma*P1/T1;
73
74 // Dirichlet on momentum
75 V2 = (Cell2.density()/rho2)*Cell2.velocity();
76 V1 = (Cell1.density()/rho1)*Cell1.velocity();
77
78 // Dirichlet on partial mass
79 if (ScalarEqNb == 1)
80 {
81 Y2 = Cell2.average(Dimension+3)/Cell2.
density();
82 Y1 = Cell1.average(Dimension+3)/Cell1.
density();
83 }
84
85 // Compute energies
86 e2 = P2/((Gamma-1.)*rho2) + 0.5*N2(V2);
87 e1 = P1/((Gamma-1.)*rho1) + 0.5*N2(V1);
88
89 // Correct C1 and C2
90 C2.setAverage(1, rho2);
91 C1.setAverage(1, rho1);

```

```

92         for (i=1; i<=Dimension; i++)
93         {
94             C2.setAverage(i+1, rho2*V2.value(i));
95             C1.setAverage(i+1, rho1*V1.value(i));
96         }
97         C2.setAverage(Dimension+2, rho2*e2);
98         C1.setAverage(Dimension+2, rho1*e1);
99
100        if (ScalarEqNb == 1)
101        {
102            C2.setAverage(Dimension+3, rho2*Y2);
103            C1.setAverage(Dimension+3, rho1*Y1);
104        }
105        break;
106
107        // OUTFLOW : use the old value of the neighbour
108        case 2:
109            C2.setAverage(Cell3.oldAverage());
110            C1.setAverage(Cell3.oldAverage());
111
112            // Also change the values in the boundary
113            Cell2.setAverage(C2.average());
114            Cell1.setAverage(C1.average());
115            break;
116
117        // FREE-SLIP WALL : Neuman on all quantities
118        case 3:
119
120            C2 = Cell3;
121            C1 = Cell4;
122            break;
123
124        // ADIABATIC WALL
125        case 4:
126
127            // Dirichlet on velocity
128            V2 = Cell2.velocity();
129            V1 = Cell1.velocity();
130
131            // Neuman on temperature
132            T2 = Cell3.temperature();
133            T1 = Cell4.temperature();
134
135            // Neuman on pressure
136            P2 = Cell3.pressure();
137            P1 = Cell4.pressure();
138
139            // Extrapolate pressure
140            //P2 = 2*Cell3.pressure()-Cell4.pressure();
141            //P1 = P2;
142
143            // Compute densities
144            rho2 = Gamma*Ma*Ma*P2/T2;
145            rho1 = Gamma*Ma*Ma*P1/T1;
146
147            // Compute energies
148            e2 = P2/((Gamma-1.)*rho2) + 0.5*N2(V2);
149            e1 = P1/((Gamma-1.)*rho1) + 0.5*N2(V1);
150
151            // Correct C1 and C2
152            C2.setAverage(1, rho2);
153            C1.setAverage(1, rho1);
154            for (i=1; i<=Dimension; i++)
155            {
156                C2.setAverage(i+1, rho2*V2.value(i));
157                C1.setAverage(i+1, rho1*V1.value(i));
158            }
159            C2.setAverage(Dimension+2, rho2*e2);
160            C1.setAverage(Dimension+2, rho1*e1);
161
162            // Neuman on partial mass
163            if (ScalarEqNb == 1)
164            {
165                C2.setAverage(Dimension+3, Cell3.average(
Dimension+3));
166                C1.setAverage(Dimension+3, Cell4.average(
Dimension+3));
167            }
168
169            break;
170
171        // ISOTHERMAL WALL
172        case 5:
173
174            // Dirichlet on velocity
175            V2 = Cell2.velocity();
176            V1 = Cell1.velocity();

```

```

177
178 // Dirichlet on temperature
179 T2 = Cell2.temperature();
180 T1 = Cell1.temperature();
181
182 // Neuman on pressure
183 P2 = Cell3.pressure();
184 P1 = Cell4.pressure();
185
186 // Extrapolate pressure
187 //P2 = 2*Cell3.pressure()-Cell4.pressure();
188 //P1 = P2;
189
190 // Compute densities
191 rho2 = Gamma*Ma*Ma*P2/T2;
192 rho1 = Gamma*Ma*Ma*P1/T1;
193
194 // Compute energies
195 e2 = P2/((Gamma-1.)*rho2) + 0.5*N2(V2);
196 e1 = P1/((Gamma-1.)*rho1) + 0.5*N2(V1);
197
198 // Correct C1 and C2
199 C2.setAverage(1, rho2);
200 C1.setAverage(1, rho1);
201 for (i=1; i<=Dimension; i++)
202 {
203     C2.setAverage(i+1, rho2*V2.value(i));
204     C1.setAverage(i+1, rho1*V1.value(i));
205 }
206 C2.setAverage(Dimension+2, rho2*e2);
207 C1.setAverage(Dimension+2, rho1*e1);
208
209 // Neuman on partial mass
210 if (ScalarEqNb == 1)
211 {
212     C2.setAverage(Dimension+3, Cell3.average(
213 Dimension+3));
214     C1.setAverage(Dimension+3, Cell4.average(
215 Dimension+3));
216 }
217 break;
218 };
219 return;
220 }
221 // --- Cell1 IN THE BOUNDARY, Cell2 IN THE FLUID -----
222
223 if (InCell1 != 0 && InCell2 == 0)
224 {
225     switch(InCell1)
226     {
227         // INFLOW
228         case 1:
229             // Dirichlet on temperature
230             T1 = Cell1.temperature();
231
232             // Extrapolate pressure from old value
233             P1 = Cell2.oldPressure();
234
235             // Compute density
236             rho1 = Gamma*Ma*Ma*P1/T1;
237
238             // Dirichlet on momentum
239             V1 = (Cell1.density()/rho1)*Cell1.velocity();
240
241             // Dirichlet on partial mass
242             if (ScalarEqNb == 1)
243                 Y1 = Cell1.average(Dimension+3)/Cell1.
244 density();
245
246             // Compute energies
247             e1 = P1/((Gamma-1.)*rho1) + 0.5*N2(V1);
248
249             // Correct C1
250             C1.setAverage(1, rho1);
251             for (i=1; i<=Dimension; i++)
252                 C1.setAverage(i+1, rho1*V1.value(i));
253             C1.setAverage(Dimension+2, rho1*e1);
254
255             if (ScalarEqNb == 1)
256                 C1.setAverage(Dimension+3, rho1*Y1);
257             break;
258
259             // OUTFLOW : Get old value of the neighbour
260             case 2:

```

```

261         C1.setAverage(Cell2.oldAverage());
262         break;
263
264     // FREE-SLIP WALL : Neuman on all quantities
265     case 3:
266
267         C1 = Cell2;
268         break;
269
270     // ADIABATIC WALL
271     case 4:
272
273         // Dirichlet on velocity
274         V1 = Cell1.velocity();
275
276         // Neuman on temperature
277         T1 = Cell2.temperature();
278
279         // Neuman on pressure
280         P1 = Cell2.pressure();
281
282         // Extrapolate pressure
283         //P1 = 2*Cell2.pressure()-Cell3.pressure();
284
285         // Compute density
286         rho1 = Gamma*Ma*Ma*P1/T1;
287
288         // Compute energy
289         e1 = P1/((Gamma-1.)*rho1) + 0.5*N2(V1);
290
291         // Correct C1
292         C1.setAverage(1, rho1);
293         for (i=1; i<=Dimension; i++)
294             C1.setAverage(i+1, rho1*V1.value(i));
295         C1.setAverage(Dimension+2, rho1*e1);
296
297         // Neuman on partial mass
298         if (ScalarEqNb == 1)
299             C1.setAverage(Dimension+3, Cell2.average(
Dimension+3));
300
301         break;
302
303     // ISOTHERMAL WALL
304     case 5:
305
306         // Dirichlet on velocity
307         V1 = Cell1.velocity();
308
309         // Dirichlet on temperature
310         T1 = Cell1.temperature();
311
312         // Neuman on pressure
313         P1 = Cell2.pressure();
314
315         // Extrapolate pressure
316         //P1 = 2*Cell2.pressure()-Cell3.pressure();
317
318         // Compute density
319         rho1 = Gamma*Ma*Ma*P1/T1;
320
321         // Compute energies
322         e1 = P1/((Gamma-1.)*rho1) + 0.5*N2(V1);
323
324         // Correct C1
325         C1.setAverage(1, rho1);
326         for (i=1; i<=Dimension; i++)
327             C1.setAverage(i+1, rho1*V1.value(i));
328         C1.setAverage(Dimension+2, rho1*e1);
329
330         // Neuman on partial mass
331         if (ScalarEqNb == 1)
332             C1.setAverage(Dimension+3, Cell2.average(
Dimension+3));
333
334         break;
335     };
336     return;
337 }
338 // --- Cell13 IN THE BOUNDARY, Cell12 IN THE FLUID -----
339
340 if (InCell13 !=0 && InCell12 == 0)
341 {
342     switch(InCell13)
343     {
344         // INFLOW
345         case 1:

```

```

346         // Dirichlet on temperature
347         T3 = Cell13.temperature();
348         T4 = Cell14.temperature();
349
350         // Extrapolate pressure from old value
351         P3 = Cell12.oldPressure();
352         P4 = P3;
353         //P4 = 2*P3 - Cell12.pressure();
354
355         // Compute densities
356         rho3 = Gamma*Ma*Ma*P3/T3;
357         rho4 = Gamma*Ma*Ma*P4/T4;
358
359         // Dirichlet on momentum
360         V3 = (Cell13.density()/rho3)*Cell13.velocity();
361         V4 = (Cell14.density()/rho4)*Cell14.velocity();
362
363         // Dirichlet on partial mass
364         if (ScalarEqNb == 1)
365         {
366             Y3 = Cell13.average(Dimension+3)/Cell13.
density();
367             Y4 = Cell14.average(Dimension+3)/Cell14.
density();
368         }
369
370         // Compute energies
371         e3 = P3/((Gamma-1.)*rho3) + 0.5*N2(V3);
372         e4 = P4/((Gamma-1.)*rho4) + 0.5*N2(V4);
373
374         // Correct C1 and C2
375         C3.setAverage(1, rho3);
376         C4.setAverage(1, rho4);
377         for (i=1; i<=Dimension; i++)
378         {
379             C3.setAverage(i+1, rho3*V3.value(i));
380             C4.setAverage(i+1, rho4*V4.value(i));
381         }
382         C3.setAverage(Dimension+2, rho3*e3);
383         C4.setAverage(Dimension+2, rho4*e4);
384
385         if (ScalarEqNb == 1)
386         {
387             C3.setAverage(Dimension+3, rho3*Y3);
388             C4.setAverage(Dimension+3, rho4*Y4);
389         }
390         break;
391
392         // OUTFLOW
393         case 2:
394
395             C3.setAverage(Cell12.oldAverage());
396             C4.setAverage(Cell12.oldAverage());
397             //C4.setAverage(2*C3.average()-Cell12.average());
398
399             // Also change the values in the boundary
400             Cell13.setAverage(C3.average());
401             Cell14.setAverage(C4.average());
402             break;
403
404         // FREE-SLIP WALL : Neuman on all quantities
405         case 3:
406
407             C3 = Cell12;
408             C4 = Cell11;
409             break;
410
411         // ADIABATIC WALL
412         case 4:
413
414             // Dirichlet on velocity
415             V3 = Cell13.velocity();
416             V4 = Cell14.velocity();
417
418             // Neuman on temperature
419             T3 = Cell12.temperature();
420             T4 = Cell11.temperature();
421
422             // Neuman on pressure
423             P3 = Cell12.pressure();
424             P4 = Cell11.pressure();
425
426             // Extrapolate pressure
427             //P3 = 2*Cell12.pressure()-Cell11.pressure();
428             //P4 = P3;
429
430             // Compute densities

```



```

431         rho3 = Gamma*Ma*Ma*P3/T3;
432         rho4 = Gamma*Ma*Ma*P4/T4;
433
434         // Compute energies
435         e3 = P3/((Gamma-1.)*rho3) + 0.5*N2(V3);
436         e4 = P4/((Gamma-1.)*rho4) + 0.5*N2(V4);
437
438         // Correct C3 and C4
439         C3.setAverage(1, rho3);
440         C4.setAverage(1, rho4);
441         for (i=1; i<=Dimension; i++)
442         {
443             C3.setAverage(i+1, rho3*V3.value(i));
444             C4.setAverage(i+1, rho4*V4.value(i));
445         }
446         C3.setAverage(Dimension+2, rho3*e3);
447         C4.setAverage(Dimension+2, rho4*e4);
448
449         // Neuman on partial mass
450         if (ScalarEqNb == 1)
451         {
452             C3.setAverage(Dimension+3, Cell2.average(
Dimension+3));
453             C4.setAverage(Dimension+3, Cell1.average(
Dimension+3));
454         }
455
456         break;
457
458         // ISOTHERMAL WALL
459         case 5:
460
461             // Dirichlet on velocity
462             V3 = Cell3.velocity();
463             V4 = Cell4.velocity();
464
465             // Dirichlet on temperature
466             T3 = Cell3.temperature();
467             T4 = Cell4.temperature();
468
469             // Neuman on pressure
470             P3 = Cell2.pressure();
471             P4 = Cell1.pressure();
472
473             // Extrapolate pressure
474             //P3 = 2*Cell2.pressure()-Cell1.pressure();
475             //P4 = P3;
476
477             // Compute densities
478             rho3 = Gamma*Ma*Ma*P3/T3;
479             rho4 = Gamma*Ma*Ma*P4/T4;
480
481             // Compute energies
482             e3 = P3/((Gamma-1.)*rho3) + 0.5*N2(V3);
483             e4 = P4/((Gamma-1.)*rho4) + 0.5*N2(V4);
484
485             // Correct C3 and C4
486             C3.setAverage(1, rho3);
487             C4.setAverage(1, rho4);
488             for (i=1; i<=Dimension; i++)
489             {
490                 C3.setAverage(i+1, rho3*V3.value(i));
491                 C4.setAverage(i+1, rho4*V4.value(i));
492             }
493             C3.setAverage(Dimension+2, rho3*e3);
494             C4.setAverage(Dimension+2, rho4*e4);
495
496             // Neuman on partial mass
497             if (ScalarEqNb == 1)
498             {
499                 C3.setAverage(Dimension+3, Cell2.average(
Dimension+3));
500                 C4.setAverage(Dimension+3, Cell1.average(
Dimension+3));
501             }
502
503             break;
504         };
505         return;
506     }
507
508     // --- Cell14 IN THE BOUNDARY, Cell13 IN THE FLUID -----
509
510     if (InCell14 != 0 && InCell13 == 0)
511     {
512         switch(InCell14)
513         {

```

```

514         // INFLOW
515         case 1:
516             // Dirichlet on temperature
517             T4 = Cell4.temperature();
518
519             // Extrapolate pressure from old value
520             P4 = Cell3.oldPressure();
521
522             // Compute density
523             rho4 = Gamma*Ma*Ma*P4/T4;
524
525             // Dirichlet on momentum
526             V4 = (Cell4.density()/rho4)*Cell4.velocity();
527
528             // Dirichlet on partial mass
529             if (ScalarEqNb == 1)
530                 Y4 = Cell4.average(Dimension+3)/Cell4.
density();
531
532             // Compute energies
533             e4 = P4/((Gamma-1.)*rho4) + 0.5*N2(V4);
534
535             // Correct C4
536             C4.setAverage(1, rho4);
537             for (i=1; i<=Dimension; i++)
538                 C4.setAverage(i+1, rho4*V4.value(i));
539             C4.setAverage(Dimension+2, rho4*e4);
540
541             if (ScalarEqNb == 1)
542                 C4.setAverage(Dimension+3, rho4*Y4);
543             break;
544
545         // OUTFLOW : Use old cell-average values of the neighbour
546         case 2:
547
548             C4.setAverage(Cell3.oldAverage());
549             break;
550
551         // FREE-SLIP WALL : Neuman on all quantities
552         case 3:
553
554             C4 = Cell3;
555             break;
556
557         // ADIABATIC WALL
558         case 4:
559
560             // Dirichlet on velocity
561             V4 = Cell4.velocity();
562
563             // Neuman on temperature
564             T4 = Cell3.temperature();
565
566             // Neuman on pressure
567             P4 = Cell3.pressure();
568
569             // Extrapolate pressure
570             //P4 = 2*Cell3.pressure()-Cell2.pressure();
571
572             // Compute density
573             rho4 = Gamma*Ma*Ma*P4/T4;
574
575             // Compute energy
576             e4 = P4/((Gamma-1.)*rho4) + 0.5*N2(V4);
577
578             // Correct C4
579             C4.setAverage(1, rho4);
580             for (i=1; i<=Dimension; i++)
581                 C4.setAverage(i+1, rho4*V4.value(i));
582             C4.setAverage(Dimension+2, rho4*e4);
583
584             // Neuman on partial mass
585             if (ScalarEqNb == 1)
586                 C4.setAverage(Dimension+3, Cell3.average(
Dimension+3));
587
588             break;
589
590         // ISOTHERMAL WALL
591         case 5:
592
593             // Dirichlet on velocity
594             V4 = Cell4.velocity();
595
596             // Dirichlet on temperature
597             T4 = Cell4.temperature();
598

```

```

599         // Neuman on pressure
600         P4 = Cell13.pressure();
601
602         // Extrapolate pressure
603         //P4 = 2*Cell13.pressure()-Cell12.pressure();
604
605         // Compute density
606         rho4 = Gamma*Ma*Ma*P4/T4;
607
608         // Compute energies
609         e4 = P4/((Gamma-1.)*rho4) + 0.5*N2(V4);
610
611         // Correct C4
612         C4.setAverage(1, rho4);
613         for (i=1; i<=Dimension; i++)
614             C4.setAverage(i+1, rho4*V4.value(i));
615         C4.setAverage(Dimension+2, rho4*e4);
616
617         // Neuman on partial mass
618         if (ScalarEqNb == 1)
619             C4.setAverage(Dimension+3, Cell13.average(
Dimension+3));
620
621         break;
622     };
623     return;
624 }
625
626 }
    
```

Here is the caller graph for this function:



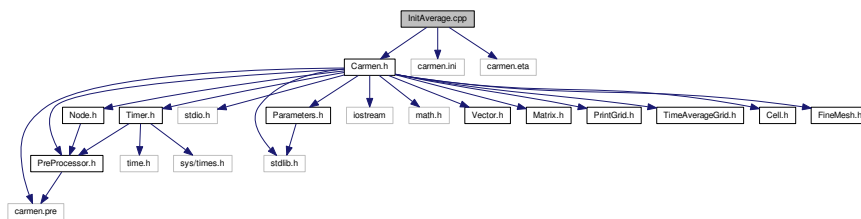
## 6.19 InitAverage.cpp File Reference

Fill the variables vector with the initial condition.

```

#include "Carmen.h"
#include "carmen.ini"
#include "carmen.eta"
    
```

Include dependency graph for InitAverage.cpp:



### Functions

- [Vector InitAverage](#) (real x, real y, real z)  
Returns the initial condition in (x, y, z) form the one defined in carmen.ini.
- [real InitResistivity](#) (real x, real y, real z)  
Returns the initial resistivity condition in (x, y, z) form the one defined in carmen.eta.

## 6.19.1 Detailed Description

Fill the variables vector with the initial condition.

## 6.19.2 Function Documentation

### 6.19.2.1 Vector InitAverage ( real x, real y = 0., real z = 0. )

Returns the initial condition in  $(x, y, z)$  form the one defined in *carmen.ini*.

#### Parameters

|          |                             |
|----------|-----------------------------|
| <i>x</i> | Position x                  |
| <i>y</i> | Position y. Defaults to 0.. |
| <i>z</i> | Position z. Defaults to 0.. |

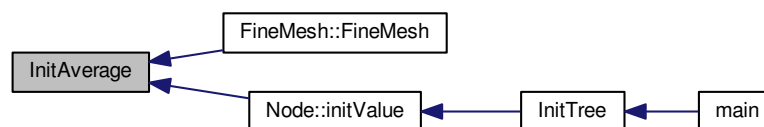
#### Returns

Vector

```

23 {
24     // --- Local variables ---
25
26     Vector Result(QuantityNb);
27     real    *Q;
28     Q = new real [QuantityNb+1];
29     int     n;
30
31     // --- Init Q ---
32
33     for (n = 1; n <= QuantityNb; n++)
34         Q[n]=0.;
35
36     // --- Use definition of initial Q contained in file 'initial' ---
37
38     #include "carmen.ini"
39
40     // --- Fill vector Result and return it ---
41
42     for (n = 1; n <= QuantityNb; n++)
43         Result.setValue(n, Q[n]);
44
45     delete[] Q;
46
47     return Result;
48 }
```

Here is the caller graph for this function:



### 6.19.2.2 real InitResistivity ( real x, real y = 0., real z = 0. )

Returns the initial resistivity condition in  $(x, y, z)$  form the one defined in *carmen.eta*.

## Parameters

|   |                             |
|---|-----------------------------|
| x | Position x                  |
| y | Position y. Defaults to 0.. |
| z | Position z. Defaults to 0.. |

## Returns

double

```

51 {
52     // --- Local variables ---
53
54     real Result=0.;
55     real Res = 0.;
56
57     #include "carmen.eta"
58
59     Result = Res;
60
61     return Result;
62 }

```

## 6.20 InitResistivity.cpp File Reference

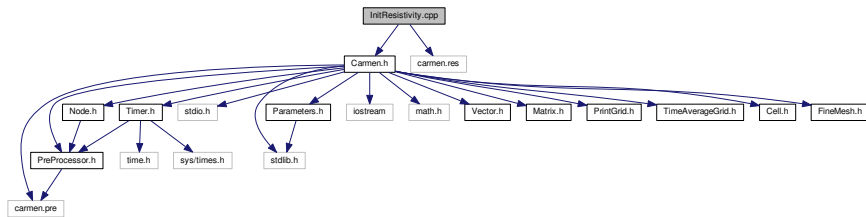
Fill the magnetic resistivity parameter (x,y,z)

```

#include "Carmen.h"
#include "carmen.res"

```

Include dependency graph for InitResistivity.cpp:



## Functions

- `real InitResistivity (real x, real y, real z)`

Returns the initial resistivity condition in (x, y, z) form the one defined in *carmen.eta*.

## 6.20.1 Detailed Description

Fill the magnetic resistivity parameter (x,y,z)

## 6.20.2 Function Documentation

6.20.2.1 `real InitResistivity ( real x, real y = 0 . , real z = 0 . )`

Returns the initial resistivity condition in (x, y, z) form the one defined in *carmen.eta*.

## Parameters

|   |                             |
|---|-----------------------------|
| x | Position x                  |
| y | Position y. Defaults to 0.. |
| z | Position z. Defaults to 0.. |

## Returns

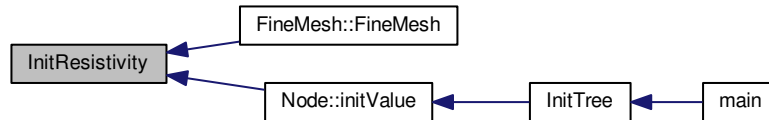
double

```

24 {
25     // --- Local variables ---
26
27     real Result=0.;
28     real Res = 0.;
29     #include "carmen.res"
30
31     Result = Res;
32
33     return Result;
34 }

```

Here is the caller graph for this function:

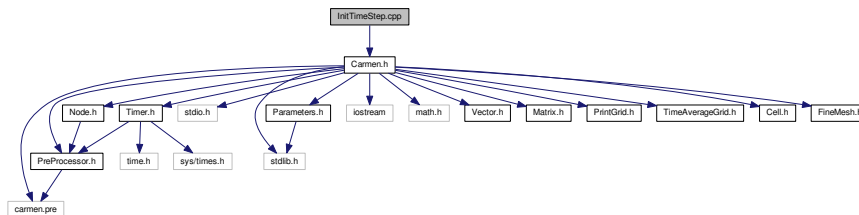


## 6.21 InitTimeStep.cpp File Reference

Compute the timestep of the very first iteration.

```
#include "Carmen.h"
```

Include dependency graph for InitTimeStep.cpp:



## Functions

- void [InitTimeStep](#) ()  
*Initns time step and all the parameters which depend on it.*

### 6.21.1 Detailed Description

Compute the timestep of the very first iteration.

## 6.21.2 Function Documentation

### 6.21.2.1 void InitTimeStep ( )

Initializes time step and all the parameters which depend on it.

#### Returns

void

```

23 {
24
25     // --- Init TimeStep -----
26
27     if (TimeStep == 0)
28     {
29         if (Resistivity) TimeStep = CFL*SpaceStep/(
Eigenvalue + eta);
30         else TimeStep = CFL*SpaceStep/Eigenvalue;
31     }
32
33     // --- Compute number of iterations -----
34
35     if (PhysicalTime != 0. && IterationNb == 0)
36         IterationNb = (int)(ceil(PhysicalTime/TimeStep));
37
38     // --- Compute Refresh -----
39
40     if (Refresh == 0)
41         Refresh = (int)(ceil(IterationNb/(RefreshNb*1.)));
42
43     // --- Compute PrintEvery -----
44
45     if ((PrintEvery == 0)&&(ImageNb != 0))
46         PrintEvery = (int)(ceil(IterationNb/(ImageNb*1.)));
47
48     // --- Compute iterations for print -----
49
50     if (PrintTime1 != 0.)
51         PrintIt1 = (int)(ceil(PrintTime1/TimeStep));
52
53     if (PrintTime2 != 0.)
54         PrintIt2 = (int)(ceil(PrintTime2/TimeStep));
55
56     if (PrintTime3 != 0.)
57         PrintIt3 = (int)(ceil(PrintTime3/TimeStep));
58
59     if (PrintTime4 != 0.)
60         PrintIt4 = (int)(ceil(PrintTime4/TimeStep));
61
62     if (PrintTime5 != 0.)
63         PrintIt5 = (int)(ceil(PrintTime5/TimeStep));
64
65     if (PrintTime6 != 0.)
66         PrintIt6 = (int)(ceil(PrintTime6/TimeStep));
67
68     // --- Compute FV reference time -----
69
70     if (Multiresolution)
71         FVTimeRef = CPUTimeRef(IterationNbRef,
ScaleNbRef);
72
73 }
```

Here is the caller graph for this function:

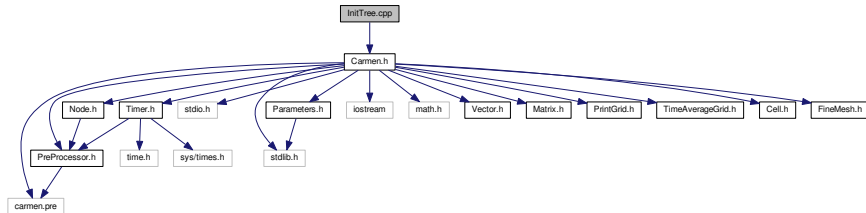


## 6.22 InitTree.cpp File Reference

Init graded tree (only for multiresolution solver)

```
#include "Carmen.h"
```

Include dependency graph for InitTree.cpp:



### Functions

- void [InitTree](#) (Node \*Root)

*Init tree structure from initial condition, starting from the node Root. Only for multiresolution computations.*

#### 6.22.1 Detailed Description

Init graded tree (only for multiresolution solver)

#### 6.22.2 Function Documentation

##### 6.22.2.1 void InitTree ( Node \* Root )

Init tree structure from initial condition, starting from the node *Root*. Only for multiresolution computations.

##### Parameters

|             |      |
|-------------|------|
| <i>Root</i> | Root |
|-------------|------|

##### Returns

void

```

23 {
24     // --- Local variables ---
25
26     int l; // Counter on levels
27
28     // --- Init cell-average value in root and split it ---
29
30     if (Recovery && UseBackup)
31
32         Root->restore();
33
34     else
35     {
36         Root->initValue();
37
38         // --- Add and init nodes in different levels, when necessary ---
39
40         for (l=1; l <= ScaleNb; l++)
41             Root->addLevel();
42     }
43
44     // -- Check if tree is graded ---
45

```



```

46     if (debug) Root->checkGradedTree();
47
48 }

```

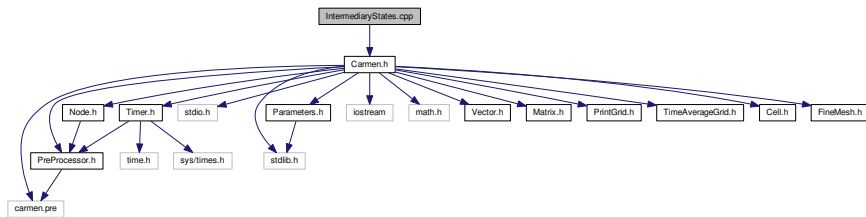
Here is the caller graph for this function:



## 6.23 IntermediaryStates.cpp File Reference

```
#include "Carmen.h"
```

Include dependency graph for IntermediaryStates.cpp:



### Functions

- **Matrix stateUstar** (const [Vector](#) &AvgL, const [Vector](#) &AvgR, const [real](#) prel, const [real](#) prer, [real](#) &slopeLeft, [real](#) &slopeRight, [real](#) &slopeM, [real](#) &slopeLeftStar, [real](#) &slopeRightStar, int AxisNo)

*Returns the intermediary states of HLLD numerical flux for MHD equations.*

### 6.23.1 Function Documentation

**6.23.1.1 Matrix stateUstar** ( const [Vector](#) & AvgL, const [Vector](#) & AvgR, const [real](#) prel, const [real](#) prer, [real](#) & slopeLeft, [real](#) & slopeRight, [real](#) & slopeM, [real](#) & slopeLeftStar, [real](#) & slopeRightStar, int AxisNo )

Returns the intermediary states of HLLD numerical flux for MHD equations.

#### Parameters

|             |                      |
|-------------|----------------------|
| <i>AvgL</i> | Left average vector  |
| <i>AvgR</i> | Right average vector |
| <i>prel</i> | Left pressure        |
| <i>prer</i> | Right pressure       |

|                       |                                        |
|-----------------------|----------------------------------------|
| <i>slopeLeft</i>      | Left slope                             |
| <i>slopeRight</i>     | Right slope                            |
| <i>slopeM</i>         | Slope value computed for HLLD          |
| <i>slopeLeftStar</i>  | Left star slope (intermediary states)  |
| <i>slopeRightStar</i> | Right star slope (intermediary states) |
| <i>AxisNo</i>         | Axis of interest                       |

## Returns

### Matrix

variables U-star

variables U-star-star

```

13 {
14   real rhol=0., rhor=0.;
15   real psil=0., psir=0.;
16   real vxl=0., vxr=0., vyl=0., vyr=0., vzl=0., vzr=0.;
17   real Bxr=0., Bxl=0., Byl=0., Byr=0., Bzl=0., Bzr=0.;
18   real vl=0., vr=0., mf=0.;
19   real Bl=0., Br=0.;
20   real pTl=0., pTr=0.;
21   real vBl=0., vBr=0.;
22   real el=0., er=0.;
23   real rhols=0., rhors=0.;
24   real vxls=0., vxrs=0., vyls=0., vyrs=0., vzls=0., vzrs=0.;
25   real Bxls=0., Bxrs=0., Byls=0., Byrs=0., Bzls=0., Bzrs=0.;
26   real pTs=0.;
27   real vBls=0., vBrs=0.;
28   real Els=0., Ers=0.;
29   real rholss=0., rhorss=0.;
30   real vxlss=0., vxrss=0., vylss=0., vyrrs=0., vzlss=0., vzrss=0.;
31   real Bxlss=0., Bxrss=0., Bylss=0., Byrrs=0., Bzlss=0., Bzrss=0.;
32   real vBlss=0., vBrrs=0.;
33   real Elss=0., Erss=0.;
34   int Bsign=0;
35   real half=0.5;
36   real epsilon2=1e-16;
37   real auxl=0., auxr=0., Sqrtrhols=0.;
38
39   Matrix U(QuantityNb,4);
40
41   //Variables
42   rhol = AvgL.value(1);
43   vxl = AvgL.value(2)/rhol;
44   vyl = AvgL.value(3)/rhol;
45   vzl = AvgL.value(4)/rhol;
46   el = AvgL.value(5);
47   psil = AvgL.value(6);
48   Bxl = AvgL.value(7);
49   Byl = AvgL.value(8);
50   Bzl = AvgL.value(9);
51
52   rhor = AvgR.value(1);
53   vxr = AvgR.value(2)/rhor;
54   vyr = AvgR.value(3)/rhor;
55   vzr = AvgR.value(4)/rhor;
56   er = AvgR.value(5);
57   psir = AvgR.value(6);
58   Bxr = AvgR.value(7);
59   Byr = AvgR.value(8);
60   Bzr = AvgR.value(9);
61
62   //v_x and v_y
63   vl = AvgL.value(AxisNo+1)/rhol;
64   vr = AvgR.value(AxisNo+1)/rhor;
65
66   // Average B value
67   mf = (AvgL.value(AxisNo+6)+AvgR.value(AxisNo+6))/(double (2.0));
68
69   if(AxisNo==1)
70   {
71     //|B|
72     Bl = sqrt( mf*mf + Byl*Byl + Bzl*Bzl );
73     Br = sqrt( mf*mf + Byr*Byr + Bzr*Bzr );
74     //Inner product v.B
75     vBl = vxl*mf + vyl*Byl + vzl*Bzl;
76     vBr = vxr*mf + vyr*Byr + vzr*Bzr;
77   }else if(AxisNo==2){

```

```

78         //|B|
79         B1 = sqrt( Bx1*Bx1 + mf*mf + Bz1*Bz1 );
80         Br = sqrt( Bxr*Bxr + mf*mf + Bzr*Bzr );
81         //Inner product v.B
82         vB1 = vx1*Bx1 + vyl*mf + vz1*Bz1;
83         vBr = vxr*Bxr + vyr*mf + vzr*Bzr;
84     }else{
85         //|B|
86         B1 = sqrt( Bx1*Bx1 + By1*By1 + mf*mf );
87         Br = sqrt( Bxr*Bxr + Byr*Byr + mf*mf );
88         //Inner product v.B
89         vB1 = vx1*Bx1 + vyl*By1 + vz1*mf;
90         vBr = vxr*Bxr + vyr*Byr + vzr*mf;
91     }
92
93     //Total Pressure
94     pTl = prel + half*B1*B1;
95     pTr = prer + half*Br*Br;
96
97     // S_M - Equation 38
98     slopeM = ((slopeRight - vr)*rhor*vr - (slopeLeft - vl)*rhol*vl - pTr +pTl)/
99             ((slopeRight - vr)*rhor - (slopeLeft - vl)*rhol);
100
101     //Sign function
102     if(mf > 0) Bsign = 1;
103     else      Bsign = -1;
104
105
106     //density - Equation 43
107     rhols = rhol*(slopeLeft-vl)/(slopeLeft-slopeM);
108     rhors = rhor*(slopeRight-vr)/(slopeRight-slopeM);
109
110     if(AxisNo==1){
111         //velocities
112         //Equation 39
113         vxls = slopeM;
114         vxrs = vxls;
115         //B_x
116         Bxls = mf;
117         Bxrs = Bxls;
118
119         auxl= (rhol*(slopeLeft - vl)*(slopeLeft - slopeM)-mf*mf);
120         auxr= (rhor*(slopeRight - vr)*(slopeRight - slopeM)-mf*mf);
121
122         if( fabs(auxl)<epsilon2 ||
123            ( ( ( fabs(slopeM -vl) ) < epsilon2 )  &&
124              ( ( fabs(By1) + fabs(Bz1) ) < epsilon2 )  &&
125              ( ( mf*mf ) > (Gamma*prel) ) ) ) ) {
126             vyls = vyl;
127             vzls = vz1;
128             Byls = By1;
129             Bzls = Bz1;
130         }else{
131             //Equation 44
132             vyls = vyl - mf*By1*(slopeM-vl)/auxl;
133             //Equation 46
134             vzls = vz1 - mf*Bz1*(slopeM-vl)/auxl;
135             //Equation 45
136             Byls = By1*(rhol*(slopeLeft-vl)*(slopeLeft-vl) - mf*mf)/auxl;
137             //Equation 47
138             Bzls = Bz1*(rhol*(slopeLeft-vl)*(slopeLeft-vl) - mf*mf)/auxl;
139         }
140         if( fabs(auxr)<epsilon2 ||
141            ( ( ( fabs(slopeM -vr) ) < epsilon2 )  &&
142              ( ( fabs(Byr) + fabs(Bzr) ) < epsilon2 )  &&
143              ( ( mf*mf ) > (Gamma*prer) ) ) ) ) {
144             vyrs = vyr;
145             vzrs = vzr;
146             Byrs = Byr;
147             Bzrs = Bzr;
148         }else{
149             //Equation 44
150             vyrs = vyr - mf*Byr*(slopeM-vr)/auxr;
151             //Equation 46
152             vzrs = vzr - mf*Bzr*(slopeM-vr)/auxr;
153             //Equation 45;
154             Byrs = Byr*(rhor*(slopeRight-vr)*(slopeRight-vr) - mf*mf)/auxr;
155             //Equation 47
156             Bzrs = Bzr*(rhor*(slopeRight-vr)*(slopeRight-vr) - mf*mf)/auxr;
157         }
158     }
159
160     }else if(AxisNo==2){
161         //velocities
162         //Equation 39
163         vyls = slopeM;
164         vyrs = vyls;
165         //B_y

```

```

166         Byls = mf;
167         Byrs = Byls;
168
169         auxl= (rho1*(slopeLeft - vl)*(slopeLeft - slopeM)-mf*mf);
170         auxr= (rho1*(slopeRight - vr)*(slopeRight - slopeM)-mf*mf);
171
172         if( fabs(auxl)<epsilon2 ||
173             ( ( ( fabs(slopeM -vl) ) < epsilon2 )  &&
174               ( ( fabs(Bxl) + fabs(Bzl) ) < epsilon2 )  &&
175               ( ( mf*mf ) > (Gamma*prel) ) ) ) ) {
176             vxls = vxl;
177             vzls = vzl;
178             Bxls = Bxl;
179             Bzls = Bzl;
180         }else{
181             //Equation 44
182             vxls = vxl - mf*Bxl*(slopeM-vl)/auxl;
183             //Equation 46
184             vzls = vzl - mf*Bzl*(slopeM-vl)/auxl;
185             //Equation 45
186             Bxls = Bxl*(rho1*(slopeLeft-vl)*(slopeLeft-vl) - mf*mf)/auxl;
187             //Equation 47
188             Bzls = Bzl*(rho1*(slopeLeft-vl)*(slopeLeft-vl) - mf*mf)/auxl;
189         }
190
191         if( fabs(auxr)<epsilon2 ||
192             ( ( ( fabs(slopeM -vr) ) < epsilon2 )  &&
193               ( ( fabs(Bxr) + fabs(Bzr) ) < epsilon2 )  &&
194               ( ( mf*mf ) > (Gamma*prer) ) ) ) ) {
195             vxrs = vxr;
196             vzrs = vzr;
197             Bxrs = Bxr;
198             Bzrs = Bzr;
199         }else{
200             //Equation 44
201             vxrs = vxr - mf*Bxr*(slopeM-vr)/auxr;
202             //Equation 46
203             vzrs = vzr - mf*Bzr*(slopeM-vr)/auxr;
204             //Equation 45
205             Bxrs = Bxr*(rho1*(slopeRight-vr)*(slopeRight-vr) - mf*mf)/auxr;
206             //Equation 47
207             Bzrs = Bzr*(rho1*(slopeRight-vr)*(slopeRight-vr) - mf*mf)/auxr;
208         }
209     }else{
210         //velocities
211         //Equation 39
212         vzls = slopeM;
213         vzrs = vzls;
214         //B_z
215         Bzls = mf;
216         Bzrs = Bzls;
217
218         auxl= (rho1*(slopeLeft - vl)*(slopeLeft - slopeM)-mf*mf);
219         auxr= (rho1*(slopeRight - vr)*(slopeRight - slopeM)-mf*mf);
220
221         if( fabs(auxl)<epsilon2 ||
222             ( ( ( fabs(slopeM -vl) ) < epsilon2 )  &&
223               ( ( fabs(Bxl) + fabs(Byl) ) < epsilon2 )  &&
224               ( ( mf*mf ) > (Gamma*prel) ) ) ) ) {
225             vxls = vxl;
226             vyls = vyl;
227             Bxls = Bxl;
228             Byls = Byl;
229         }else{
230             //Equation 44
231             vxls = vxl - mf*Bxl*(slopeM-vl)/auxl;
232             //Equation 46
233             vyls = vyl - mf*Byl*(slopeM-vl)/auxl;
234             //Equation 45
235             Bxls = Bxl*(rho1*(slopeLeft-vl)*(slopeLeft-vl) - mf*mf)/auxl;
236             //Equation 47
237             Byls = Byl*(rho1*(slopeLeft-vl)*(slopeLeft-vl) - mf*mf)/auxl;
238         }
239
240         if( fabs(auxr)<epsilon2 ||
241             ( ( ( fabs(slopeM -vr) ) < epsilon2 )  &&
242               ( ( fabs(Bxr) + fabs(Byr) ) < epsilon2 )  &&
243               ( ( mf*mf ) > (Gamma*prer) ) ) ) ) {
244             vxrs = vxr;
245             vyrs = vyr;
246             Bxrs = Bxr;
247             Byrs = Byr;
248         }else{
249             //Equation 44
250             vxrs = vxr - mf*Bxr*(slopeM-vr)/auxr;
251             //Equation 46
252             vyrs = vyr - mf*Byr*(slopeM-vr)/auxr;

```

```

253         //Equation 45
254         Bxrs = Bxr*(rhor*(slopeRight-vr)*(slopeRight-vr) - mf*mf)/auxr;
255         //Equation 47
256         Byrs = Byr*(rhor*(slopeRight-vr)*(slopeRight-vr) - mf*mf)/auxr;
257     }
258 }
259
260 //total pressure - Equation 41
261 pTs = pTl + rhol*(slopeLeft - vl)*(slopeM - vl);
262
263 //inner product vs*Bs
264 vBls = vxls*Bxls + vylys*Byls + vzls*Bzls;
265 vBrs = vxrs*Bxrs + vyrs*Byrs + vzrs*Bzrs;
266
267 //energy - Equation 48
268 Els = ((slopeLeft - vl)*el - pTl*vl+pTs*slopeM+mf*(vBl - vBls))/(slopeLeft - slopeM);
269 Ers = ((slopeRight-vr)*er - pTr*vr+pTs*slopeM+mf*(vBr - vBrs))/(slopeRight - slopeM);
270
271 //U-star variables
272 //Left
273 U.setValue(1,1,rhols);
274 U.setValue(2,1,rhols*vxls);
275 U.setValue(3,1,rhols*vylys);
276 U.setValue(4,1,rhols*vzls);
277 U.setValue(5,1,Els);
278 U.setValue(6,1,psil);
279 U.setValue(7,1,Bxls);
280 U.setValue(8,1,Byls);
281 U.setValue(9,1,Bzls);
282 //Right
283 U.setValue(1,2,rhors);
284 U.setValue(2,2,rhors*vxrs);
285 U.setValue(3,2,rhors*vyrs);
286 U.setValue(4,2,rhors*vzrs);
287 U.setValue(5,2,Ers);
288 U.setValue(6,2,psir);
289 U.setValue(7,2,Bxrs);
290 U.setValue(8,2,Byrs);
291 U.setValue(9,2,Bzrs);
292
293
294
295
296
297 Sqrtrhols = (sqrt(rhols)+sqrt(rhors));
298
299 if((mf*mf/min((Bl*Bl),(Br*Br))) < epsilon2){
300 for(int k=1;k<=QuantityNb;k++){
301     U.setValue(k,3, U.value(k,1));
302     U.setValue(k,4, U.value(k,2));
303 }
304 }else{
305     //density - Equation 49
306     rholss = rhols;
307     rhorss = rhors;
308
309     if(AxisNo==1){
310         //velocities
311         //Equation 39
312         vxlss = slopeM;
313         vxrss = slopeM;
314         //Equation 59
315         vylss = (sqrt(rhols)*vylys + sqrt(rhors)*vyrs + (Byrs - Byls)*Bsign)/Sqrtrhols;
316         vyrrs = vylss;
317         //Equation 60
318         vzlss = (sqrt(rhols)*vzls + sqrt(rhors)*vzrs + (Bzrs - Bzls)*Bsign)/Sqrtrhols;
319         vzrrs = vzlss;
320
321         //Magnetic Field components
322         //B_x
323         Bxlls = mf;
324         Bxrss = mf;
325         //Equation 61
326         Bylls = (sqrt(rhols)*Byrs + sqrt(rhors)*Byls + sqrt(rhols*rhors)*(vyrs - vyls)*Bsign)/
Sqrtrhols;
327         Byrrs = Bylls;
328         //Equation 62
329         Bzlls = (sqrt(rhols)*Bzrs + sqrt(rhors)*Bzls + sqrt(rhols*rhors)*(vzrs - vzls)*Bsign)/
Sqrtrhols;
330         Bzrrs = Bzlls;
331     }
332     else if(AxisNo==2){
333         //velocities
334         //Equation 59
335         vxlss = (sqrt(rhols)*vxls + sqrt(rhors)*vxrs + (Bxrs - Bxls)*Bsign)/Sqrtrhols;
336         vxrrs = vxlss;
337         //Equation 39
338         vylss = slopeM;

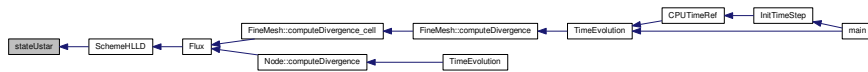
```

```

339         vyrss = slopeM;
340         //Equation 60
341         vzlss = (sqrt(rhols)*vzls + sqrt(rhors)*vzrs + (Bzrs - Bzls)*Bsign)/Sqrtrhols;
342         vzrss = vzlss;
343
344         //Magnetic Field components
345         //Equation 61
346         Bxlss = (sqrt(rhols)*Bxrs + sqrt(rhors)*Bxls + sqrt(rhols*rhors)*(vxrs - vxls)*Bsign)/
Sqrtrhols;
347         Bxrss = Bxlss;
348         //B_y
349         Bylss = mf;
350         Byrss = mf;
351         //Equation 62
352         Bzlss = (sqrt(rhols)*Bzrs + sqrt(rhors)*Bzls + sqrt(rhols*rhors)*(vzrs - vzls)*Bsign)/
Sqrtrhols;
353         Bzrss = Bzlss;
354
355     }else{
356         //velocities
357         //Equation 59
358         vxlss = (sqrt(rhols)*vxls + sqrt(rhors)*vxrs + (Bxrs - Bxls)*Bsign)/Sqrtrhols;
359         vxrss = vxlss;
360         //Equation 60
361         vylss = (sqrt(rhols)*vyls + sqrt(rhors)*vyrs + (Byrs - Byls)*Bsign)/Sqrtrhols;
362         vyrss = vylss;
363         //Equation 39
364         vzlss = slopeM;
365         vzrss = slopeM;
366
367         //Magnetic Field components
368         //Equation 61
369         Bxlss = (sqrt(rhols)*Bxrs + sqrt(rhors)*Bxls + sqrt(rhols*rhors)*(vxrs - vxls)*Bsign)/
Sqrtrhols;
370         Bxrss = Bxlss;
371         //Equation 62
372         Bylss = (sqrt(rhols)*Byrs + sqrt(rhors)*Byls + sqrt(rhols*rhors)*(vyrs - vyls)*Bsign)/
Sqrtrhols;
373         Byrss = Bylss;
374         //B_y
375         Bzlss = mf;
376         Bzrss = mf;
377
378     }
379
380
381     //inner product vss*Bss
382     vBlss = vxlss*Bxlss + vylss*Bylss + vzlss*Bzlss;
383     vBrss = vxrss*Bxrss + vyrss*Byrss + vzrss*Bzrss;
384
385     //Energy - Equation 63
386     Elss = Els - sqrt(rhols)*(vBlss - vBlss)*Bsign;
387     Erss = Ers + sqrt(rhors)*(vBrss - vBrss)*Bsign;
388
389
390
391
392     //U-star-star variables
393     //Left
394     U.setValue(1,3,rholss);
395     U.setValue(2,3,rholss*vxlss);
396     U.setValue(3,3,rholss*vylss);
397     U.setValue(4,3,rholss*vzlss);
398     U.setValue(5,3,Elss);
399     U.setValue(6,3,psil);
400     U.setValue(7,3,Bxlss);
401     U.setValue(8,3,Bylss);
402     U.setValue(9,3,Bzlss);
403     //Right
404     U.setValue(1,4,rhorss);
405     U.setValue(2,4,rhorss*vxrss);
406     U.setValue(3,4,rhorss*vyrss);
407     U.setValue(4,4,rhorss*vzrss);
408     U.setValue(5,4,Erss);
409     U.setValue(6,4,psir);
410     U.setValue(7,4,Bxrss);
411     U.setValue(8,4,Byrss);
412     U.setValue(9,4,Bzrss);
413     }
414     //Equation 61 - S-star left and S-star right
415     slopeLeftStar = slopeM - Abs(mf)/sqrt(rhols);
416     slopeRightStar = slopeM + Abs(mf)/sqrt(rhors);
417
418     return U;
419 }

```

Here is the caller graph for this function:

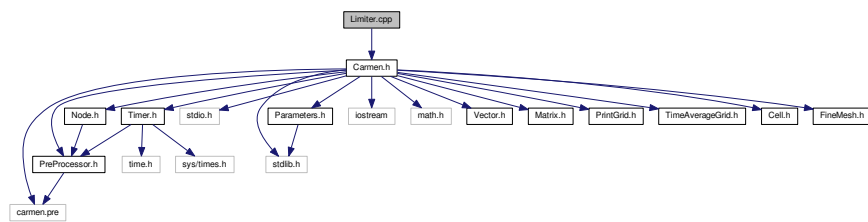


## 6.24 Limiter.cpp File Reference

Limiter functions for the conservative variables.

```
#include "Carmen.h"
```

Include dependency graph for Limiter.cpp:



### Functions

- [real Limiter](#) (const [real](#) r)  
Returns the value of slope limiter from a real value x.
- [Vector Limiter](#) (const [Vector](#) u, const [Vector](#) v)  
Returns the value of the slope limiter between the slopes u and v.

#### 6.24.1 Detailed Description

Limiter functions for the conservative variables.

#### 6.24.2 Function Documentation

##### 6.24.2.1 [real Limiter](#) ( const [real](#) x )

Returns the value of slope limiter from a real value x.

##### Parameters

|   |     |
|---|-----|
| X | ... |
|---|-----|

##### Returns

double

```

23 {
24   real Result = 0.;
25
26   switch (LimiterNo)
27   {
28     case 1: // Min-Mod
  
```

```

29     Result = Max(0., Min(1., r));
30     break;
31
32     case 2: // Van Albada
33         Result = (r<=0.)? 0.: (r*r+r)/(r*r+1.);
34         break;
35
36     case 3: // Van Leer
37         Result = (r<=0.) ? 0.: (r+Abs(r))/(1.+Abs(r));
38         break;
39
40     case 4: // Superbee
41         Result = (r<=0.) ? 0.: Max(0., Max(Min(2.*r,1.), Min(r,2.)));
42         break;
43     case 5: // Monotonized Central
44         Result = max(0.0, min(min(2*r, 0.5*(1+r)), 2.0));
45         break;
46
47 };
48
49     return Result;
50 }

```

### 6.24.2.2 Vector Limiter ( const Vector *u*, const Vector *v* )

Returns the value of the slope limiter between the slopes *u* and *v*.

#### Parameters

|          |        |
|----------|--------|
| <i>u</i> | Vector |
| <i>v</i> | Vector |

#### Returns

Vector

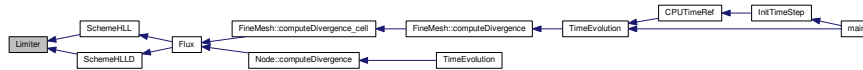
```

57 {
58     // Min Mod limiter
59
60     int LimiterNo = 3;
61
62     Vector Result(u.dimension());
63     int i;
64     real x, y; // slopes
65
66     for (i=1; i<=u.dimension(); i++)
67     {
68         x = u.value(i);
69         y = v.value(i);
70
71         switch(LimiterNo)
72         {
73             // MIN-MOD
74             case 1:
75                 if (x == y)
76                     Result.setValue(i, 0.);
77                 else
78                     Result.setValue(i, Min(1., fabs(x)/fabs(x-y)));
79                 break;
80
81             // VAN LEER
82             case 3:
83             default:
84                 if ((fabs(x) + fabs(y)) == 0.)
85                     Result.setValue(i, 0.);
86                 else
87                     Result.setValue(i, fabs(x) / (fabs(x)+fabs(y)));
88                 break;
89         };
90     }
91
92     return Result;
93 }

```



Here is the caller graph for this function:

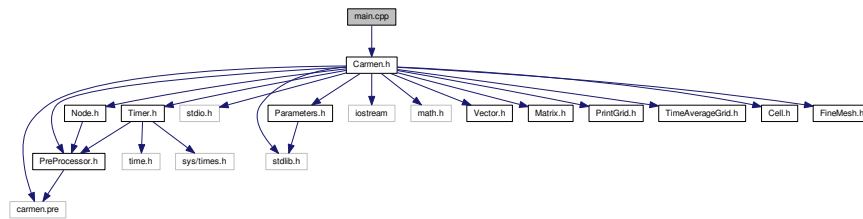


## 6.25 main.cpp File Reference

Main function.

```
#include "Carmen.h"
```

Include dependency graph for main.cpp:



### Functions

- int main (int argc, char \*argv[])

#### 6.25.1 Detailed Description

Main function.

#### 6.25.2 Function Documentation

##### 6.25.2.1 int main ( int argc, char \* argv[] )

```

107 {
108     // --- Init Cluster variable -----
109
110     // carmen 0 => local execution i.e. show time on screen
111     // carmen 1 => cluster execution i.e. refresh Performance.dat (default)
112
113     #if defined PARMPI
114         // --- MPI Runtime system initialization
115         // size - total number of processors
116         // rnak - current CPU
117         MPI_Init(&argc,&argv);
118         MPI_Comm_size(MPI_COMM_WORLD, &size);
119         MPI_Comm_rank(MPI_COMM_WORLD, &rank);
120     #else
121         size=1;
122         rank=0;
123     #endif
124
125     if (argc == 2)
126         Cluster = atoi(argv[1]);
127
128     // --- Print messages on screen-----
129
130     cout << "carmen: begin execution.\n\n";
131     printf("Carmen %4.2f \n",CarmenVersion);

```

```

132     cout << "Copyright (C) 2000-2005 by Olivier Roussel.\n";
133     cout << "All rights reserved.\n\n";
134
135 #if defined PARMPI
136     //Synchronize all parallel branches
137     MPI_Barrier(MPI_COMM_WORLD);
138 #endif
139
140     CPUTime.start();
141
142     // --- Create first node of the tree structure -----
143
144     Node *Mesh=0;
145     FineMesh *FMesh=0;
146
147
148     // --- Init global values (See Parameters.h and Parameters.cpp) -----
149
150     cout << "carmen: init computation ...\n";
151     InitParameters();
152
153     // --- Debug output information for parallel execution -----
154
155 #if defined PARMPI
156     if (Multiresolution)
157     {
158         printf("\nParallel Multiresolution solver not implemented yet!\n");
159         exit(0);
160     }
161
162     printf("My Rank=%d\n",rank);
163
164     // --- Each CPU print his coordinates in the virtual processor cart -----
165     printf("Cart_i = %d;    Cart_j = %d;    Cart_k = %d;\n",coords[0],
166         coords[1],coords[2]);
167
168     // --- Each CPU print his computation domain
169     printf("Xmin = %lf;    XMax = %lf;\n",XMin[1],XMax[1]);
170     printf("Ymin = %lf;    YMax = %lf;\n",XMin[2],XMax[2]);
171     printf("Zmin = %lf;    ZMax = %lf;\n",XMin[3],XMax[3]);
172
173     // --- And the local scale number -----
174     printf("ScaleNb = %d\n",ScaleNb);
175 #endif
176
177     // --- Allocate -----
178
179     if (Multiresolution)
180         Mesh = new Node;
181     else
182         FMesh = new FineMesh;
183
184     // --- Init tree structure -----
185
186     if (Multiresolution)
187     {
188         InitTree(Mesh);
189         RefreshTree(Mesh);
190     }
191
192     // --- Compute initial integral values and init time step -----
193
194     if (Multiresolution)
195         Mesh->computeIntegral();
196     else
197         FMesh->computeIntegral();
198
199     // -- Write integral values --
200
201     // -- Compute initial time step --
202     InitTimeStep();
203
204     if (rank==0) PrintIntegral("Integral.dat");
205
206     // --- Save initial values into files -----
207
208     if (PrintEvery == 0)
209     {
210         if (Multiresolution)
211             View(Mesh, "Tree_0.dat", "Mesh_0.dat", "Average_0.vtk");
212         else
213             View(FMesh,"Average_0.vtk");
214     }
215
216     // --- When PrintEvery != 0, save initial values into specific name format ---
217

```

```

218     if (PrintEvery != 0)
219     {
220         if (Multiresolution)
221             ViewEvery(Mesh, 0);
222         else
223             ViewEvery(FMesh, 0);
224     }
225
226     // --- Parallel execution only -----
227     // --- Save to disk DX header for ouput files -----
228     // --- This file is needed for the external postprocessing (merging files from the different
    processors)
229
230 #if defined PARMPI
231
232     real    tempXMin[4];
233     real    tempXMax[4];
234
235     // --- Save original task parameters for the parallel execution
236     int tempScaleNb=ScaleNb;
237
238     // --- Simulate sequential running
239     ScaleNb=AllTaskScaleNb;
240
241     for (int i=0;i<4;i++)
242     {
243         tempXMin[i]=XMin[i];
244         tempXMax[i]=XMax[i];
245         // --- Simulate sequential running
246         XMin[i]=AllXMin[i];
247         XMax[i]=AllXMax[i];
248     }
249
250     // --- Write header with parameters, as we have run sequential code
251     if (rank==0) FMesh->writeHeader("header.txt");
252
253     // Restore variables
254     for (int i=0;i<4;i++)
255     {
256         XMin[i]=tempXMin[i];
257         XMax[i]=tempXMax[i];
258     }
259
260     ScaleNb=tempScaleNb;
261 #endif
262 #endif
263
264     // --- Done ---
265
266     cout << "carmen: done.\n";
267
268     // --- Write solver type ---
269
270     if (Multiresolution)
271         cout << "carmen: multiresolution (MR) solver.\n";
272     else
273         cout << "carmen: finite volume (FV) solver.\n";
274
275     // --- Write number of iterations ---
276
277     if (IterationNb == 1)
278         cout << "carmen: compute 1 iteration ...\n";
279     else
280         cout << "carmen: compute " << IterationNb << " iterations ...\n";
281
282     printf("\n\n");
283
284     // --- Begin time iteration -----
285
286     for (IterationNo = 1; IterationNo <= IterationNb;
    IterationNo++)
287     {
288         // initializing eigenvalues - slopes
289         EigenvalueX = 0.;
290         EigenvalueY = 0.;
291         EigenvalueZ = 0.;
292         DIVBMax = 0.;
293
294         // --- Time evolution procedure ---
295         if (Multiresolution)
296             TimeEvolution(Mesh);
297         else
298             TimeEvolution(FMesh);
299
300         // --- Remesh ---
301         if (Multiresolution) Remesh(Mesh);
302

```

```

303     // --- Check CPU Time ---
304     CPUTime.check();
305
306     // --- Write information every (Refresh) iteration ---
307     if ((IterationNo-1)%Refresh == 0)
308     {
309         // - Write integral values -
310         if (rank==0) PrintIntegral("Integral.dat");
311
312         if (Cluster == 0)
313             ShowTime(CPUTime); // Show time on screen
314     //else
315         if (rank==0) Performance("carmen.prf"); // Refresh file "carmen.prf"
316     }
317
318     // --- Backup data every (10*Refresh) iteration ---
319     if ((IterationNo-1)%(10*Refresh) == 0 && UseBackup)
320     {
321         if (Multiresolution)
322             Backup(Mesh);
323         else
324             Backup(FMesh);
325     }
326
327     // --- Print solution if IterationNo = PrintIt1 to PrintIt6 ---
328     if (Multiresolution)
329         ViewIteration(Mesh);
330     else
331         ViewIteration(FMesh);
332
333     // --- Print solution if IterationNo is a multiple of PrintEvery ---
334     if (PrintEvery != 0)
335     {
336         {
337             if (IterationNo%PrintEvery == 0)
338             {
339                 if (Multiresolution)
340                     ViewEvery(Mesh, IterationNo);
341                 else
342                     ViewEvery(FMesh, IterationNo);
343             }
344         }
345
346         //if(ElapsedTime>=PhysicalTime)break;
347
348         // --- End time iteration -----
349     }
350
351     // --- Backup final data -----
352
353     IterationNo--;
354
355     if (UseBackup)
356     {
357         if (Multiresolution)
358             Backup(Mesh);
359         else
360             Backup(FMesh);
361     }
362
363     // --- Write integral values -----
364
365     if (rank==0) PrintIntegral("Integral.dat");
366
367     IterationNo++;
368
369     // --- Save values into file -----
370
371     if (Multiresolution)
372         View(Mesh, "Tree.dat", "Mesh.dat", "Average.vtk");
373     else
374         View(FMesh, "Average.vtk");
375
376     cout << "\ncarmen: done.\n";
377
378     // --- Analyse performance and save it into file -----
379
380     if (rank==0) Performance("carmen.prf");
381
382     // --- End -----
383
384     if (Multiresolution)
385         delete Mesh;
386     else
387         delete FMesh;
388
389

```

```

390 #if defined PARMPI
391
392 //free memory for the MPI runtime variables
393 delete[] disp;
394 delete[] blocklen;
395 int sz;
396 MPI_Buffer_detach(&MPIbuffer, &sz);
397 // for (int i = 0; i < 4*Dimension; i++) MPI_Request_free(&req[i]);
398 MPI_Finalize();
399
400 #endif
401
402 cout <<"carmen: end execution.\n";
403 return EXIT_SUCCESS;
404 }

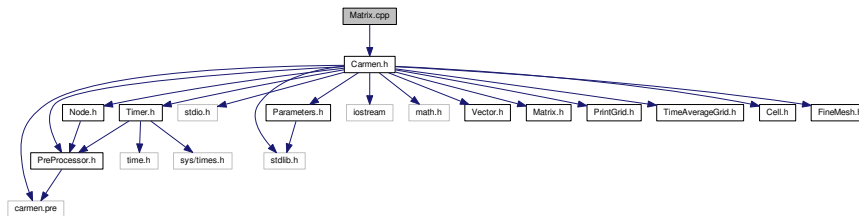
```

## 6.26 Matrix.cpp File Reference

Construct the data structures.

```
#include "Carmen.h"
```

Include dependency graph for Matrix.cpp:



## Functions

- **Matrix operator\*** (const real a, const Matrix &M)  
*Returns the product of the current matrix and a real a.*
- **ostream & operator<<** (ostream &out, const Matrix &M)  
*Writes the components of the matrix M on screen.*

### 6.26.1 Detailed Description

Construct the data structures.

### 6.26.2 Function Documentation

#### 6.26.2.1 Matrix operator\* ( const real a, const Matrix & M )

Returns the product of the current matrix and a real a.

Example :

```

#include "Matrix.h"
Matrix M(5, 3);
Matrix P;
real b = 2.;
...

```

$P = b * M;$

The operation  $P = M * b$  can also be done. See [Matrix Matrix::operator\\*\(const real a\) const](#).

Parameters

|     |                        |
|-----|------------------------|
| $a$ | Real value             |
| $M$ | <a href="#">Matrix</a> |

Returns

[Matrix](#)

```
1041 {
1042     return M*a;
1043 }
```

### 6.26.2.2 ostream& operator<< ( ostream & out, const Matrix & M )

Writes the components of the matrix  $M$  on screen.

Parameters

|       |                        |
|-------|------------------------|
| $out$ |                        |
| $M$   | <a href="#">Matrix</a> |

Returns

ostream&

```
1054 {
1055     int n;
1056     int m;
1057
1058     for (n = 1; n <= M.lines(); n++)
1059     {
1060         for (m = 1; m <= M.columns(); m++)
1061         {
1062             out<<n<<"<<m<<": "<<M.value(n,m)<<endl;
1063         }
1064     }
1065     return out;
1066 }
```

## 6.27 Matrix.h File Reference

This graph shows which files directly or indirectly include this file:



### Classes

- class [Matrix](#)  
*Standard class for every matrix in Carmen.*

### Functions

- [Matrix operator\\*](#) (const [real](#) a, const [Matrix](#) &M)

Returns the product of the current matrix and a real  $a$ .

- ostream & operator<< (ostream &out, const Matrix &M)

Writes the components of the matrix  $M$  on screen.

## 6.27.1 Function Documentation

### 6.27.1.1 Matrix operator\* ( const real $a$ , const Matrix & $M$ )

Returns the product of the current matrix and a real  $a$ .

Example :

```
#include "Matrix.h"
Matrix M(5,3);
Matrix P;
real b = 2.;
...
P = b*M;
```

The operation  $P = M*b$  can also be done. See [Matrix Matrix::operator\\*\(const real a\) const](#).

Parameters

|     |                        |
|-----|------------------------|
| $a$ | Real value             |
| $M$ | <a href="#">Matrix</a> |

Returns

[Matrix](#)

```
1041 {
1042     return M*a;
1043 }
```

### 6.27.1.2 ostream& operator<< ( ostream & out, const Matrix & $M$ )

Writes the components of the matrix  $M$  on screen.

Parameters

|       |                        |
|-------|------------------------|
| $out$ |                        |
| $M$   | <a href="#">Matrix</a> |

Returns

ostream&

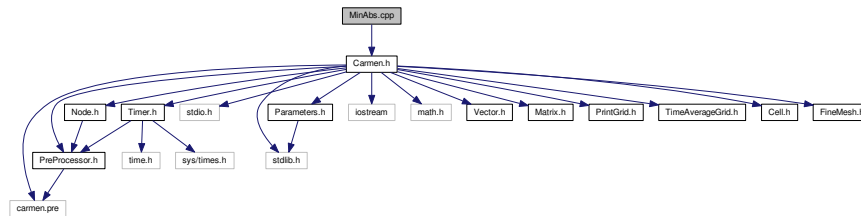
```
1054 {
1055     int n;
1056     int m;
1057
1058     for (n = 1; n <= M.lines(); n++)
1059     {
1060         for (m = 1; m <= M.columns(); m++)
1061         {
1062             out<<n<<" "<<m<<" : "<<M.value(n,m)<<endl;
1063         }
1064     }
1065     return out;
1066 }
```

## 6.28 MinAbs.cpp File Reference

Computes the minimal value between 2 numbers.

```
#include "Carmen.h"
```

Include dependency graph for MinAbs.cpp:



### Functions

- [real MinAbs](#) (const [real](#) a, const [real](#) b)  
*Returns the minimum in module of a and b.*

#### 6.28.1 Detailed Description

Computes the minimal value between 2 numbers.

#### 6.28.2 Function Documentation

##### 6.28.2.1 [real MinAbs](#) ( const [real](#) a, const [real](#) b )

Returns the minimum in module of *a* and *b*.

##### Parameters

|          |            |
|----------|------------|
| <i>a</i> | Real value |
| <i>b</i> | Real value |

##### Returns

double

```

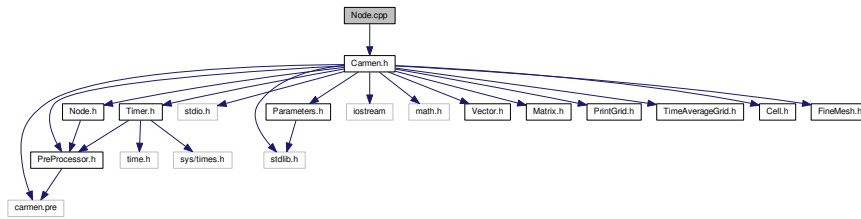
23 {
24     return (fabs(a) <= fabs(b)) ? a : b;
25 }
```

## 6.29 Node.cpp File Reference

Constructs the tree structure and computes the MHD multiresolution approach.



```
#include "Carmen.h"
Include dependency graph for Node.cpp:
```

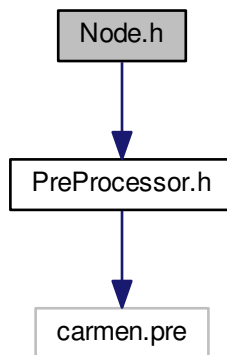


### 6.29.1 Detailed Description

Constructs the tree structure and computes the MHD multiresolution approach.

## 6.30 Node.h File Reference

```
#include "PreProcessor.h"
Include dependency graph for Node.h:
```



This graph shows which files directly or indirectly include this file:



### Classes

- class [Node](#)

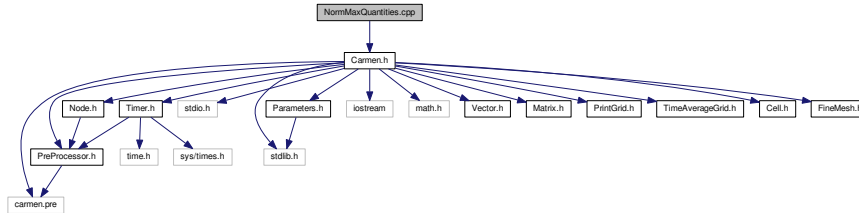
An object [Node](#) is an element of a graded tree structure, used for multiresolution computations. Its contains the following informations:

## 6.31 NormMaxQuantities.cpp File Reference

Compute the Linf norm of a vector containing the physical quantities divided by their characteristic value.

```
#include "Carmen.h"
```

Include dependency graph for NormMaxQuantities.cpp:



### Functions

- [real NormMaxQuantities](#) (const [Vector](#) &V)

*Returns the Max-norm of the vector where every quantity is divided by its characteristic value.*

#### 6.31.1 Detailed Description

Compute the Linf norm of a vector containing the physical quantities divided by their characteristic value.

#### Author

Anna Karina Fontes Gomes

#### Date

January-2017

#### 6.31.2 Function Documentation

##### 6.31.2.1 [real NormMaxQuantities](#) ( const [Vector](#) & V )

Returns the Max-norm of the vector where every quantity is divided by its characteristic value.

#### Parameters

|   |                        |
|---|------------------------|
| V | <a href="#">Vector</a> |
|---|------------------------|

#### Returns

double

```

26 {
27     Vector W(QuantityNb);
28     int AxisNo=1;
29     real MomentumMax=0.;
30     real MagMax=0.;
31
32
33
34     W.setZero();
35
36     /*
37     // Density
  
```

```

38     W.setValue(1, V.value(1)/QuantityMax.value(1));
39
40     // Momentum
41     W.setValue(2, V.value(2)/QuantityMax.value(2) );
42     W.setValue(3, V.value(3)/QuantityMax.value(3) );
43     W.setValue(4, V.value(4)/QuantityMax.value(4) );
44
45     // Energy
46     W.setValue(5, V.value(5)/QuantityMax.value(5) );
47
48     // psi
49     // W.setValue(6, V.value(6)/QuantityMax.value(6));
50
51     // Magnetic Field
52     W.setValue(7, V.value(7)/QuantityMax.value(7) );
53     W.setValue(8, V.value(8)/QuantityMax.value(8) );
54     W.setValue(9, V.value(9)/QuantityMax.value(9) );
55 */
56
57 // --- Compute Linf norm ---
58
59 W.setValue(1, (V.value(1))/QuantityMax.value(1));
60 W.setValue(5, (V.value(5))/QuantityMax.value(5));
61 //W.setValue(6, (V.value(6))/QuantityMax.value(6));
62
63
64 for (AxisNo = 1; AxisNo <= Dimension; AxisNo++)
65 {
66     MomentumMax = Max( MomentumMax, QuantityMax.value(AxisNo+1) );
67     MagMax       = Max( MagMax,      QuantityMax.value(AxisNo+6) );
68     W.setValue(2, W.value(2) + V.value(AxisNo+1)*V.value(AxisNo+1));
69     W.setValue(7, W.value(7) + V.value(AxisNo+6)*V.value(AxisNo+6));
70 }
71
72 if(Dimension==2){
73     W.setValue(4, (V.value(4))/QuantityMax.value(4));
74     W.setValue(9, (V.value(9))/QuantityMax.value(9));
75 }
76
77 W.setValue(2, sqrt(W.value(2))/MomentumMax);
78 W.setValue(7, sqrt(W.value(7))/MagMax );
79
80 if(IterationNo==0) return NMax(V);
81 return NMax(W);
82 }

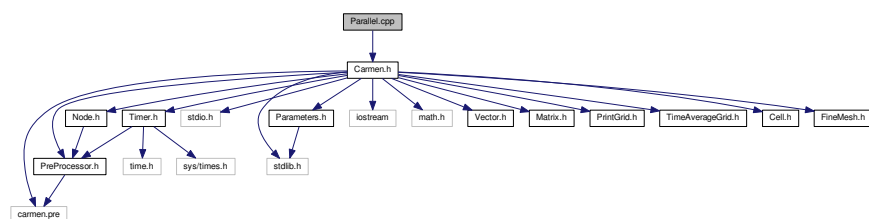
```

## 6.32 Parallel.cpp File Reference

Parallel implementation (not working yet)

```
#include "Carmen.h"
```

Include dependency graph for Parallel.cpp:



## Functions

- void [CreateMPITopology](#) ()  
*Parallel function DOES NOT WORK!*
- void [FillCellAddr](#) (Cell \*Mesh4MPI, int d, int &n)
- void [FillNbAddr](#) (Cell \*\*\*Nb, int l, int i, int j, int &n)
- void [CreateMPIType](#) (FineMesh \*Root)

- void `FreeMPIType` ()  
*Parallel function DOES NOT WORK!*
- void `CreateMPILinks` ()  
*Parallel function DOES NOT WORK!*
- void `CPUExchange` (`FineMesh *Root`, int WS)  
*Parallel function DOES NOT WORK!*
- void `ReduceIntegralValues` ()  
*Parallel function DOES NOT WORK!*

### 6.32.1 Detailed Description

Parallel implementation (not working yet)

### 6.32.2 Function Documentation

#### 6.32.2.1 void `CPUExchange` ( `FineMesh * Root`, int )

Parallel function DOES NOT WORK!

Parameters

|             |           |
|-------------|-----------|
| <i>Root</i> | Fine mesh |
|-------------|-----------|

Returns

void

```

350                                     {
351     CommTimer.start ();
352     #if defined PARMPI
353     int i,k;
354     int exNb=0;
355
356     WhatSend=WS;
357     CellElementsNb=0;
358
359     for (i=0;i<16;i++) {
360         k=1<<i;
361         if ((WS & k) != 0) CellElementsNb++;
362     }
363
364     static bool ft=true;
365     // if (ft==true) {
366     CreateMPIType(Root);
367     // CreateMPILinks();
368     // ft=false;
369     // }
370
371     // MPI_Startall(4*Dimension,req);
372
373
374     //Send
375     switch (MPISendType) {
376     case 0:
377         MPI_Ibsend(MPI_BOTTOM, 1, MPItypeSiL, rank_il, 100, comm_cart ,&req[exNb++]);
378         MPI_Ibsend(MPI_BOTTOM, 1, MPItypeSiU, rank_iu, 200, comm_cart, &req[exNb++]);
379         break;
380
381     case 10:
382         MPI_Isend(MPI_BOTTOM, 1, MPItypeSiL, rank_il, 100, comm_cart,&req[exNb++]);
383         MPI_Isend(MPI_BOTTOM, 1, MPItypeSiU, rank_iu, 200, comm_cart,&req[exNb++]);
384         break;
385
386     case 20:
387         MPI_Issend(MPI_BOTTOM, 1, MPItypeSiL, rank_il, 100, comm_cart,&req[exNb++]);
388         MPI_Issend(MPI_BOTTOM, 1, MPItypeSiU, rank_iu, 200, comm_cart,&req[exNb++]);
389         break;
390     }
391
392     if (Dimension >= 2) {
393         switch (MPISendType) {

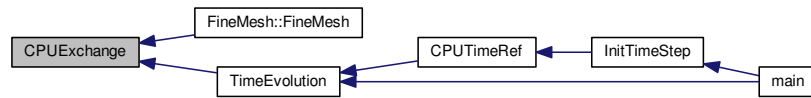
```

```

394     case 0:
395         MPI_Ibsend(MPI_BOTTOM, 1, MPItypeSjL, rank_jl, 300, comm_cart,&req[exNb++]);
396         MPI_Ibsend(MPI_BOTTOM, 1, MPItypeSjU, rank_ju, 400, comm_cart,&req[exNb++]);
397         break;
398
399     case 10:
400         MPI_Isend(MPI_BOTTOM, 1, MPItypeSjL, rank_jl, 300, comm_cart,&req[exNb++]);
401         MPI_Isend(MPI_BOTTOM, 1, MPItypeSjU, rank_ju, 400, comm_cart,&req[exNb++]);
402         break;
403
404     case 20:
405         MPI_Issend(MPI_BOTTOM, 1, MPItypeSjL, rank_jl, 300, comm_cart,&req[exNb++]);
406         MPI_Issend(MPI_BOTTOM, 1, MPItypeSjU, rank_ju, 400, comm_cart,&req[exNb++]);
407         break;
408     }
409 }
410
411 if (Dimension == 3) {
412     switch (MPISendType) {
413     case 0:
414         MPI_Ibsend(MPI_BOTTOM, 1, MPItypeSkL, rank_kl, 500, comm_cart,&req[exNb++]);
415         MPI_Ibsend(MPI_BOTTOM, 1, MPItypeSkU, rank_ku, 600, comm_cart,&req[exNb++]);
416         break;
417
418     case 10:
419         MPI_Isend(MPI_BOTTOM, 1, MPItypeSkL, rank_kl, 500, comm_cart,&req[exNb++]);
420         MPI_Isend(MPI_BOTTOM, 1, MPItypeSkU, rank_ku, 600, comm_cart,&req[exNb++]);
421         break;
422
423     case 20:
424         MPI_Issend(MPI_BOTTOM, 1, MPItypeSkL, rank_kl, 500, comm_cart,&req[exNb++]);
425         MPI_Issend(MPI_BOTTOM, 1, MPItypeSkU, rank_ku, 600, comm_cart,&req[exNb++]);
426         break;
427     }
428 }
429
430 //Recv
431
432 if (MPIRecvType==0) {
433     MPI_Recv(MPI_BOTTOM, 1, MPItypeRiL, rank_il, 200, comm_cart, &st[6]);
434     MPI_Recv(MPI_BOTTOM, 1, MPItypeRiU, rank_iu, 100, comm_cart, &st[7]);
435 } else
436 {
437     MPI_Irecv(MPI_BOTTOM, 1, MPItypeRiL, rank_il, 200, comm_cart, &req[exNb++]);
438     MPI_Irecv(MPI_BOTTOM, 1, MPItypeRiU, rank_iu, 100, comm_cart, &req[exNb++]);
439 }
440
441 if (Dimension >= 2) {
442     if (MPIRecvType==0) {
443         MPI_Recv(MPI_BOTTOM, 1, MPItypeRjL, rank_jl, 400, comm_cart, &st[8]);
444         MPI_Recv(MPI_BOTTOM, 1, MPItypeRjU, rank_ju, 300, comm_cart, &st[9]);
445     } else
446     {
447         MPI_Irecv(MPI_BOTTOM, 1, MPItypeRjL, rank_jl, 400, comm_cart, &req[exNb++]);
448         MPI_Irecv(MPI_BOTTOM, 1, MPItypeRjU, rank_ju, 300, comm_cart, &req[exNb++]);
449     }
450 }
451
452 if (Dimension == 3) {
453     if (MPIRecvType==0) {
454         MPI_Recv(MPI_BOTTOM, 1, MPItypeRkL, rank_kl, 600, comm_cart, &st[10]);
455         MPI_Recv(MPI_BOTTOM, 1, MPItypeRkU, rank_ku, 500, comm_cart, &st[11]);
456     } else
457     {
458         MPI_Irecv(MPI_BOTTOM, 1, MPItypeRkL, rank_kl, 600, comm_cart, &req[exNb++]);
459         MPI_Irecv(MPI_BOTTOM, 1, MPItypeRkU, rank_ku, 500, comm_cart, &req[exNb++]);
460     }
461 }
462
463 FreeMPIType();
464 #endif
465 CommTimer.stop();
466 }

```

Here is the caller graph for this function:



### 6.32.2.2 void CreateMPLinks ( )

Parallel function DOES NOT WORK!

Returns

void

```

271     {
272     int exNb;
273     exNb=0;
274     #if defined PARMPI
275
276     //Send
277
278     switch (MPISendType) {
279     case 0:
280         MPI_Bsend_init(MPI_BOTTOM, 1, MPItypeSiL, rank_il, 100, comm_cart, &req[exNb++]);
281         MPI_Bsend_init(MPI_BOTTOM, 1, MPItypeSiU, rank_iu, 200, comm_cart, &req[exNb++]);
282         break;
283
284     case 10:
285         MPI_Send_init(MPI_BOTTOM, 1, MPItypeSiL, rank_il, 100, comm_cart, &req[exNb++]);
286         MPI_Send_init(MPI_BOTTOM, 1, MPItypeSiU, rank_iu, 200, comm_cart, &req[exNb++]);
287         break;
288
289     case 20:
290         MPI_Ssend_init(MPI_BOTTOM, 1, MPItypeSiL, rank_il, 100, comm_cart, &req[exNb++]);
291         MPI_Ssend_init(MPI_BOTTOM, 1, MPItypeSiU, rank_iu, 200, comm_cart, &req[exNb++]);
292         break;
293     }
294
295     if (Dimension >= 2) {
296         switch (MPISendType) {
297         case 0:
298             MPI_Bsend_init(MPI_BOTTOM, 1, MPItypeSjL, rank_jl, 300, comm_cart, &req[exNb++]);
299             MPI_Bsend_init(MPI_BOTTOM, 1, MPItypeSjU, rank_ju, 400, comm_cart, &req[exNb++]);
300             break;
301
302         case 10:
303             MPI_Send_init(MPI_BOTTOM, 1, MPItypeSjL, rank_jl, 300, comm_cart, &req[exNb++]);
304             MPI_Send_init(MPI_BOTTOM, 1, MPItypeSjU, rank_ju, 400, comm_cart, &req[exNb++]);
305             break;
306
307         case 20:
308             MPI_Ssend_init(MPI_BOTTOM, 1, MPItypeSjL, rank_jl, 300, comm_cart, &req[exNb++]);
309             MPI_Ssend_init(MPI_BOTTOM, 1, MPItypeSjU, rank_ju, 400, comm_cart, &req[exNb++]);
310             break;
311         }
312     }
313
314     if (Dimension == 3) {
315         switch (MPISendType) {
316         case 0:
317             MPI_Bsend_init(MPI_BOTTOM, 1, MPItypeSkL, rank_kl, 500, comm_cart, &req[exNb++]);
318             MPI_Bsend_init(MPI_BOTTOM, 1, MPItypeSkU, rank_ku, 600, comm_cart, &req[exNb++]);
319             break;
320
321         case 10:
322             MPI_Send_init(MPI_BOTTOM, 1, MPItypeSkL, rank_kl, 500, comm_cart, &req[exNb++]);
323             MPI_Send_init(MPI_BOTTOM, 1, MPItypeSkU, rank_ku, 600, comm_cart, &req[exNb++]);
324             break;
325
326         case 20:
327             MPI_Ssend_init(MPI_BOTTOM, 1, MPItypeSkL, rank_kl, 500, comm_cart, &req[exNb++]);
328             MPI_Ssend_init(MPI_BOTTOM, 1, MPItypeSkU, rank_ku, 600, comm_cart, &req[exNb++]);
  
```

```

329         break;
330     }
331 }
332
333 //Recv
334
335 MPI_Recv_init(MPI_BOTTOM, 1, MPItypeRiL, rank_il, 200, comm_cart, &req[exNb++]);
336 MPI_Recv_init(MPI_BOTTOM, 1, MPItypeRiU, rank_iu, 100, comm_cart, &req[exNb++]);
337
338 if (Dimension >= 2) {
339     MPI_Recv_init(MPI_BOTTOM, 1, MPItypeRjL, rank_jl, 400, comm_cart, &req[exNb++]);
340     MPI_Recv_init(MPI_BOTTOM, 1, MPItypeRjU, rank_ju, 300, comm_cart, &req[exNb++]);
341 }
342
343 if (Dimension == 3) {
344     MPI_Recv_init(MPI_BOTTOM, 1, MPItypeRkL, rank_kl, 600, comm_cart, &req[exNb++]);
345     MPI_Recv_init(MPI_BOTTOM, 1, MPItypeRkU, rank_ku, 500, comm_cart, &req[exNb++]);
346 }
347 #endif
348 }

```

### 6.32.2.3 void CreateMPITopology ( )

Parallel function DOES NOT WORK!

#### Returns

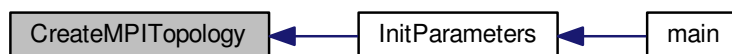
void

```

22     {
23 #if defined PARMPI
24     int src;
25     int periods[]={1,1,1};
26     CartDims[0]=CartDims[1]=CartDims[2]=0;
27
28     MPI_Dims_create(size,Dimension,CartDims);
29     MPI_Cart_create(MPI_COMM_WORLD,Dimension,CartDims,periods,1,&comm_cart);
30     MPI_Comm_rank(comm_cart, &rank);
31     MPI_Cart_coords(comm_cart,rank,Dimension,coords);
32
33     MPI_Cart_shift(comm_cart, 0, -1, &src, &rank_il);
34     MPI_Cart_shift(comm_cart, 0, 1, &src, &rank_iu);
35
36     if (Dimension >= 2) {
37         MPI_Cart_shift(comm_cart, 1, -1, &src, &rank_jl);
38         MPI_Cart_shift(comm_cart, 1, 1, &src, &rank_ju);
39     }
40
41     if (Dimension == 3) {
42         MPI_Cart_shift(comm_cart, 2, -1, &src, &rank_kl);
43         MPI_Cart_shift(comm_cart, 2, 1, &src, &rank_ku);
44     }
45 #endif
46 }

```

Here is the caller graph for this function:



### 6.32.2.4 void CreateMPIType ( FineMesh \* Root )

```

121     {
122 #if defined PARMPI

```

```

123 int i,j,k;
124 int n,d,l;
125
126 Cell *MeshCell;
127 MeshCell=Root->MeshCell;
128
129 n=0;
130 for (l=0;l<NeighbourNb;l++)
131     for (j=0;j<one_D;j++)
132         for (k=0;k<two_D;k++) FillNbAddr (Root->Neighbour_iL,l,j,k,n);
133 MPI_Type_hindexed (CellElementsNb*NeighbourNb*one_D*two_D,blocklen,disp,
MPI_Type, &MPItypeRiL);
134 MPI_Type_commit (&MPItypeRiL);
135
136 n=0;
137 for (l=0;l<NeighbourNb;l++)
138     for (j=0;j<one_D;j++)
139         for (k=0;k<two_D;k++) FillNbAddr (Root->Neighbour_iU,l,j,k,n);
140 MPI_Type_hindexed (CellElementsNb*NeighbourNb*one_D*two_D,blocklen,disp,
MPI_Type, &MPItypeRiU);
141 MPI_Type_commit (&MPItypeRiU);
142
143 n=0;
144 for (l=0;l<NeighbourNb;l++)
145     for (j=0;j<one_D;j++)
146         for (k=0;k<two_D;k++) {
147             i=1;
148             d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
149             FillCellAddr (MeshCell,d,n);
150         }
151 MPI_Type_hindexed (CellElementsNb*NeighbourNb*one_D*two_D,blocklen,disp,
MPI_Type, &MPItypeSiL);
152 MPI_Type_commit (&MPItypeSiL);
153
154 n=0;
155 for (l=0;l<NeighbourNb;l++)
156     for (j=0;j<one_D;j++)
157         for (k=0;k<two_D;k++) {
158             i=(1<<ScaleNb)-NeighbourNb+1;
159             d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
160             FillCellAddr (MeshCell,d,n);
161         }
162 MPI_Type_hindexed (CellElementsNb*NeighbourNb*one_D*two_D,blocklen,disp,
MPI_Type, &MPItypeSiU);
163 MPI_Type_commit (&MPItypeSiU);
164
165 if (Dimension >= 2) {
166     n=0;
167     for (l=0;l<NeighbourNb;l++)
168         for (i=0;i<one_D;i++)
169             for (k=0;k<two_D;k++) FillNbAddr (Root->
Neighbour_jL,l,i,k,n);
170 MPI_Type_hindexed (CellElementsNb*NeighbourNb*one_D*two_D,blocklen,disp,
MPI_Type, &MPItypeRjL);
171 MPI_Type_commit (&MPItypeRjL);
172
173     n=0;
174     for (l=0;l<NeighbourNb;l++)
175         for (i=0;i<one_D;i++)
176             for (k=0;k<two_D;k++) FillNbAddr (Root->
Neighbour_jU,l,i,k,n);
177 MPI_Type_hindexed (CellElementsNb*NeighbourNb*one_D*two_D,blocklen,disp,
MPI_Type, &MPItypeRjU);
178 MPI_Type_commit (&MPItypeRjU);
179
180     n=0;
181     for (l=0;l<NeighbourNb;l++)
182         for (i=0;i<one_D;i++)
183             for (k=0;k<two_D;k++) {
184                 j=1;
185                 d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
186                 FillCellAddr (MeshCell,d,n);
187             }
188 MPI_Type_hindexed (CellElementsNb*NeighbourNb*one_D*two_D,blocklen,disp,
MPI_Type, &MPItypeSjL);
189 MPI_Type_commit (&MPItypeSjL);
190
191     n=0;
192     for (l=0;l<NeighbourNb;l++)
193         for (i=0;i<one_D;i++)
194             for (k=0;k<two_D;k++) {
195                 j=(1<<ScaleNb)-NeighbourNb+1;
196                 d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
197                 FillCellAddr (MeshCell,d,n);
198             }
199 MPI_Type_hindexed (CellElementsNb*NeighbourNb*one_D*two_D,blocklen,disp,
MPI_Type, &MPItypeSjU);

```



```

200     MPI_Type_commit (&MPITypeSjU);
201 }
202
203
204 if (Dimension == 3) {
205     n=0;
206     for (l=0;l<NeighbourNb;l++)
207         for (i=0;i<one_D;i++)
208             for (j=0;j<two_D;j++) FillNbAddr (Root->
Neighbour_kL,l,i,j,n);
209     MPI_Type_hindexed (CellElementsNb*NeighbourNb*one_D*two_D,blocklen,disp,
MPI_Type,&MPITypeRkL);
210     MPI_Type_commit (&MPITypeRkL);
211
212     n=0;
213     for (l=0;l<NeighbourNb;l++)
214         for (i=0;i<one_D;i++)
215             for (j=0;j<two_D;j++) FillNbAddr (Root->
Neighbour_kU,l,i,j,n);
216     MPI_Type_hindexed (CellElementsNb*NeighbourNb*one_D*two_D,blocklen,disp,
MPI_Type,&MPITypeRkU);
217     MPI_Type_commit (&MPITypeRkU);
218
219     n=0;
220     for (l=0;l<NeighbourNb;l++)
221         for (i=0;i<one_D;i++)
222             for (j=0;j<two_D;j++) {
223                 k=1;
224                 d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
225                 FillCellAddr (MeshCell,d,n);
226             }
227     MPI_Type_hindexed (CellElementsNb*NeighbourNb*one_D*two_D,blocklen,disp,
MPI_Type,&MPITypeSkL);
228     MPI_Type_commit (&MPITypeSkL);
229
230     n=0;
231     for (l=0;l<NeighbourNb;l++)
232         for (i=0;i<one_D;i++)
233             for (j=0;j<two_D;j++) {
234                 k=(1<<ScaleNb)-NeighbourNb+1;
235                 d=i + (1<<ScaleNb)*(j + (1<<ScaleNb)*k);
236                 FillCellAddr (MeshCell,d,n);
237             }
238     MPI_Type_hindexed (CellElementsNb*NeighbourNb*one_D*two_D,blocklen,disp,
MPI_Type,&MPITypeSkU);
239     MPI_Type_commit (&MPITypeSkU);
240 }
241
242 #endif
243 }

```

Here is the caller graph for this function:



### 6.32.2.5 void FillCellAddr ( Cell \* Mesh4MPI, int d, int & n )

```

49     {
50 #if defined PARMPI
51
52     if ((WhatSend & SendQ) != 0) {
53         MPI_Address (Mesh4MPI[d].Q.U, &disp[n]);
54         blocklen[n++] = Mesh4MPI[d].Q.dimension();
55     }
56
57     if ((WhatSend & SendQs) != 0) {
58         MPI_Address (Mesh4MPI[d].Qs.U, &disp[n]);
59         blocklen[n++] = Mesh4MPI[d].Qs.dimension();
60     }
61
62     if ((WhatSend & SendX) != 0) {
63         MPI_Address (Mesh4MPI[d].X.U, &disp[n]);

```

```

64     blocklen[n++]=Mesh4MPI[d].X.dimension();
65 }
66
67 if ((WhatSend & SenddX) != 0) {
68     MPI_Address(Mesh4MPI[d].dX.U, &disp[n]);
69     blocklen[n++]=Mesh4MPI[d].dX.dimension();
70 }
71
72 if ((WhatSend & SendD) != 0) {
73     MPI_Address(Mesh4MPI[d].D.U, &disp[n]);
74     blocklen[n++]=Mesh4MPI[d].D.dimension();
75 }
76
77 if ((WhatSend & SendGrad) != 0) {
78     MPI_Address(Mesh4MPI[d].Grad.U, &disp[n]);
79     blocklen[n++]=Mesh4MPI[d].Grad.columns()*Mesh4MPI[d].Grad.
lines();
80 }
81 #endif
82 }

```

Here is the caller graph for this function:



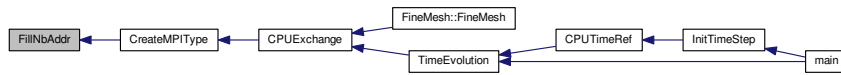
### 6.32.2.6 void FillNbAddr ( Cell \*\*\* Nb, int l, int i, int j, int & n )

```

85                                     {
86 #if defined PARMPI
87     if ((WhatSend & SendQ) != 0) {
88         MPI_Address(Nb[l][i][j].Q.U, &disp[n]);
89         blocklen[n++]=Nb[l][i][j].Q.dimension();
90     }
91
92     if ((WhatSend & SendQs) != 0) {
93         MPI_Address(Nb[l][i][j].Qs.U, &disp[n]);
94         blocklen[n++]=Nb[l][i][j].Qs.dimension();
95     }
96
97     if ((WhatSend & SendX) != 0) {
98         MPI_Address(Nb[l][i][j].X.U, &disp[n]);
99         blocklen[n++]=Nb[l][i][j].X.dimension();
100    }
101
102    if ((WhatSend & SenddX) != 0) {
103        MPI_Address(Nb[l][i][j].dX.U, &disp[n]);
104        blocklen[n++]=Nb[l][i][j].dX.dimension();
105    }
106
107    if ((WhatSend & SendD) != 0) {
108        MPI_Address(Nb[l][i][j].D.U, &disp[n]);
109        blocklen[n++]=Nb[l][i][j].D.dimension();
110    }
111
112
113    if ((WhatSend & SendGrad) != 0) {
114        MPI_Address(Nb[l][i][j].Grad.U, &disp[n]);
115        blocklen[n++]=Nb[l][i][j].Grad.columns()*Nb[l][i][j].Grad.
lines();
116    }
117 #endif
118 }

```

Here is the caller graph for this function:



### 6.32.2.7 void FreeMPIType ( )

Parallel function DOES NOT WORK!

#### Returns

void

```

247     {
248     #if defined PARMPI
249     MPI_Type_free (&MPItypeSiL);
250     MPI_Type_free (&MPItypeSiU);
251     MPI_Type_free (&MPItypeRiL);
252     MPI_Type_free (&MPItypeRiU);
253
254     if (Dimension >= 2) {
255     MPI_Type_free (&MPItypeSjL);
256     MPI_Type_free (&MPItypeSjU);
257     MPI_Type_free (&MPItypeRjL);
258     MPI_Type_free (&MPItypeRjU);
259     }
260
261     if (Dimension == 3) {
262     MPI_Type_free (&MPItypeSkL);
263     MPI_Type_free (&MPItypeSkU);
264     MPI_Type_free (&MPItypeRkL);
265     MPI_Type_free (&MPItypeRkU);
266     }
267     #endif
268     }
    
```

Here is the caller graph for this function:



### 6.32.2.8 void ReduceIntegralValues ( )

Parallel function DOES NOT WORK!

#### Returns

void

```

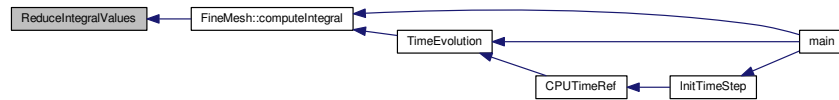
469     {
470     real rb; //Recieve Buffer
471     rb=0.0;
472     CommTimer.start ();
473     #if defined PARMPI
474     MPI_Reduce (&ErrorMax, &rb, 1, MPI_Type, MPI_MAX, 0, MPI_COMM_WORLD);
475     ErrorMax=rb;
476     }
    
```

```

477 MPI_Reduce (&ErrorMid, &rb, 1, MPI_Type, MPI_SUM, 0, MPI_COMM_WORLD);
478 ErrorMid=rb/size;
479
480 MPI_Reduce (&ErrorL2, &rb, 1, MPI_Type, MPI_SUM, 0, MPI_COMM_WORLD);
481 ErrorL2=rb/size;
482
483 MPI_Reduce (&ErrorNb, &rb, 1, MPI_Type, MPI_SUM, 0, MPI_COMM_WORLD);
484 ErrorNb=rb;
485
486 MPI_Allreduce (&FlameVelocity, &rb, 1, MPI_Type, MPI_SUM, MPI_COMM_WORLD);
487 FlameVelocity=rb;
488
489 MPI_Allreduce (&GlobalMomentum, &rb, 1, MPI_Type, MPI_SUM, MPI_COMM_WORLD);
490 GlobalMomentum=rb;
491
492 MPI_Allreduce (&GlobalEnergy, &rb, 1, MPI_Type, MPI_SUM, MPI_COMM_WORLD);
493 GlobalEnergy=rb;
494
495 MPI_Reduce (&ExactMomentum, &rb, 1, MPI_Type, MPI_SUM, 0, MPI_COMM_WORLD);
496 ExactMomentum=rb;
497
498 MPI_Reduce (&ExactEnergy, &rb, 1, MPI_Type, MPI_SUM, 0, MPI_COMM_WORLD);
499 ExactEnergy=rb;
500
501 MPI_Allreduce (&GlobalReactionRate, &rb, 1, MPI_Type, MPI_SUM, MPI_COMM_WORLD);
502 GlobalReactionRate=rb;
503
504 MPI_Allreduce (&EigenvalueMax, &rb, 1, MPI_Type, MPI_MAX, MPI_COMM_WORLD);
505 EigenvalueMax=rb;
506
507 #endif
508 CommTimer.stop ();
509 }

```

Here is the caller graph for this function:



## 6.33 Parameters.cpp File Reference

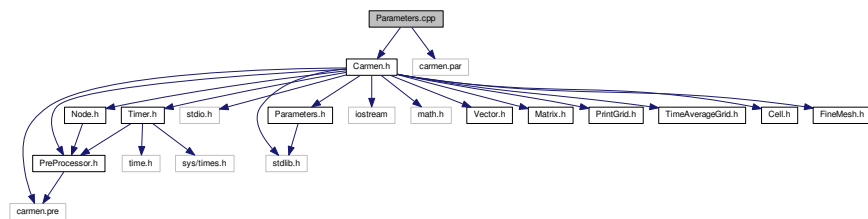
User parameters.

```

#include "Carmen.h"
#include "carmen.par"

```

Include dependency graph for Parameters.cpp:



## Functions

- void [InitParameters](#) ()

*Inites parameters from file carmen.par. If a parameter is not mentioned in this file, the default value is used.*

## Variables

- `real CarmenVersion` =2.483
- `int StepNb` =2
- `int IterationNb` =0
- `int IterationNbRef` =1000
- `real CFL` =0.5
- `real TimeStep` =0.
- `real PhysicalTime` =0.
- `int Refresh` =0
- `int RefreshNb` =200
- `real PrintTime1` =0.
- `real PrintTime2` =0.
- `real PrintTime3` =0.
- `real PrintTime4` =0.
- `real PrintTime5` =0.
- `real PrintTime6` =0.
- `int PrintIt1` =0
- `int PrintIt2` =0
- `int PrintIt3` =0
- `int PrintIt4` =0
- `int PrintIt5` =0
- `int PrintIt6` =0
- `int PrintEvery` =0
- `int ImageNb` =0
- `bool UseBackup` = true
- `bool Recovery` =false
- `bool TimeAveraging` =false
- `real StartTimeAveraging` =0.
- `bool ComputeCPUTimeRef` =false
- `int EquationType` =7
- `int SchemeNb` =1
- `int LimiterNo` =5
- `int ScalarEqNb` =0
- `int DivClean` =2
- `int Dimension` =1
- `int Coordinate` =1
- `real XMin` [4] ={-1.,-1.,-1.,-1.}
- `real XMax` [4] ={1.,1.,1.,1.}
- `int CMin` [4] ={3,3,3,3}
- `int CMax` [4] ={3,3,3,3}
- `bool UseBoundaryRegions` = false
- `bool Multiresolution` =true
- `bool TimeAdaptivity` =false
- `int TimeAdaptivityFactor` = 1
- `int ScaleNb` =5
- `int ScaleNbRef` =2
- `int IcNb` = 0
- `int PrintMoreScales` =0
- `real Tolerance` =0.
- `real ToleranceScale` =1.
- `real PenalizeFactor` =1.
- `bool ConstantTimeStep` =true
- `real ElapsedTime` =0.
- `real ExpectedCompression` = 0.

- bool `CVS` =false
- bool `LES` =false
- real `SmoothCoeff` =0.
- int `ThresholdNorm` =0
- real `Celerity` = 1.
- real `Viscosity` = 1.
- real `Re` =100.
- real `Pr` =0.71
- real `Gamma` =1.4
- real `Ma` =0.3
- real `Fr` =0.
- real `TRef` =273.
- real `Circulation` =10.
- real `ForceX` =0.
- real `ForceY` =0.
- real `ForceZ` =0.
- real `ModelConstant` =0.1
- real `ThermalConduction` =0.
- real `ConstantForce` =true
- bool `Resistivity` =false
- bool `Diffusivity` =false
- bool `ComputeTemp` =false
- real `eta` =0.
- real `chi` =0.
- real `Alpha` =0.64
- real `Ze` =10.
- real `Le` =1.
- real `Sigma` =5.E-02
- real `DIVB` =0.
- real `DIVBMax` =0.
- real `Bdivergence` =0.
- real `PsiGrad` =0.
- real `ch` =0.0
- real `auxvar` =0.
- bool `debug` =false
- bool `FluxCorrection` =true
- int `PostProcessing` =2
- bool `DatalsBinary` =true
- bool `ZipData` =true
- bool `WriteAsPoints` =false
- real `SpaceStep` =0.
- real `Eigenvalue` =0.
- real `EigenvalueX` =0.
- real `EigenvalueY` =0.
- real `EigenvalueZ` =0.
- real `EigenvalueMax` =0.
- Vector `QuantityMax`
- Vector `QuantityAverage`
- real `pi` =acos(-1.)
- int `StepNo` =1
- int `IterationNo` =0
- Timer `CPUTime`
- int `Cluster` =1
- int `QuantityNb` =2
- double `FVTimeRef` =0.

- int CellNb =0
- int LeafNb =0
- real TotalLeafNb =0.
- real TotalCellNb =0.
- real ErrorMax =0.
- real ErrorGlobalMax =0.
- real ErrorMid =0.
- real ErrorGlobalMid =0.
- real ErrorL2 =0.
- real ErrorGlobalL2 =0.
- int ErrorNb =0
- int ErrorGlobalNb =0
- real RKFEError =0.
- real RKFAccuracyFactor =1.E-03
- real RKFSafetyFactor =0.01
- real cr =0.
- real Helicity =0.
- real GlobalMomentum =0.
- real GlobalEnergy =0.
- real GlobalEnstrophy =0.
- real ExactMomentum =0.
- real ExactEnergy =0.
- real FlameVelocity =0.
- real GlobalMomentumOld =0.
- real GlobalVolume =0.
- real GlobalReactionRate =0.
- real AverageRadius =0.
- real PreviousAverageRadius =0.
- real PreviousAverageRadius2 =0.
- real XCenter [4] ={0.,0.,0.,0.}
- real ReactionRateMax =0.
- FILE \* GlobalFile
- int ChildNb =0
- real IntVorticity =0.
- real IntDensity =0.
- Vector IntMomentum
- real IntEnergy =0.
- real BaroclinicEffect =0.
- int rank
- int size
- int CPUScales
- real AllXMin [4]
- real AllXMax [4]
- int AllTaskScaleNb
- int NeighbourNb
- int coords [3]
- int CartDims [3]
- int CellElementsNb
- int MaxCellElementsNb
- int one\_D
- int two\_D
- int MPISendType
- int MPIRecvType
- Timer CommTimer
- int rank\_il

- int `rank_iu`
- int `rank_jl`
- int `rank_ju`
- int `rank_kl`
- int `rank_ku`
- int `WhatSend`
- int `SendD` = 1 << 0
- int `SendGrad` = 1 << 1
- int `SendQ` = 1 << 2
- int `SendQs` = 1 << 3
- int `SendX` = 1 << 4
- int `SenddX` = 1 << 5
- char `BackupName` [255]

### 6.33.1 Detailed Description

User parameters.

### 6.33.2 Function Documentation

#### 6.33.2.1 void InitParameters ( )

Initializes parameters from file `carmen.par`. If a parameter is not mentioned in this file, the default value is used.

#### Returns

void

— Compute ch -----

```

284 {
285     // --- Local variables -----
286
287     int i;          // Counter
288
289     // --- Set global variables from file "carmen.par" -----
290
291     #include "carmen.par"
292
293     // --- Adapt IterationNbRef to the dimension -----
294
295     IterationNbRef=(int) (exp((4.-Dimension)*log(10.)));
296
297     // --- Compute the number of children of a given parent cell ---
298
299     ChildNb = (1<<Dimension);
300
301     #if defined PARMPI
302
303     AllTaskScaleNb=ScaleNb;
304     for (i=0;i<4;i++)
305     {
306         AllXMax[i]=XMax[i];
307         AllXMin[i]=XMin[i];
308     }
309
310     //some combinations give deadlock...
311     MPISendType = 10;    //0 - Ibsend; 10 - Isend; 20 - Issend;
312     MPIRecvType = 1;    //0 - Recv; 1 - Irecv;
313
314     CPUScales=0;
315     int tmp=size;
316     while ((tmp=(tmp>>1))>0) CPUScales++;
317     ScaleNb=CPUScales/Dimension;
318
319     one_D=1; two_D=1;
320     if (Dimension >= 2) one_D=1<<ScaleNb;
321     if (Dimension == 3) two_D=1<<ScaleNb;

```



```

322
323 // #if defined PARMPI
324
325     NeighbourNb=2;
326     MaxCellElementsNb=6;
327
328     // -- Create memory arrays that are needs for the MPI Type creation ---
329     disp = new MPI_Aint[NeighbourNb*MaxCellElementsNb*
one_D*two_D];
330     blocklen = new int [NeighbourNb*MaxCellElementsNb*
one_D*two_D];
331
332     // --- Allocate additional memory for MPI buffer send---
333     Cell tc;
334     int CellElNb,bufsize;
335     CellElNb=tc.size().dimension()+tc.center().dimension()+tc.average().dimension()+tc.
tempAverage().dimension()+tc.divergence().dimension();
336
337     if (EquationType==6)
338         CellElNb += tc.gradient().lines()*tc.gradient().columns();
339
340     bufsize=(CellElNb*one_D*two_D*NeighbourNb+MPI_BSEND_OVERHEAD)*2*
Dimension+1024;
341     MPIbuffer=new real[bufsize];
342     MPI_Buffer_attach(MPIbuffer,bufsize*sizeof(real));
343
344 #else
345     NeighbourNb=0;
346 #endif
347
348 #if defined PARMPI
349     CreateMPITopology();
350
351     // --- Compute domain coordinates for the processors ---
352     XMin[1] = AllXMin[1] + coords[0]*(AllXMax[1]-AllXMin[1])/
CartDims[0];
353     XMax[1] = AllXMin[1] + (coords[0]+1)*(AllXMax[1]-
AllXMin[1])/CartDims[0];
354
355     if (Dimension >= 2)
356     {
357         XMin[2] = AllXMin[2] + coords[1]*(AllXMax[2]-
AllXMin[2])/CartDims[1];
358         XMax[2] = AllXMin[2] + (coords[1]+1)*(AllXMax[2]-
AllXMin[2])/CartDims[1];
359     }
360
361     if (Dimension == 3)
362     {
363         XMin[3] = AllXMin[3] + coords[2]*(AllXMax[3]-
AllXMin[3])/CartDims[2];
364         XMax[3] = AllXMin[3] + (coords[2]+1)*(AllXMax[3]-
AllXMin[3])/CartDims[2];
365     }
366
367     // --- Set the backup file name for the current processor
368     sprintf(BackupName,"%d_%d_%d_%s",coords[0],coords[1],
coords[2],"carmen.bak");
369 #else
370     sprintf(BackupName,"%s","carmen.bak");
371 #endif
372
373
374     // --- Use CVS only if Dimension > 1 -----
375
376     if (Dimension == 1)
377         CVS = false;
378
379     // --- TimeAveraging always false if not Navier-Stokes -----
380
381     if (EquationType != 6)
382         TimeAveraging = false;
383
384     // --- If there is no file "carmen.bak", set Recovery=false -----
385
386     if (!fopen(BackupName,"r"))
387         Recovery = false;
388
389     // --- If PrintMoreScales != 0 or 1 with Multiresolution = false, print error and stop ---
390
391     if (!Multiresolution && (!(PrintMoreScales == 0 ||
PrintMoreScales == -1)))
392     {
393         cout << "Parameters.cpp: In method 'void InitParameters()':\n";
394         cout << "Parameters.cpp: value of PrintMoreScales incompatible with FV computations\n";
395         cout << "Parameters.cpp: must be 0 or -1\n";
396         cout << "carmen: *** [Parameters.o] Execution error\n";

```

```

397     cout << "carmen: abort execution.\n";
398     exit(1);
399 }
400
401 // --- Compute global volume -----
402
403 GlobalVolume = fabs(XMax[1]-XMin[1]);
404
405 if (Dimension > 1)
406     GlobalVolume *= fabs(XMax[2]-XMin[2]);
407
408 if (Dimension > 2)
409     GlobalVolume *= fabs(XMax[3]-XMin[3]);
410
411 // --- Compute PostProcessing and DataIsBinary -----
412
413 // In 1D, use Gnuplot instead of Data Explorer
414
415 if (Dimension == 1 && PostProcessing == 2)
416     PostProcessing = 1;
417
418 // In 2D-3D, use Data Explorer instead of Gnuplot
419
420 if (Dimension != 1 && PostProcessing == 1)
421     PostProcessing = 2;
422
423 // --- Compute number of conservative quantities -----
424
425 QuantityNb = 9;
426
427 // --- Set the dimension of QuantityMax to QuantityNb -----
428
429 QuantityMax.setDimension(QuantityNb);
430
431 // --- Set the dimension of QuantityAverage to 4 (pressure, vorticity, entropy, volume)
432
433 QuantityAverage.setDimension(4);
434
435 // --- Set the dimension of IntMomentum to dimension -----
436
437 IntMomentum.setDimension(Dimension);
438
439 // --- Compute minimal space step -----
440
441 SpaceStep = fabs(XMax[1]-XMin[1]);
442
443 for (i = 2; i <= Dimension; i++)
444     SpaceStep = Min(SpaceStep, fabs(XMax[i]-XMin[i]));
445
446 SpaceStep /= (1<<ScaleNb);
447
448 // --- Compute time step from CFL if TimeStep = 0 -----
449
450 if (TimeStep == 0.)
451 {
452     if (fabs(Eigenvalue)>0.0e-20)
453         TimeStep = CFL*SpaceStep/Eigenvalue;
454     else
455         TimeStep = 0.0001;
456 }
457 else
458     ConstantTimeStep = true;
459
460 ch = CFL*SpaceStep/TimeStep;
461
462
463 }

```

Here is the caller graph for this function:



### 6.33.3 Variable Documentation

#### 6.33.3.1 int AllTaskScaleNb

Global Scale number

#### 6.33.3.2 real AllXMax[4]

#### 6.33.3.3 real AllXMin[4]

Global domain parameters

#### 6.33.3.4 real Alpha =0.64

Temperature ratio

#### 6.33.3.5 real auxvar =0.

Auxiliar variable

#### 6.33.3.6 real AverageRadius =0.

Average radius of the flame ball

#### 6.33.3.7 char BackupName[255]

#### 6.33.3.8 real BaroclinicEffect =0.

Intensity of the baroclinic effects

#### 6.33.3.9 real Bdivergence =0.

Auxilian divergence variable

#### 6.33.3.10 real CarmenVersion =2.483

Version release

#### 6.33.3.11 int CartDims[3]

#### 6.33.3.12 real Celerity = 1.

Advection-diffusion celerity (0, 1, -1)

#### 6.33.3.13 int CellElementsNb

#### 6.33.3.14 int CellNb =0

Number of cells

6.33.3.15 **real CFL =0.5**

Courant-Friedrich-Levy number

6.33.3.16 **real ch =0.0**

Divergence cleaning ch parameter

6.33.3.17 **real chi =0.**

Artificial diffusion constant

6.33.3.18 **int ChildNb =0**

Number of children for a given parent (equal to 2\*\*Dimension)

6.33.3.19 **real Circulation =10.**

Circulation parameter

6.33.3.20 **int Cluster =1**

0 for local execution, 1 for cluster

6.33.3.21 **int CMax[4] ={3,3,3,3}**

Max. boundary condition (1 = Boundary, 2 = Symetric, 3 = Periodic)

6.33.3.22 **int CMin[4] ={3,3,3,3}**

Min. boundary condition (1 = Boundary, 2 = Symetric, 3 = Periodic)

6.33.3.23 **Timer CommTimer**

Communication timer for performance analyse

6.33.3.24 **bool ComputeCPUTimeRef =false**

True = the reference CPU time is being computed

6.33.3.25 **bool ComputeTemp =false**

6.33.3.26 **real ConstantForce =true**

False = adapt force to maintain constant energy

6.33.3.27 **bool ConstantTimeStep =true**

true = constant TimeStep

6.33.3.28 int Coordinate =1

1 = Cartesian, 2 = Spherical in x

6.33.3.29 int coords[3]

Current CPU coordinates in the virtual CPU processors cart

6.33.3.30 int CPUScales

CPUScales=log2(Number of processors)

6.33.3.31 Timer CPUTime

Timer for CPU time

6.33.3.32 real cr =0.

Alpha parameter divergence cleaning

6.33.3.33 bool CVS =false

True = use Donoho thresholding to perform CVS.

6.33.3.34 bool DatalBinary =true

true = write data in binary format, false = write data in ASCII format

6.33.3.35 bool debug =false

true = check if tree is graded

6.33.3.36 bool Diffusivity =false

True = artificial diffusion. False = no artificial diffusion

6.33.3.37 int Dimension =1

Dimension (1,2,3)

6.33.3.38 real DIVB =0.

Divergence of B

6.33.3.39 real DIVBMax =0.

Maximum divergence of B

6.33.3.40 int DivClean =2

Divergence cleaning: 1-EGLM 2-GLM

6.33.3.41 real Eigenvalue =0.

Maximal eigenvalue

6.33.3.42 real EigenvalueMax =0.

Maximal eigenvalue

6.33.3.43 real EigenvalueX =0.

Eigenvalue at x direction

6.33.3.44 real EigenvalueY =0.

Eigenvalue at y direction

6.33.3.45 real EigenvalueZ =0.

Eigenvalue at z direction

6.33.3.46 real ElapsedTime =0.

ElapsedTime

6.33.3.47 int EquationType =7

Type of equation

6.33.3.48 real ErrorGlobalL2 =0.

Global L2 error on space and time

6.33.3.49 real ErrorGlobalMax =0.

Global error max on space and time

6.33.3.50 real ErrorGlobalMid =0.

Global mean error on space and time

6.33.3.51 int ErrorGlobalNb =0

Number of points for the computation of the global mean error

6.33.3.52 `real ErrorL2 =0.`

L2 error on the grid

6.33.3.53 `real ErrorMax =0.`

Error Max on the grid

6.33.3.54 `real ErrorMid =0.`

Mean error on the grid

6.33.3.55 `int ErrorNb =0`

Number of points for the computation of the mean error

6.33.3.56 `real eta =0.`

Resistivity function

6.33.3.57 `real ExactEnergy =0.`

Global energy for the exact solution (only for EquationType = 1 or 2)

6.33.3.58 `real ExactMomentum =0.`

Global momentum for the exact solution (only for EquationType = 1 or 2)

6.33.3.59 `real ExpectedCompression = 0.`

Expected memory compression

6.33.3.60 `real FlameVelocity =0.`

Flame velocity

6.33.3.61 `bool FluxCorrection =true`

true = conservative flux correction

6.33.3.62 `real ForceX =0.`

6.33.3.63 `real ForceY =0.`

6.33.3.64 `real ForceZ =0.`

6.33.3.65 `real Fr =0.`

Froude number

6.33.3.66 `double FVTimeRef =0.`

FV reference CPU time for 1 iteration

6.33.3.67 `real Gamma =1.4`

Adiabatic function

6.33.3.68 `real GlobalEnergy =0.`

Global energy (only for EquationType = 1 or 2)

6.33.3.69 `real GlobalEnstrophy =0.`

Global enstrophy (only for EquationType = 6)

6.33.3.70 `FILE* GlobalFile`

Global file

6.33.3.71 `real GlobalMomentum =0.`

Global momentum (only for EquationType = 1 or 2)

6.33.3.72 `real GlobalMomentumOld =0.`

Old global momentum (only for EquationType = 6)

6.33.3.73 `real GlobalReactionRate =0.`

Global reaction rate

6.33.3.74 `real GlobalVolume =0.`

Global volume

6.33.3.75 `real Helicity =0.`

Time derivative of helicity (must be zero)

6.33.3.76 `int IcNb = 0`

Initial condition suavization

6.33.3.77 `int ImageNb =0`

Print ImageNb images



6.33.3.78 `real IntDensity =0.`

Integral of the density

6.33.3.79 `real IntEnergy =0.`

Integral of the energy

6.33.3.80 `Vector IntMomentum`

Integral of the modulus of the momentum

6.33.3.81 `real IntVorticity =0.`

Integral of the modulus of the vorticity

6.33.3.82 `int IterationNb =0`

Number of iterations

6.33.3.83 `int IterationNbRef =1000`

Number of iterations for the FV reference computation

6.33.3.84 `int IterationNo =0`

Current iteration number

6.33.3.85 `real Le =1.`

Lewis number

6.33.3.86 `int LeafNb =0`

Number of leaves

6.33.3.87 `bool LES =false`

True = use eddy-viscosity.

6.33.3.88 `int LimiterNo =5`

Limiter number

6.33.3.89 `real Ma =0.3`

Mach number

6.33.3.90 int MaxCellElementsNb

6.33.3.91 real ModelConstant =0.1

Constant used in the turbulence model

6.33.3.92 int MPIRecvType

6.33.3.93 int MPISendType

Type of calling MPI communication functions. See [Parameters.cpp](#) for the more information

6.33.3.94 bool Multiresolution =true

true = Multiresolution, false = FV on fine mesh

6.33.3.95 int NeighbourNb

Important parameter: The deep of the inter-CPU domain overlapping

6.33.3.96 int one\_D

6.33.3.97 real PenalizeFactor =1.

Factor of penalization (obsolete)

6.33.3.98 real PhysicalTime =0.

Physical elapsed time

6.33.3.99 real pi =acos(-1.)

Pi constant

6.33.3.100 int PostProcessing =2

1 = Gnuplot, 2 = Data Explorer, 3 = Tecplot

6.33.3.101 real Pr =0.71

Prandtl number

6.33.3.102 real PreviousAverageRadius =0.

Previous average radius

6.33.3.103 real PreviousAverageRadius2 =0.

Previous average radius

6.33.3.104 int PrintEvery =0

Print every PrintEvery iteration

6.33.3.105 int PrintIt1 =0

Iteration for print 1

6.33.3.106 int PrintIt2 =0

Iteration for print 2

6.33.3.107 int PrintIt3 =0

Iteration for print 3

6.33.3.108 int PrintIt4 =0

Iteration for print 4

6.33.3.109 int PrintIt5 =0

Iteration for print 5

6.33.3.110 int PrintIt6 =0

Iteration for print 6

6.33.3.111 int PrintMoreScales =0

More scales to print

6.33.3.112 real PrintTime1 =0.

Time for print 1

6.33.3.113 real PrintTime2 =0.

Time for print 2

6.33.3.114 real PrintTime3 =0.

Time for print 3

6.33.3.115 real PrintTime4 =0.

Time for print 4

6.33.3.116 **real** PrintTime5 =0.

Time for print 5

6.33.3.117 **real** PrintTime6 =0.

Time for print 6

6.33.3.118 **real** PsiGrad =0.

Psi gradient

6.33.3.119 **Vector** QuantityAverage

**Vector** containing the average quantities

6.33.3.120 **Vector** QuantityMax

**Vector** containing the maximal quantities

6.33.3.121 **int** QuantityNb =2

Number of conservative quantities

6.33.3.122 **int** rank

Current CPU

6.33.3.123 **int** rank\_il

axis X, direction - low

6.33.3.124 **int** rank\_iu

axis X, direction - high

6.33.3.125 **int** rank\_jl

axis Y, direction - low

6.33.3.126 **int** rank\_ju

6.33.3.127 **int** rank\_kl

6.33.3.128 **int** rank\_ku

6.33.3.129 **real** Re =100.

Reynolds number

6.33.3.130 `real ReactionRateMax =0.`

Maximum of the reaction rate

6.33.3.131 `bool Recovery =false`

true = restore data from previous computation

6.33.3.132 `int Refresh =0`

/ Refresh rate

6.33.3.133 `int RefreshNb =200`

Number of refreshments

6.33.3.134 `bool Resistivity =false`

True = resistive model. False = ideal model

6.33.3.135 `real RKFAccuracyFactor =1.E-03`

Desired value of RKFEError (only if TimeAdaptivity=true)

6.33.3.136 `real RKFEError =0.`

Maximum of the relative errors between RK2 and RK3 (only if TimeAdaptivity=true)

6.33.3.137 `real RKFSafetyFactor =0.01`

Safety factor for the computation of the time step (only if TimeAdaptivity=true)

6.33.3.138 `int ScalarEqNb =0`

Number of additional scalar equations

6.33.3.139 `int ScaleNb =5`

Maximal number of scales allowed

6.33.3.140 `int ScaleNbRef =2`

Number of scales for the FV reference computation

6.33.3.141 `int SchemeNb =1`

Scheme number

6.33.3.142 int SendD = 1 << 0

6.33.3.143 int SenddX = 1 << 5

6.33.3.144 int SendGrad = 1 << 1

6.33.3.145 int SendQ = 1 << 2

6.33.3.146 int SendQs = 1 << 3

6.33.3.147 int SendX = 1 << 4

6.33.3.148 real Sigma =5.E-02

Radiation coefficient

6.33.3.149 int size

Number of processors

6.33.3.150 real SmoothCoeff =0.

Smoothing coefficient

6.33.3.151 real SpaceStep =0.

Space step

6.33.3.152 real StartTimeAveraging =0.

Time where the time-averaging procedure must start

6.33.3.153 int StepNb =2

Number of steps for the time integration

6.33.3.154 int StepNo =1

Current step number for the time integration

6.33.3.155 real ThermalConduction =0.

Dimensionless thermal conduction

6.33.3.156 int ThresholdNorm =0

Normalization of the wavelet basis for the threshold

6.33.3.157 bool TimeAdaptivity =false

true = use time adaptivity (only when Multiresolution = true, dummy else)

6.33.3.158 `int TimeAdaptivityFactor = 1`

The factor of time step between two scales is  $2^{\text{TimeAdaptivityFactor}}$

6.33.3.159 `bool TimeAveraging = false`

true = use a time-averaging grid (only for turbulence)

6.33.3.160 `real TimeStep = 0.`

Time step

6.33.3.161 `real Tolerance = 0.`

Prescribed tolerance

6.33.3.162 `real ToleranceScale = 1.`

Scale factor for tolerance (when Tolerance = 0).

6.33.3.163 `real TotalCellNb = 0.`

Total number of cells for all iterations

6.33.3.164 `real TotalLeafNb = 0.`

Total number of leaves for all iterations

6.33.3.165 `real TRef = 273.`

Reference temperature for Sutherland's law

6.33.3.166 `int two_D`

6.33.3.167 `bool UseBackup = true`

true = use Backup procedure.

6.33.3.168 `bool UseBoundaryRegions = false`

true = use file carmen.bc

6.33.3.169 `real Viscosity = 1.`

0 or 1. 0 means no viscosity.

6.33.3.170 `int WhatSend`

6.33.3.171 `bool WriteAsPoints =false`

true = write data as point-values, false = write data as cell-averages

6.33.3.172 `real XCenter[4] ={0.,0.,0.,0.}`

Coordinate of the center of the flame ball

6.33.3.173 `real XMax[4] ={1.,1.,1.,1.}`

Maximal values of coordinates;

6.33.3.174 `real XMin[4] ={-1.,-1.,-1.,-1.}`

Minimal values of coordinates;

6.33.3.175 `real Ze =10.`

Equivalent to Beta

6.33.3.176 `bool ZipData =true`

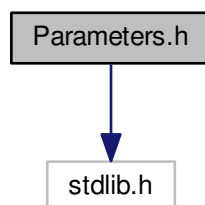
true = zip data files

## 6.34 Parameters.h File Reference

This header contains all parameters as global variables.

```
#include <stdlib.h>
```

Include dependency graph for Parameters.h:



This graph shows which files directly or indirectly include this file:





## Variables

- int Cluster
- int StepNb
- int StepNo
- int IterationNb
- int IterationNbRef
- int IterationNo
- real CFL
- real TimeStep
- real PhysicalTime
- Timer CPUTime
- double FVTimeRef
- int Refresh
- int RefreshNb
- real PrintTime1
- real PrintTime2
- real PrintTime3
- real PrintTime4
- real PrintTime5
- real PrintTime6
- int PrintIt1
- int PrintIt2
- int PrintIt3
- int PrintIt4
- int PrintIt5
- int PrintIt6
- int PrintEvery
- int ImageNb
- bool TimeAveraging
- real StartTimeAveraging
- bool Recovery
- bool UseBackup
- bool ConstantTimeStep
- real ElapsedTime
- real SpaceStep
- bool ComputeCPUTimeRef
- int SchemeNb
- int IcNb
- int LimiterNo
- int QuantityNb
- int EquationType
- int ScalarEqNb
- int DivClean
- int Dimension
- int ChildNb
- int Coordinate
- real XMin [4]
- real XMax [4]
- int CMin [4]
- int CMax [4]
- bool UseBoundaryRegions
- bool Multiresolution
- bool TimeAdaptivity
- int TimeAdaptivityFactor

- int CellNb
- int LeafNb
- real TotalCellNb
- real TotalLeafNb
- int ScaleNb
- int ScaleNbRef
- int PrintMoreScales
- real Tolerance
- real ToleranceScale
- real ToleranceInit
- real PenalizeFactor
- real ExpectedCompression
- bool CVS
- bool LES
- int ThresholdNorm
- real SmoothCoeff
- real eta
- real Gamma
- bool Resistivity
- bool Diffusivity
- real chi
- real Celerity
- real Viscosity
- real Re
- real Pr
- real Ma
- real Fr
- real TRef
- real ModelConstant
- real ThermalConduction
- real ConstantForce
- real Circulation
- bool ComputeTemp
- real Alpha
- real Ze
- real Le
- real Sigma
- real ErrorMax
- real ErrorGlobalMax
- real ErrorMid
- real ErrorGlobalMid
- real ErrorL2
- real ErrorGlobalL2
- int ErrorNb
- int ErrorGlobalNb
- real cr
- real Helicity
- real GlobalMomentum
- real GlobalEnergy
- real GlobalEnstrophy
- real ExactMomentum
- real ExactEnergy
- real GlobalMomentumOld
- real GlobalVolume
- real EigenvalueZ

- real EigenvalueY
- real EigenvalueX
- real Eigenvalue
- real EigenvalueMax
- Vector QuantityMax
- Vector QuantityAverage
- real RKFEError
- real RKFAccuracyFactor
- real RKFSafetyFactor
- real FlameVelocity
- real GlobalReactionRate
- real AverageRadius
- real PreviousAverageRadius
- real PreviousAverageRadius2
- real XCenter [4]
- real ReactionRateMax
- real IntVorticity
- real IntDensity
- Vector IntMomentum
- real IntEnergy
- real BaroclinicEffect
- real DIVB
- real DIVBMax
- real Bdivergence
- real PsiGrad
- real ch
- real auxvar
- real pi
- bool debug
- bool FluxCorrection
- real CarmenVersion
- FILE \* GlobalFile
- int PostProcessing
- bool DatsBinary
- bool ZipData
- bool WriteAsPoints
- int AllTaskScaleNb
- int CPUScales
- int size
- int rank
- real AllXMin [4]
- real AllXMax [4]
- int NeighbourNb
- int coords [3]
- int CartDims [3]
- int CellElementsNb
- int MaxCellElementsNb
- int one\_D
- int two\_D
- int MPISendType
- int MPIRecvType
- Timer CommTimer
- int rank\_il
- int rank\_iu
- int rank\_jl

- int [rank\\_ju](#)
- int [rank\\_kl](#)
- int [rank\\_ku](#)
- int [WhatSend](#)
- int [SendD](#)
- int [SendGrad](#)
- int [SendQ](#)
- int [SendQs](#)
- int [SendX](#)
- int [SenddX](#)
- char [BackupName](#) [255]

### 6.34.1 Detailed Description

This header contains all parameters as global variables.

### 6.34.2 Variable Documentation

#### 6.34.2.1 int AllTaskScaleNb

Global Scale number

#### 6.34.2.2 real AllXMax[4]

#### 6.34.2.3 real AllXMin[4]

Global domain parameters

#### 6.34.2.4 real Alpha

Temperature ratio

#### 6.34.2.5 real auxvar

Auxiliar variable

#### 6.34.2.6 real AverageRadius

Average radius of the flame ball

#### 6.34.2.7 char BackupName[255]

#### 6.34.2.8 real BaroclinicEffect

Intensity of the baroclinic effects

#### 6.34.2.9 real Bdivergence

Auxilian divergence variable

6.34.2.10 real CarmenVersion

Version release

6.34.2.11 int CartDims[3]

6.34.2.12 real Celerity

Advection-diffusion celerity (0, 1, -1)

6.34.2.13 int CellElementsNb

6.34.2.14 int CellNb

Number of cells

6.34.2.15 real CFL

Courant-Friedrich-Levy number

6.34.2.16 real ch

Divergence cleaning ch parameter

6.34.2.17 real chi

Artificial diffusion constant

6.34.2.18 int ChildNb

Number of children for a given parent (equal to 2\*\*Dimension)

6.34.2.19 real Circulation

Circulation parameter

6.34.2.20 int Cluster

0 for local execution, 1 for cluster

6.34.2.21 int CMax[4]

Max. boundary condition (1 = Boundary, 2 = Symetric, 3 = Periodic)

6.34.2.22 int CMin[4]

Min. boundary condition (1 = Boundary, 2 = Symetric, 3 = Periodic)

**6.34.2.23 Timer CommTimer**

Communication timer for performance analyse

**6.34.2.24 bool ComputeCPUTimeRef**

True = the reference CPU time is being computed

**6.34.2.25 bool ComputeTemp****6.34.2.26 real ConstantForce**

False = adapt force to maintain constant energy

**6.34.2.27 bool ConstantTimeStep**

true = constant TimeStep

**6.34.2.28 int Coordinate**

1 = Cartesian, 2 = Spherical in x

**6.34.2.29 int coords[3]**

Current CPU coordinates in the virtual CPU processors cart

**6.34.2.30 int CPUScales**

$CPUScales = \log_2(\text{Number of processors})$

**6.34.2.31 Timer CPUTime**

[Timer](#) for CPU time

**6.34.2.32 real cr**

Alpha parameter divergence cleaning

**6.34.2.33 bool CVS**

True = use Donoho thresholding to perform CVS.

**6.34.2.34 bool DatalsBinary**

true = write data in binary format, false = write data in ASCII format

**6.34.2.35 bool debug**

true = check if tree is graded

**6.34.2.36 bool Diffusivity**

True = artificial diffusion. False = no artificial diffusion

**6.34.2.37 int Dimension**

Dimension (1,2,3)

**6.34.2.38 real DIVB**

Divergence of B

**6.34.2.39 real DIVBMax**

Maximum divergence of B

**6.34.2.40 int DivClean**

Divergence cleaning: 1-EGLM 2-GLM

**6.34.2.41 real Eigenvalue**

Maximal eigenvalue

**6.34.2.42 real EigenvalueMax**

Maximal eigenvalue

**6.34.2.43 real EigenvalueX**

Eigenvalue at x direction

**6.34.2.44 real EigenvalueY**

Eigenvalue at y direction

**6.34.2.45 real EigenvalueZ**

Eigenvalue at z direction

**6.34.2.46 real ElapsedTime**

ElapsedTime

**6.34.2.47 int EquationType**

Type of equation

**6.34.2.48 real ErrorGlobalL2**

Global L2 error on space and time

**6.34.2.49 real ErrorGlobalMax**

Global error max on space and time

**6.34.2.50 real ErrorGlobalMid**

Global mean error on space and time

**6.34.2.51 int ErrorGlobalNb**

Number of points for the computation of the global mean error

**6.34.2.52 real ErrorL2**

L2 error on the grid

**6.34.2.53 real ErrorMax**

Error Max on the grid

**6.34.2.54 real ErrorMid**

Mean error on the grid

**6.34.2.55 int ErrorNb**

Number of points for the computation of the mean error

**6.34.2.56 real eta**

Resistivity function

**6.34.2.57 real ExactEnergy**

Global energy for the exact solution (only for EquationType = 1 or 2)

**6.34.2.58 real ExactMomentum**

Global momentum for the exact solution (only for EquationType = 1 or 2)

**6.34.2.59 real ExpectedCompression**

Expected memory compression



**6.34.2.60 real FlameVelocity**

Flame velocity

**6.34.2.61 bool FluxCorrection**

true = conservative flux correction

**6.34.2.62 real Fr**

Froude number

**6.34.2.63 double FVTimeRef**

FV reference CPU time for 1 iteration

**6.34.2.64 real Gamma**

Adiabatic function

**6.34.2.65 real GlobalEnergy**

Global energy (only for EquationType = 1 or 2)

**6.34.2.66 real GlobalEnstrophy**

Global enstrophy (only for EquationType = 6)

**6.34.2.67 FILE\* GlobalFile**

Global file

**6.34.2.68 real GlobalMomentum**

Global momentum (only for EquationType = 1 or 2)

**6.34.2.69 real GlobalMomentumOld**

Old global momentum (only for EquationType = 6)

**6.34.2.70 real GlobalReactionRate**

Global reaction rate

**6.34.2.71 real GlobalVolume**

Global volume

**6.34.2.72 real Helicity**

Time derivative of helicity (must be zero)

**6.34.2.73 int IcNb**

Initial condition suavization

**6.34.2.74 int ImageNb**

Print ImageNb images

**6.34.2.75 real IntDensity**

Integral of the density

**6.34.2.76 real IntEnergy**

Integral of the energy

**6.34.2.77 Vector IntMomentum**

Integral of the modulus of the momentum

**6.34.2.78 real IntVorticity**

Integral of the modulus of the vorticity

**6.34.2.79 int IterationNb**

Number of iterations

**6.34.2.80 int IterationNbRef**

Number of iterations for the FV reference computation

**6.34.2.81 int IterationNo**

Current iteration number

**6.34.2.82 real Le**

Lewis number

**6.34.2.83 int LeafNb**

Number of leaves

**6.34.2.84 bool LES**

True = use eddy-viscosity.

**6.34.2.85 int LimiterNo**

Limiter number

**6.34.2.86 real Ma**

Mach number

**6.34.2.87 int MaxCellElementsNb****6.34.2.88 real ModelConstant**

Constant used in the turbulence model

**6.34.2.89 int MPIRecvType****6.34.2.90 int MPISendType**

Type of calling MPI communication functions. See [Parameters.cpp](#) for the more information

**6.34.2.91 bool Multiresolution**

true = Multiresolution, false = FV on fine mesh

**6.34.2.92 int NeighbourNb**

Important parameter: The deep of the inter-CPU domain overlapping

**6.34.2.93 int one\_D****6.34.2.94 real PenalizeFactor**

Factor of penalization (obsolete)

**6.34.2.95 real PhysicalTime**

Physical elapsed time

**6.34.2.96 real pi**

Pi constant

**6.34.2.97 int PostProcessing**

1 = Gnuplot, 2 = Data Explorer, 3 = Tecplot

6.34.2.98 **real** Pr

Prandtl number

6.34.2.99 **real** PreviousAverageRadius

Previous average radius

6.34.2.100 **real** PreviousAverageRadius2

Previous average radius

6.34.2.101 **int** PrintEvery

Print every PrintEvery iteration

6.34.2.102 **int** PrintIt1

Iteration for print 1

6.34.2.103 **int** PrintIt2

Iteration for print 2

6.34.2.104 **int** PrintIt3

Iteration for print 3

6.34.2.105 **int** PrintIt4

Iteration for print 4

6.34.2.106 **int** PrintIt5

Iteration for print 5

6.34.2.107 **int** PrintIt6

Iteration for print 6

6.34.2.108 **int** PrintMoreScales

More scales to print

6.34.2.109 **real** PrintTime1

Time for print 1

6.34.2.110 **real** PrintTime2

Time for print 2

6.34.2.111 **real** PrintTime3

Time for print 3

6.34.2.112 **real** PrintTime4

Time for print 4

6.34.2.113 **real** PrintTime5

Time for print 5

6.34.2.114 **real** PrintTime6

Time for print 6

6.34.2.115 **real** PsiGrad

Psi gradient

6.34.2.116 **Vector** QuantityAverage

**Vector** containing the average quantities

6.34.2.117 **Vector** QuantityMax

**Vector** containing the maximal quantities

6.34.2.118 **int** QuantityNb

Number of conservative quantites

6.34.2.119 **int** rank

Current CPU

6.34.2.120 **int** rank\_il

axis X, direction - low

6.34.2.121 **int** rank\_iu

axis X, direction - high

6.34.2.122 int rank\_jl

axis Y, direction - low

6.34.2.123 int rank\_ju

6.34.2.124 int rank\_kl

6.34.2.125 int rank\_ku

6.34.2.126 real Re

Reynolds number

6.34.2.127 real ReactionRateMax

Maximum of the reaction rate

6.34.2.128 bool Recovery

true = restore data from previous computation

6.34.2.129 int Refresh

/ Refresh rate

6.34.2.130 int RefreshNb

Number of refreshments

6.34.2.131 bool Resistivity

True = resistive model. False = ideal model

6.34.2.132 real RKFAccuracyFactor

Desired value of RKFEError (only if TimeAdaptivity=true)

6.34.2.133 real RKFEError

Maximum of the relative errors between RK2 and RK3 (only if TimeAdaptivity=true)

6.34.2.134 real RKFSafetyFactor

Safety factor for the computation of the time step (only if TimeAdaptivity=true)

6.34.2.135 int ScalarEqNb

Number of additional scalar equations

6.34.2.136 int ScaleNb

Maximal number of scales allowed

6.34.2.137 int ScaleNbRef

Number of scales for the FV reference computation

6.34.2.138 int SchemeNb

Scheme number

6.34.2.139 int SendD

6.34.2.140 int SenddX

6.34.2.141 int SendGrad

6.34.2.142 int SendQ

6.34.2.143 int SendQs

6.34.2.144 int SendX

6.34.2.145 real Sigma

Radiation coefficient

6.34.2.146 int size

Number of processors

6.34.2.147 real SmoothCoeff

Smoothing coefficient

6.34.2.148 real SpaceStep

Space step

6.34.2.149 real StartTimeAveraging

Time where the time-averaging procedure must start

6.34.2.150 int StepNb

Number of steps for the time integration

6.34.2.151 int StepNo

Current step number for the time integration

**6.34.2.152 real ThermalConduction**

Dimensionless thermal conduction

**6.34.2.153 int ThresholdNorm**

Normalization of the wavelet basis for the threshold

**6.34.2.154 bool TimeAdaptivity**

true = use time adaptivity (only when Multiresolution = true, dummy else)

**6.34.2.155 int TimeAdaptivityFactor**

The factor of time step between two scales is  $2^{\text{TimeAdaptivityFactor}}$

**6.34.2.156 bool TimeAveraging**

true = use a time-averaging grid (only for turbulence)

**6.34.2.157 real TimeStep**

Time step

**6.34.2.158 real Tolerance**

Prescribed tolerance

**6.34.2.159 real ToleranceInit**

Prescribed tolerance for initial condition (obsolete)

**6.34.2.160 real ToleranceScale**

Scale factor for tolerance (when Tolerance = 0).

**6.34.2.161 real TotalCellNb**

Total number of cells for all iterations

**6.34.2.162 real TotalLeafNb**

Total number of leaves for all iterations

**6.34.2.163 real TRef**

Reference temperature for Sutherland's law



6.34.2.164 `int two_D`

6.34.2.165 `bool UseBackup`

true = use Backup procedure.

6.34.2.166 `bool UseBoundaryRegions`

true = use file carmen.bc

6.34.2.167 `real Viscosity`

0 or 1. 0 means no viscosity.

6.34.2.168 `int WhatSend`

6.34.2.169 `bool WriteAsPoints`

true = write data as point-values, false = write data as cell-averages

6.34.2.170 `real XCenter[4]`

Coordinate of the center of the flame ball

6.34.2.171 `real XMax[4]`

Maximal values of coordinates;

6.34.2.172 `real XMin[4]`

Minimal values of coordinates;

6.34.2.173 `real Ze`

Equivalent to Beta

6.34.2.174 `bool ZipData`

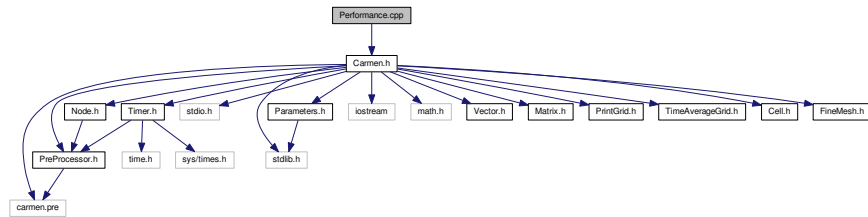
true = zip data files

## 6.35 Performance.cpp File Reference

Simulation information.

```
#include "Carmen.h"
```

Include dependency graph for Performance.cpp:



## Functions

- void [Performance](#) (const char \*FileName)  
*Computes the performance of the computation and, for cluster computations, write it into file FileName.*

### 6.35.1 Detailed Description

Simulation information.

### 6.35.2 Function Documentation

#### 6.35.2.1 void Performance ( const char \* FileName )

Computes the performance of the computation and, for cluster computations, write it into file *FileName*.

#### Parameters

| <i>FileName</i> | Name of the file. |
|-----------------|-------------------|
|-----------------|-------------------|

#### Returns

void

```

23 {
24     // --- Local variables ---
25
26     bool EndComputation;           // True if end computation
27     int FineCellNb;               // Number of cells on fine grid
28     int CellPVirt;
29     FILE *output;                 // Pointer to output file
30
31     double realtimefull;          //full real time
32     double ftime;                 // real time
33     double ctime;                 // CPU time
34     unsigned int ttime, rtime;    // total and remaining real time (in seconds)
35     unsigned int tctime=0, rctime=0; // total and remaining CPU time (in seconds)
36     unsigned int rest;
37     int day, hour, min, sec;
38
39     // --- Init EndComputation
40
41     EndComputation = (IterationNo > IterationNb);
42
43     // --- Compute FineCellNb ---
44
45     FineCellNb = 1<<(ScaleNb*Dimension);
46     CellPVirt = 1<<(ScaleNb*(Dimension-1));
47     CellPVirt = CellPVirt*2*Dimension;
48     // --- Write in file ---
49
50 /*
51     char CPUFileName[255];
  
```

```

52 #if defined PARMPI
53     sprintf(CPUFileName, "%d_%d_%d_%s", coords[0], coords[1], coords[2], FileName);
54 // strcpy(CPUFileName, FileName);
55 #else
56     strcpy(CPUFileName, FileName);
57 #endif
58 */
59
60     if ((output = fopen(FileName, "w")))
61     {
62
63         realtimefull=ftime = CPUTime.realTime();
64         ctime = CPUTime.CPUTime();
65
66         if (!EndComputation)
67         {
68             ttime = (unsigned int)((ftime*IterationNb)/IterationNo);
69             rtime = (unsigned int)((ftime*(IterationNb-IterationNo))/
IterationNo);
70             tctime = (unsigned int)((ctime*IterationNb)/IterationNo);
71             rctime = (unsigned int)((ctime*(IterationNb-IterationNo))/
IterationNo);
72         }
73
74         fprintf(output, "Dimension           : %12i\n", Dimension);
75
76         if (EndComputation)
77             fprintf(output, "Iterations           : %12i\n", IterationNb);
78         else
79         {
80             fprintf(output, "Iterations (total)   : %12i\n", IterationNb);
81             fprintf(output, "Iterations (elapsed) : %12i\n", IterationNo);
82             fprintf(output, "In progress         : %13.6f %%\n", 100.*
IterationNo/(1.*IterationNb));
83         }
84
85         fprintf(output, "Scales (max)        : %12i\n", ScaleNb);
86         fprintf(output, "Cells (max)         : %12i\n", (1<<(ScaleNb*Dimension)));
87
88         if (Multiresolution)
89             fprintf(output, "Solver              :          MR\n");
90         else
91             fprintf(output, "Solver              :          FV\n");
92
93         //fprintf(output, "Time integration    :          explicit\n");
94         fprintf(output, "Time accuracy order : %12i\n", StepNb);
95         fprintf(output, "Time step           : %13.6e s\n", TimeStep);
96         fprintf(output, "Threshold parameter : %13.6e \n", Tolerance);
97         fprintf(output, "Threshold norm      : %12i\n", ThresholdNorm);
98         fprintf(output, "CFL                 : %13.6e \n", CFL);
99
100        if(Resistivity)
101            fprintf(output, "Eta                 : %13.6e \n", eta);
102        if(Diffusivity)
103            fprintf(output, "Chi                 : %13.6e \n", chi);
104
105        if (EndComputation)
106        {
107            fprintf(output, "Physical time       : %13.6e s\n", ElapsedTime); //TimeStep *
IterationNb);
108            fprintf(output, "CPU time (s)       : %13.6e s\n", ctime);
109
110            if (Multiresolution)
111                fprintf(output, "CPU time / it. x pt : %13.6e s\n", ctime/
TotalLeafNb);
112            else
113                fprintf(output, "CPU time / it. x pt : %13.6e s\n", ctime/((1<<(
ScaleNb*Dimension))*IterationNb));
114
115            if (Multiresolution)
116            {
117
118                fprintf(output, "Leaf compression   : %13.6f %% \n", (100.*
TotalLeafNb)/(1.0*IterationNb*FineCellNb));
119                //fprintf(output, "Memory compression : %13.6f %% \n",
(100.*TotalCellNb)/(1.0*IterationNb*(FineCellNb)));
120                fprintf(output, "Memory compression : %13.6f %% \n", (100.*
TotalCellNb)/(1.0*IterationNb*(FineCellNb + CellPVirt)));
121                fprintf(output, "CPU compression    : %13.6f %% \n", (100.*ctime)/(IterationNb*
FVTimeRef));
122            }
123            else
124            {
125                fprintf(output, "Leaf compression   : %13.6f %% \n", 100.);
126                fprintf(output, "Memory compression : %13.6f %% \n", 100.);
127                fprintf(output, "CPU compression    : %13.6f %% \n", 100.);
128            }

```

```

129     }
130     else
131     {
132         fprintf(output, "Total physical time   :%13.6e s\n", TimeStep * IterationNb);
133         fprintf(output, "Elapsed physical time :%13.6e s\n", TimeStep *
IterationNb);
134         if (Multiresolution)
135         {
136             fprintf(output, "Leaf compression   :%13.6f %% \n", (100.*
TotalLeafNb)/(1.0*IterationNb*FineCellNb));
137             fprintf(output, "Memory compression  :%13.6f %% \n", (100.*
TotalCellNb)/(1.0*IterationNb*FineCellNb));
138             fprintf(output, "CPU compression    :%13.6f %% \n", (100.*ctime)/(
IterationNb*FVTimeRef));
139         }
140         else
141         {
142             fprintf(output, "Leaf compression   :%13.6f %% \n", 100.);
143             fprintf(output, "Memory compression  :%13.6f %% \n", 100.);
144             fprintf(output, "CPU compression    :%13.6f %% \n", 100.);
145         }
146     }
147     fprintf(output, "\n");
148
149     if (EndComputation)
150     {
151         // --- Print final time -----
152
153         rest = (unsigned int) (ctime);
154         day  = rest/86400;
155         rest %= 86400;
156         hour = rest/3600;
157         rest %= 3600;
158         min  = rest/60;
159         rest %= 60;
160         sec  = rest;
161         rest = (unsigned int) (ctime);
162
163         if (rest >= 86400)
164             fprintf(output, "CPU time : %5d day %2d h %2d min %2d s\n", day, hour, min, sec);
165
166         if ((rest < 86400)&&(rest >= 3600))
167             fprintf(output, "CPU time : %2d h %2d min %2d s\n", hour, min, sec);
168
169         if ((rest < 3600)&&(rest >= 60))
170             fprintf(output, "CPU time : %2d min %2d s\n", min, sec);
171
172         if (rest < 60)
173             fprintf(output, "CPU time : %2d s\n", sec);
174     }
175     else
176     {
177         // --- Print total time -----
178
179         rest = tctime;
180         day  = rest/86400;
181         rest %= 86400;
182         hour = rest/3600;
183         rest %= 3600;
184         min  = rest/60;
185         rest %= 60;
186         sec  = rest;
187
188         if (tctime >= 86400)
189             fprintf(output, "Total CPU time (estimation) : %5d day %2d h %2d min %2d s\n", day,
hour, min, sec);
190
191         if ((tctime < 86400)&&(tctime >= 3600))
192             fprintf(output, "Total CPU time (estimation) : %2d h %2d min %2d s\n", hour, min, sec);
193
194         if ((tctime < 3600)&&(tctime >= 60))
195             fprintf(output, "Total CPU time (estimation) : %2d min %2d s\n", min, sec);
196
197         if (tctime < 60)
198             fprintf(output, "Total CPU time (estimation) : %2d s\n", sec);
199
200         // --- Print remaining time -----
201
202         rest = rctime;
203         day  = rest/86400;
204         rest %= 86400;
205         hour = rest/3600;
206         rest %= 3600;
207         min  = rest/60;
208         rest %= 60;
209         sec  = rest;
210

```

```

211
212     if (rctime >= 86400)
213         fprintf(output, "Remaining CPU time (estimation) : %5d day %2d h %2d min %2d s\n", day,
hour, min, sec);
214
215     if ((rctime < 86400)&&(rctime >= 3600))
216         fprintf(output, "Remaining CPU time (estimation) : %2d h %2d min %2d s\n", hour, min, sec);
217
218     if ((rctime < 3600)&&(rctime >= 60))
219         fprintf(output, "Remaining CPU time (estimation) : %2d min %2d s\n", min, sec);
220
221     if (rctime < 60)
222         fprintf(output, "Remaining CPU time (estimation) : %2d s\n", sec);
223
224 }
225
226
227 #if defined PARMPI
228     fprintf(output, "\n");
229     fprintf(output, "Real time (time() function) :%lf\n", realtimefull);
230     fprintf(output, "clock() function :%lf\n", ctime);
231     fprintf(output, "\nCommunications real timer: %lf\n", CommTimer.
realTime());
232     fprintf(output, "Communications clock():%lf\n", CommTimer.
CPUTime());
233 #endif
234
235     fprintf(output, "\n");
236     fclose (output);
237 }
238 else
239 {
240     cout << "Performance.cpp: In method `void Performance(Node*, char*)':\n";
241     cout << "Performance.cpp: cannot open file " << FileName << '\n';
242     cout << "carmen: *** [Performance.o] Execution error\n";
243     cout << "carmen: abort execution.\n";
244     exit(1);
245 }
246 }

```

Here is the caller graph for this function:

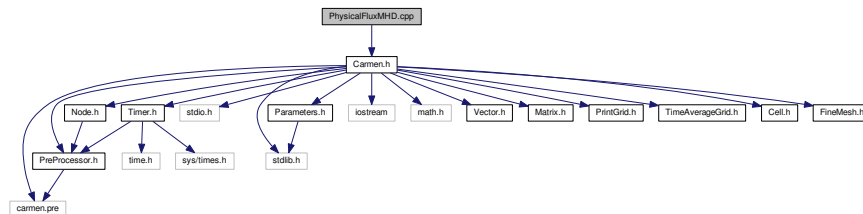


## 6.36 PhysicalFluxMHD.cpp File Reference

Computes the MHD physical flux.

#include "Carmen.h"

Include dependency graph for PhysicalFluxMHD.cpp:



## Functions

- **Vector FluxX** (const **Vector** &Avg)  
*Returns the physical flux of MHD equations in X direction.*
- **Vector FluxY** (const **Vector** &Avg)  
*Returns the physical flux of MHD equations in Y direction.*
- **Vector FluxZ** (const **Vector** &Avg)  
*Returns the physical flux of MHD equations in Z direction.*

### 6.36.1 Detailed Description

Computes the MHD physical flux.

#### Author

Anna Karina Fontes Gomes

#### Version

2.0

### 6.36.2 Function Documentation

#### 6.36.2.1 Vector FluxX ( const Vector & Avg )

Returns the physical flux of MHD equations in X direction.

#### Parameters

|            |                 |
|------------|-----------------|
| <i>Avg</i> | Average vector. |
|------------|-----------------|

#### Returns

**Vector**

```

9 {
10   real rho;
11   real vx, vy, vz;
12   real pre, e;
13   real Bx, By, Bz;
14   real Bx2, By2, Bz2, B2;
15   real vx2, vy2, vz2, v2;
16   real half = 0.5;
17   Vector F(QuantityNb);
18
19   //Variables
20   rho = Avg.value(1);
21   vx = Avg.value(2)/rho;
22   vy = Avg.value(3)/rho;
23   vz = Avg.value(4)/rho;
24   e = Avg.value(5);
25   Bx = Avg.value(7);
26   By = Avg.value(8);
27   Bz = Avg.value(9);
28
29   Bx2 = Bx*Bx;
30   By2 = By*By;
31   Bz2 = Bz*Bz;
32   B2 = half*(Bz2+Bx2+By2);
33
34   vx2 = vx*vx;
35   vy2 = vy*vy;
36   vz2 = vz*vz;
37   v2 = half*(vz2+vx2+vy2);
38
39   //pressure
40   pre = (Gamma -1.)*(e - rho*v2 - B2);
41

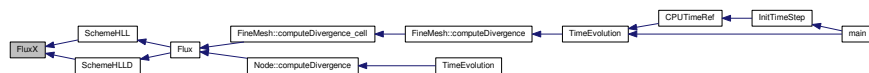
```

```

42 //Physical flux - x-direction
43 F.setValue(1,rho*vx);
44 F.setValue(2,rho*vx2 + pre + half*(Bz2+By2-Bx2));
45 F.setValue(3,rho*vx*vy - Bx*By);
46 F.setValue(4,rho*vx*vz - Bx*Bz);
47 F.setValue(5,(e + pre + B2)*vx - Bx*(vx*Bx + vy*By + vz*Bz) );
48 F.setValue(6,0.0);
49 F.setValue(7,0.0);
50 F.setValue(8,vx*By - vy*Bx);
51 F.setValue(9,vx*Bz - vz*Bx);
52
53
54 return F;
55 }

```

Here is the caller graph for this function:



### 6.36.2.2 Vector FluxY ( const Vector & Avg )

Returns the physical flux of MHD equations in Y direction.

Parameters

|            |                 |
|------------|-----------------|
| <b>Avg</b> | Average vector. |
|------------|-----------------|

Returns

**Vector**

```

59 {
60 real rho;
61 real vx, vy, vz;
62 real pre, e;
63 real Bx, By, Bz;
64 real Bx2, By2, Bz2, B2;
65 real vx2, vy2, vz2, v2;
66 real half = 0.5;
67
68 Vector G(QuantityNb);
69
70 //Variables
71 rho = Avg.value(1);
72 vx = Avg.value(2)/rho;
73 vy = Avg.value(3)/rho;
74 vz = Avg.value(4)/rho;
75 e = Avg.value(5);
76 Bx = Avg.value(7);
77 By = Avg.value(8);
78 Bz = Avg.value(9);
79
80 Bx2 = Bx*Bx;
81 By2 = By*By;
82 Bz2 = Bz*Bz;
83 B2 = half*(Bz2+Bx2+By2);
84
85 vx2 = vx*vx;
86 vy2 = vy*vy;
87 vz2 = vz*vz;
88 v2 = half*(vz2+vx2+vy2);
89
90 //pressure
91 pre = (Gamma -1.)*(e - rho*v2 - B2);
92
93 //Physical flux - y-direction
94 G.setValue(1,rho*vy);
95 G.setValue(2,rho*vx*vy - Bx*By);
96 G.setValue(3,rho*vy2 + pre + half*(Bx2+Bz2-By2));
97 G.setValue(4,rho*vy*vz - By*Bz);
98 G.setValue(5,(e + pre + B2)*vy - By*(vx*Bx + vy*By + vz*Bz));

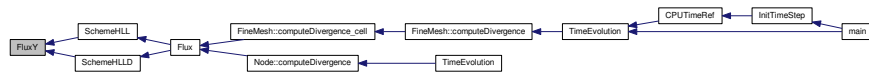
```

```

99     G.setValue(6,0.0);
100    G.setValue(7,vy*Bx - vx*By);
101    G.setValue(8,0.0);
102    G.setValue(9,vy*Bz - vz*By);
103    return G;
104 }

```

Here is the caller graph for this function:



### 6.36.2.3 Vector FluxZ ( const Vector & Avg )

Returns the physical flux of MHD equations in Z direction.

#### Parameters

|            |                 |
|------------|-----------------|
| <i>Avg</i> | Average vector. |
|------------|-----------------|

#### Returns

##### Vector

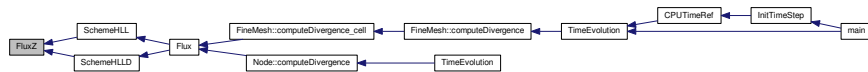
```

107 {
108     real rho;
109     real vx, vy, vz;
110     real pre, e;
111     real Bx, By, Bz;
112     real Bx2, By2, Bz2, B2;
113     real vx2, vy2, vz2, v2;
114     real half = 0.5;
115
116     Vector H(QuantityNb);
117
118     //Variables
119     rho = Avg.value(1);
120     vx = Avg.value(2)/rho;
121     vy = Avg.value(3)/rho;
122     vz = Avg.value(4)/rho;
123     e = Avg.value(5);
124     Bx = Avg.value(7);
125     By = Avg.value(8);
126     Bz = Avg.value(9);
127
128     Bx2 = Bx*Bx;
129     By2 = By*By;
130     Bz2 = Bz*Bz;
131     B2 = half*(Bz2+Bx2+By2);
132
133     vx2 = vx*vx;
134     vy2 = vy*vy;
135     vz2 = vz*vz;
136     v2 = half*(vz2+vx2+vy2);
137
138     //pressure
139     pre = (Gamma - 1.)*(e - rho*v2 - B2);
140
141     //Physical flux - y-direction
142     H.setValue(1, rho*vz);
143     H.setValue(2, rho*vz*vx - Bz*Bx);
144     H.setValue(3, rho*vz*vy - Bz*By);
145     H.setValue(4, rho*vz2 + pre + half*(Bx2+By2-Bz2));
146     H.setValue(5, (e + pre + B2)*vz - Bz*(vx*Bx + vy*By + vz*Bz));
147     H.setValue(6, 0.0);
148     H.setValue(7, vz*Bx - vx*Bz);
149     H.setValue(8, vz*By - vy*Bz);
150     H.setValue(9, 0.0);
151
152     return H;
153 }

```



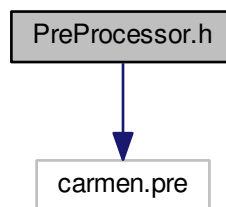
Here is the caller graph for this function:



## 6.37 PreProcessor.h File Reference

```
#include "carmen.pre"
```

Include dependency graph for PreProcessor.h:



This graph shows which files directly or indirectly include this file:



### Macros

- #define `real` `double`
- #define `FORMAT` `"%23.16e "`
- #define `TXTFORMAT` `"%-23s "`
- #define `REAL` `"double"`
- #define `BACKUP_FILE_FORMAT` `"%lf"`
- #define `byte` `unsigned char`
- #define `MPI_Type` `MPI_DOUBLE`

### 6.37.1 Macro Definition Documentation

6.37.1.1 #define `BACKUP_FILE_FORMAT` `"%lf"`

6.37.1.2 #define `byte` `unsigned char`

6.37.1.3 #define `FORMAT` `"%23.16e "`

6.37.1.4 #define `MPI_Type` `MPI_DOUBLE`

6.37.1.5 `#define real double`

6.37.1.6 `#define REAL "double"`

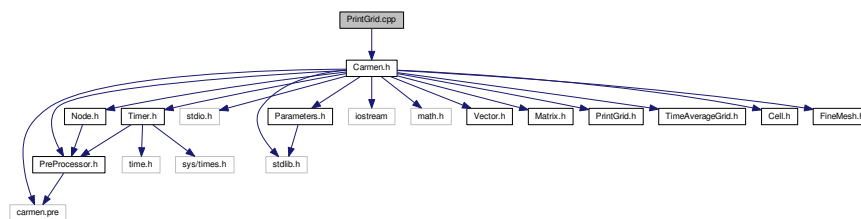
6.37.1.7 `#define TXTFORMAT "%-23s "`

## 6.38 PrintGrid.cpp File Reference

Functions to print every variable of the MHD model.

```
#include "Carmen.h"
```

Include dependency graph for PrintGrid.cpp:



### 6.38.1 Detailed Description

Functions to print every variable of the MHD model.

## 6.39 PrintGrid.h File Reference

This graph shows which files directly or indirectly include this file:



### Classes

- class [PrintGrid](#)

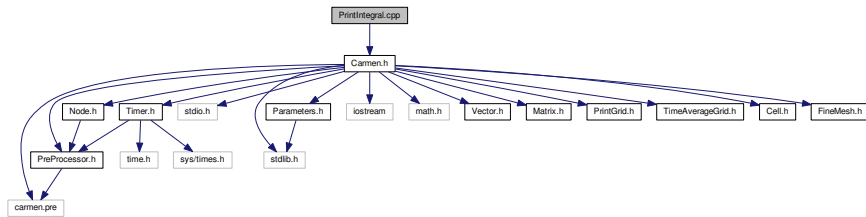
An object [PrintGrid](#) is a special regular grid created to write tree-structured data into an output file.

## 6.40 PrintIntegral.cpp File Reference

Print integral values into file "FileName".

```
#include "Carmen.h"
```

Include dependency graph for PrintIntegral.cpp:



## Functions

- void [PrintIntegral](#) (const char \*FileName)  
Writes the integral values, like e.g flame velocity, global error, into file *FileName*.

### 6.40.1 Detailed Description

Print integral values into file "FileName".

### 6.40.2 Function Documentation

#### 6.40.2.1 void PrintIntegral ( const char \* *FileName* )

Writes the integral values, like e.g flame velocity, global error, into file *FileName*.

#### Parameters

| <i>FileName</i> | Name of the file |
|-----------------|------------------|
|-----------------|------------------|

#### Returns

void

```

31 {
32     // --- Local variables ---
33
34     real    t;           // time
35     FILE   *output;    // output file
36     int    i;          // counter
37     real   Volume=1.;  // Total volume
38
39     // --- Open file ---
40
41     if ( (IterationNo == 0) ? (output = fopen(FileName,"w")) : (output = fopen(FileName,"a")) )
42     {
43         // HEADER
44
45         if (IterationNo == 0)
46         {
47
48             fprintf(output, "#");
49             fprintf(output, TXTFORMAT, " Time");
50             fprintf(output, TXTFORMAT, " CFL");
51             fprintf(output, TXTFORMAT, " Energy");
52             fprintf(output, TXTFORMAT, " Div B Max");
53             fprintf(output, TXTFORMAT, " ch");
54             fprintf(output, TXTFORMAT, " Helicity");
55             fprintf(output, TXTFORMAT, " DivB Max norm");
56 /*
57             if (Multiresolution)
58             {
59                 fprintf(output, TXTFORMAT, "Memory comp.");

```

```

60         fprintf(output, TXTFORMAT, "CPU comp.");
61         if (ExpectedCompression != 0. || CVS)
62             fprintf(output, TXTFORMAT, "Tolerance");
63         // if (CVS)
64         //     fprintf(output, TXTFORMAT, "Av. Pressure");
65
66     }
67 */
68     if (!ConstantTimeStep)
69     {
70         if (StepNb == 3) fprintf(output, TXTFORMAT, "RKF Error");
71         fprintf(output, TXTFORMAT, "Next time step");
72         fprintf(output, "%13s ", "IterationNo");
73         fprintf(output, "%13s ", "IterationNb");
74     }
75
76     fprintf(output, "\n");
77
78 }
79
80 if (ConstantTimeStep)
81     t=IterationNo*TimeStep;
82 else
83     t = ElapsedTime;
84
85 fprintf(output, FORMAT, t);
86
87 // --- Compute total volume ---
88
89 for (i=1; i<= Dimension; i++)
90     Volume *= fabs(XMax[i]-XMin[i]);
91
92 // Print CFL
93 fprintf(output, FORMAT, Eigenvalue*TimeStep/SpaceStep);
94
95 // Print momentum and energy
96 //fprintf(output, FORMAT, GlobalMomentum);
97 fprintf(output, FORMAT, GlobalEnergy);
98 fprintf(output, FORMAT, DIVBMax);
99 fprintf(output, FORMAT, ch);
100    fprintf(output, FORMAT, Helicity);
101    fprintf(output, FORMAT, DIVB);
102
103
104 /*
105     if (Multiresolution)
106     {
107         fprintf(output, FORMAT, (1.*CellNb)/(1<<(ScaleNb*Dimension)));
108         fprintf(output, FORMAT, CPUTime.CPUTime()/(IterationNo*FVTimeRef));
109
110         if (ExpectedCompression != 0.)
111             fprintf(output, FORMAT, Tolerance);
112
113         // if (CVS)
114         //{
115         //     fprintf(output, FORMAT, ComputedTolerance(ScaleNb));
116         //     fprintf(output, FORMAT, QuantityAverage.value(1));
117         //}
118     }
119 */
120     if (!ConstantTimeStep)
121     {
122         if (StepNb == 3) fprintf(output, FORMAT, RKFError);
123         fprintf(output, FORMAT, TimeStep);
124         fprintf(output, "%13i ", IterationNo);
125         fprintf(output, "%13i ", IterationNb);
126     }
127
128     fprintf(output, "\n");
129     fclose(output);
130 }
131 else
132 {
133     cout << "PrintIntegral.cpp: In method `void PrintIntegral(Node*, char*)':\n";
134     cout << "PrintIntegral.cpp: cannot open file " << FileName << '\n';
135     cout << "carmen: *** [PrintIntegral.o] Execution error\n";
136     cout << "carmen: abort execution.\n";
137     exit(1);
138 }
139 }

```

Here is the caller graph for this function:

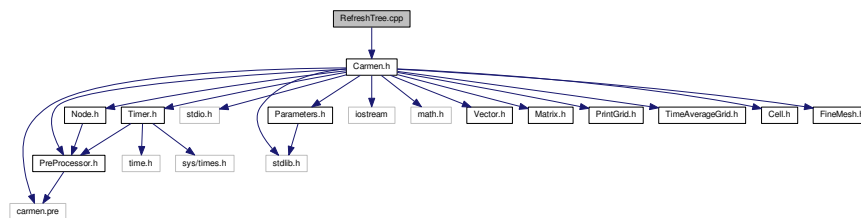


## 6.41 RefreshTree.cpp File Reference

Refresh the tree structure.

```
#include "Carmen.h"
```

Include dependency graph for RefreshTree.cpp:



### Functions

- void [RefreshTree](#) (Node \*Root)

*Refresh the tree structure, i.e. compute the cell-averages of the internal nodes by projection and those of the virtual leaves by prediction. The root node is Root. Only for multiresolution computations.*

#### 6.41.1 Detailed Description

Refresh the tree structure.

#### 6.41.2 Function Documentation

##### 6.41.2.1 void RefreshTree ( Node \* Root )

Refresh the tree structure, i.e. compute the cell-averages of the internal nodes by projection and those of the virtual leaves by prediction. The root node is *Root*. Only for multiresolution computations.

##### Parameters

|             |      |
|-------------|------|
| <i>Root</i> | Root |
|-------------|------|

**Returns**

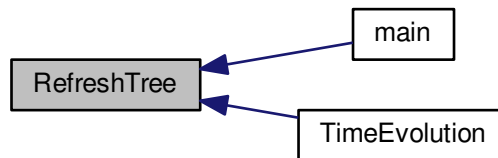
void

```

23 {
24     // --- Project : compute cell-average values in all nodes ---
25     Root->project();
26
27     // --- Fill virtual children with predicted values ---
28     Root->fillVirtualChildren();
29 }

```

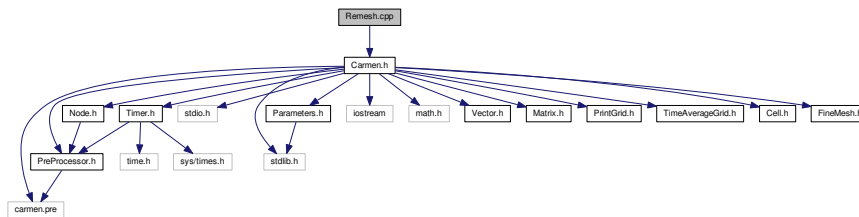
Here is the caller graph for this function:

**6.42 Remesh.cpp File Reference**

Remesh the mesh.

```
#include "Carmen.h"
```

Include dependency graph for Remesh.cpp:

**Functions**

- void [Remesh](#) (Node \*Root)

*Remesh the tree structure after a time evolution. The root node is Root. Only for multiresolution computations.*

**6.42.1 Detailed Description**

Remesh the mesh.

**6.42.2 Function Documentation**

#### 6.42.2.1 void Remesh ( Node \* Root )

Remesh the tree structure after a time evolution. The root node is *Root*. Only for multiresolution computations.

## Parameters

|             |      |
|-------------|------|
| <i>Root</i> | Root |
|-------------|------|

## Returns

void

```

23 {
24     // --- Refresh tree structure ---
25     // RefreshTree(Root);
26
27     // --- Check if tree is graded ---
28     if (debug) Root->checkGradedTree();
29
30     // --- Adapt : depending on details, refine or combine ---
31     Root->adapt();
32
33     // --- Check if tree is graded ---
34     if (debug) Root->checkGradedTree();
35 }

```

Here is the caller graph for this function:

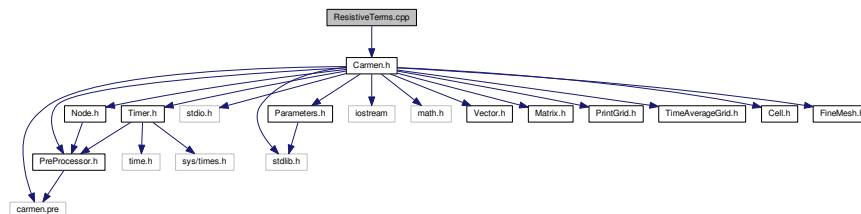


## 6.43 ResistiveTerms.cpp File Reference

This computes the resistive terms of energy and magnetic field Equations.

```
#include "Carmen.h"
```

Include dependency graph for ResistiveTerms.cpp:



## Functions

- [Vector ResistiveTerms](#) ([Cell &Cell1](#), [Cell &Cell2](#), [Cell &Cell3](#), [Cell &Cell4](#), int AxisNo)  
*Returns the resistive source terms in the cell UserCell.*

### 6.43.1 Detailed Description

This computes the resistive terms of energy and magnetic field Equations.



**Author**

Anna Karina Fontes Gomes

**Version**

2.0

**Date**

Sep-2016

**6.43.2 Function Documentation****6.43.2.1 Vector ResistiveTerms ( Cell & Cell1, Cell & Cell2, Cell & Cell3, Cell & Cell4, int AxisNo )**Returns the resistive source terms in the cell *UserCell*.**Parameters**

|               |                  |
|---------------|------------------|
| <i>Cell1</i>  | Cell 1           |
| <i>Cell2</i>  | Cell 2           |
| <i>Cell3</i>  | Cell 3           |
| <i>Cell4</i>  | Cell 4           |
| <i>AxisNo</i> | Axis of interest |

**Returns**[Vector](#)

X - direction

2D

Y - direction

2D

Z - direction

3D

```

12 {
13     // --- Local variables ---
14     Vector B(3), Bi(3), Bj(3), Bk(3);
15     Vector Result(QuantityNb);
16     Vector Bavg(3);
17     real Jx = 0., Jy = 0., Jz = 0.;
18     real dx, dy, dz;
19     real ResX= 0., ResY= 0., ResZ= 0., ResE= 0.;
20     real eta0=0., etai=0., etaj=0., etak=0., etaR=0.;
21
22     dx = Cell2.size(1);
23     dy = Cell2.size(2);
24     dz = Cell2.size(3);
25
26     eta0 = Cell1.Res;
27     etai = Cell2.Res;
28     etaj = Cell3.Res;
29     etak = Cell4.Res;
30
31     for(int i=1; i <= 3; i++){
32         B.setValue(i, Cell1.average(i+6));
33         Bi.setValue(i, Cell2.average(i+6));
34         Bj.setValue(i, Cell3.average(i+6));
35         Bk.setValue(i, Cell4.average(i+6));
36     }
37
38
39     if(AxisNo == 1){

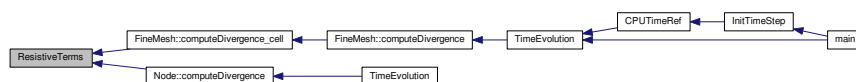
```

```

41     etaR = (eta0 + etai)/2.;
42
43     Bavg.setValue(2, 0.5*(B.value(2) + Bi.value(2)));
44     Bavg.setValue(3, 0.5*(B.value(3) + Bi.value(3)));
45
46     Jy = -(B.value(3) - Bi.value(3))/dx;
47     Jz = (B.value(2) - Bi.value(2))/dx;
48
49     Jz = Jz - (B.value(1) - Bj.value(1))/dy;
50
51     if(Dimension==3){
52         Jy = Jy + (B.value(1) - Bk.value(1))/dz;
53     }
54
55     ResE = etaR*(Bavg.value(2)*Jz - Bavg.value(3)*Jy);
56     ResX = 0.;
57     ResY = etaR*Jz;
58     ResZ = -etaR*Jy;
59
60
61 }else if(AxisNo == 2){
62     etaR = (eta0 + etaj)/2.;
63
64     Bavg.setValue(1, 0.5*(B.value(1) + Bj.value(1)));
65     Bavg.setValue(3, 0.5*(B.value(3) + Bj.value(3)));
66
67     Jx = (B.value(3) - Bj.value(3))/dy;
68     Jz = -(B.value(1) - Bj.value(1))/dy;
69
70     Jz = Jz + (B.value(2) - Bi.value(2))/dx;
71
72     if(Dimension==3){
73         Jx = Jx + (B.value(2) - Bk.value(2))/dz;
74     }
75
76     ResE = etaR*(Bavg.value(3)*Jx - Bavg.value(1)*Jz);
77     ResX = -etaR*Jz;
78     ResY = 0.;
79     ResZ = etaR*Jx;
80
81 }else{
82     etaR = (eta0 + etak)/2.;
83
84     Bavg.setValue(1, 0.5*(B.value(1) + Bk.value(1)));
85     Bavg.setValue(2, 0.5*(B.value(2) + Bk.value(2)));
86
87     Jx = -(B.value(2) - Bk.value(2))/dz;
88     Jy = (B.value(1) - Bk.value(1))/dz;
89
90     Jx = Jx + (B.value(3) - Bj.value(3))/dy;
91     Jy = Jy - (B.value(3) - Bi.value(3))/dx;
92
93     ResE = etaR*(Bavg.value(1)*Jy - Bavg.value(2)*Jx);
94     ResX = etaR*Jy;
95     ResY = -etaR*Jx;
96     ResZ = 0.;
97 }
98
99 Result.setZero();
100
101 // These values will be added to the numerical flux
102 Result.setValue(5, ResE);
103 Result.setValue(7, ResX);
104 Result.setValue(8, ResY);
105 Result.setValue(9, ResZ);
106
107 return Result;
108
109 }

```

Here is the caller graph for this function:

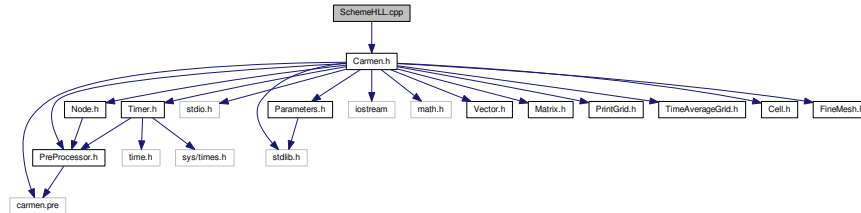


## 6.44 SchemeHLL.cpp File Reference

Computes the HLL Riemann solver.

```
#include "Carmen.h"
```

Include dependency graph for SchemeHLL.cpp:



### Functions

- **Vector SchemeHLL** (const [Cell](#) &Cell1, const [Cell](#) &Cell2, const [Cell](#) &Cell3, const [Cell](#) &Cell4, int AxisNo)
 

Returns the HLL numerical flux for MHD equations. The scheme uses four cells to estimate the flux at the interface. Cell2 and Cell3 are the first neighbours on the left and right sides. Cell1 and Cell4 are the second neighbours on the left and right sides.

#### 6.44.1 Detailed Description

Computes the HLL Riemann solver.

##### Author

Anna Karina Fontes Gomes

##### Version

4.0

##### Date

July-2016

#### 6.44.2 Function Documentation

##### 6.44.2.1 Vector SchemeHLL ( const [Cell](#) & Cell1, const [Cell](#) & Cell2, const [Cell](#) & Cell3, const [Cell](#) & Cell4, const int AxisNo )

Returns the HLL numerical flux for MHD equations. The scheme uses four cells to estimate the flux at the interface. Cell2 and Cell3 are the first neighbours on the left and right sides. Cell1 and Cell4 are the second neighbours on the left and right sides.

##### Parameters

---

|               |                                    |
|---------------|------------------------------------|
| <i>Cell1</i>  | second neighbour on the left side  |
| <i>Cell2</i>  | first neighbour on the left side   |
| <i>Cell3</i>  | first neighbour on the right side  |
| <i>Cell4</i>  | second neighbour on the right side |
| <i>AxisNo</i> | Axis of interest.                  |

## Returns

### Vector

```

12 {
13
14     // General variables
15
16     Vector    LeftAverage(QuantityNb); //
17     Vector    RightAverage(QuantityNb); // Conservative quantities
18     Vector    Result(QuantityNb);      // MHD numerical flux
19     int aux=0;
20     // Variables for the HLL scheme
21     Vector    FL(QuantityNb), FR(QuantityNb); //Left and right physical fluxes
22     Vector    VL(3), VR(3); // Left and right velocities
23     Vector    BL(3), BR(3); // Left and right velocities
24     real     rhoL=0., rhoR=0.; // Left and right densities
25     real     eL=0., eR=0.; // Left and right energies
26     real     preL=0., preR=0.;
27     real     bkL=0., bkR=0.;
28     real     aL=0., aR=0.;
29     real     bL=0., bR=0.;
30     real     cfL=0., cfR=0.;
31     real     SL=0., SR=0.;
32     real     dx=0.;
33     dx = Cell2.size(AxisNo);
34     real     r, Limit, LeftSlope = 0., RightSlope = 0.; // Left and right slopes
35     int i;
36
37 // --- Limiter function -----
38
39 for (i=1; i<=QuantityNb; i++)
40 {
41     // --- Compute left cell-average value ---
42
43     if (Cell2.average(i) != Cell1.average(i))
44     {
45         RightSlope = Cell3.average(i)-Cell2.average(i);
46         LeftSlope = Cell2.average(i)-Cell1.average(i);
47         r = RightSlope/LeftSlope;
48         Limit = Limiter(r);
49         LeftAverage.setValue(i, Cell2.average(i) + 0.5*Limit*LeftSlope);
50         aux = 1;
51     }
52     else
53         LeftAverage.setValue(i, Cell2.average(i));
54
55     // --- Compute right cell-average value ---
56
57     if (Cell3.average(i) != Cell2.average(i))
58     {
59         RightSlope = Cell4.average(i)-Cell3.average(i);
60         LeftSlope = Cell3.average(i)-Cell2.average(i);
61         r = RightSlope/LeftSlope;
62         Limit = Limiter(r);
63         RightAverage.setValue(i, Cell3.average(i) - 0.5*Limit*LeftSlope);
64         aux = 1;
65     }
66     else
67         RightAverage.setValue(i, Cell3.average(i));
68 }
69
70
71 // --- HLL scheme -----
72
73 // --- Conservative variables ---
74
75 // Left and right densities
76 rhoL = LeftAverage.value(1);
77 rhoR = RightAverage.value(1);
78
79 // Left and right momentum and magnetic field
80 for (int i=1;i<=3;i++)
81 {
82     VL.setValue( i, LeftAverage.value(i+1));
83     VR.setValue( i, RightAverage.value(i+1));

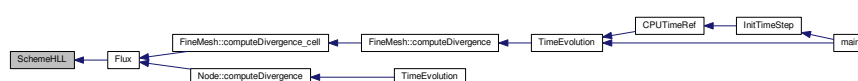
```

```

84     BL.setValue( i, LeftAverage.value(i+6));
85     BR.setValue( i, RightAverage.value(i+6));
86 }
87
88 // Left and right energies
89 eL = LeftAverage.value(5);
90 eR = RightAverage.value(5);
91
92 // Left and right pressures
93 preL = (Gamma -1.)*(eL - 0.5*(VL*VL)/rhoL - 0.5*(BL*BL));
94 preR = (Gamma -1.)*(eR - 0.5*(VR*VR)/rhoR - 0.5*(BR*BR));
95
96 // --- Magnetoacoustic waves calculations ---
97
98 bkL = power2(BL.value(AxisNo))/rhoL;
99 bkR = power2(BR.value(AxisNo))/rhoR;
100
101 aL = Gamma*preL/rhoL;
102 aR = Gamma*preR/rhoR;
103
104 bL = (BL*BL)/rhoL;
105 bR = (BR*BR)/rhoR;
106
107 // Left and Right fast speeds
108 cfL = sqrt(0.5*(aL + bL + sqrt(power2(aL + bL) - 4.0*aL*bkL)));
109 cfR = sqrt(0.5*(aR + bR + sqrt(power2(aR + bR) - 4.0*aR*bkR)));
110
111 // Left and right slopes
112 SL = Min(Min(VL.value(AxisNo)/rhoL - cfL, VR.value(AxisNo)/rhoR - cfR),0.0);
113 SR = Max(Max(VL.value(AxisNo)/rhoL + cfL, VR.value(AxisNo)/rhoR + cfR),0.0);
114
115 // --- Physical flux ---
116 if(AxisNo ==1){
117     EigenvalueX = Max(Max(Abs(SL),Abs(SR)),
EigenvalueX);
118     FL = FluxX(LeftAverage);
119     FR = FluxX(RightAverage);
120 }else if(AxisNo ==2){
121     EigenvalueY = Max(Max(Abs(SL),Abs(SR)),
EigenvalueY);
122     FL = FluxY(LeftAverage);
123     FR = FluxY(RightAverage);
124 }else{
125     EigenvalueZ = Max(Max(Abs(SL),Abs(SR)),
EigenvalueZ);
126     FL = FluxZ(LeftAverage);
127     FR = FluxZ(RightAverage);
128 }
129
130
131 // --- HLL Riemann Solver ---
132
133 for(int i=1;i<=QuantityNb;i++)
134 {
135     Result.setValue(i, (SR*FL.value(i) - SL*FR.value(i) + SR*SL*(RightAverage.value(i) - LeftAverage.
value(i)))/(SR-SL));
136 }
137
138 // Parabolic-Hyperbolic divergence Cleaning (Dedner, 2002)
139 fluxCorrection(Result, LeftAverage, RightAverage, AxisNo);
140
141 // Artificial diffusion terms
142 if(Diffusivity && aux==1) Result = Result - ArtificialViscosity(
LeftAverage,RightAverage,dx,AxisNo);
143
144 return Result;
145
146 }

```

Here is the caller graph for this function:

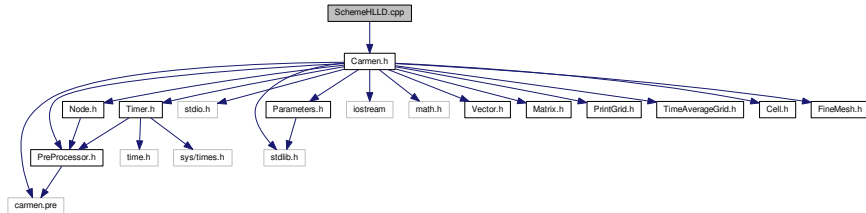


## 6.45 SchemeHLLD.cpp File Reference

Computes the HLLD Riemann solver.

```
#include "Carmen.h"
```

Include dependency graph for SchemeHLLD.cpp:



### Functions

- **Vector SchemeHLLD** (const [Cell](#) &Cell1, const [Cell](#) &Cell2, const [Cell](#) &Cell3, const [Cell](#) &Cell4, int AxisNo)
 

*Returns the HLLD numerical flux for MHD equations. The scheme uses four cells to estimate the flux at the interface. Cell2 and Cell3 are the first neighbours on the left and right sides. Cell1 and Cell4 are the second neighbours on the left and right sides.*

#### 6.45.1 Detailed Description

Computes the HLLD Riemann solver.

##### Author

Anna Karina Fontes Gomes

##### Version

4.0

##### Date

July-2016

#### 6.45.2 Function Documentation

##### 6.45.2.1 Vector SchemeHLLD ( const *Cell* & *Cell1*, const *Cell* & *Cell2*, const *Cell* & *Cell3*, const *Cell* & *Cell4*, const int *AxisNo* )

Returns the HLLD numerical flux for MHD equations. The scheme uses four cells to estimate the flux at the interface. *Cell2* and *Cell3* are the first neighbours on the left and right sides. *Cell1* and *Cell4* are the second neighbours on the left and right sides.

##### Parameters

---

|               |                                    |
|---------------|------------------------------------|
| <i>Cell1</i>  | second neighbour on the left side  |
| <i>Cell2</i>  | first neighbour on the left side   |
| <i>Cell3</i>  | first neighbour on the right side  |
| <i>Cell4</i>  | second neighbour on the right side |
| <i>AxisNo</i> | Axis of interest.                  |

## Returns

### Vector

```

13 {
14
15     // General variables
16
17     Vector    LeftAverage(QuantityNb); //
18     Vector    RightAverage(QuantityNb); // Conservative quantities
19     Vector    Result(QuantityNb);      // MHD numerical flux
20     int aux=0;
21
22     // Variables for the HLL scheme
23     Vector    FL(QuantityNb), FR(QuantityNb); //Left and right physical fluxes
24     Vector    VL(3), VR(3); // Left and right velocities
25     Vector    BL(3), BR(3); // Left and right velocities
26     real     rhoL=0., rhoR=0.; // Left and right densities
27     real     eL=0., eR=0.; // Left and right energies
28     real     preL=0., preR=0.;
29     real     bkL=0., bkR=0.;
30     real     aL=0., aR=0.;
31     real     bL=0., bR=0.;
32     real     cfL=0., cfR=0.;
33     real     SL=0., SR=0.;
34     real     SLS=0., SRS=0.;
35     real     SM=0.;
36     Matrix    U(QuantityNb,4);
37     Matrix    F(QuantityNb,2);
38     real     dx=0.;
39     dx = Cell2.size(AxisNo);
40     real     r, Limit, LeftSlope = 0., RightSlope = 0.; // Left and right slopes
41     int i;
42
43 // --- Limiter function -----
44
45     for (i=1; i<=QuantityNb; i++)
46     {
47         // --- Compute left cell-average value ---
48
49         if (Cell2.average(i) != Cell1.average(i))
50         {
51             RightSlope = Cell3.average(i)-Cell2.average(i);
52             LeftSlope = Cell2.average(i)-Cell1.average(i);
53             r = RightSlope/LeftSlope;
54             Limit = Limiter(r);
55             LeftAverage.setValue(i, Cell2.average(i) + 0.5*Limit*LeftSlope);
56             aux = 1;
57         }
58         else
59             LeftAverage.setValue(i, Cell2.average(i));
60
61         // --- Compute right cell-average value ---
62
63         if (Cell3.average(i) != Cell2.average(i))
64         {
65             RightSlope = Cell4.average(i)-Cell3.average(i);
66             LeftSlope = Cell3.average(i)-Cell2.average(i);
67             r = RightSlope/LeftSlope;
68             Limit = Limiter(r);
69             RightAverage.setValue(i, Cell3.average(i) - 0.5*Limit*LeftSlope);
70             aux = 1;
71         }
72         else
73             RightAverage.setValue(i, Cell3.average(i));
74     }
75
76 // --- HLLD scheme -----
77
78 // --- Conservative variables ---
79
80 // Left and right densities
81 rhoL = LeftAverage.value(1);
82 rhoR = RightAverage.value(1);
83
84 // Left and right momentum and magnetic field

```

```

85     for (int i=1;i<=3;i++)
86     {
87         VL.setValue( i, LeftAverage.value (i+1));
88         VR.setValue( i, RightAverage.value (i+1));
89         BL.setValue( i, LeftAverage.value (i+6));
90         BR.setValue( i, RightAverage.value (i+6));
91     }
92
93     // Left and right energies
94     eL = LeftAverage.value(5);
95     eR = RightAverage.value(5);
96
97     // Left and right pressures
98     preL = (Gamma -1.)*(eL - 0.5*(VL*VL)/rhoL - 0.5*(BL*BL));
99     preR = (Gamma -1.)*(eR - 0.5*(VR*VR)/rhoR - 0.5*(BR*BR));
100
101     // --- Magnetoacoustic waves computation ---
102     bkL = power2(BL.value(AxisNo))/rhoL;
103     bkR = power2(BR.value(AxisNo))/rhoR;
104
105     aL = Gamma*preL/rhoL;
106     aR = Gamma*preR/rhoR;
107
108     bL = (BL*BL)/rhoL;
109     bR = (BR*BR)/rhoR;
110
111     // Left and Right fast speeds
112     cfL = sqrt(0.5*(aL + bL + sqrt(power2(aL + bL) - 4.0*aL*bkL)));
113     cfR = sqrt(0.5*(aR + bR + sqrt(power2(aR + bR) - 4.0*aR*bkR)));
114
115     // Left and Right slopes
116     SL = Min((VL.value(AxisNo))/rhoL, (VR.value(AxisNo))/rhoR) - Max(cfL,cfR);
117     SR = Max((VL.value(AxisNo))/rhoL, (VR.value(AxisNo))/rhoR) + Max(cfL,cfR);
118
119     // --- Physical flux ---
120     if(AxisNo ==1){
121         EigenvalueX = Max(Max(Abs(SL),Abs(SR)),
EigenvalueX);
122         FL = FluxX(LeftAverage);
123         FR = FluxX(RightAverage);
124     }else if(AxisNo ==2){
125         EigenvalueY = Max(Max(Abs(SL),Abs(SR)),
EigenvalueY);
126         FL = FluxY(LeftAverage);
127         FR = FluxY(RightAverage);
128     }else{
129         EigenvalueZ = Max(Max(Abs(SL),Abs(SR)),
EigenvalueZ);
130         FL = FluxZ(LeftAverage);
131         FR = FluxZ(RightAverage);
132     }
133
134     // Intermediary states U* and U**
135     U = stateUstar(LeftAverage, RightAverage, preL, preR, SL, SR, SM, SLS, SRS,AxisNo);
136
137     // --- HLLD Riemann Solver ---
138
139     for(int i=1;i<=QuantityNb;i++)
140     {
141         //Flux Function - Equation 66
142         //F_L
143         if(SL>=0.)
144             Result.setValue(i, FL.value(i));
145         //F-star left // FL=FLstar
146         else if(SLS>=0. && SL<0.)
147             Result.setValue(i, FL.value(i) + SL*(U.value(i,1) - LeftAverage.value(i)));
148         //F-star-star left
149         else if(SM>=0. && SLS<0.)
150             Result.setValue(i, FL.value(i) + SLS*U.value(i,3) - (SLS - SL)*U.value(i,1) - SL*
LeftAverage.value(i));
151         //F-star-star right
152         else if(SRS>=0. && SM<0.)
153             Result.setValue(i, FR.value(i) + SRS*U.value(i,4) - (SRS - SR)*U.value(i,2) - SR*
RightAverage.value(i));
154         //F-star right
155         else if(SR>=0. && SRS<0.)
156             Result.setValue(i, FR.value(i) + SR*(U.value(i,2) - RightAverage.value(i)));
157         //F_R
158         else
159             Result.setValue(i, FR.value(i));
160     }
161     // Parabolic-Hyperbolic divergence Cleaning (Dedner, 2002)
162     //fluxCorrection(Result, Cell2.average(), Cell3.average(), AxisNo);
163     fluxCorrection(Result, LeftAverage, RightAverage, AxisNo);
164
165     // Artificial diffusion terms
166     if(Diffusivity && aux==1) Result = Result - ArtificialViscosity(

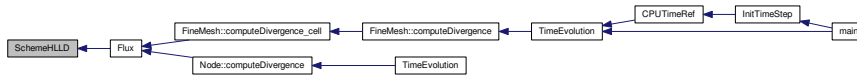
```



```

167     LeftAverage, RightAverage, dx, AxisNo) ;
168     return Result;
169 }
170 }
    
```

Here is the caller graph for this function:

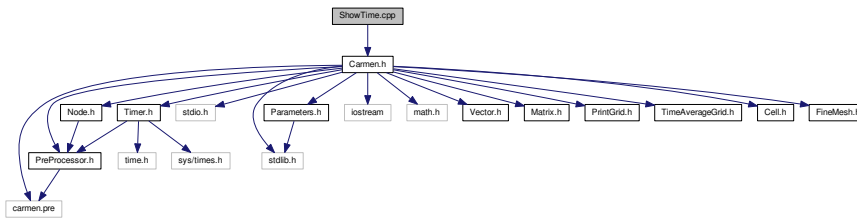


## 6.46 ShowTime.cpp File Reference

Computes the CPU Time.

```
#include "Carmen.h"
```

Include dependency graph for ShowTime.cpp:



### Functions

- void [ShowTime](#) (Timer arg)  
Writes on screen the estimation of total and remaining CPU times. These informations are stored in the timer arg.

#### 6.46.1 Detailed Description

Computes the CPU Time.

#### 6.46.2 Function Documentation

##### 6.46.2.1 void ShowTime ( Timer arg )

Writes on screen the estimation of total and remaining CPU times. These informations are stored in the timer arg.

##### Parameters

|     |          |
|-----|----------|
| arg | Argument |
|-----|----------|

## Returns

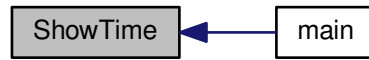
void

```

24 {
25 // double ftime; // real time
26 double ctime; // CPU time
27 // unsigned int ttime, rtime; // total and remaining real time (in seconds)
28 unsigned int tctime, rctime; // total and remaining CPU time (in seconds)
29
30 int day, hour, min, sec;
31 unsigned int rest;
32
33 // --- Write total and remaining estimated time -----
34
35 // ftime = arg.GetRealTime();
36 ctime = arg.CPUTime();
37 // ttime = (unsigned int)((ftime*IterationNb)/IterationNo);
38 // rtime = (unsigned int)((ftime*(IterationNb-IterationNo))/IterationNo);
39 tctime = (unsigned int)((ctime*IterationNb)/IterationNo);
40 rctime = (unsigned int)((ctime*(IterationNb-IterationNo))/
IterationNo);
41
42 // --- Show total time -----
43
44 rest = tctime;
45 day = rest/86400;
46 rest %= 86400;
47 hour = rest/3600;
48 rest %= 3600;
49 min = rest/60;
50 rest %= 60;
51 sec = rest;
52
53 printf("\033[1A\033[1A");
54
55 if (tctime >= 86400)
56 printf("Total CPU time (estimation) : %5d day %2d h %2d min %2d s\n", day, hour, min, sec);
57
58 if ((tctime < 86400)&&(tctime >= 3600))
59 printf("Total CPU time (estimation) : %2d h %2d min %2d s \n", hour, min, sec);
60
61 if ((tctime < 3600)&&(tctime >= 60))
62 printf("Total CPU time (estimation) : %2d min %2d s \n", min, sec);
63
64 if (tctime < 60)
65 printf("Total CPU time (estimation) : %2d s \n", sec);
66
67 // --- Show remaining time -----
68
69 rest = rctime;
70 day = rest/86400;
71 rest %= 86400;
72 hour = rest/3600;
73 rest %= 3600;
74 min = rest/60;
75 rest %= 60;
76 sec = rest;
77
78 if (rctime >= 86400)
79 printf("Remaining CPU time (estimation) : %5d day %2d h %2d min %2d s\n", day, hour, min, sec);
80
81 if ((rctime < 86400)&&(rctime >= 3600))
82 printf("Remaining CPU time (estimation) : %2d h %2d min %2d s \n", hour, min, sec);
83
84 if ((rctime < 3600)&&(rctime >= 60))
85 printf("Remaining CPU time (estimation) : %2d min %2d s \n", min, sec);
86
87 if (rctime < 60)
88 printf("Remaining CPU time (estimation) : %2d s \n", sec);
89
90 }

```

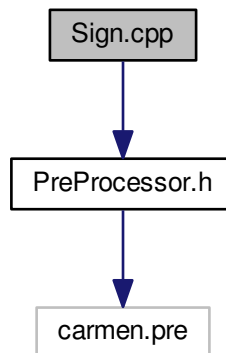
Here is the caller graph for this function:



## 6.47 Sign.cpp File Reference

Sign function.

```
#include "PreProcessor.h"  
Include dependency graph for Sign.cpp:
```



### Functions

- int [Sign](#) (const [real](#) a)  
*Returns 1 if a is non-negative, -1 elsewhere.*

#### 6.47.1 Detailed Description

Sign function.

#### 6.47.2 Function Documentation

##### 6.47.2.1 int Sign ( const real a )

Returns 1 if a is non-negative, -1 elsewhere.

## Parameters

|          |            |
|----------|------------|
| <i>a</i> | Real value |
|----------|------------|

## Returns

int

```

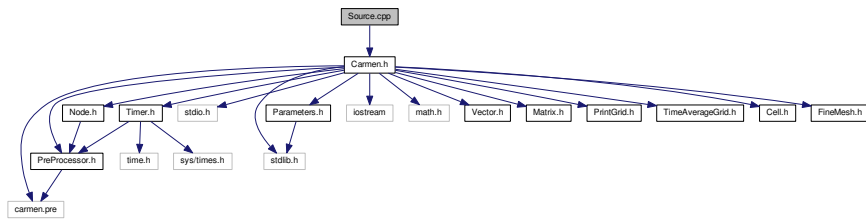
23 {
24     if (a >= 0)
25         return 1;
26     else
27         return -1;
28 }
```

## 6.48 Source.cpp File Reference

Computes the source terms of the system.

```
#include "Carmen.h"
```

Include dependency graph for Source.cpp:



## Functions

- [Vector Source \(Cell & UserCell\)](#)  
*Returns the source term in the cell UserCell.*

### 6.48.1 Detailed Description

Computes the source terms of the system.

### 6.48.2 Function Documentation

#### 6.48.2.1 Vector Source ( Cell & UserCell )

Returns the source term in the cell *UserCell*.

## Parameters

|                 |                            |
|-----------------|----------------------------|
| <i>UserCell</i> | <a href="#">Cell</a> value |
|-----------------|----------------------------|

## Returns

[Vector](#)

Gravity vector

```

24 {
25     // --- Local variables ---
26
27     Vector Force(Dimension);
28     Vector Result(QuantityNb);
29     Result.setZero();
30
31     Vector V(3);
32     real Gx=0., Gy=0., Gz=0., rho=0.;
33     for(int i=1;i<=3;i++)
34         V.setValue(i,UserCell.average(i+1));
35     rho = UserCell.density();
36     Gz = 0.2;
37     Result.setValue(2, rho*Gx);
38     Result.setValue(3, rho*Gy);
39     Result.setValue(4, rho*Gz);
40     Result.setValue(5, rho*(Gx*V.value(1) + Gy*V.value(2) + Gz*V.value(3)));
41
42     Result.setZero();
43     return Result;
44 }

```

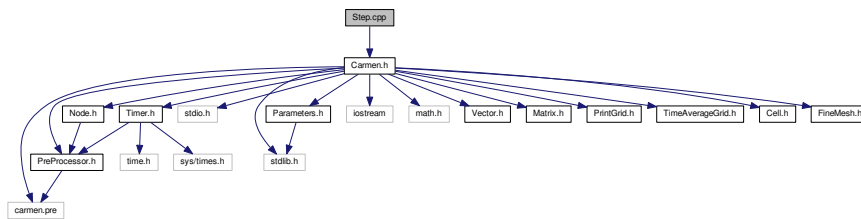
Here is the caller graph for this function:



## 6.49 Step.cpp File Reference

This function returns  $u(x) = 1$  if  $x < 0$  or  $u(x) = 1$  if  $x < 0$  or  $0$  if  $x > 0$  or  $1/2$  if  $x = 0$ .

#include "Carmen.h"  
 Include dependency graph for Step.cpp:



### Functions

- **real Step** (real x)  
 Returns a step (1 if  $x < 0$ , 0 if  $x > 0$ , 0.5 if  $x=0$ )

#### 6.49.1 Detailed Description

This function returns  $u(x) = 1$  if  $x < 0$  or  $u(x) = 1$  if  $x < 0$  or  $0$  if  $x > 0$  or  $1/2$  if  $x = 0$ .

#### 6.49.2 Function Documentation

**6.49.2.1 real Step ( real x )**

Returns a step (1 if  $x < 0$ , 0 if  $x > 0$ , 0.5 if  $x=0$ )

## Parameters

|   |            |
|---|------------|
| x | Real value |
|---|------------|

## Returns

double

```

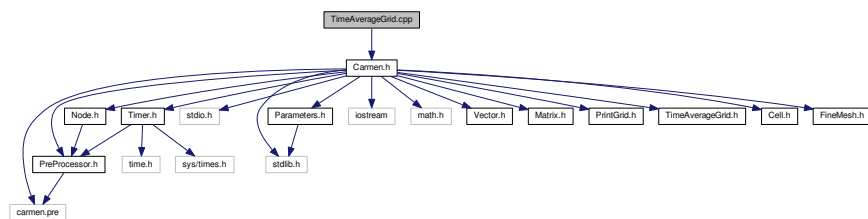
25 {
26     if (x < 0.)
27         return 1.;
28     else if (x > 0.)
29         return 0.;
30     else
31         return .5;
32 }
```

## 6.50 TimeAverageGrid.cpp File Reference

Averages the grid over time.

```
#include "Carmen.h"
```

Include dependency graph for TimeAverageGrid.cpp:



### 6.50.1 Detailed Description

Averages the grid over time.

## 6.51 TimeAverageGrid.h File Reference

This graph shows which files directly or indirectly include this file:



## Classes

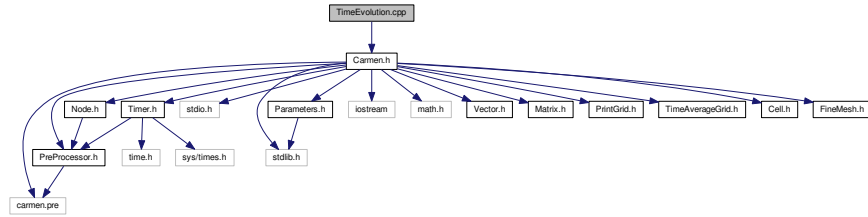
- class [TimeAverageGrid](#)  
*Time Average Grid.*

## 6.52 TimeEvolution.cpp File Reference

Time evolution for finite volume with multiresolution.

```
#include "Carmen.h"
```

Include dependency graph for TimeEvolution.cpp:



## Functions

- void `TimeEvolution (Node *Root)`  
*Computes a time evolution on the tree structure, the root node being Root. Only for multiresolution computations.*
- void `TimeEvolution (FineMesh *Root)`  
*Computes a time evolution on the regular fine mesh Root. Only for finite volume computations.*

### 6.52.1 Detailed Description

Time evolution for finite volume with multiresolution.

### 6.52.2 Function Documentation

#### 6.52.2.1 void TimeEvolution ( Node \* Root )

Computes a time evolution on the tree structure, the root node being *Root*. Only for multiresolution computations.

#### Parameters

|             |           |
|-------------|-----------|
| <i>Root</i> | Root node |
|-------------|-----------|

#### Returns

void

```

20 {
21
22     // --- Smooth data ---
23
24     if ( SmoothCoeff != 0.)
25         Root->smooth();
26
27     // --- Store cell-average values of leaves ---
28     Root->store();
29
30     // --- Refresh tree structure ---
31     RefreshTree(Root);
32
33     for (StepNo = 1; StepNo <= StepNb; StepNo++)
34     {
35         // --- Compute divergence ---
36         Root->computeDivergence();
37         // --- Runge-Kutta step ---
38         Root->RungeKutta();
39         // --- Divergence cleaning source-terms
40         Root->computeCorrection();
41
42     }
43
44     // --- Refresh tree structure ---
  
```



```

45     RefreshTree (Root);
46
47
48
49     // --- Check stability ---
50     Root->checkStability();
51
52     // --- Compute integral values ---
53     Root->computeIntegral();
54
55     // --- Compute total number of cells and leaves ---
56
57     TotalCellNb += CellNb;
58     TotalLeafNb += LeafNb;
59     //cout<<"eigen= "<<Eigenvalue<<endl;
60     // --- Compute elapsed time and adapt time step ---
61     Eigenvalue = Max(EigenvalueX,Max(EigenvalueY,
EigenvalueZ));
62     ElapsedTime += TimeStep;
63     if (!ConstantTimeStep) AdaptTimeStep();
64
65     // --- Compute divergence-free correction constant
66     //ch = CFL*SpaceStep/TimeStep;
67     ch = Max(CFL*SpaceStep/TimeStep, Eigenvalue);
68
69 }

```

### 6.52.2.2 void TimeEvolution ( FineMesh \* Root )

Computes a time evolution on the regular fine mesh *Root*. Only for finite volume computations.

#### Parameters

|             |           |
|-------------|-----------|
| <i>Root</i> | Fine mesh |
|-------------|-----------|

#### Returns

void

```

78 {
79
80     // --- Store cell-average values into temporary ---
81     Root->store();
82
83     for (StepNo = 1; StepNo <= StepNb; StepNo++)
84     {
85         // --- Compute divergence for neighbour cells ---
86         //The same conception with background computations, see upper...
87         Root->computeDivergence(1);
88         // --- Runge-Kutta step for neighbour cells ---
89         Root->RungeKutta(1);
90         // --- Divergence cleaning source term
91         Root->computeCorrection(1);
92         // --- Start inter-CPU exchanges ---
93         CPUExchange(Root, SendQ);
94         // --- Compute divergence for internal cells ---
95         Root->computeDivergence(0);
96         // --- Runge-Kutta step for internal cells ---
97         Root->RungeKutta(0);
98         // --- Divergence cleaning source term
99         Root->computeCorrection(0);
100
101 #if defined PARMPI
102     CommTimer.start(); //Communication Timer Start
103     //Waiting while inter-CPU exchanges are finished
104     if (MPIRecvType == 1) //for nonblocking receive...
105         MPI_Waitall(4*Dimension, req, st);
106     CommTimer.stop();
107 #endif
108
109     }
110
111     // --- Check stability ---
112     Root->checkStability();
113
114     // --- Compute integral values ---
115     Root->computeIntegral();
116
117     // --- Compute elapsed time and adapt time step ---

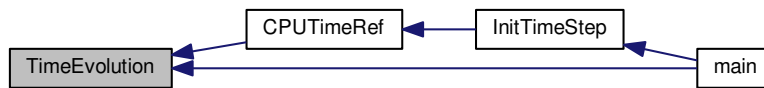
```

```

118
119     if (!ComputeCPUTimeRef)
120     {
121         Eigenvalue = Max(EigenvalueX,Max(
EigenvalueY,EigenvalueZ));
122         ElapsedTime += TimeStep;
123         if (!ConstantTimeStep) AdaptTimeStep();
124         // --- Compute divergence-free correction constant
125         //ch = CFL*SpaceStep/TimeStep;
126         ch = Max(CFL*SpaceStep/TimeStep, Eigenvalue);
127     }
128
129     // --- Compute time-average values ---
130
131     if (TimeAveraging)
132         Root->computeTimeAverage();
133
134 }

```

Here is the caller graph for this function:



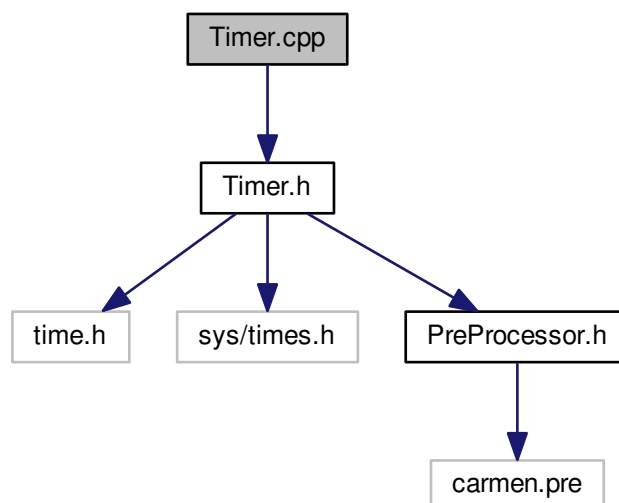
## 6.53 Timer.cpp File Reference

Computes time.

```
#include "Timer.h"

```

Include dependency graph for Timer.cpp:

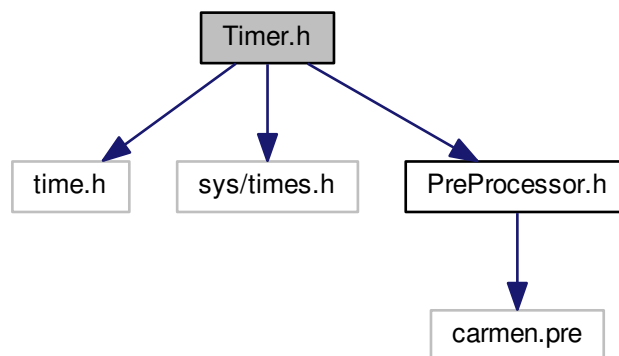


### 6.53.1 Detailed Description

Computes time.

## 6.54 Timer.h File Reference

```
#include <time.h>
#include <sys/times.h>
#include "PreProcessor.h"
Include dependency graph for Timer.h:
```



This graph shows which files directly or indirectly include this file:



### Classes

- class [Timer](#)

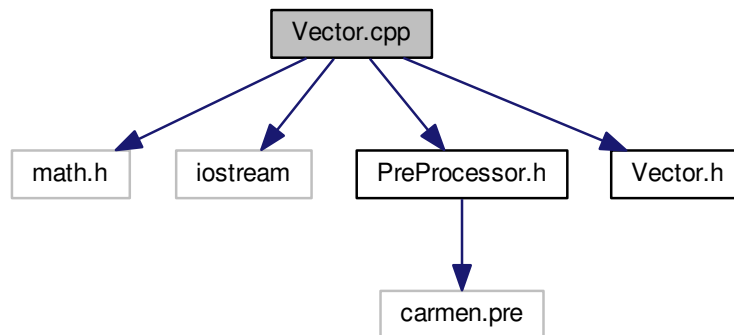
*An object [Timer](#) gives information on the CPU time of long-time computations.*

## 6.55 Vector.cpp File Reference

Creates vector structure.

```
#include <math.h>
#include <iostream>
#include "PreProcessor.h"
#include "Vector.h"
```

Include dependency graph for Vector.cpp:



## Functions

- **Vector operator\*** (const **real** a, const **Vector** &V)  
*Returns the product of the current vector and a real a.*
- **int dim** (const **Vector** &V)  
*Returns the dimension of the vector. Similar to **int Vector::dimension()**.*
- **Vector abs** (const **Vector** &V)  
*Returns the absolute value term by term of the vector.*
- **real N1** (const **Vector** &V)  
*Returns the L1-norm of the vector.*
- **real N2** (const **Vector** &V)  
*Returns the L2-norm of the vector.*
- **real NMax** (const **Vector** &V)  
*Returns the Max-norm of the vector.*
- **ostream & operator<<** (ostream &out, const **Vector** &V)  
*Writes the components of the vector V on screen.*

### 6.55.1 Detailed Description

Creates vector structure.

### 6.55.2 Function Documentation

#### 6.55.2.1 Vector abs ( const Vector & V )

Returns the absolute value term by term of the vector.

#### Parameters

---

|   |        |
|---|--------|
| V | Vector |
|---|--------|

**Returns**

Vector

```

1485 {
1486     int n;
1487     real a;
1488     Vector result( dim(V) );
1489
1490     for (n = 1; n <= dim(V); n++)
1491     {
1492         a = V.value(n);
1493         if (a < 0.)
1494             result.setValue(n, -a);
1495         else
1496             result.setValue(n, a);
1497     }
1498     return result;
1499 }

```

**6.55.2.2 int dim ( const Vector & V )**

Returns the dimension of the vector. Similar to `int Vector::dimension()`.

**Parameters**

|   |        |
|---|--------|
| V | Vector |
|---|--------|

**Returns**

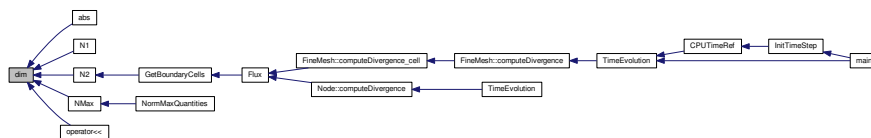
int

```

1474 {
1475     return V.dimension();
1476 }

```

Here is the caller graph for this function:

**6.55.2.3 real N1 ( const Vector & V )**

Returns the L1-norm of the vector.

**Parameters**

|   |        |
|---|--------|
| V | Vector |
|---|--------|

## Returns

real

```

1508 {
1509     int n;
1510     real result;
1511
1512     result = 0.;
1513     for (n = 1; n <= dim(V); n++)
1514         result += fabs(V.value(n));
1515
1516     return result;
1517 }

```

## 6.55.2.4 real N2 ( const Vector &amp; V )

Returns the L2-norm of the vector.

## Parameters

|   |        |
|---|--------|
| V | Vector |
|---|--------|

## Returns

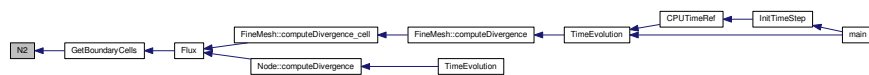
real

```

1526 {
1527     int n;
1528     real result;
1529
1530     result = 0.;
1531     for (n = 1; n <= dim(V); n++)
1532         result += V.value(n)*V.value(n);
1533
1534     return sqrt(result);
1535 }

```

Here is the caller graph for this function:



## 6.55.2.5 real NMax ( const Vector &amp; V )

Returns the Max-norm of the vector.

## Parameters

|   |        |
|---|--------|
| V | Vector |
|---|--------|

## Returns

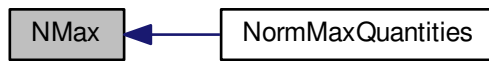
real

```

1544 {
1545     int n;
1546     real result;
1547
1548     result = 0.;
1549     for (n = 1; n <= dim(V); n++)
1550         if (result < fabs(V.value(n))) result = fabs(V.value(n));
1551
1552     return result;
1553 }

```

Here is the caller graph for this function:



### 6.55.2.6 Vector operator\*( const real a, const Vector & V )

Returns the product of the current vector and a real *a*.

Example :

```

#include "Vector.h"
Vector V(1.,0.,0.);
Vector W;
real x = 2.;
...
W = x*V;
  
```

The operation  $W = V*x$  can also be done. See [Vector Vector::operator\\*\(const real a\) const](#).

Parameters

|          |                        |
|----------|------------------------|
| <i>a</i> | Real value             |
| <i>V</i> | <a href="#">Vector</a> |

Returns

[Vector](#)

```

1463 {
1464     return V*a;
1465 }
  
```

### 6.55.2.7 ostream& operator<< ( ostream & out, const Vector & V )

Writes the components of the vector *V* on screen.

Parameters

|            |                        |
|------------|------------------------|
| <i>out</i> |                        |
| <i>V</i>   | <a href="#">Vector</a> |

Returns

ostream&

```

1563 {
1564     int n;
1565
1566     for (n = 1; n <= dim(V); n++)
  
```

```

1567     {
1568         out<<n<<": "<<V.value(n)<<endl;
1569     }
1570     return out;
1571 }

```

## 6.56 Vector.h File Reference

This graph shows which files directly or indirectly include this file:



### Classes

- class [Vector](#)  
*Standard class for every vector in Carmen.*

### Functions

- [Vector operator\\*](#) (const [real](#) a, const [Vector](#) &V)  
*Returns the product of the current vector and a real a.*
- [Vector abs](#) (const [Vector](#) &V)  
*Returns the absolute value term by term of the vector.*
- int [dim](#) (const [Vector](#) &V)  
*Returns the dimension of the vector. Similar to int [Vector::dimension\(\)](#).*
- [real N1](#) (const [Vector](#) &V)  
*Returns the L1-norm of the vector.*
- [real N2](#) (const [Vector](#) &V)  
*Returns the L2-norm of the vector.*
- [real NMax](#) (const [Vector](#) &V)  
*Returns the Max-norm of the vector.*
- ostream & [operator<<](#) (ostream &out, const [Vector](#) &V)  
*Writes the components of the vector V on screen.*

### 6.56.1 Function Documentation

#### 6.56.1.1 Vector abs ( const Vector & V )

Returns the absolute value term by term of the vector.

#### Parameters

|   |                        |
|---|------------------------|
| V | <a href="#">Vector</a> |
|---|------------------------|

#### Returns

[Vector](#)

```

1485 {
1486     int n;
1487     real a;
1488     Vector result( dim(V) );
1489
1490     for (n = 1; n <= dim(V); n++)
1491     {

```



```

1492     a = V.value(n);
1493     if (a < 0.)
1494         result.setValue(n, -a);
1495     else
1496         result.setValue(n, a);
1497     }
1498     return result;
1499 }

```

### 6.56.1.2 int dim ( const Vector & V )

Returns the dimension of the vector. Similar to `int Vector::dimension()`.

Parameters

|   |        |
|---|--------|
| V | Vector |
|---|--------|

Returns

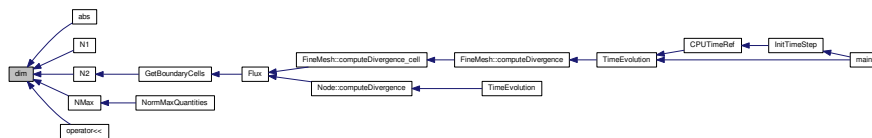
int

```

1474 {
1475     return V.dimension();
1476 }

```

Here is the caller graph for this function:



### 6.56.1.3 real N1 ( const Vector & V )

Returns the L1-norm of the vector.

Parameters

|   |        |
|---|--------|
| V | Vector |
|---|--------|

Returns

real

```

1508 {
1509     int n;
1510     real result;
1511     result = 0.;
1512     for (n = 1; n <= dim(V); n++)
1513         result += fabs(V.value(n));
1514     return result;
1515 }

```

### 6.56.1.4 real N2 ( const Vector & V )

Returns the L2-norm of the vector.

## Parameters

|   |        |
|---|--------|
| V | Vector |
|---|--------|

## Returns

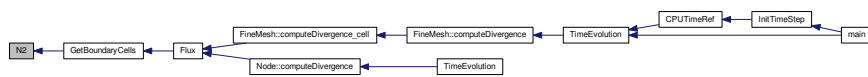
real

```

1526 {
1527     int n;
1528     real result;
1529
1530     result = 0.;
1531     for (n = 1; n <= dim(V); n++)
1532         result += V.value(n)*V.value(n);
1533
1534     return sqrt(result);
1535 }

```

Here is the caller graph for this function:



## 6.56.1.5 real NMax ( const Vector &amp; V )

Returns the Max-norm of the vector.

## Parameters

|   |        |
|---|--------|
| V | Vector |
|---|--------|

## Returns

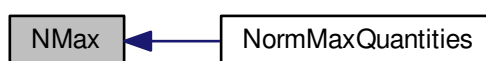
real

```

1544 {
1545     int n;
1546     real result;
1547
1548     result = 0.;
1549     for (n = 1; n <= dim(V); n++)
1550         if (result < fabs(V.value(n))) result = fabs(V.value(n));
1551
1552     return result;
1553 }

```

Here is the caller graph for this function:



**6.56.1.6 Vector operator\*( const real a, const Vector & V )**

Returns the product of the current vector and a real *a*.

Example :

```
#include "Vector.h"
Vector V(1.,0.,0.);
Vector W;
real x = 2.;
...
W = x*V;
```

The operation  $W = V*x$  can also be done. See [Vector Vector::operator\\*\(const real a\) const](#).

Parameters

|          |                        |
|----------|------------------------|
| <i>a</i> | Real value             |
| <i>V</i> | <a href="#">Vector</a> |

Returns

[Vector](#)

```
1463 {
1464     return V*a;
1465 }
```

**6.56.1.7 ostream& operator<< ( ostream & out, const Vector & V )**

Writes the components of the vector *V* on screen.

Parameters

|            |                        |
|------------|------------------------|
| <i>out</i> |                        |
| <i>V</i>   | <a href="#">Vector</a> |

Returns

[ostream&](#)

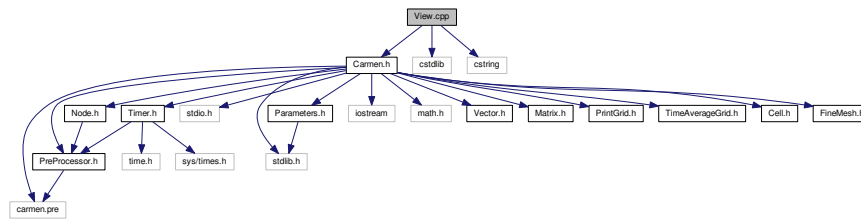
```
1563 {
1564     int n;
1565
1566     for (n = 1; n <= dim(V); n++)
1567     {
1568         out<<n<<": "<<V.value(n)<<endl;
1569     }
1570     return out;
1571 }
```

**6.57 View.cpp File Reference**

Visualization for multiresolution.

```
#include "Carmen.h"
#include <cstdlib>
#include <cstring>
```

Include dependency graph for View.cpp:



## Functions

- void `View (Node *Root, const char *TreeFileName, const char *MeshFileName, const char *AverageFileName)`  
Writes the data of the tree structure into files *TreeFileName* (tree structure), *MeshFileName* (mesh) and *AverageFileName* (cell-averages). The root node is *Root*. Only for multiresolution computations.
- void `View (FineMesh *Root, const char *AverageFileName)`  
Writes the current cell-averages of the fine mesh *Root* into file *AverageFileName*. Only for finite volume computations.

### 6.57.1 Detailed Description

Visualization for multiresolution.

### 6.57.2 Function Documentation

#### 6.57.2.1 void View ( Node \* Root, const char \* TreeFileName, const char \* MeshFileName, const char \* AverageFileName )

Writes the data of the tree structure into files *TreeFileName* (tree structure), *MeshFileName* (mesh) and *AverageFileName* (cell-averages). The root node is *Root*. Only for multiresolution computations.

#### Parameters

|                        |                   |
|------------------------|-------------------|
| <i>Root</i>            | Root node         |
| <i>TreeFileName</i>    | Tree file name    |
| <i>MeshFileName</i>    | Mesh file name    |
| <i>AverageFileName</i> | Average file name |

#### Returns

void

```

36 {
37     char buf[256];
38     int iaux;
39
40     // write tree (debugging only)
41     if (debug) Root->writeTree(TreeFileName);
42
43     // Root->computeCorrection();
44
45     // write mesh for graphic visualisation
46     if (Dimension != 1)
47     {
48         Root->writeHeader(MeshFileName);
49         Root->writeAverage(MeshFileName);
50
51         // Compress data (if parameter ZipData is true)
  
```

```

52     if (ZipData)
53     {
54         sprintf(buf,"gzip %s",MeshFileName);
55         iaux=system(buf);
56     }
57 }
58 else
59     Root->writeMesh(MeshFileName);
60
61
62 // write cell-averages in multiresolution representation (1D) or on fine grid (2-3D)
63 if (Dimension != 1)
64 {
65     Root->writeFineGrid(AverageFileName,ScaleNb+
PrintMoreScales);
66
67     // Compress data
68     if (ZipData)
69     {
70         sprintf(buf,"gzip %s",AverageFileName);
71         iaux=system(buf);
72     }
73 }
74 else
75 {
76     Root->writeHeader(AverageFileName);
77     Root->writeAverage(AverageFileName);
78 }
79 }

```

### 6.57.2.2 void View ( FineMesh \* Root, const char \* AverageFileName )

Writes the current cell-averages of the fine mesh *Root* into file *AverageFileName*. Only for finite volume computations.

#### Parameters

|                        |           |
|------------------------|-----------|
| <i>Root</i>            | Fine mesh |
| <i>AverageFileName</i> | File name |

#### Returns

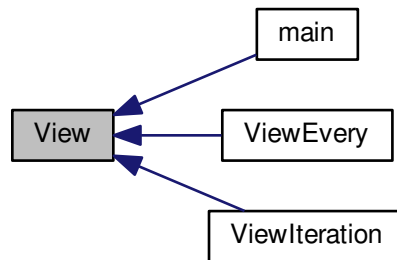
void

```

88 {
89     char buf[256];
90     int iaux;
91
92
93     char CPUFileName[255];
94 #if defined PARMPI
95     sprintf(CPUFileName,"%d_%d_%d_%s",coords[0],coords[1],coords[2],AverageFileName);
96 #else
97     strcpy(CPUFileName, AverageFileName);
98 #endif
99
100     // write header for graphic visualization
101     Root->writeHeader(CPUFileName);
102
103     // write cell-average values for graphic visualization
104     Root->writeAverage(CPUFileName);
105
106     // Compress data
107     if (Dimension != 1)
108     {
109         if (ZipData)
110         {
111             sprintf(buf,"gzip %s",CPUFileName);
112             iaux=system(buf);
113         }
114     }
115
116     // --- Write time-average values into file ---
117
118     if (TimeAveraging)
119         Root->writeTimeAverage("TimeAverage.dat");
120
121 }

```

Here is the caller graph for this function:

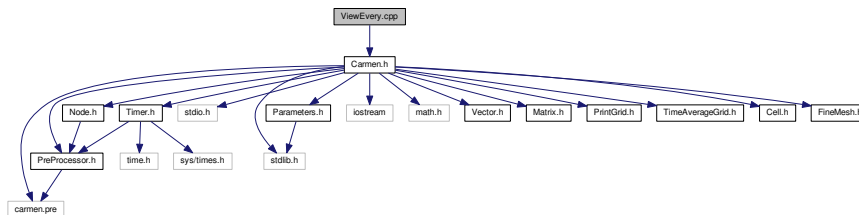


## 6.58 ViewEvery.cpp File Reference

Print solution every PrintEvery iteration.

```
#include "Carmen.h"
```

Include dependency graph for ViewEvery.cpp:



## Functions

- void [ViewEvery](#) (Node \*Root, int arg)

*Writes into file the data of the tree structure at iteration arg. The output file names are AverageNNN.dat and MeshNNN.dat, NNN being the iteration in an accurate format. The root node is Root. Only for multiresolution computations.*

- void [ViewEvery](#) (FineMesh \*Root, int arg)

*Same as previous for a fine mesh Root. Only for finite volume computations.*

### 6.58.1 Detailed Description

Print solution every PrintEvery iteration.

### 6.58.2 Function Documentation

#### 6.58.2.1 void ViewEvery ( Node \* Root, int arg )

Writes into file the data of the tree structure at iteration arg. The output file names are AverageNNN.dat and MeshNNN.dat, NNN being the iteration in an accurate format. The root node is Root. Only for multiresolution computations.

## Parameters

|             |           |
|-------------|-----------|
| <i>Root</i> | Root node |
| <i>arg</i>  | Argument  |

## Returns

void

```

33 {
34   char AverageName[256];      // File name for AverageNNN.dat
35   char MeshName[256];       // File name for MeshNNN.dat
36   char AverageFormat[256];  // File format for AverageNNN.dat
37   char MeshFormat[256];    // File format for MeshNNN.dat
38
39   sprintf(AverageFormat, "Average%s0%i.vtk", "%", DigitNumber(
  IterationNb));
40   sprintf(AverageName, AverageFormat, arg);
41   sprintf(MeshFormat, "Mesh%s0%i.dat", "%", DigitNumber(
  IterationNb));
42   sprintf(MeshName, MeshFormat, arg);
43
44   View(Root, "Tree.dat", MeshName, AverageName);
45 }
```

## 6.58.2.2 void ViewEvery ( FineMesh \* Root, int arg )

Same as previous for a fine mesh *Root*. Only for finite volume computations.

## Parameters

|             |           |
|-------------|-----------|
| <i>Root</i> | Fine mesh |
| <i>arg</i>  | argument  |

## Returns

void

```

54 {
55   char AverageName[256];      // File name for AverageNNN.dat
56   char AverageFormat[256];  // File format for AverageNNN.dat
57
58   sprintf(AverageFormat, "Average%s0%i.vtk", "%", DigitNumber(
  IterationNb));
59   sprintf(AverageName, AverageFormat, arg);
60
61   View(Root, AverageName);
62 }
```

Here is the caller graph for this function:

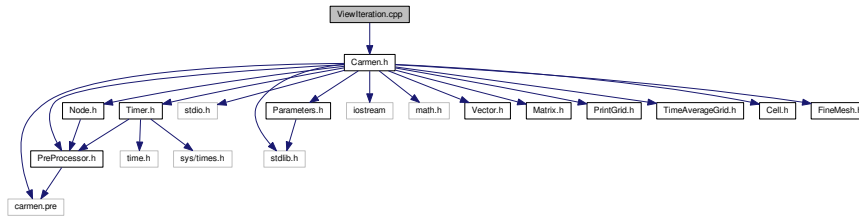


## 6.59 ViewIteration.cpp File Reference

Print solution if IterationNo = PrintIt1 to PrintIt6.

```
#include "Carmen.h"
```

Include dependency graph for ViewIteration.cpp:



## Functions

- void `ViewIteration (Node *Root)`  
Writes into file the data of the tree structure from physical time `PrintTime1` to physical time `PrintTime6`. The output file names are `Average_N.dat` and `Mesh_N.dat`, `N` being between 1 and 6. The root node is `Root`. Only for multiresolution computations.
- void `ViewIteration (FineMesh *Root)`  
Same as previous for a fine mesh `Root`. Only for finite volume computations.

### 6.59.1 Detailed Description

Print solution if `IterationNo = PrintIt1` to `PrintIt6`.

### 6.59.2 Function Documentation

#### 6.59.2.1 void ViewIteration ( Node \* Root )

Writes into file the data of the tree structure from physical time `PrintTime1` to physical time `PrintTime6`. The output file names are `Average_N.dat` and `Mesh_N.dat`, `N` being between 1 and 6. The root node is `Root`. Only for multiresolution computations.

#### Parameters

|             |           |
|-------------|-----------|
| <i>Root</i> | Root node |
|-------------|-----------|

#### Returns

void

```

34 {
35     if (IterationNo == PrintIt1)
36         View(Root, "Tree.dat", "Mesh_1.dat", "Average_1.vtk");
37
38     if (IterationNo == PrintIt2)
39         View(Root, "Tree.dat", "Mesh_2.dat", "Average_2.vtk");
40
41     if (IterationNo == PrintIt3)
42         View(Root, "Tree.dat", "Mesh_3.dat", "Average_3.vtk");
43
44     if (IterationNo == PrintIt4)
45         View(Root, "Tree.dat", "Mesh_4.dat", "Average_4.vtk");
46
47     if (IterationNo == PrintIt5)
48         View(Root, "Tree.dat", "Mesh_5.dat", "Average_5.vtk");
49
50     if (IterationNo == PrintIt6)
51         View(Root, "Tree.dat", "Mesh_6.dat", "Average_6.vtk");
52 }
  
```



### 6.59.2.2 void ViewIteration ( FineMesh \* Root )

Same as previous for a fine mesh *Root*. Only for finite volume computations.

## Parameters

|             |           |
|-------------|-----------|
| <i>Root</i> | Fine mesh |
|-------------|-----------|

## Returns

void

```
61 {  
62     if (IterationNo == PrintIt1)  
63         View(Root, "Average_1.vtk");  
64  
65     if (IterationNo == PrintIt2)  
66         View(Root, "Average_2.vtk");  
67  
68     if (IterationNo == PrintIt3)  
69         View(Root, "Average_3.vtk");  
70  
71     if (IterationNo == PrintIt4)  
72         View(Root, "Average_4.vtk");  
73  
74     if (IterationNo == PrintIt5)  
75         View(Root, "Average_5.vtk");  
76  
77     if (IterationNo == PrintIt6)  
78         View(Root, "Average_6.vtk");  
79 }
```

Here is the caller graph for this function:



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