Guacho 3D V1.1

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Contents

1	GUA	ACHO-3	D Docume	entati	on													1
	1.1	Introdu	iction							 	 	 	 	 				1
	1.2	release	e.notes .							 	 	 	 					1
	1.3	require	ements .							 	 ٠.	 	 					1
2	Mod	lules Inc	dex															3
	2.1	Module	es List							 	 		 	 				3
3	File	Index																5
	3.1	File Lis	st							 	 	 	 	 				5
4	Mod	lule Dod	umentati	on														7
	4.1	bound	aries Modu	ule Re	eferenc	e .				 	 		 	 				7
		4.1.1	Detailed	Desc	ription					 	 	 	 	 				7
		4.1.2	Function	/Subr	outine	Docu	ımer	ntatio	n .	 	 		 	 				7
			4.1.2.1	bou	ndaryi					 	 	 	 	 				7
			4.1.2.2	bou	ndaryii	٠.				 	 	 	 	 				7
	4.2	colden	s_utilities	Modu	le Refe	erenc	е.			 	 		 	 				7
		4.2.1	Detailed	Desc	ription					 	 	 	 	 				8
		4.2.2	Function	/Subr	outine	Docu	ımer	ntatio	n .	 	 	 	 	 				8
			4.2.2.1	fill_r	map .					 	 	 	 	 				8
			4.2.2.2	getx	xyz .					 	 	 	 	 				9
			4.2.2.3	init_	_colden	ıs .				 	 	 	 					9
			4.2.2.4	read	d_data					 	 		 	 				9
			4.2.2.5	rota	ition_x					 	 		 	 				10
			4.2.2.6	rota	ıtion_y					 	 		 					10
			4.2.2.7	rota	.tion_z					 	 		 					10
			4.2.2.8	write	e_map					 	 		 					11
	4.3	consta	nts Modul	e Ref	erence					 	 	 	 					11
	4.4	cooling	g_chi Modu	ule Re	eferenc	е.				 	 	 	 	 				12
		4.4.1	Detailed	Desc	ription					 	 	 	 	 				12
		4.4.2	Function	/Subr	outine	Docu	ımer	ntatio	n .	 	 	 	 	 				12

iv CONTENTS

		4.4.2.1	coolchi	12
		4.4.2.2	coolingchi	12
4.5	cooling	_dmc Mod	dule Reference	13
	4.5.1	Detailed	Description	13
	4.5.2	Function	/Subroutine Documentation	13
		4.5.2.1	cooldmc	13
		4.5.2.2	coolingdmc	13
		4.5.2.3	read_table	13
4.6	cooling	_h Module	e Reference	13
	4.6.1	Detailed	Description	14
	4.6.2	Function	/Subroutine Documentation	14
		4.6.2.1	aloss	14
		4.6.2.2	alpha	15
		4.6.2.3	alpha1	16
		4.6.2.4	atomic	16
		4.6.2.5	betah	16
		4.6.2.6	colf	17
		4.6.2.7	coolingh	17
4.7	difrad I	Module Re	eference	17
	4.7.1	Detailed	Description	18
	4.7.2	Function	/Subroutine Documentation	18
		4.7.2.1	diffuse_rad	18
		4.7.2.2	emdiff	18
		4.7.2.3	init_rand	19
		4.7.2.4	photons	19
		4.7.2.5	progress	19
		4.7.2.6	radbounds	19
		4.7.2.7	random_versor	20
		4.7.2.8	starsource	20
4.8	globals	Module F	Reference	20
	4.8.1	Detailed	Description	21
4.9	h_alph	a_utilities	Module Reference	21
	4.9.1	Detailed	Description	22
	4.9.2	Function	/Subroutine Documentation	22
		4.9.2.1	fill_map	22
		4.9.2.2	getxyz	23
		4.9.2.3	init_ha	23
		4.9.2.4	read_data	23
		4.9.2.5	rotation_x	24
		4.9.2.6	rotation_y	24

CONTENTS

		4.9.2.7 rotation_z	24
		4.9.2.8 write_ha	25
		4.9.2.9 write_rg	25
4.10	hll Mod	ule Reference	25
	4.10.1	Detailed Description	25
	4.10.2	Function/Subroutine Documentation	25
		4.10.2.1 hllfluxes	26
		4.10.2.2 prim2fhll	27
4.11	hllc Mo	dule Reference	28
	4.11.1	Detailed Description	28
	4.11.2	Function/Subroutine Documentation	28
		4.11.2.1 hllcfluxes	28
		4.11.2.2 prim2fhllc	29
4.12	hlld Mo	dule Reference	29
	4.12.1	Detailed Description	30
	4.12.2	Function/Subroutine Documentation	30
		4.12.2.1 hlldfluxes	30
		4.12.2.2 prim2fhlld	30
4.13	hlle Mo	dule Reference	31
	4.13.1	Detailed Description	31
	4.13.2	Function/Subroutine Documentation	31
		4.13.2.1 hllefluxes	31
		4.13.2.2 prim2fhlle	32
4.14	hydro_c	core Module Reference	32
	4.14.1	Detailed Description	33
	4.14.2	Function/Subroutine Documentation	33
		4.14.2.1 calcprim	33
		4.14.2.2 cfast	34
		4.14.2.3 cfastx	34
		4.14.2.4 csound	34
		4.14.2.5 get_timestep	34
		4.14.2.6 limiter	35
		4.14.2.7 prim2f	36
		4.14.2.8 prim2u	36
		4.14.2.9 swapy	36
		4.14.2.10 swapz	36
		4.14.2.11 u2prim	36
4.15	hydro_s	solver Module Reference	37
	4.15.1	Detailed Description	37
	4.15.2	Function/Subroutine Documentation	37

vi CONTENTS

		4.15.2.1 step	37
		4.15.2.2 tstep	38
		4.15.2.3 viscosity	38
4.16	init Mod	dule Reference	38
	4.16.1	Detailed Description	38
	4.16.2	Function/Subroutine Documentation	38
		4.16.2.1 initflow	38
		4.16.2.2 initmain	38
4.17	lyman_	alpha_utilities Module Reference	39
	4.17.1	Detailed Description	39
	4.17.2	Function/Subroutine Documentation	39
		4.17.2.1 fill_map	39
		4.17.2.2 getxyz	40
		4.17.2.3 init_la	40
		4.17.2.4 phigauss	41
		4.17.2.5 read_data	41
		4.17.2.6 rotation_x	41
		4.17.2.7 rotation_y	41
		4.17.2.8 rotation_z	41
		4.17.2.9 write_la	42
4.18	_	n_module Module Reference	42
		Detailed Description	42
	4.18.2	Function/Subroutine Documentation	42
		4.18.2.1 write_bin	42
		4.18.2.2 write_header	43
4.19		p_module Module Reference	43
		Detailed Description	43
	4.19.2	Function/Subroutine Documentation	43
		4.19.2.1 outputsilo	43
		4.19.2.2 writeblocks	44
		4.19.2.3 writemaster	44
4.20	_	module Module Reference	44
		Detailed Description	44
	4.20.2	Function/Subroutine Documentation	45
		4.20.2.1 write_vtk	45
4.21		Module Reference	46
		Detailed Description	46
	4.21.2	Function/Subroutine Documentation	46
4.00		4.21.2.1 write_output	46
4.22	sources	s Module Reference	47

CONTENTS vii

	4.22.1	Detailed Description
	4.22.2	Function/Subroutine Documentation
		4.22.2.1 divbcorr_source
		4.22.2.2 divergence_b
		4.22.2.3 getpos
		4.22.2.4 grav_source
		4.22.2.5 radpress_source
		4.22.2.6 source
4.23	therma	I_cond Module Reference
	4.23.1	Detailed Description
	4.23.2	Function/Subroutine Documentation
		4.23.2.1 get_dt_cond
		4.23.2.2 heatfluxes
		4.23.2.3 init_thermal_cond
		4.23.2.4 ksp
		4.23.2.5 ksp_parl
		4.23.2.6 ksp_perp
		4.23.2.7 mhd_heatfluxes
		4.23.2.8 progress
		4.23.2.9 st_steps
		4.23.2.10 substep
		4.23.2.11 superstep
		4.23.2.12 thermal_bounds
		4.23.2.13 thermal_conduction
File	Docume	entation 5
5.1		esquivel/Desktop/Guacho-Working/doc/mainpage.h File Reference
5.2		esquivel/Desktop/Guacho-Working/src/boundaries.f90 File Reference
	5.2.1	Detailed Description
5.3		esquivel/Desktop/Guacho-Working/src/coldens.f90 File Reference
	5.3.1	Detailed Description
	5.3.2	Function/Subroutine Documentation
		5.3.2.1 coldens
5.4	/Users/	esquivel/Desktop/Guacho-Working/src/constants.f90 File Reference
	5.4.1	Detailed Description
5.5	/Users/	/esquivel/Desktop/Guacho-Working/src/cooling_chi.f90 File Reference
	5.5.1	Detailed Description
5.6	/Users/	esquivel/Desktop/Guacho-Working/src/cooling_dmc.f90 File Reference
	5.6.1	Detailed Description
5.7	/Users/	esquivel/Desktop/Guacho-Working/src/cooling_h.f90 File Reference

5

viii CONTENTS

	5.7.1 Detailed Description	60
5.8	/Users/esquivel/Desktop/Guacho-Working/src/difrad.f90 File Reference	60
	5.8.1 Detailed Description	61
5.9	/Users/esquivel/Desktop/Guacho-Working/src/globals.f90 File Reference	61
	5.9.1 Detailed Description	63
5.10	/Users/esquivel/Desktop/Guacho-Working/src/h_alpha_proj.f90 File Reference	63
	5.10.1 Detailed Description	63
	5.10.2 Function/Subroutine Documentation	64
	5.10.2.1 h_alpha_proj	64
5.11	/Users/esquivel/Desktop/Guacho-Working/src/hll.f90 File Reference	64
	5.11.1 Detailed Description	64
5.12	/Users/esquivel/Desktop/Guacho-Working/src/hllc.f90 File Reference	65
	5.12.1 Detailed Description	65
5.13	/Users/esquivel/Desktop/Guacho-Working/src/hlld.f90 File Reference	65
	5.13.1 Detailed Description	66
5.14	/Users/esquivel/Desktop/Guacho-Working/src/hlle.f90 File Reference	66
	5.14.1 Detailed Description	66
5.15	/Users/esquivel/Desktop/Guacho-Working/src/hydro_core.f90 File Reference	66
	5.15.1 Detailed Description	67
5.16	/Users/esquivel/Desktop/Guacho-Working/src/hydro_solver.f90 File Reference	67
	5.16.1 Detailed Description	68
5.17	/Users/esquivel/Desktop/Guacho-Working/src/init.f90 File Reference	68
	5.17.1 Detailed Description	68
5.18	/Users/esquivel/Desktop/Guacho-Working/src/lyman_alpha_tau.f90 File Reference	68
	5.18.1 Detailed Description	69
	5.18.2 Function/Subroutine Documentation	70
	5.18.2.1 lyman_alpha_tau	70
5.19	/Users/esquivel/Desktop/Guacho-Working/src/main.f90 File Reference	70
	5.19.1 Detailed Description	71
5.20	/Users/esquivel/Desktop/Guacho-Working/src/Out_BIN_Module.f90 File Reference	71
	5.20.1 Detailed Description	71
5.21	/Users/esquivel/Desktop/Guacho-Working/src/Out_Silo_Module.f90 File Reference	71
	5.21.1 Detailed Description	72
5.22	/Users/esquivel/Desktop/Guacho-Working/src/Out_VTK_Module.f90 File Reference	72
	5.22.1 Detailed Description	72
5.23	/Users/esquivel/Desktop/Guacho-Working/src/output.f90 File Reference	72
	5.23.1 Detailed Description	73
5.24	/Users/esquivel/Desktop/Guacho-Working/src/sources.f90 File Reference	73
	5.24.1 Detailed Description	73
5.25	/Users/esquivel/Desktop/Guacho-Working/src/thermal_cond.f90 File Reference	74

CONTENTS		i
5.25.1	1 Detailed Description	
Index		77

Chapter 1

GUACHO-3D Documentation

Authors

Alejandro Esquivel et al.

1.1 Introduction

Documentation of the Guacho code

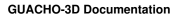
1.2 release.notes

This program is distributed in the hope that it will be useful, but WITHOUT ANY WARRANTY; without even the implied warranty of MERCHANTABILITY or FITNESS FOR A PARTICULAR PURPOSE. See the GNU General Public License for more details.

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1.3 requirements

Fortran 90/95 compiler with C preprocessor, Message Passing Interface (optional), gmake.



Chapter 2

Modules Index

2.1 Modules List

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

4 Modules Index

out_vtk_	<u>module</u>	
	Output in VTK format	44
output		
	Writes output	46
sources		
	Adds source terms	47
thermal_	_cond	
	Adds thermal conducion	50

Chapter 3

File Index

3.1 File List

е	re is a list of all documented files with brief descriptions:	
	/Users/esquivel/Desktop/Guacho-Working/doc/mainpage.h	
	Webpage frontend	55
	/Users/esquivel/Desktop/Guacho-Working/src/boundaries.f90	
	Boundary conditions	55
	/Users/esquivel/Desktop/Guacho-Working/src/coldens.f90	
	Column density projection	55
	/Users/esquivel/Desktop/Guacho-Working/src/constants.f90	
	Constants module	57
	/Users/esquivel/Desktop/Guacho-Working/src/cooling_chi.f90	
	Cooling module with CHIANTI generated cooling curves	58
	/Users/esquivel/Desktop/Guacho-Working/src/cooling_dmc.f90	
	Cooling module with Digarno Mac Cray coronal cooling curve	59
	/Users/esquivel/Desktop/Guacho-Working/src/cooling h.f90	
	Cooling with hydrogen rate parametrized cooling	59
	/Users/esquivel/Desktop/Guacho-Working/src/difrad.f90	
	Diffuse radiation module	60
	/Users/esquivel/Desktop/Guacho-Working/src/globals.f90	
	Global variables	61
	/Users/esquivel/Desktop/Guacho-Working/src/h alpha proj.f90	
	H alpha projection	63
	/Users/esquivel/Desktop/Guacho-Working/src/hll.f90	
	HLL approximate Riemann solver module	64
	/Users/esquivel/Desktop/Guacho-Working/src/hllc.f90	
	HLLC approximate Riemann solver module	65
	/Users/esquivel/Desktop/Guacho-Working/src/hlld.f90	
	HLLD approximate Riemann solver module	65
	/Users/esquivel/Desktop/Guacho-Working/src/hlle.f90	
	HLLE approximate Riemann solver module	66
	/Users/esquivel/Desktop/Guacho-Working/src/hydro core.f90	
	Hydrodynamical and Magnetohidrodynamocal bacic module	66
	/Users/esquivel/Desktop/Guacho-Working/src/hydro_solver.f90	
	Hydrodynamical and Magnetohidrodynamocal solver module	67
	/Users/esquivel/Desktop/Guacho-Working/src/init.f90	
	Guacho-3D initialization module	68
	/Users/esquivel/Desktop/Guacho-Working/src/lyman_alpha_tau.f90	-
	Lyman alpha utilities	68
	/Users/esquivel/Desktop/Guacho-Working/src/main.f90	
	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	

6 File Index

/Users/esquivel/Desktop/Guacho-Working/src/Out_BIN_Module.f90	
Output in BIN Format	71
/Users/esquivel/Desktop/Guacho-Working/src/Out_Silo_Module.f90	
Output in Silo Format	71
/Users/esquivel/Desktop/Guacho-Working/src/Out_VTK_Module.f90	
Output in VTK Format	72
/Users/esquivel/Desktop/Guacho-Working/src/output.f90	
Writes Output	72
/Users/esquivel/Desktop/Guacho-Working/src/sources.f90	
Adds source terms	73
/Users/esquivel/Desktop/Guacho-Working/src/thermal_cond.f90	
Thermal conduction module	74

Chapter 4

Module Documentation

4.1 boundaries Module Reference

Boundary conditions.

Functions/Subroutines

• subroutine boundaryi ()

Boundary conditions for 1st order half timestep.

• subroutine boundaryii ()

Boundary conditions for 2nd order half timestep.

4.1.1 Detailed Description

Sets boundary conditions, the type of boundaries is set in the Makefile

4.1.2 Function/Subroutine Documentation

4.1.2.1 subroutine boundaries::boundaryi ()

Boundary conditions for 1st order half timestep

The conditions only are imposed at the innermost ghost cell, on the u (unstepped) variables

Definition at line 48 of file boundaries.f90.

4.1.2.2 subroutine boundaries::boundaryii ()

Boundary conditions for 2nd order half timestep

The conditions only are imposed in two ghost cells on the up (stepped) variables

Definition at line 259 of file boundaries.f90.

4.2 coldens_utilities Module Reference

Column densirt projection.

Functions/Subroutines

• subroutine init_coldens ()

Initializes data.

• subroutine read_data (u, itprint, filepath)

reads data from file

• subroutine getxyz (i, j, k, x, y, z)

gets position of a cell

• subroutine rotation_x (theta, x, y, z, xn, yn, zn)

Rotation around the X axis.

• subroutine rotation_y (theta, x, y, z, xn, yn, zn)

Rotation around the Y axis.

subroutine rotation_z (theta, x, y, z, xn, yn, zn)

Rotation around the Z axis.

• subroutine fill map (nxmap, nymap, u, map, dxT, dyT, theta x, theta y, theta z)

Fill target map.

• subroutine write_map (fileout, nxmap, nymap, map)

Writes projection to file.

4.2.1 Detailed Description

Utilities to compute a column density map

4.2.2 Function/Subroutine Documentation

4.2.2.1 subroutine coldens_utilities::fill_map (integer, intent(in) nxmap, integer, intent(in) nymap, real, dimension(neq,nxmin:nxmax,nymin:nymax, nzmin:nzmax), intent(in) u, real, dimension(nxmap,nymap), intent(out) map, real, intent(in) dxT, real, intent(in) dyT, real, intent(in) theta_x, real, intent(in) theta_y, real, intent(in) theta_z)

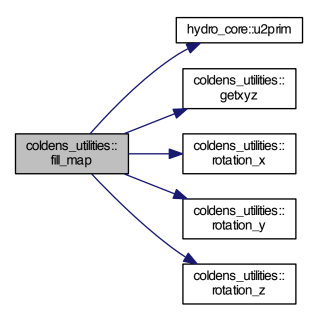
Fills the target map of one MPI block

Parameters

integer	integer [in] nxmap : Number of X cells in target	
integer	[in] nymap : Number of Y cells in target	
real	[in] u(neq,nxmin:nxmax,nymin:nymax, nzmin:nzmax) : conserved variables	
real	[out] map(nxmap,mymap) : Target map	
real	[in] dxT: target pixel width	
real	[in] dyT: target pixel height	
real	[in] thetax : Rotation around X	
real [in] thetay: Rotation around Y		
real	[in] thetaz : Rotation around Z	

Definition at line 286 of file coldens.f90.

Here is the call graph for this function:



4.2.2.2 subroutine coldens_utilities::getxyz (integer, intent(in) *i*, integer, intent(in) *j*, integer, intent(in) *k*, real, intent(out) *x*, real, intent(out) *y*, real, intent(out) *z*)

Returns the position and spherical radius calculated with respect to the center of the grid

Parameters

integer	[in] i : cell index in the x direction
integer [in] j : cell index in the y direction	
integer	[in] k : cell index in the z direction
real	[in] x : x position in the grid
real	[in] y: y position in the grid
real	[in] z : z position in the grid

Definition at line 188 of file coldens.f90.

4.2.2.3 subroutine coldens_utilities::init_coldens()

Initializes data, MPI and other stuff

Definition at line 36 of file coldens.f90.

4.2.2.4 subroutine coldens_utilities::read_data (real, dimension(neq,nxmin:nxmax,nymin:nymax,nzmin:nzmax), intent(out) u, integer, intent(in) itprint, character (len=128), intent(in) filepath)

reads data from file

Parameters

real [out] u(neq,nxmin:nxmax,nymin:nymax,nzmin:nzmax) : conserved variables	
integer	[in] itprint : number of output
string	[in] filepath: path where the output is

Definition at line 135 of file coldens.f90.

4.2.2.5 subroutine coldens_utilities::rotation_x (real, intent(in) theta, real, intent(in) x, real, intent(in) y, real, intent(in) z, real, intent(out) xn, real, intent(out) yn, real, intent(out) zn)

Does a rotation around the x axis

Parameters

real	[in], theta: Angle of rotation (in radians)
real	[in], x : original x position in the grid
real	[in], y : original y position in the grid
real	[in], x : original z position in the grid
real	[out], x : final x position in the grid
real	[out], y: final y position in the grid
real	[out], x : final z position in the grid

Definition at line 214 of file coldens.f90.

4.2.2.6 subroutine coldens_utilities::rotation_y (real, intent(in) theta, real, intent(in) x, real, intent(in) y, real, intent(in) z, real, intent(out) xn, real, intent(out) yn, real, intent(out) zn)

Does a rotation around the x axis

Parameters

real	[in], theta: Angle of rotation (in radians)
real	[in], x : original x position in the grid
real	[in], y : original y position in the grid
real	[in], x : original z position in the grid
real	[out], x : final x position in the grid
real	[out], y: final y position in the grid
real	[out], x : final z position in the grid

Definition at line 238 of file coldens.f90.

4.2.2.7 subroutine coldens_utilities::rotation_z (real, intent(in) *theta*, real, intent(in) *x*, real, intent(in) *y*, real, intent(in) *y*, real, intent(out) *yn*, real, intent(out) *yn*, real, intent(out) *zn*)

Does a rotation around the x axis

Parameters

real	[in], theta: Angle of rotation (in radians)
real	[in], x : original x position in the grid
real	[in], y : original y position in the grid
real	[in], x : original z position in the grid
real	[out], x : final x position in the grid
real	[out], y: final y position in the grid

,	
real	[out], x : final z position in the grid
rour	[[odf], X : midi z position in the grid

Definition at line 260 of file coldens.f90.

4.2.2.8 subroutine coldens_utilities::write_map (character (len=128), intent(in) *fileout*, integer, intent(in) *nxmap*, integer, intent(in) *nymap*, real, dimension(nxmap,nymap), intent(in) *map*)

Writes projection to file

Parameters

integer	[in] itprint : number of output
string	[in] fileout : file where to write
integer	[in] nxmap : Number of X cells in target
integer	[in] nymap : Number of Y cells in target
real	[in] map(nxmap,mymap) : Target map

Definition at line 340 of file coldens.f90.

4.3 constants Module Reference

real, parameter pi =acos(-1.)

Module containing physical and asronomical constants.

Variables

```
• real, parameter amh =1.66e-24
     hydrogen mass
• real, parameter mu =0.5
     mean atomic mass
• real, parameter kb =1.38e-16
     Boltzmann constant (cgs)
• real, parameter rg =8.3145e7
     Gas constant (cgs)
• real, parameter ggrav =6.67259e-8
     Gravitational constant (cgs)
• real, parameter clight =2.99E10
     speed of light in vacuum (cgs)
• real, parameter msun =1.99E33
     solar radius (cgs)
• real, parameter rsun =6.955e10
     solar mass (cgs)
• real, parameter mjup =1.898E30
     Jupiter mass (cgs)
• real, parameter rjup =7.1492E9
     Jupiter radius (cgs)
• real, parameter au =1.496e13
```

1AU in cm

1pc in cm

• real, parameter pc =3.0857E18

real, parameter kpc =3.0857E21

1Kpc in cm

• real, parameter hr =3600.

1hr in seconds

• real, parameter day =86400.

1day in seconds

• real, parameter yr =3.1536E7

1yr in seconds

• real, parameter myr =3.1536E13

1Myr in seconds

4.4 cooling_chi Module Reference

Cooling module with CHIANTI generated cooling curves.

Functions/Subroutines

• subroutine read_table ()

Reads the cooling curve table.

• real(kind=8) function coolchi (T)

Returns the cooling coefficient interpolating the table.

subroutine coolingchi ()

High level wrapper to apply cooling with CHIANTI tables.

Variables

• real(kind=8), dimension(2, 41) cooltab

4.4.1 Detailed Description

Cooling module with CHIANTI generated cooling curves
The location of the tables is assumed to be in src/CHIANTIIib/coolingCHIANTI.tab

4.4.2 Function/Subroutine Documentation

4.4.2.1 real (kind=8) function cooling_chi::coolchi (real, intent(in) T)

Parameters

```
real [in] T : Temperature K
```

Definition at line 75 of file cooling_chi.f90.

4.4.2.2 subroutine cooling_chi::coolingchi()

High level wrapper to apply cooling with CHIANTI tables cooling is applied in the entire domain and updates both the conserved and primitive variables

Definition at line 102 of file cooling_chi.f90.

Here is the call graph for this function:

4.5 cooling_dmc Module Reference

Cooling module with Dalgarno McCray coronal cooling curve.

Functions/Subroutines

• subroutine read_table ()

Reads the cooling curve table.

• real(kind=8) function cooldmc (T)

Returns the cooling coefficient interpolating the table.

• subroutine coolingdmc ()

High level wrapper to apply cooling with DMC table.

Variables

• real(kind=8), dimension(2, 41) cooltab

4.5.1 Detailed Description

Cooling module with Dalgarno McCray coronal cooling curve
The location of the tables is assumed to be in src/DMClib/coolingDMC.tab, it is read by init subroutine

4.5.2 Function/Subroutine Documentation

4.5.2.1 real (kind=8) function cooling_dmc::cooldmc (real, intent(in) T)

Parameters

```
real | [in] T : Temperature K
```

Definition at line 77 of file cooling dmc.f90.

4.5.2.2 subroutine cooling_dmc::coolingdmc ()

High level wrapper to apply cooling with DMC table cooling is applied in the entire domain and updates both the conserved and primitive variables

Definition at line 103 of file cooling_dmc.f90.

Here is the call graph for this function:

4.5.2.3 subroutine cooling_dmc::read_table ()

Reads the Dalgarno McCray cooling courve the location is assumed in src/DMClib/coolingDMC.tab, it is read by init subroutine

Definition at line 45 of file cooling dmc.f90.

4.6 cooling_h Module Reference

Cooling with parametrized cooling and H rate equation.

Functions/Subroutines

• subroutine coolingh ()

High level wrapper to apply cooling.

• real(kind=8) function alpha (T)

calculates the recombination rate (case B)

• real(kind=8) function alpha1 (T)

calculates the recombination rate to level 1

real(kind=8) function colf (T)

calculates the collisional ionization rate

real(kind=8) function betah (T)

betaH(T)

real(kind=8) function aloss (X1, X2, DT, DEN, DH0, TE0)

Non equilibrium cooling.

• subroutine atomic (dt, uu, tau, radphi)

Updates the ionization fraction and applpies cooling.

4.6.1 Detailed Description

Cooling with parametrized cooling and H rate equation

4.6.2 Function/Subroutine Documentation

4.6.2.1 real (kind=8) function cooling_h::aloss (real (kind=8), intent(in) X1, real (kind=8), intent(in) X2, real, intent(in) DT, real (kind=8), intent(in) DEN, real (kind=8), intent(in) DEN, real (kind=8), intent(in) TEO)

Non-equilibrium energy loss for low temperatures considering the collisional excitation of [O I] and [O II] lines and radiative recombination of H. This cooling rate is multiplied by a factor of 7.033 so that it has the same value as the "coronal equilibrium" cooling rate at a temperature of 44770 K (at temperatures higher than this value, the equilibrium cooling rate is used). The collisional ionization of H and excitation of Lyman-alpha are computed separately, and added to the cooling rate.

Parameters

real8	[in] x1 : initial H ionization fraction
real8 [in] x2 : final H ionization fraction	
real	[in] dt : timestep
real8	[in] den: total density of hydrogen
real8	[in] dh0 : density of neutral hydrogen
real8	[in] Te0 : Temperature

Definition at line 164 of file cooling_h.f90.

Here is the call graph for this function:



4.6.2.2 real (kind=8) function cooling_h::alpha (real (kind=8), intent(in) $\it T$)

calculates the recombination rate (case B)

Parameters

real8	[in] T : Temperature K	

Definition at line 80 of file cooling_h.f90.

4.6.2.3 real (kind=8) function cooling_h::alpha1 (real (kind=8), intent(in) T)

calculates the recombination rate to level 1

Parameters

real8	[in] T : Temperature K
-------	------------------------

Definition at line 97 of file cooling_h.f90.

4.6.2.4 subroutine cooling_h::atomic (real, intent(in) *dt*, real, dimension(neq), intent(out) *uu*, real, intent(in) *tau*, real, intent(in) *radphi*)

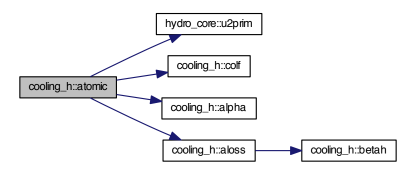
Calculates the new ionization state and energy density using a time dependent ionization calculation and an approximate time dependent cooling calculation

Parameters

real	[in] dt : timestep (seconds)
real	[in] uu(neq) : conserved variablas in one cell
real	[in] tau : optical depth (not in use)
real	[in] radphi : photoionizing rate

Definition at line 264 of file cooling_h.f90.

Here is the call graph for this function:



4.6.2.5 real (kind=8) function cooling_h::betah (real (kind=8), intent(in) T)

 $\beta_H(T)$

Parameters

real 8[in] T : Temperature K

Definition at line 130 of file cooling_h.f90.

4.6.2.6 real (kind=8) function cooling_h::colf (real (kind=8), intent(in) T)

calculates the collisional ionization rate

Parameters

real8[in] T: Temperature K

Definition at line 113 of file cooling_h.f90.

4.6.2.7 subroutine cooling_h::coolingh()

High level wrapper to apply cooling parametrized cooling curve, uses the ionization state of hydrogen and ties the O I and II to it

Definition at line 42 of file cooling_h.f90.

Here is the call graph for this function:

4.7 difrad Module Reference

Ray tracing Radiative Trasnport.

Functions/Subroutines

• subroutine init_rand ()

initializes random number generation

• subroutine emdiff (emax)

calculates the diffuse fotoionization emissivity

• subroutine random_versor (xd, yd, zd)

returns the 3 components of a random versor

• subroutine starsource (srad, x0, y0, z0, x, y, z, xd, yd, zd)

Place photon packets at a "star" surface.

• subroutine photons (xl0, yl0, zl0, xd, yd, zd, f)

Photon trajectories.

• subroutine radbounds ()

follows the rays across MPI boundaries

• subroutine progress (j, tot)

Progress bar.

• subroutine diffuse_rad ()

Diffuse radiation driver.

Variables

• real, parameter a0 =6.3e-18

Fotoionization cross section.

• integer, parameter nrays =1000000

Number of rays.

real, dimension(:,:,:), allocatable ph

Photoionizing rate.

• real, dimension(:,:,:), allocatable em

Photoionizing emissivity.

real, dimension(:,:,:), allocatable photl
 Auxiliary buffer for MPI.

• real, dimension(:,:,:), allocatable photr

Auxiliary buffer for MPI.

real, dimension(:,:,:), allocatable photb

Auxiliary buffer for MPI.

• real, dimension(:,:,:), allocatable phott

Auxiliary buffer for MPI.

real, dimension(:,:,:), allocatable photo

Auxiliary buffer for MPI.

• real, dimension(:,:,:), allocatable photi

Auxiliary buffer for MPI.

• integer, dimension(6) buffersize

Auxiliary buffer for MPI.

4.7.1 Detailed Description

Ray tracing Radiative Trasnport

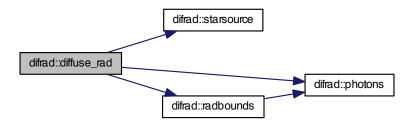
4.7.2 Function/Subroutine Documentation

4.7.2.1 subroutine difrad::diffuse_rad ()

Upper level wrapper to compute the diffuse photoionization rate

Definition at line 657 of file difrad.f90.

Here is the call graph for this function:



4.7.2.2 subroutine difrad::emdiff (real, intent(out) emax)

calculates the diffuse fotoionization emissivity in the entire domain

Parameters

real	[out] emax : maximum emissivity in the entire grid
------	--

Definition at line 98 of file difrad.f90.

Here is the call graph for this function:



4.7.2.3 subroutine difrad::init_rand()

initializes random number generation

Definition at line 56 of file difrad.f90.

4.7.2.4 subroutine difrad::photons (real, intent(in) x10, real, intent(in) y10, real, intent(in) z10, real, intent(in) xd, real, intent(in) xd, real, intent(in) t) yd, real, intent(in) zd, real, intent(in) t)

Launches a photon from cell (xc,yc,zc) in the (xd,yd,zd) direction, with f and ionizing photons, and updates the photoionizing rate

Parameters

real	[in] xl0 : Initial X position
real	[in] yl0 : Initial Y position
real	[in] zl0 : Initial Z position
real	[in] xd : Direction in X
real	[in] yd : Direction in Y
real	[in] zd : Direction in Z
real	[in] f : NUmber of photoionizong photons

Definition at line 252 of file difrad.f90.

4.7.2.5 subroutine difrad::progress (integer(kind=4) j, integer(kind=4), intent(in) tot)

Progress bar (only tested with Fortran compiler) takes a number between 1 and tot

Parameters

integer	[in] j : current iteration
integer	[in] tot : total number of iterartions

Definition at line 635 of file difrad.f90.

4.7.2.6 subroutine difrad::radbounds ()

follows the rays across MPI boundaries

Definition at line 455 of file difrad.f90.

Here is the call graph for this function:



4.7.2.7 subroutine difrad::random_versor (real, intent(out) xd, real, intent(out) yd, real, intent(out) zd)

returns the 3 components of a random versor (unit magnitude)

Parameters

real	[out] xd : x component
real	[out] yd : y component
real	[out] zd : z component

Definition at line 149 of file difrad.f90.

4.7.2.8 subroutine difrad::starsource (real, intent(in) *srad*, real, intent(in) *x0*, real, intent(in) *y0*, real, intent(in) *z0*, real, intent(out) *x*, real, intent(out) *y*, real, intent(out) *yd*, real, intent(out) *yd*, real, intent(out) *zd*)

returns the random location and direction at a star surface, if the direction goes into the star, the direction is inverted Parameters

real	[in] Srad : radius of the "star"
real	[in] x0 : X position of the center of the star
real	[in] y0 : Y position of the center of the star
real	[in] y0 : Z position of the center of the star
real	[out] x : random X position at the star surface
real	[out] y : random Y position at the star surface
real	[out] z : random Z position at the star surface
real	[out] xd : random X direction
real	[out] yd : random Y direction
real	[out] zd : random Z direction

Definition at line 187 of file difrad.f90.

4.8 globals Module Reference

Module containing global variables.

Variables

- real, dimension(:,:,:,:), allocatable u
 conserved varibles
- real, dimension(:,:,:,:), allocatable up conserved varibles after 1/2 timestep

```
• real, dimension(:,:,:,:), allocatable primit
      primitive varibles
• real, dimension(:,:,:,:), allocatable f
      X fluxes.
• real, dimension(:,:,:), allocatable g
      Y fluxes.
• real, dimension(:,:,:), allocatable h
      Z fluxes.

 real dx

      grid spacing in X

 real dy

      grid spacing in Y

 real dz

      grid spacing in Z
• integer, dimension(0:2) coords
      position of neighboring MPI blocks
· integer left
      MPI neighbor in the -x direction.
· integer right
      MPI neighbor in the +x direction.

    integer top

      MPI neighbor in the -y direction.

    integer bottom

      MPI neighbor in the +y direction.
· integer out
      MPI neighbor in the -z direction.

    integer in

      MPI neighbor in the +z direction.

    integer rank

      MPI rank.
· integer comm3d
      Cartessian MPI comunicator.
· real time
      Current time.

    real dt_cfl

      Current CFL $ t$.
• integer currentiteration
      Current iteration.

    real, dimension(:,:,:), allocatable temp

      Temperature array [K].
```

4.8.1 Detailed Description

This mudules contains variables that are treated as global in the code

4.9 h_alpha_utilities Module Reference

H alpha projection.

Functions/Subroutines

• subroutine init_ha ()

Initializes data.

• subroutine read_data (u, itprint, filepath)

reads data from file

• subroutine getxyz (i, j, k, x, y, z)

gets position of a cell

• subroutine rotation_x (theta, x, y, z, xn, yn, zn)

Rotation around the X axis.

• subroutine rotation_y (theta, x, y, z, xn, yn, zn)

Rotation around the Y axis.

• subroutine rotation_z (theta, x, y, z, xn, yn, zn)

Rotation around the Z axis.

• subroutine fill map (nxmap, nymap, u, map, dxT, dyT, theta x, theta y, theta z)

Fill target map.

• subroutine write_ha (fileout, nxmap, nymap, map)

Writes projection to file.

• subroutine write_rg (fileout, nxmap, nymap, map)

Writes projection to file in rg format.

4.9.1 Detailed Description

Utilities to compute an H alpha map

4.9.2 Function/Subroutine Documentation

4.9.2.1 subroutine h_alpha_utilities::fill_map (integer, intent(in) nxmap, integer, intent(in) nymap, real, dimension(neq,nxmin:nxmax,nymin:nymax, nzmin:nzmax), intent(in) u, real, dimension(nxmap,nymap), intent(out) map, real, intent(in) dxT, real, intent(in) dyT, real, intent(in) theta_x, real, intent(in) theta_y, real, intent(in) theta_z)

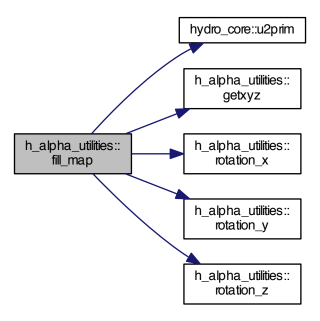
Fills the target map of one MPI block

Parameters

integer	[in] nxmap : Number of X cells in target
integer	[in] nymap : Number of Y cells in target
real	[in] u(neq,nxmin:nxmax,nymin:nymax, nzmin:nzmax) : conserved variables
real	[out] map(nxmap,mymap) : Target map
real	[in] dxT: target pixel width
real	[in] dyT: target pixel height
real	[in] thetax : Rotation around X
real	[in] thetay: Rotation around Y
real	[in] thetaz : Rotation around Z

Definition at line 285 of file h_alpha_proj.f90.

Here is the call graph for this function:



4.9.2.2 subroutine h_alpha_utilities::getxyz (integer, intent(in) *i*, integer, intent(in) *j*, integer, intent(in) *k*, real, intent(out) *x*, real, intent(out) *y*, real, intent(out) *z*)

Returns the position and spherical radius calculated with respect to the center of the grid

Parameters

integer	[in] i : cell index in the x direction
integer	[in] j : cell index in the y direction
integer	[in] k : cell index in the z direction
real	[in] x : x position in the grid
real	[in] y: y position in the grid
real	[in] z : z position in the grid

Definition at line 187 of file h_alpha_proj.f90.

 $4.9.2.3 \quad subroutine \ h_alpha_utilities::init_ha\ (\quad)$

Initializes data, MPI and other stuff

Definition at line 35 of file h_alpha_proj.f90.

4.9.2.4 subroutine h_alpha_utilities::read_data (real, dimension(neq,nxmin:nxmax,nymin:nymax,nzmin:nzmax), intent(out) *u*, integer, intent(in) *itprint*, character (len=128), intent(in) *filepath*)

reads data from file

Parameters

real	[out] u(neq,nxmin:nxmax,nymin:nymax,nzmin:nzmax) : conserved variables
integer	[in] itprint : number of output
string	[in] filepath : path where the output is

Definition at line 134 of file h_alpha_proj.f90.

4.9.2.5 subroutine h_alpha_utilities::rotation_x (real, intent(in) *theta,* real, intent(in) *x,* real, intent(in) *y,* real, intent(in) *z,* real, intent(out) *xn,* real, intent(out) *yn,* real, intent(out) *zn*)

Does a rotation around the x axis

Parameters

real	[in], theta: Angle of rotation (in radians)
real	[in], x : original x position in the grid
real	[in], y : original y position in the grid
real	[in], x : original z position in the grid
real	[out], x : final x position in the grid
real	[out], y: final y position in the grid
real	[out], x : final z position in the grid

Definition at line 213 of file h_alpha_proj.f90.

4.9.2.6 subroutine h_alpha_utilities::rotation_y (real, intent(in) theta, real, intent(in) x, real, intent(in) y, real, intent(in) z, real, intent(out) xn, real, intent(out) yn, real, intent(out) zn)

Does a rotation around the x axis

Parameters

real	[in], theta : Angle of rotation (in radians)
real	[in], x : original x position in the grid
real	[in], y : original y position in the grid
real	[in], x : original z position in the grid
real	[out], x : final x position in the grid
real	[out], y: final y position in the grid
real	[out], x : final z position in the grid

Definition at line 237 of file h_alpha_proj.f90.

4.9.2.7 subroutine h_alpha_utilities::rotation_z (real, intent(in) *theta*, real, intent(in) *x*, real, intent(in) *y*, real, intent(in) *y*, real, intent(in) *y*, real, intent(out) *yn*, real, intent(out) *zn*)

Does a rotation around the x axis

Parameters

real	[in], theta: Angle of rotation (in radians)
real	[in], x : original x position in the grid
real	[in], y : original y position in the grid
real	[in], x : original z position in the grid
real	[out], x : final x position in the grid
real	[out], y: final y position in the grid

4.10 hll Module Reference 25

real	[out], x : final z position in the grid

Definition at line 259 of file h_alpha_proj.f90.

4.9.2.8 subroutine h_alpha_utilities::write_ha (character (len=128), intent(in) *fileout*, integer, intent(in) *nxmap*, integer, intent(in) *nymap*, real, dimension(nxmap,nymap), intent(in) *map*)

Writes projection to file

Parameters

integer	[in] itprint : number of output
string	[in] fileout : file where to write
integer	[in] nxmap : Number of X cells in target
integer	[in] nymap : Number of Y cells in target
real	[in] map(nxmap,mymap) : Target map

Definition at line 362 of file h_alpha_proj.f90.

4.9.2.9 subroutine h_alpha_utilities::write_rg (character (len=128), intent(in) *fileout*, integer, intent(in) *nxmap*, integer, intent(in) *nymap*, real, dimension(nxmap,nymap), intent(in) *map*)

Writes projection to file

Parameters

integer	[in] itprint : number of output
string	[in] fileout : file where to write
integer	[in] nxmap : Number of X cells in target
integer	[in] nymap : Number of Y cells in target
real	[in] map(nxmap,mymap) : Target map

Definition at line 391 of file h_alpha_proj.f90.

4.10 hll Module Reference

HLL approximate Riemann solver module.

Functions/Subroutines

- subroutine prim2fhl (prim1, primr, ff)
 - Solves the Riemann problem at the interface PL,PR using the HLL solver.
- subroutine hllfluxes (choice)

Calculates HLL fluxes from the primitive variables on all the domain.

4.10.1 Detailed Description

The module contains the routines needed to Solve the Riemann problem in the entire domain and return the physical fluxes in x,y,z with the HLL solver

4.10.2 Function/Subroutine Documentation

4.10.2.1 subroutine hll::hllfluxes (integer, intent(in) choice)

Calculates HLL fluxes from the primitive variables on all the domain

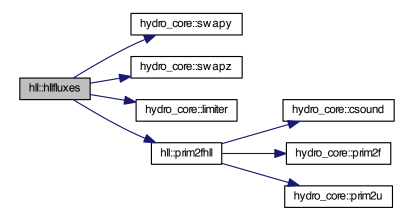
4.10 hll Module Reference 27

Parameters

integer	[in] choice: 1, uses primit for the 1st half of timestep (first order)
	2 uses primit for second order timestep

Definition at line 93 of file hll.f90.

Here is the call graph for this function:



4.10.2.2 subroutine hll::prim2fhll (real, dimension(neq), intent(in) *priml*, real, dimension(neq), intent(in) *primr*, real, dimension(neq), intent(inout) *ff*)

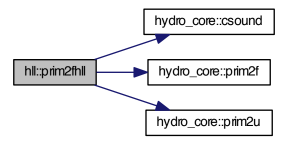
Solves the Riemann problem at the interface betweem PL and PR using the HLL solver The fluxes are computed in the X direction, to obtain the y ans z directions a swap is performed

Parameters

real	[in] primL : primitives at the Left state
real	[in] primR : primitives at the Right state
real	[out] ff : fluxes at the interface ($F_{i+1/2}$)

Definition at line 48 of file hll.f90.

Here is the call graph for this function:



4.11 hllc Module Reference

HLLC approximate Riemann solver module.

Functions/Subroutines

• subroutine prim2fhllc (priml, primr, ff)

Solves the Riemann problem at the interface PL,PR using the HLLC solver.

• subroutine hllcfluxes (choice)

Calculates HLLC fluxes from the primitive variables on all the domain.

4.11.1 Detailed Description

The module contains the routines needed to Solve the Riemann problem in the entire domain and return the physical fluxes in x,y,z with the HLLC solver

4.11.2 Function/Subroutine Documentation

4.11.2.1 subroutine hllc::hllcfluxes (integer, intent(in) choice)

Calculates HLLC fluxes from the primitive variables on all the domain

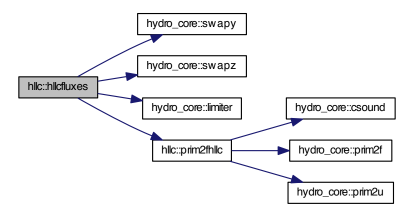
Parameters

integer	[in] choice: 1, uses primit for the 1st half of timestep (first order)
	2 uses primit for second order timestep

Definition at line 144 of file hllc.f90.

4.12 hlld Module Reference 29

Here is the call graph for this function:



4.11.2.2 subroutine hllc::prim2fhllc (real, dimension(neq), intent(in) *priml*, real, dimension(neq), intent(in) *primr*, real, dimension(neq), intent(inout) *ff*)

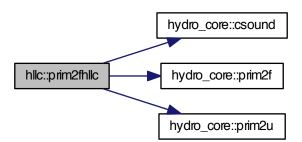
Solves the Riemann problem at the interface betweem PL and PR using the HLLC solver The fluxes are computed in the X direction, to obtain the y ans z directions a swap is performed

Parameters

rea	[in] primL : primitives at the Left state
rea	[in] primR : primitives at the Right state
rea	[out] ff : fluxes at the interface ($F_{i+1/2}$)

Definition at line 47 of file hllc.f90.

Here is the call graph for this function:



4.12 hlld Module Reference

HLLD approximate Riemann solver module.

Functions/Subroutines

• subroutine prim2fhlld (priml, primr, ff)

Solves the Riemann problem at the interface PL,PR using the HLLD solver.

subroutine hlldfluxes (choice)

Calculates HLLD fluxes from the primitive variables on all the domain.

4.12.1 Detailed Description

The module contains the routines needed to Solve the Riemann problem in the entire domain and return the physical fluxes in x,y,z with the HLLD solver

4.12.2 Function/Subroutine Documentation

4.12.2.1 subroutine hlld::hlldfluxes (integer, intent(in) choice)

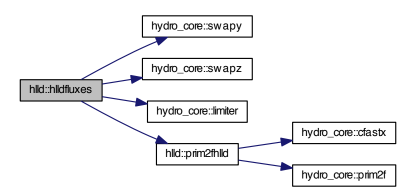
Calculates HLLD fluxes from the primitive variables on all the domain

Parameters

integer	[in] choice: 1, uses primit for the 1st half of timestep (first order)
	2 uses primit for second order timestep

Definition at line 328 of file hlld.f90.

Here is the call graph for this function:



4.12.2.2 subroutine hlld::prim2fhlld (real, dimension(neq), intent(in) *priml*, real, dimension(neq), intent(in) *primr*, real, dimension(neq), intent(inout) *ff*)

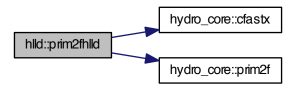
Solves the Riemann problem at the interface betweem PL and PR using the HLLD solver The fluxes are computed in the X direction, to obtain the y ans z directions a swap is performed 4.13 hlle Module Reference 31

Parameters

real	[in] primL : primitives at the Left state
real	[in] primR: primitives at the Right state
real	[out] ff : fluxes at the interface ($F_{i+1/2}$)

Definition at line 49 of file hlld.f90.

Here is the call graph for this function:



4.13 hlle Module Reference

HLLE approximate Riemann solver module.

Functions/Subroutines

• subroutine prim2fhlle (priml, primr, ff)

Solves the Riemann problem at the interface PL,PR using the HLLE solver.

• subroutine hllefluxes (choice)

Calculates HLLE fluxes from the primitive variables on all the domain.

4.13.1 Detailed Description

The module contains the routines needed to Solve the Riemann problem in the entire domain and return the physical fluxes in x,y,z with the HLLE solver

4.13.2 Function/Subroutine Documentation

4.13.2.1 subroutine hlle::hllefluxes (integer, intent(in) choice)

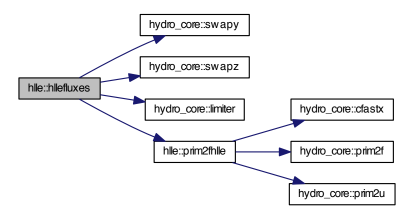
Calculates HLLE fluxes from the primitive variables on all the domain

Parameters

integer	[in] choice: 1, uses primit for the 1st half of timestep (first order)
	2 uses primit for second order timestep

Definition at line 94 of file hlle.f90.

Here is the call graph for this function:



4.13.2.2 subroutine hlle::prim2fhlle (real, dimension(neq), intent(in) *priml*, real, dimension(neq), intent(in) *primr*, real, dimension(neq), intent(inout) *ff*)

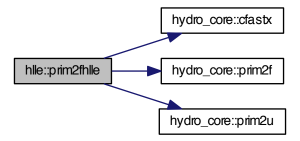
Solves the Riemann problem at the interface betweem PL and PR using the HLLE solver The fluxes are computed in the X direction, to obtain the y ans z directions a swap is performed

Parameters

real	[in] primL : primitives at the Left state
real	[in] primR: primitives at the Right state
real	[out] ff: fluxes at the interface ($F_{i+1/2}$)

Definition at line 49 of file hlle.f90.

Here is the call graph for this function:



4.14 hydro_core Module Reference

Basic hydro (and MHD) subroutines utilities.

Functions/Subroutines

subroutine u2prim (uu, prim, T)

Computes the primitive variables and temperature from conserved variables on a single cell.

• subroutine calcprim (u, primit)

Updated the primitives, using the conserved variables in the entire domain.

• subroutine prim2u (prim, uu)

Computes the conserved conserved variables from the primitives in a single cell.

• subroutine prim2f (prim, ff)

Computes the Euler Fluxes in one cell.

• subroutine swapy (var, neq)

Swaps the x and y components in a cell.

subroutine swapz (var, neq)

Swaps the x and z components in a cell.

• subroutine csound (p, d, cs)

Computes the sound speed.

subroutine cfast (p, d, bx, by, bz, cfx, cfy, cfz)

Computes the fast magnetosonic speeds in the 3 coordinates.

• subroutine cfastx (prim, cfX)

Computes the fast magnetosonic speed in the x direction.

subroutine get timestep (current iter, n iter, current time, tprint, dt, dump flag)

Otains the timestep allowed by the CFL condition in the entire.

• subroutine limiter (PLL, PL, PR, PRR, neq)

Performs a linear reconstruction of the primitive variables.

4.14.1 Detailed Description

This module contains subroutines and utilities that are the core of the hydro (and MHD) that are common to most implementations and will be used for the different specific solvers

4.14.2 Function/Subroutine Documentation

4.14.2.1 subroutine hydro_core::calcprim (real, dimension(neq,nxmin:nxmax,nymin:nymax,nzmin:nzmax), intent(in) *u*, real, dimension(neq,nxmin:nxmax,nymin:nymax,nzmin:nzmax), intent(out) *primit*)

Updated the primitives, using the conserved variables in the entire domain

Parameters

real	[in] u(neq,nxmin:nxmax,nymin:nymax,nzmin:nzmax) : conserved variables
real	[out] prim(neq,nxmin:nxmax,nymin:nymax,nzmin:nzmax) : primitive variables

Definition at line 119 of file hydro_core.f90.

Here is the call graph for this function:



4.14.2.2 subroutine hydro_core::cfast (real, intent(in) p, real, intent(in) d, real, intent(in) bx, real, intent(in) by, real, intent(in) bz, real, intent(out) cfx, real, intent(out) cfy, real, intent(out) cfz

Computes the fast magnetosonic speeds in the 3 coordinates

Parameters

real	[in] p : value of pressure
real	[in] d : value of density
real	[in] Bx : value of the x component of the magnetic field
real	[in] By : value of the y component of the magnetic field
real	[in] Bz : value of the z component of the magnetic field
real	[out] csx : fast magnetisonic speed in x
real	[out] csy : fast magnetisonic speed in y
real	[out] csz : fast magnetisonic speed in z

Definition at line 327 of file hydro_core.f90.

4.14.2.3 subroutine hydro_core::cfastx (real, dimension(neq), intent(in) prim, real, intent(out) cfX)

Computes the fast magnetosonic speed in the x direction

Parameters

real	[in] prim(neq) : vector with the primitives in one cell

Definition at line 352 of file hydro_core.f90.

4.14.2.4 subroutine hydro_core::csound (real, intent(in) p, real, intent(in) d, real, intent(out) cs)

Computes the sound speed

Parameters

real	[in] p : value of pressure
real	[in] d : value of density
real	[out] cs : sound speed

Definition at line 301 of file hydro_core.f90.

4.14.2.5 subroutine hydro_core::get_timestep (integer, intent(in) *current_iter*, integer, intent(in) *n_iter*, real, intent(in) *current_time*, real, intent(in) *tprint*, real, intent(out) *dt*, logical, intent(out) *dump_flag*)

Otains the timestep allowed by the CFL condition in the entire domain using the global primitives, and sets logical variable to dump output

Parameters

integer	[in] current_iter: Current iteration, it starts with a small but increasing CFL in the first N_trans
	iterarions
integer	[in] n_iter: Number of iterations to go from a small CFL to the final CFL (in parameters.f90)
real	[in] current_time : Current (global) simulation time
real	[in] tprint : time for the next programed disk dump
real	[out] : Δt allowed by the CFL condition
logical	[out] dump_flag : Flag to write to disk

Definition at line 384 of file hydro_core.f90.

Here is the call graph for this function:

4.14.2.6 subroutine hydro_core::limiter (real, dimension(neq), intent(in) *PLL*, real, dimension(neq), intent(inout) *PL*, real, dimension(neq), intent(in) *PRR*, integer, intent(in) *neq*)

returns a linear reconstruction of the variables at the interface beteen the primitives PLL, PL, PR, PRR The reconstruction is made with a slope limiter chosen at compilation time (i.e. set on the Makefile)

Parameters

real	[in] : primitives at the left of the left state
real	[inout] : primitives at the left state
real	[inout] : primitives at the right state
real	[in] : primitives at the right of the right state
real	[in] : number of equations

Definition at line 462 of file hydro_core.f90.

4.14.2.7 subroutine hydro_core::prim2f (real, dimension(neq), intent(in) prim, real, dimension(neq), intent(out) ff)

Computes the Euler Fluxes in one cell, using the primitices

It returns the flux in the x direction (i.e. F), the y and z fluxes can be obtained swaping the respective entries (see swapy and swapz subroutines)

Parameters

real	[in] prim(neq) : primitives in one cell
real	[out] ff(neq) : Euler Fluxes (x direction)

Definition at line 199 of file hydro_core.f90.

4.14.2.8 subroutine hydro_core::prim2u (real, dimension(neq), intent(in) prim, real, dimension(neq), intent(out) uu)

Computes the conserved conserved variables from the primitives in a single cell

Parameters

real	[in] prim(neq) : primitives in one cell
real	[out] uu(neq) : conserved varibles in one cell

Definition at line 158 of file hydro_core.f90.

4.14.2.9 subroutine hydro_core::swapy (real, dimension(neq), intent(inout) var, integer, intent(in) neq)

Swaps the x and y components in a cell.

Parameters

real	[inout] var(neq) : variable to be swapped
real	[in] neq : number of equations in the code

Definition at line 249 of file hydro_core.f90.

4.14.2.10 subroutine hydro_core::swapz (real, dimension(neq), intent(inout) var, integer, intent(in) neq)

Swaps the x and z components in a cell.

Parameters

real	[inout] var(neq) : variable to be swapped
real	[in] neq : number of equations in the code

Definition at line 275 of file hydro_core.f90.

4.14.2.11 subroutine hydro_core::u2prim (real, dimension(neq), intent(in) *uu*, real, dimension(neq), intent(out) *prim*, real, intent(out) *T*)

Computes the primitive variables and temperature from conserved variables on a single cell

Parameters

real	[in] uu(neq) : conserved variables in one cell
real	[out] prim(neq) : primitives in one cell
real	[out] T : Temperature [K]

Definition at line 44 of file hydro_core.f90.

4.15 hydro_solver Module Reference

Advances the simulation one timestep.

Functions/Subroutines

• subroutine viscosity ()

Adds artificial viscosity to the conserved variables.

• subroutine step (dt)

Upwind timestep.

• subroutine tstep ()

High level wrapper to advancce the simulation.

4.15.1 Detailed Description

Advances the solution from t to $t+\Delta t$

4.15.2 Function/Subroutine Documentation

4.15.2.1 subroutine hydro_solver::step (real, intent(in) dt)

Performs the upwind timestep according to

$$U_i^{n+1} = U_i^n - \frac{\Delta t}{\Delta x} \left[F_{i+1/2}^{n+1/2} - F_{i-1/2}^{n+1/2} \right]$$

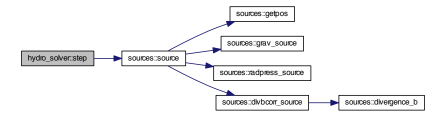
(in 3D), it takes U^{n+1} =up from the global variables and U^n =u

Parameters

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Definition at line 82 of file hydro_solver.f90.

Here is the call graph for this function:



4.15.2.2 subroutine hydro_solver::tstep ()

High level wrapper to advancce the simulation

The variables are taken from the globals module.

Definition at line 124 of file hydro_solver.f90.

Here is the call graph for this function:

```
4.15.2.3 subroutine hydro_solver::viscosity ( )
```

Adds artificial viscosity to the conserved variables

Takes the variables from the globals module and it assumes that the up are the stepped variables, while u are unstepped

Definition at line 52 of file hydro_solver.f90.

4.16 init Module Reference

Guacho-3D initialization.

Functions/Subroutines

• subroutine initmain (tprint, itprint)

Main initialization routine.

subroutine initflow (itprint)

Initializes the conserved variables, in the globals module.

4.16.1 Detailed Description

This module contains the routines needed to initializa the code, it also initializes all the modules set by the user.

4.16.2 Function/Subroutine Documentation

4.16.2.1 subroutine init::initflow (integer, intent(inout) itprint)

Initializes the conserved variables, in the globals module

Parameters

real [inout] itprint : number of current output

Definition at line 435 of file init.f90.

4.16.2.2 subroutine init::initmain (real, intent(out) tprint, integer, intent(out) itprint)

This subsroutine initializes all the variables in the globals module, MPI, cooling and user_mod routines; and outputs to screen the main parameters used in the run

real	[out] tprint : time of next output
integer	[out] itprint : number of next output

Definition at line 41 of file init.f90.

Here is the call graph for this function:

4.17 lyman_alpha_utilities Module Reference

Lyman_alpha_utilities.

Functions/Subroutines

• subroutine init_la ()

Initializes data.

• subroutine read_data (u, itprint, filepath)

reads data from file

• subroutine getxyz (i, j, k, x, y, z)

gets position of a cell

• subroutine rotation_x (theta, x, y, z, xn, yn, zn)

Rotation around the X axis.

• subroutine rotation_y (theta, x, y, z, xn, yn, zn)

Rotation around the Y axis.

• subroutine rotation_z (theta, x, y, z, xn, yn, zn)

Rotation around the Z axis.

- subroutine fill_map (nxmap, nymap, nvmap, vmin, vmax, u, map, dxT, dyT, theta_x, theta_y, theta_z) Fill target map.
- subroutine write_la (itprint, filepath, nxmap, nymap, nvmap, map)

Writes projection to file.

• subroutine phigauss (T, vzn, vmin, vmax, nvmap, profile)

This routine computes a gaussian line profile.

4.17.1 Detailed Description

Utilities to compute the Lyman-

4.17.2 Function/Subroutine Documentation

4.17.2.1 subroutine lyman_alpha_utilities::fill_map (integer, intent(in) nxmap, integer, intent(in) nymap, integer, intent(in) nvmap, real, intent(in) vmin, real, intent(in) vmax, real, dimension(neq,nxmin:nxmax,nymin:nymax, nzmin:nzmax), intent(in) u, real, dimension(nxmap,nymap,nvmap), intent(out) map, real, intent(in) dxT, real, intent(in) dyT, real, intent(in) theta_x, real, intent(in) theta_y, real, intent(in) theta_z)

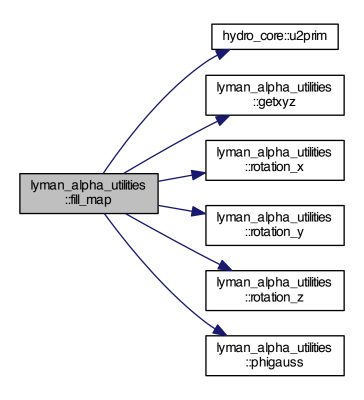
Fills the target map of one MPI block

integer	[in] nxmap : Number of X cells in target

integer	[in] nymap : Number of Y cells in target
real	[in] u(neq,nxmin:nxmax,nymin:nymax, nzmin:nzmax) : conserved variables
real	[out] map(nxmap,mymap) : Target map
real	[in] dxT: target pixel width
real	[in] dyT: target pixel height
real	[in] thetax : Rotation around X
real	[in] thetay: Rotation around Y
real	[in] thetaz : Rotation around Z

Definition at line 285 of file lyman_alpha_tau.f90.

Here is the call graph for this function:



4.17.2.2 subroutine lyman_alpha_utilities::getxyz (integer, intent(in) *i*, integer, intent(in) *j*, integer, intent(in) *k*, real, intent(out) *x*, real, intent(out) *y*, real, intent(out) *z*)

Returns the position and spherical radius calculated with respect to the center of the grid

integer	[in] i : cell index in the x direction
integer	[in] j : cell index in the y direction
integer	[in] k : cell index in the z direction
real	[in] x : x position in the grid
real	[in] y: y position in the grid
real	[in] z : z position in the grid

Definition at line 186 of file lyman_alpha_tau.f90.

4.17.2.3 subroutine lyman_alpha_utilities::init_la ()

Initializes data, MPI and other stuff

Definition at line 36 of file lyman_alpha_tau.f90.

4.17.2.4 subroutine lyman_alpha_utilities::phigauss (real, intent(in) *T*, real, intent(in) *vzn*, real, intent(in) *vmax*, integer, intent(in) *nvmap*, real, dimension(nvmap), intent(out) *profile*)

This routine computes a gaussian line profile

Definition at line 386 of file lyman_alpha_tau.f90.

4.17.2.5 subroutine lyman_alpha_utilities::read_data (real, dimension(neq,nxmin:nxmax,nymin:nymax,nzmin:nzmax), intent(out) u, integer, intent(in) itprint, character (len=128), intent(in) filepath)

reads data from file

Parameters

real	[out] u(neq,nxmin:nxmax,nymin:nymax,nzmin:nzmax) : conserved variables
integer	[in] itprint : number of output
string	[in] filepath: path where the output is

Definition at line 136 of file lyman_alpha_tau.f90.

4.17.2.6 subroutine lyman_alpha_utilities::rotation_x (real, intent(in) *theta*, real, intent(in) *x*, real, intent(in) *y*, real, intent(out) *yn*, real, intent(out) *yn*, real, intent(out) *zn*)

Does a rotation around the x axis

Parameters

real	[in], theta : Angle of rotation (in radians)
real	[in], x : original x position in the grid
real	[in], y : original y position in the grid
real	[in], x : original z position in the grid
real	[out], x : final x position in the grid
real	[out], y: final y position in the grid
real	[out], x : final z position in the grid

Definition at line 212 of file lyman_alpha_tau.f90.

4.17.2.7 subroutine lyman_alpha_utilities::rotation_y (real, intent(in) theta, real, intent(in) x, real, intent(in) y, real, intent(in) z, real, intent(out) xn, real, intent(out) yn, real, intent(out) zn)

Does a rotation around the x axis

real	[in], theta: Angle of rotation (in radians)
real	[in], x : original x position in the grid
real	[in], y : original y position in the grid
real	[in], x : original z position in the grid
real	[out], x : final x position in the grid
real	[out], y: final y position in the grid
real	[out], x : final z position in the grid

Definition at line 236 of file lyman_alpha_tau.f90.

4.17.2.8 subroutine lyman_alpha_utilities::rotation_z (real, intent(in) *theta*, real, intent(in) *x*, real, intent(in) *y*, real, intent(out) *yn*, real, intent(out) *yn*, real, intent(out) *zn*)

Does a rotation around the x axis

Parameters

real	[in], theta: Angle of rotation (in radians)
real	[in], x : original x position in the grid
real	[in], y : original y position in the grid
real	[in], x : original z position in the grid
real	[out], x : final x position in the grid
real	[out], y: final y position in the grid
real	[out], x : final z position in the grid

Definition at line 258 of file lyman alpha tau.f90.

4.17.2.9 subroutine lyman_alpha_utilities::write_la (integer, intent(in) *itprint*, character (len=128), intent(in) *filepath*, integer, intent(in) *nxmap*, integer, intent(in) *nvmap*, real, dimension(nxmap,nymap,nvmap), intent(in) *map*)

Writes projection to file

Parameters

integer	[in] itprint : number of output
string	[in] filepath: path where to write
integer	[in] nxmap : Number of X cells in target
integer	[in] nymap : Number of Y cells in target
integer	[in] nvmap : Number of velocity channels
real	[in] map(nxmap,mymap) : Target map

Definition at line 361 of file lyman_alpha_tau.f90.

4.18 out_bin_module Module Reference

Output in BIN format.

Functions/Subroutines

• subroutine write_header (unit, neq_out, nghost_out)

Writes header.

• subroutine write_bin (itprint)

Writes Data, one file per processor.

4.18.1 Detailed Description

This module writes the ouput in BIN format

4.18.2 Function/Subroutine Documentation

4.18.2.1 subroutine out_bin_module::write_bin (integer, intent(in) itprint)

Writes Data in BIN format one file per processor

Parameters

integer	[in] itprint : number of output
---------	---------------------------------

Definition at line 111 of file Out_BIN_Module.f90.

Here is the call graph for this function:



4.18.2.2 subroutine out_bin_module::write_header (integer, intent(in) *unit*, integer, intent(in) *neq_out*, integer, intent(in) *nghost_out*)

Writes header for binary input

Parameters

integer	[in] unit : number of logical unit

Definition at line 43 of file Out_BIN_Module.f90.

4.19 out_silo_module Module Reference

Output in Silo (+HDF5) Format.

Functions/Subroutines

• subroutine writeblocks (itprint)

Writes Data, one file per processor.

• subroutine writemaster (itprint)

Writes the Master File.

• subroutine outputsilo (itprint)

Upper level wrapper.

4.19.1 Detailed Description

This module writes the ouput in SILO (HDF5) format

4.19.2 Function/Subroutine Documentation

4.19.2.1 subroutine out_silo_module::outputsilo (integer, intent(in) itprint)

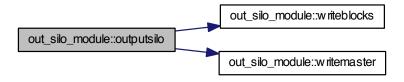
Upper level wrapper for the SILO output

Parameters

integer	[in] itprint : number of output

Definition at line 347 of file Out_Silo_Module.f90.

Here is the call graph for this function:



4.19.2.2 subroutine out_silo_module::writeblocks (integer, intent(in) itprint)

Writes Data in silo format one file per processor

Parameters

integer	[in] itprint : number of output

Definition at line 44 of file Out_Silo_Module.f90.

4.19.2.3 subroutine out_silo_module::writemaster (integer, intent(in) itprint)

Writes the master file with the metadata and multivars

Parameters

integer	[in] itprint : number of output

Definition at line 198 of file Out_Silo_Module.f90.

4.20 out_vtk_module Module Reference

Output in VTK format.

Functions/Subroutines

• subroutine write_vtk (itprint)

Writes Data, one file per processor.

4.20.1 Detailed Description

This module writes the ouput in VTK format

4.20.2 Function/Subroutine Documentation

4.20.2.1 subroutine out_vtk_module::write_vtk (integer, intent(in) itprint)

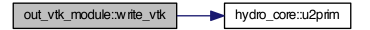
Writes Data in VTK format one file per processor

Parameters

integer	[in] itprint : number of output
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Definition at line 43 of file Out_VTK_Module.f90.

Here is the call graph for this function:



4.21 output Module Reference

Writes output.

Functions/Subroutines

• subroutine write_output (itprint)

Writes output.

4.21.1 Detailed Description

This module writes the ouput in the formats specified in the makefile

4.21.2 Function/Subroutine Documentation

4.21.2.1 subroutine output::write_output (integer, intent(in) itprint)

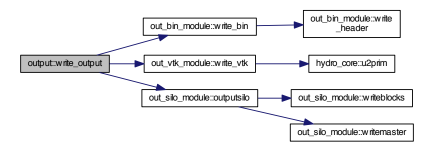
Writes output, the format is chosen in makefile Supported formats are *.bin and VTK (both BINARY), Silo (+hdf5)

Parameters

integer	[in] itprint : number of output

Definition at line 42 of file output.f90.

Here is the call graph for this function:



4.22 sources Module Reference

Adds source terms.

Functions/Subroutines

• subroutine getpos (i, j, k, x, y, z, r)

Gets position in the grid.

• subroutine grav_source (xc, yc, zc, pp, s)

Gravity due to point sources.

• subroutine radpress_source (i, j, k, xc, yc, zc, rc, pp, s)

Radiation pressure force.

• subroutine divergence_b (i, j, k, d)

Computes div(B)

• subroutine divbcorr_source (i, j, k, pp, s)

8 Wave source terms for div(B) correction

• subroutine source (i, j, k, prim, s)

Upper level wrapper for sources.

4.22.1 Detailed Description

This module adds the source terms from gravity, radiation pressure (not fully tested), and div(B) cleaning if the 8 wave scheme is used

4.22.2 Function/Subroutine Documentation

4.22.2.1 subroutine sources::divbcorr_source (integer, intent(in) *i*, integer, intent(in) *j*, integer, intent(in) *k*, real, dimension(neq), intent(in) *pp*, real, dimension(neq), intent(inout) *s*)

Adds terms proportional to div B in Faraday's Law, momentum equationand energy equation as propoes in Powell et al. 1999

Parameters

integer	[in] i : cell index in the X direction
integer	[in] j : cell index in the Y direction
integer	[in] k : cell index in the Z direction
real	[in] pp(neq) : vector of primitive variables
real	[out] s(neq) : vector with source terms

Definition at line 201 of file sources.f90.

Here is the call graph for this function:



4.22.2.2 subroutine sources::divergence_b (integer, intent(in) i, integer, intent(in) j, integer, intent(in) k, real, intent(out) d)

Computes div(B)

Parameters

integer	[in] i : cell index in the X direction
integer	[in] j : cell index in the Y direction
integer	[in] k : cell index in the Z direction
real	[out] d :: div(B)

Definition at line 178 of file sources.f90.

4.22.2.3 subroutine sources::getpos (integer, intent(in) *i*, integer, intent(in) *j*, integer, intent(in) *k*, real, intent(out) *x*, real, intent(out) *y*, real, intent(out) *z*, real, intent(out) *r*)

Gets the position and spherical radius calculated with respect to the center of the grid

Parameters

integer	[in] i : index in the X direction
integer	[in] j : index in the Y direction
integer	[in] k : index in the Z direction
real	[out] x : X position form the center of the grid (code units)
real	[out] y: Y position form the center of the grid (code units)
real	[out] z : Z position form the center of the grid (code units)
real	[out] r : Spherical radius form the center of the grid (code units)

Definition at line 55 of file sources.f90.

4.22.2.4 subroutine sources::grav_source (real, intent(in) xc, real, intent(in) yc, real, intent(in) zc, real, dimension(neq), intent(in) pp, real, dimension(neq), intent(inout) s)

Adds the gravitational force due to point particles, at this moment is fixed to two point sources (exoplanet)

Parameters

real	[in] xc : X position of the cell
real	[in] yc : Y position of the cell
real	[in] zc : Z position of the cell
real	[in] pp(neq) : vector of primitive variables
real	[out] s(neq) : vector with source terms

Definition at line 82 of file sources.f90.

4.22.2.5 subroutine sources::radpress_source (integer, intent(in) i, integer, intent(in) j, integer, intent(in) k, real, intent(in) xc, real, intent(in) yc, real, intent(in) zc, real, intent(in) rc, real, dimension(neq), intent(in) pp, real, dimension(neq), intent(inout) s)

Adds the radiaiton pressure force due to photo-ionization

Parameters

integer	[in] i : cell index in the X direction
integer	[in] j : cell index in the Y direction
integer	[in] k : cell index in the Z direction
real	[in] xc : X position of the cell
real	[in] yc : Y position of the cell
real	[in] zc : Z position of the cell
reak	[in] rc : $\sqrt{x^2 + y^2 + z^2}$
real	[in] pp(neq) : vector of primitive variables
real	[out] s(neq) : vector with source terms

Definition at line 140 of file sources.f90.

4.22.2.6 subroutine sources::source (integer, intent(in) *i,* integer, intent(in) *j,* integer, intent(in) *k,* real, dimension(neq), intent(in) *prim,* real, dimension(neq), intent(out) *s*)

Upper level wrapper for sources

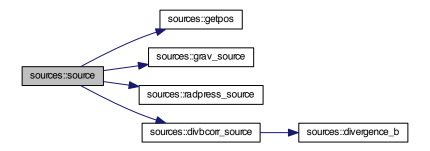
Main driver, this is called from the upwind stepping

Parameters

integer	[in] i : cell index in the X direction
integer	[in] j : cell index in the Y direction
integer	[in] k : cell index in the Z direction
real	[in] prim(neq): vector of primitive variables
real	[out] s(neq) : vector with source terms'

Definition at line 240 of file sources.f90.

Here is the call graph for this function:



4.23 thermal_cond Module Reference

Adds thermal conducion.

Functions/Subroutines

subroutine init_thermal_cond ()

Intializes Temperature array.

• subroutine get_dt_cond (dt)

computes conduction timescale

• subroutine progress (j, tot)

Progress bar.

• real function ksp (T)

Spitzer conductivity.

real function ksp_parl (xtemp)

Spitzer parallel conductivity.

• real function ksp_perp (xtemp, xdens, B2)

Spitzer perpendicular conductivity.

• subroutine heatfluxes ()

Returns Heat Fluxes.

• subroutine mhd_heatfluxes ()

Returns Heat Fluxes with anisotropic thermal conduction.

• subroutine thermal_bounds ()

Exchanges ghost cells for energy only.

• real function superstep (N, snu)

Length of superstep.

• real function substep (j, N, nu)

Size of substep j.

• subroutine st_steps (fs, Ns, fstep)

Returns the number of Supersteps.

• subroutine thermal_conduction ()

Upper level wrapper for thermal conduction.

Variables

• real, parameter ph =0.4

Parameter for the sturated regime in McKee.

• real, parameter nu =0.01

Super-stepping daMPI_NBg factor.

• real, parameter snu =sqrt(nu)

Sqrt of damping factor.

• integer, parameter max_iter = 100

Maximum number of iterations.

• real, parameter tstep_red_factor =0.25

timestep reduction factor for the conduction

real dt_cond

conduction timestep

integer tc_log

loical unit to write TC log

4.23.1 Detailed Description

Adds a thermal conduction term, affects both the primitive and conserved variables

4.23.2 Function/Subroutine Documentation

4.23.2.1 subroutine thermal_cond::get_dt_cond (real, intent(out) dt)

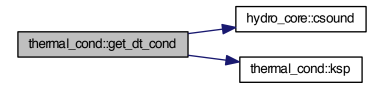
computes conduction timescale (in seconds)

Parameters

real	[out] dt :: conduction timescale

Definition at line 83 of file thermal cond.f90.

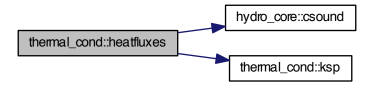
Here is the call graph for this function:



4.23.2.2 subroutine thermal_cond::heatfluxes ()

Heat flux, if saturation enabled it takes minimum of the Spitzer and the saturated value The result is stored in the 5th component of global the F,G,H fluxes (in cgs, conversion is done in dt product) Definition at line 194 of file thermal_cond.f90.

Here is the call graph for this function:



4.23.2.3 subroutine thermal_cond::init_thermal_cond ()

Intializes Temperature array (to resolve dependencies it was moved to the globals module)

Definition at line 55 of file thermal_cond.f90.

4.23.2.4 real function thermal_cond::ksp (real, intent(in) T)

Computes the Spitzer conductivity

Parameters

real [in] T : temperature [K]

Definition at line 147 of file thermal_cond.f90.

4.23.2.5 real function thermal_cond::ksp_parl (real, intent(in) xtemp)

Computes the Spitzer conductivity parallel to B

Parameters

real [in] T : temperature [K]

Definition at line 162 of file thermal_cond.f90.

4.23.2.6 real function thermal_cond::ksp_perp (real, intent(in) xtemp, real, intent(in) xdens, real, intent(in) B2)

Computes the Spitzer conductivity perpendicular to B

Parameters

real [in] T : temperature [K]

Definition at line 177 of file thermal_cond.f90.

4.23.2.7 subroutine thermal_cond::mhd_heatfluxes ()

Heat flux, if sturation enabled takes minimum of the Spitzer and the saturated value

The result is stored in the 5th component of global the F,G,H fluxes (in cgs, conversion is done in dt product)

Definition at line 285 of file thermal_cond.f90.

Here is the call graph for this function:

4.23.2.8 subroutine thermal_cond::progress (integer(kind=4) j, integer(kind=4), intent(in) tot)

Progress bar (only tested with intel Fortran compiler) takes a number between 1 and tot

Parameters

integer	[in] j : current iteration
integer	[in] tot: total number of iterartions

Definition at line 125 of file thermal cond.f90.

4.23.2.9 subroutine thermal_cond::st_steps (real, intent(in) fs, integer, intent(out) Ns, real, intent(out) fstep)

Returns the number of Supersteps

Parameters

real	fs : ratio of dtcond/dthydro
integer	Ns : Number of Supersteps
real	fstep : Number of supersteps (float)

Definition at line 674 of file thermal cond.f90.

Here is the call graph for this function:



4.23.2.10 real function thermal_cond::substep (integer, intent(in) j, integer, intent(in) N, real, intent(in) nu)

Returns the size of substep j of N

Parameters

integer	[in] j : index of current step
integer	[in] N : Total number of substeps
real	[in] nu : daMPI_NBg factor

Definition at line 656 of file thermal_cond.f90.

4.23.2.11 real function thermal_cond::superstep (integer N, real, intent(in) snu)

Returns the length of the superstep with N inner substeps

integer	[in] N : Nunber of inner substeps
real	[in] snu : sqrt of daMPI_NBg factor

Definition at line 635 of file thermal_cond.f90.

4.23.2.12 subroutine thermal_cond::thermal_bounds ()

Exchanges one layer of boundaries, only the equation that corresponds to the energy

Definition at line 508 of file thermal_cond.f90.

4.23.2.13 subroutine thermal_cond::thermal_conduction ()

This routine adds the heat conduction, receives the hydro timestep in seconds, and assumes the primitives and Temp(i,j,k) arrays are updated

Definition at line 700 of file thermal_cond.f90.

Here is the call graph for this function:

Chapter 5

File Documentation

- 5.1 /Users/esquivel/Desktop/Guacho-Working/doc/mainpage.h File Reference Webpage frontend.
- 5.2 /Users/esquivel/Desktop/Guacho-Working/src/boundaries.f90 File Reference
 Boundary conditions.

Modules

module boundaries
 Boundary conditions.

Functions/Subroutines

- subroutine boundaries::boundaryi ()
 - Boundary conditions for 1st order half timestep.
- subroutine boundaries::boundaryii ()

Boundary conditions for 2nd order half timestep.

5.2.1 Detailed Description

Author

Alejandro Esquivel

Date

2/Nov/2014

5.3 /Users/esquivel/Desktop/Guacho-Working/src/coldens.f90 File Reference

Column density projection.

58 File Documentation

Modules

• module coldens_utilities

Column densirt projection.

Functions/Subroutines

• subroutine coldens utilities::init coldens ()

Initializes data.

• subroutine coldens_utilities::read_data (u, itprint, filepath)

reads data from file

• subroutine coldens_utilities::getxyz (i, j, k, x, y, z)

gets position of a cell

• subroutine coldens_utilities::rotation_x (theta, x, y, z, xn, yn, zn)

Rotation around the X axis.

• subroutine coldens_utilities::rotation_y (theta, x, y, z, xn, yn, zn)

Rotation around the Y axis.

subroutine coldens_utilities::rotation_z (theta, x, y, z, xn, yn, zn)

Rotation around the Z axis.

• subroutine coldens_utilities::fill_map (nxmap, nymap, u, map, dxT, dyT, theta_x, theta_y, theta_z)

Fill target map.

• subroutine coldens_utilities::write_map (fileout, nxmap, nymap, map)

Writes projection to file.

· program coldens

Computes the H-alpha emission.

5.3.1 Detailed Description

Author

Alejandro Esquivel

Date

2/Nov/2014

5.3.2 Function/Subroutine Documentation

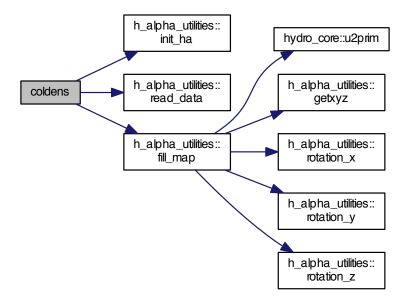
5.3.2.1 program coldens ()

Computes the H-alpha apbsorption

It rotates the data along each of the coordinates axis by an amount $\theta_x, \theta_y, \theta_z$, and projects the map along the the LOS, which is taken to be the Z axis

Definition at line 370 of file coldens.f90.

Here is the call graph for this function:



5.4 /Users/esquivel/Desktop/Guacho-Working/src/constants.f90 File Reference

Constants module.

Modules

· module constants

Module containing physical and asronomical constants.

Variables

• real, parameter constants::pi =acos(-1.)

 π

• real, parameter constants::amh =1.66e-24

hydrogen mass

• real, parameter constants::mu =0.5

mean atomic mass

• real, parameter constants::kb =1.38e-16

Boltzmann constant (cgs)

• real, parameter constants::rg =8.3145e7

Gas constant (cgs)

• real, parameter constants::ggrav =6.67259e-8

Gravitational constant (cgs)

• real, parameter constants::clight =2.99E10

speed of light in vacuum (cgs)

60 File Documentation

```
    real, parameter constants::msun =1.99E33
solar radius (cgs)
```

• real, parameter constants::rsun =6.955e10 solar mass (cgs)

• real, parameter constants::mjup =1.898E30

Jupiter mass (cgs)

• real, parameter constants::rjup =7.1492E9

Jupiter radius (cgs)

• real, parameter constants::au =1.496e13

1AU in cm

• real, parameter constants::pc =3.0857E18

1pc in cm

• real, parameter constants::kpc =3.0857E21

1Kpc in cm

• real, parameter constants::hr =3600.

1hr in seconds

• real, parameter constants::day =86400.

1day in seconds

real, parameter constants::yr =3.1536E7

1yr in seconds

• real, parameter constants::myr =3.1536E13

1Myr in seconds

5.4.1 Detailed Description

Author

Alejandro Esquivel

Date

2/Nov/2014

5.5 /Users/esquivel/Desktop/Guacho-Working/src/cooling_chi.f90 File Reference

Cooling module with CHIANTI generated cooling curves.

Modules

· module cooling_chi

Cooling module with CHIANTI generated cooling curves.

Functions/Subroutines

• subroutine cooling_chi::read_table ()

Reads the cooling curve table.

real(kind=8) function cooling_chi::coolchi (T)

Returns the cooling coefficient interpolating the table.

• subroutine cooling_chi::coolingchi ()

High level wrapper to apply cooling with CHIANTI tables.

Variables

• real(kind=8), dimension(2, 41) cooling_chi::cooltab

5.5.1 Detailed Description

Author

Alejandro Esquivel

Date

2/Nov/2014

5.6 /Users/esquivel/Desktop/Guacho-Working/src/cooling_dmc.f90 File Reference

Cooling module with Dlgarno Mac Cray coronal cooling curve.

Modules

module cooling_dmc

Cooling module with Dalgarno McCray coronal cooling curve.

Functions/Subroutines

• subroutine cooling_dmc::read_table ()

Reads the cooling curve table.

• real(kind=8) function cooling_dmc::cooldmc (T)

Returns the cooling coefficient interpolating the table.

• subroutine cooling_dmc::coolingdmc ()

High level wrapper to apply cooling with DMC table.

Variables

• real(kind=8), dimension(2, 41) cooling_dmc::cooltab

5.6.1 Detailed Description

Author

Alejandro Esquivel

Date

2/Nov/2014

5.7 /Users/esquivel/Desktop/Guacho-Working/src/cooling_h.f90 File Reference

Cooling with hydrogen rate parametrized cooling.

62 File Documentation

Modules

· module cooling h

Cooling with parametrized cooling and H rate equation.

Functions/Subroutines

```
    subroutine cooling h::coolingh ()
```

High level wrapper to apply cooling.

• real(kind=8) function cooling_h::alpha (T)

calculates the recombination rate (case B)

• real(kind=8) function cooling_h::alpha1 (T)

calculates the recombination rate to level 1

• real(kind=8) function cooling_h::colf (T)

calculates the collisional ionization rate

• real(kind=8) function cooling_h::betah (T)

betaH(T)

• real(kind=8) function cooling h::aloss (X1, X2, DT, DEN, DH0, TE0)

Non equilibrium cooling.

• subroutine cooling_h::atomic (dt, uu, tau, radphi)

Updates the ionization fraction and applpies cooling.

5.7.1 Detailed Description

Author

Alejandro Esquivel

Date

2/Nov/2014

5.8 /Users/esquivel/Desktop/Guacho-Working/src/difrad.f90 File Reference

Diffuse radiation module.

Modules

· module difrad

Ray tracing Radiative Trasnport.

Functions/Subroutines

• subroutine difrad::init rand ()

initializes random number generation

subroutine difrad::emdiff (emax)

calculates the diffuse fotoionization emissivity

• subroutine difrad::random_versor (xd, yd, zd)

returns the 3 components of a random versor

• subroutine difrad::starsource (srad, x0, y0, z0, x, y, z, xd, yd, zd)

Place photon packets at a "star" surface.

• subroutine difrad::photons (xl0, yl0, zl0, xd, yd, zd, f)

Photon trajectories.

• subroutine difrad::radbounds ()

follows the rays across MPI boundaries

• subroutine difrad::progress (j, tot)

Progress bar.

• subroutine difrad::diffuse_rad ()

Diffuse radiation driver.

Variables

• real, parameter difrad::a0 =6.3e-18

Fotoionization cross section.

• integer, parameter difrad::nrays =1000000

Number of rays.

• real, dimension(:,:,:), allocatable difrad::ph

Photoionizing rate.

• real, dimension(:,:,:), allocatable difrad::em

Photoionizing emissivity.

real, dimension(:,:,:), allocatable difrad::photl

Auxiliary buffer for MPI.

real, dimension(:,:,:), allocatable difrad::photr

Auxiliary buffer for MPI.

• real, dimension(:,:,:), allocatable difrad::photb

Auxiliary buffer for MPI.

• real, dimension(:,:,:), allocatable difrad::phott

Auxiliary buffer for MPI.

• real, dimension(:,:,:), allocatable difrad::photo

Auxiliary buffer for MPI.

• real, dimension(:,:,:), allocatable difrad::photi

Auxiliary buffer for MPI.

• integer, dimension(6) difrad::buffersize

Auxiliary buffer for MPI.

5.8.1 Detailed Description

Author

Alejandro Esquivel

Date

2/Nov/2014

5.9 /Users/esquivel/Desktop/Guacho-Working/src/globals.f90 File Reference

Global variables.

Modules

· module globals

Module containing global variables.

Variables

```
• real, dimension(:,:,:,:), allocatable globals::u
      conserved varibles

    real, dimension(:,:,:,:), allocatable globals::up

      conserved varibles after 1/2 timestep
• real, dimension(:,:,:), allocatable globals::primit
      primitive varibles
• real, dimension(:,:,:,:), allocatable globals::f
      X fluxes.
• real, dimension(:,:,:,:), allocatable globals::g
      Y fluxes.
• real, dimension(:,:,:,:), allocatable globals::h
      Z fluxes.
· real globals::dx
      grid spacing in X
· real globals::dy
      grid spacing in Y
· real globals::dz
      grid spacing in Z
• integer, dimension(0:2) globals::coords
      position of neighboring MPI blocks
· integer globals::left
      MPI neighbor in the -x direction.
· integer globals::right
      MPI neighbor in the +x direction.
· integer globals::top
      MPI neighbor in the -y direction.
• integer globals::bottom
      MPI neighbor in the +y direction.
· integer globals::out
      MPI neighbor in the -z direction.
· integer globals::in
      MPI neighbor in the +z direction.
• integer globals::rank
      MPI rank.
• integer globals::comm3d
```

Cartessian MPI comunicator.

• real, dimension(:,:,:), allocatable globals::temp

real globals::time
 Current time.

 real globals::dt_cfl

Current CFL \$ t\$.
• integer globals::currentiteration

Current iteration.

Temperature array [K].

5.9.1 Detailed Description

Author

Alejandro Esquivel

Date

2/Nov/2014

5.10 /Users/esquivel/Desktop/Guacho-Working/src/h_alpha_proj.f90 File Reference

H alpha projection.

Modules

module h_alpha_utilities
 H alpha projection.

Functions/Subroutines

• subroutine h_alpha_utilities::init_ha ()

Initializes data.

• subroutine h_alpha_utilities::read_data (u, itprint, filepath)

reads data from file

subroutine h_alpha_utilities::getxyz (i, j, k, x, y, z)

gets position of a cell

• subroutine h_alpha_utilities::rotation_x (theta, x, y, z, xn, yn, zn)

Rotation around the X axis.

• subroutine h_alpha_utilities::rotation_y (theta, x, y, z, xn, yn, zn)

Rotation around the Y axis.

• subroutine h_alpha_utilities::rotation_z (theta, x, y, z, xn, yn, zn)

Rotation around the Z axis.

• subroutine h_alpha_utilities::fill_map (nxmap, nymap, u, map, dxT, dyT, theta_x, theta_y, theta_z)

• subroutine h_alpha_utilities::write_ha (fileout, nxmap, nymap, map)

Writes projection to file.

• subroutine h_alpha_utilities::write_rg (fileout, nxmap, nymap, map)

Writes projection to file in rg format.

program h_alpha_proj

Computes the H-alpha emission.

5.10.1 Detailed Description

Author

Alejandro Esquivel

Date

2/Nov/2014

5.10.2 Function/Subroutine Documentation

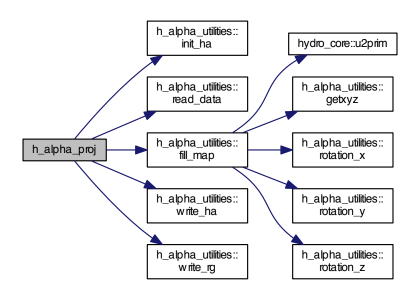
5.10.2.1 program h_alpha_proj ()

Computes the H-alpha apbsorption

It rotates the data along each of the coordinates axis by an amount $\theta_x, \theta_y, \theta_z$, and projects the map along the the LOS, which is taken to be the Z axis

Definition at line 428 of file h_alpha_proj.f90.

Here is the call graph for this function:



5.11 /Users/esquivel/Desktop/Guacho-Working/src/hll.f90 File Reference

HLL approximate Riemann solver module.

Modules

· module hll

HLL approximate Riemann solver module.

Functions/Subroutines

• subroutine hll::prim2fhll (priml, primr, ff)

Solves the Riemann problem at the interface PL,PR using the HLL solver.

subroutine hll::hllfluxes (choice)

Calculates HLL fluxes from the primitive variables on all the domain.

5.11.1 Detailed Description

Author

Alejandro Esquivel

Date

2/Nov/2014

5.12 /Users/esquivel/Desktop/Guacho-Working/src/hllc.f90 File Reference

HLLC approximate Riemann solver module.

Modules

· module hllc

HLLC approximate Riemann solver module.

Functions/Subroutines

• subroutine hllc::prim2fhllc (priml, primr, ff)

Solves the Riemann problem at the interface PL,PR using the HLLC solver.

• subroutine hllc::hllcfluxes (choice)

Calculates HLLC fluxes from the primitive variables on all the domain.

5.12.1 Detailed Description

Author

Alejandro Esquivel

Date

2/Nov/2014

5.13 /Users/esquivel/Desktop/Guacho-Working/src/hlld.f90 File Reference

HLLD approximate Riemann solver module.

Modules

module hlld

HLLD approximate Riemann solver module.

Functions/Subroutines

• subroutine hlld::prim2fhlld (priml, primr, ff)

Solves the Riemann problem at the interface PL,PR using the HLLD solver.

• subroutine hlld::hlldfluxes (choice)

Calculates HLLD fluxes from the primitive variables on all the domain.

5.13.1 Detailed Description

Author

C. Villarreal D'Angelo, A. Esquivel, M. Schneiter

Date

2/Nov/2014

5.14 /Users/esquivel/Desktop/Guacho-Working/src/hlle.f90 File Reference

HLLE approximate Riemann solver module.

Modules

· module hlle

HLLE approximate Riemann solver module.

Functions/Subroutines

• subroutine hlle::prim2fhlle (priml, primr, ff)

Solves the Riemann problem at the interface PL,PR using the HLLE solver.

• subroutine hlle::hllefluxes (choice)

Calculates HLLE fluxes from the primitive variables on all the domain.

5.14.1 Detailed Description

Author

C. Villarreal D'Angelo, A. Esquivel, M. Schneiter

Date

2/Nov/2014

5.15 /Users/esquivel/Desktop/Guacho-Working/src/hydro_core.f90 File Reference

Hydrodynamical and Magnetohidrodynamocal bacic module.

Modules

· module hydro_core

Basic hydro (and MHD) subroutines utilities.

Functions/Subroutines

• subroutine hydro core::u2prim (uu, prim, T)

Computes the primitive variables and temperature from conserved variables on a single cell.

• subroutine hydro_core::calcprim (u, primit)

Updated the primitives, using the conserved variables in the entire domain.

• subroutine hydro_core::prim2u (prim, uu)

Computes the conserved conserved variables from the primitives in a single cell.

• subroutine hydro_core::prim2f (prim, ff)

Computes the Euler Fluxes in one cell.

subroutine hydro_core::swapy (var, neq)

Swaps the x and y components in a cell.

• subroutine hydro_core::swapz (var, neq)

Swaps the x and z components in a cell.

• subroutine hydro_core::csound (p, d, cs)

Computes the sound speed.

• subroutine hydro_core::cfast (p, d, bx, by, bz, cfx, cfy, cfz)

Computes the fast magnetosonic speeds in the 3 coordinates.

• subroutine hydro_core::cfastx (prim, cfX)

Computes the fast magnetosonic speed in the x direction.

subroutine hydro_core::get_timestep (current_iter, n_iter, current_time, tprint, dt, dump_flag)

Otains the timestep allowed by the CFL condition in the entire.

• subroutine hydro core::limiter (PLL, PL, PR, PRR, neg)

Performs a linear reconstruction of the primitive variables.

• real function average (a, b)

5.15.1 Detailed Description

Author

Alejandro Esquivel

Date

2/Nov/2014

5.16 /Users/esquivel/Desktop/Guacho-Working/src/hydro_solver.f90 File Reference

Hydrodynamical and Magnetohidrodynamocal solver module.

Modules

· module hydro_solver

Advances the simulation one timestep.

Functions/Subroutines

• subroutine hydro_solver::viscosity ()

Adds artificial viscosity to the conserved variables.

subroutine hydro_solver::step (dt)

Upwind timestep.

subroutine hydro_solver::tstep ()

High level wrapper to advancce the simulation.

5.16.1 Detailed Description

Author

Alejandro Esquivel

Date

2/Nov/2014

5.17 /Users/esquivel/Desktop/Guacho-Working/src/init.f90 File Reference

Guacho-3D initialization module.

Modules

· module init

Guacho-3D initialization.

Functions/Subroutines

• subroutine init::initmain (tprint, itprint)

Main initialization routine.

• subroutine init::initflow (itprint)

Initializes the conserved variables, in the globals module.

5.17.1 Detailed Description

Author

Alejandro Esquivel

Date

2/Nov/2014

5.18 /Users/esquivel/Desktop/Guacho-Working/src/lyman_alpha_tau.f90 File Reference

Lyman_alpha_utilities.

Modules

· module lyman_alpha_utilities

Lyman_alpha_utilities.

Functions/Subroutines

• subroutine lyman_alpha_utilities::init_la ()

Initializes data.

• subroutine lyman_alpha_utilities::read_data (u, itprint, filepath)

reads data from file

• subroutine lyman_alpha_utilities::getxyz (i, j, k, x, y, z)

gets position of a cell

• subroutine lyman_alpha_utilities::rotation_x (theta, x, y, z, xn, yn, zn)

Rotation around the X axis.

• subroutine lyman_alpha_utilities::rotation_y (theta, x, y, z, xn, yn, zn)

Rotation around the Y axis.

• subroutine lyman_alpha_utilities::rotation_z (theta, x, y, z, xn, yn, zn)

Rotation around the Z axis.

• subroutine lyman_alpha_utilities::fill_map (nxmap, nymap, nvmap, vmin, vmax, u, map, dxT, dyT, theta_x, theta_y, theta_z)

Fill target map.

• subroutine lyman_alpha_utilities::write_la (itprint, filepath, nxmap, nymap, nvmap, map)

Writes projection to file.

• subroutine lyman alpha utilities::phigauss (T, vzn, vmin, vmax, nvmap, profile)

This routine computes a gaussian line profile.

• program lyman_alpha_tau

Computes the Ly-alpha apbsorption.

5.18.1 Detailed Description

Author

Alejandro Esquivel

Date

2/Nov/2014

5.18.2 Function/Subroutine Documentation

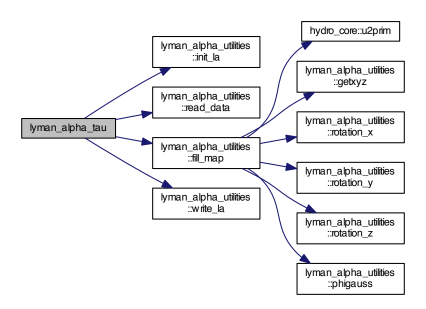
5.18.2.1 program lyman_alpha_tau ()

Computes the Ly-alpha apbsorption

It rotates the data along each of the coordinates axis by an amount $\theta_x, \theta_y, \theta_z$, and the LOS is along the Z axis

Definition at line 419 of file lyman_alpha_tau.f90.

Here is the call graph for this function:



5.19 /Users/esquivel/Desktop/Guacho-Working/src/main.f90 File Reference

Guacho-3D main program.

Functions/Subroutines

· program guacho

Guacho-3D Main Program This is the main program unit of the Guacho-3D code.

The code itegrates Euler equations in three dimensions, the choice of the integration method is set in the makefile. The flow (conserved) variables are taken to be:

ieq=

1 : rho (total)

2 : rho u

3 : rho v

4 : rho w

5 : Internal energy (thermal+kinetic)

6: bx (optional, if MHD or PMHD)

7: by (optional, if MHD or PMHD)

```
8: bz (optional, if MHD or PMHD)
additional variables advected into the flow, e.g.:
9 (6): n_HI
10 (7): n_HII
11 (8): n_HeI
12 (9): n_HeII
13 (10): n_HeIII
14 (11): rho*zbar
15 (12): ne
This can be changed bu the user according to cooling function for instance.
```

5.19.1 Detailed Description

Author

Alejandro Esquivel

Date

2/Nov/2014

5.20 /Users/esquivel/Desktop/Guacho-Working/src/Out_BIN_Module.f90 File Reference

Output in BIN Format.

Modules

module out_bin_module
 Output in BIN format.

Functions/Subroutines

subroutine out_bin_module::write_header (unit, neq_out, nghost_out)
 Writes header.

• subroutine out_bin_module::write_bin (itprint)

Writes Data, one file per processor.

5.20.1 Detailed Description

Author

Alejandro Esquivel

Date

2/Nov/2014

5.21 /Users/esquivel/Desktop/Guacho-Working/src/Out_Silo_Module.f90 File Reference

Output in Silo Format.

Modules

module out_silo_module
 Output in Silo (+HDF5) Format.

Functions/Subroutines

• subroutine out_silo_module::writeblocks (itprint)

Writes Data, one file per processor.

subroutine out_silo_module::writemaster (itprint)

Writes the Master File.

• subroutine out_silo_module::outputsilo (itprint)

Upper level wrapper.

5.21.1 Detailed Description

Author

Alejandro Esquivel

Date

2/Nov/2014

5.22 /Users/esquivel/Desktop/Guacho-Working/src/Out_VTK_Module.f90 File Reference

Output in VTK Format.

Modules

module out_vtk_module
 Output in VTK format.

Functions/Subroutines

subroutine out_vtk_module::write_vtk (itprint)
 Writes Data, one file per processor.

5.22.1 Detailed Description

Author

Alejandro Esquivel

Date

2/Nov/2014

5.23 /Users/esquivel/Desktop/Guacho-Working/src/output.f90 File Reference

Writes Output.

Modules

module output
 Writes output.

Functions/Subroutines

subroutine output::write_output (itprint)
 Writes output.

5.23.1 Detailed Description

Author

Alejandro Esquivel

Date

2/Nov/2014

5.24 /Users/esquivel/Desktop/Guacho-Working/src/sources.f90 File Reference

Adds source terms.

Modules

module sources

Adds source terms.

Functions/Subroutines

• subroutine sources::getpos (i, j, k, x, y, z, r)

Gets position in the grid.

• subroutine sources::grav_source (xc, yc, zc, pp, s)

Gravity due to point sources.

• subroutine sources::radpress_source (i, j, k, xc, yc, zc, rc, pp, s)

Radiation pressure force.

• subroutine sources::divergence_b (i, j, k, d)

Computes div(B)

• subroutine sources::divbcorr_source (i, j, k, pp, s)

8 Wave source terms for div(B) correction

• subroutine sources::source (i, j, k, prim, s)

Upper level wrapper for sources.

5.24.1 Detailed Description

Author

Alejandro Esquivel

Date

2/Nov/2014

5.25 /Users/esquivel/Desktop/Guacho-Working/src/thermal_cond.f90 File Reference

Thermal conduction module.

Modules

· module thermal cond

Adds thermal conducion.

Functions/Subroutines

• subroutine thermal_cond::init_thermal_cond ()

Intializes Temperature array.

subroutine thermal_cond::get_dt_cond (dt)

computes conduction timescale

• subroutine thermal_cond::progress (j, tot)

Progress bar.

real function thermal_cond::ksp (T)

Spitzer conductivity.

• real function thermal_cond::ksp_parl (xtemp)

Spitzer parallel conductivity.

• real function thermal_cond::ksp_perp (xtemp, xdens, B2)

Spitzer perpendicular conductivity.

• subroutine thermal_cond::heatfluxes ()

Returns Heat Fluxes.

• subroutine thermal_cond::mhd_heatfluxes ()

Returns Heat Fluxes with anisotropic thermal conduction.

• subroutine thermal cond::thermal bounds ()

Exchanges ghost cells for energy only.

• real function thermal_cond::superstep (N, snu)

Length of superstep.

• real function thermal cond::substep (j, N, nu)

Size of substep j.

• subroutine thermal_cond::st_steps (fs, Ns, fstep)

Returns the number of Supersteps.

subroutine thermal_cond::thermal_conduction ()

Upper level wrapper for thermal conduction.

Variables

• real, parameter thermal_cond::ph =0.4

Parameter for the sturated regime in McKee.

• real, parameter thermal_cond::nu =0.01

Super-stepping daMPI_NBg factor.

• real, parameter thermal_cond::snu =sqrt(nu)

Sqrt of damping factor.

• integer, parameter thermal_cond::max_iter = 100

Maximum number of iterations.

• real, parameter thermal cond::tstep red factor =0.25

timestep reduction factor for the conduction

• real thermal_cond::dt_cond

conduction timestep

• integer thermal_cond::tc_log

loical unit to write TC log

5.25.1 Detailed Description

Author

Alejandro Esquivel & Ernesto Zurbiggen

Date

07/Sep/2015

Index

/Users/esquivel/Desktop/Guacho-Working/doc/mainpage	⊹aloss
h, 55	cooling_h, 14
/Users/esquivel/Desktop/Guacho-Working/src/Out_BI ←	alpha
N_Module.f90, 71	cooling_h, 14
/Users/esquivel/Desktop/Guacho-Working/src/Out_	alpha1
Silo_Module.f90, 71	cooling_h, 16
/Users/esquivel/Desktop/Guacho-Working/src/Out_V TK_Module.f90, 72	atomic cooling_h, 16
/Users/esquivel/Desktop/Guacho-Working/src/boundaries. f90, 55	betah
/Users/esquivel/Desktop/Guacho-Working/src/coldens. ← f90, 55	cooling_h, 16 boundaries, 7
/Users/esquivel/Desktop/Guacho-Working/src/constants. ← f90, 57	boundaryi, 7 boundaryii, 7
/Users/esquivel/Desktop/Guacho-Working/src/cooling ←	boundaryi
_chi.f90, 58	boundaries, 7 boundaryii
/Users/esquivel/Desktop/Guacho-Working/src/cooling ←dmc.f90, 59	boundaries, 7
/Users/esquivel/Desktop/Guacho-Working/src/cooling ← h.f90, 59	calcprim
/Users/esquivel/Desktop/Guacho-Working/src/difrad. ←	hydro_core, 33 cfast
f90, 60	hydro_core, 33
/Users/esquivel/Desktop/Guacho-Working/src/globals. ← f90, 61	cfastx
/Users/esquivel/Desktop/Guacho-Working/src/h_←	hydro_core, 34
alpha_proj.f90, 63	coldens
/Users/esquivel/Desktop/Guacho-Working/src/hll.f90, 64	coldens.f90, 56 coldens.f90
/Users/esquivel/Desktop/Guacho-Working/src/hllc.f90,	coldens, 56
65	coldens_utilities, 7
/Users/esquivel/Desktop/Guacho-Working/src/hlld.f90, 65	fill_map, 8 getxyz, 9
/Users/esquivel/Desktop/Guacho-Working/src/hlle.f90,	init_coldens, 9
66 /Users/esquivel/Desktop/Guacho-Working/src/hydro_←	read_data, 9
core.f90, 66	rotation_x, 10 rotation_y, 10
/Users/esquivel/Desktop/Guacho-Working/src/hydro_← solver.f90, 67	rotation_z, 10
/Users/esquivel/Desktop/Guacho-Working/src/init.f90,	write_map, 11 colf
68	cooling_h, 17
/Users/esquivel/Desktop/Guacho-Working/src/lyman_← alpha_tau.f90, 68	constants, 11 coolchi
/Users/esquivel/Desktop/Guacho-Working/src/main.f90,	cooling_chi, 12 cooldmc
/Users/esquivel/Desktop/Guacho-Working/src/output.↔	cooling_dmc, 13
f90, 72	cooling_chi, 12
/Users/esquivel/Desktop/Guacho-Working/src/sources. ← f90, 73	coolchi, 12 coolingchi, 12
/Users/esquivel/Desktop/Guacho-Working/src/thermal ←	cooling_dmc, 13
cond.f90, 74	cooldmc, 13

80 INDEX

coolingdmc, 13	h_alpha_proj.f90
read table, 13	h_alpha_proj, 64
cooling_h, 13	h_alpha_utilities, 21
aloss, 14	fill_map, 22
alpha, 14	getxyz, 23
alpha1, 16	init_ha, 23
atomic, 16	read_data, 23
betah, 16	rotation_x, 24
colf, 17	rotation y, 24
coolingh, 17	rotation_z, 24
-	
coolingchi	write_ha, 25
cooling_chi, 12	write_rg, 25
coolingdmc	heatfluxes
cooling_dmc, 13	thermal_cond, 51
coolingh	hll, 25
cooling_h, 17	hllfluxes, 25
csound	prim2fhll, 27
hydro_core, 34	hllc, 28
	hllcfluxes, 28
diffuse_rad	prim2fhllc, 29
difrad, 18	hllcfluxes
difrad, 17	hllc, 28
diffuse rad, 18	hlld, 29
emdiff, 18	hlldfluxes, 30
init_rand, 19	prim2fhlld, 30
photons, 19	hlldfluxes
progress, 19	hlld, 30
radbounds, 19	hlle, 31
random_versor, 20	hllefluxes, 31
starsource, 20	prim2fhlle, 32
divbcorr_source	hllefluxes
sources, 47	hlle, 31
divergence_b	hllfluxes
sources, 48	hll, 25
	hydro_core, 32
emdiff	calcprim, 33
difrad, 18	cfast, 33
	cfastx, 34
fill_map	csound, 34
coldens_utilities, 8	get_timestep, 34
h_alpha_utilities, 22	limiter, 34
lyman_alpha_utilities, 39	prim2f, 36
	prim2u, <mark>36</mark>
get_dt_cond	swapy, 36
thermal_cond, 51	swapz, 36
get_timestep	u2prim, 36
hydro_core, 34	hydro_solver, 37
getpos	step, 37
sources, 48	•
getxyz	tstep, 37
coldens_utilities, 9	viscosity, 38
h_alpha_utilities, 23	init, 38
_ · _	initflow, 38
lyman_alpha_utilities, 40	initmain, 38
globals, 20	
grav_source	init_coldens
sources, 48	coldens_utilities, 9
h alaba and	init_ha
h_alpha_proj	h_alpha_utilities, 23
h_alpha_proj.f90, 64	init_la

INDEX 81

lyman_alpha_utilities, 40	prim2fhllc
init rand	hllc, 29
difrad, 19	prim2fhlld
init_thermal_cond	hlld, 30
thermal_cond, 52	prim2fhlle
initflow	hlle, 32
init, 38	prim2u
initmain	hydro_core, 36
init, 38	progress
init, 30	difrad, 19
ksp	thermal_cond, 53
thermal_cond, 52	thermal_cond, 55
ksp_parl	radbounds
thermal_cond, 52	difrad, 19
	radpress_source
ksp_perp	sources, 49
thermal_cond, 52	random_versor
limiter	
	difrad, 20
hydro_core, 34	read_data
lyman_alpha_tau	coldens_utilities, 9
lyman_alpha_tau.f90, 70	h_alpha_utilities, 23
lyman_alpha_tau.f90	lyman_alpha_utilities, 41
lyman_alpha_tau, 70	read_table
lyman_alpha_utilities, 39	cooling_dmc, 13
fill_map, 39	rotation_x
getxyz, 40	coldens_utilities, 10
init_la, 40	h_alpha_utilities, 24
phigauss, 40	lyman_alpha_utilities, 41
read_data, 41	rotation_y
rotation_x, 41	coldens_utilities, 10
rotation_y, 41	h_alpha_utilities, 24
rotation z, 41	lyman_alpha_utilities, 41
write_la, 42	rotation z
	coldens_utilities, 10
mhd heatfluxes	h_alpha_utilities, 24
thermal_cond, 52	lyman_alpha_utilities, 41
_ ,	,,
out bin module, 42	source
write bin, 42	sources, 49
write header, 43	sources, 47
out_silo_module, 43	divbcorr_source, 47
outputsilo, 43	divergence_b, 48
writeblocks, 44	getpos, 48
writemaster, 44	grav_source, 48
out_vtk_module, 44	radpress_source, 49
write_vtk, 45	source, 49
output, 46	st steps
·	thermal_cond, 53
write_output, 46	starsource
outputsilo	difrad, 20
out_silo_module, 43	
phiggues	step
phigauss	hydro_solver, 37
lyman_alpha_utilities, 40	substep
photons	thermal_cond, 53
difrad, 19	superstep
prim2f	thermal_cond, 53
hydro_core, 36	swapy
prim2fhll	hydro_core, 36
hll, 27	swapz

82 INDEX

```
hydro_core, 36
thermal_bounds
     thermal_cond, 54
thermal_cond, 50
     get_dt_cond, 51
     heatfluxes, 51
     init_thermal_cond, 52
     ksp, 52
     ksp_parl, 52
     ksp_perp, 52
     mhd_heatfluxes, 52
     progress, 53
     st_steps, 53
     substep, 53
     superstep, 53
     thermal\_bounds,\, \color{red} 54
     thermal_conduction, 54
thermal_conduction
     thermal cond, 54
tstep
     hydro_solver, 37
u2prim
     hydro_core, 36
viscosity
     hydro_solver, 38
write_bin
     out_bin_module, 42
write ha
     h_alpha_utilities, 25
write_header
     out bin module, 43
write la
     lyman_alpha_utilities, 42
write_map
     coldens_utilities, 11
write_output
     output, 46
write_rg
     h_alpha_utilities, 25
write_vtk
     out_vtk_module, 45
writeblocks
     out_silo_module, 44
writemaster
     out_silo_module, 44
```