# radmc3dPy

v0.25

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# **Contents**

1	Nam	espace	Index											1
	1.1	Packa	ges					 	 	 	 		 	 1
2	Clas	s Index												3
	2.1	Class	List					 	 	 	 		 	 3
3	Nam	espace	Docume	ntation										5
	3.1	radmc	3dPy Nam	espace Re	ference			 	 	 	 		 	 5
		3.1.1	Detailed	Description	1			 	 	 	 		 	 5
	3.2	radmc	3dPy.analy	ze Names	pace Ref	erence		 	 	 	 		 	 5
		3.2.1	Detailed	Description	1			 	 	 	 		 	 6
		3.2.2	Function	Document	ation .			 	 	 	 		 	 6
			3.2.2.1	getDens\	/struct			 	 	 	 		 	 6
			3.2.2.2	readData				 	 	 	 		 	 6
			3.2.2.3	readGrid				 	 	 	 		 	 7
			3.2.2.4	readOpa	C			 	 	 	 		 	 7
			3.2.2.5	readPara	ms			 	 	 	 		 	 8
			3.2.2.6	readSpec	ctrum .			 	 	 	 		 	 8
			3.2.2.7	writeDefa	ultParfile			 	 	 	 		 	 8
	3.3	radmc	3dPy.crd_f	rans Name	space R	eferenc	e	 	 	 	 		 	 8
		3.3.1	Detailed	Description	n			 	 	 	 		 	 9
		3.3.2	Function	Document	ation .			 	 	 	 		 	 9
			3.3.2.1	csrot				 	 	 	 		 	 9
			3.3.2.2	ctrans_sp	h2cart			 	 	 	 		 	 9
			3.3.2.3	ctrans_sp	oh2cyl .			 	 	 	 		 	 10
			3.3.2.4	vrot				 	 	 	 		 	 10
			3.3.2.5	vtrans_sp	oh2cart			 	 	 	 		 	 10
	3.4	radmc	3dPy.imag	e Namespa	ace Refer	rence		 	 	 	 		 	 10
		3.4.1		Description										
		3.4.2		Document										
			3.4.2.1	cmask .										
				netPSF										11

ii CONTENTS

		3.4.2.3	makeImage	. 12
		3.4.2.4	plotImage	. 12
		3.4.2.5	readImage	. 13
3.5	radmc	3dPy.mode	el_lines_nlte_lvg_1d_1 Namespace Reference	. 13
	3.5.1	Detailed	Description	. 13
	3.5.2	Function	Documentation	. 13
		3.5.2.1	getDefaultParams	. 13
		3.5.2.2	getDustDensity	. 14
		3.5.2.3	getDustTemperature	. 14
		3.5.2.4	getGasAbundance	. 14
		3.5.2.5	getGasDensity	. 14
		3.5.2.6	getGasTemperature	. 14
		3.5.2.7	getModelDesc	. 15
		3.5.2.8	getVelocity	. 15
		3.5.2.9	getVTurb	. 15
3.6	radmc	3dPy.mode	el_ppdisk Namespace Reference	. 15
	3.6.1	Detailed	Description	. 15
	3.6.2	Function	Documentation	. 16
		3.6.2.1	getDefaultParams	. 16
		3.6.2.2	getDustDensity	. 16
		3.6.2.3	getGasAbundance	. 16
		3.6.2.4	getGasDensity	. 17
		3.6.2.5	getModelDesc	. 17
		3.6.2.6	getVelocity	. 17
		3.6.2.7	getVTurb	. 17
3.7	radmc	3dPy.mode	el_simple_1 Namespace Reference	. 17
	3.7.1	Detailed	Description	. 17
	3.7.2	Function	Documentation	. 18
		3.7.2.1	getDefaultParams	. 18
		3.7.2.2	getDustDensity	. 18
		3.7.2.3	getModelDesc	. 18
3.8	radmc	3dPy.mode	el_spher1d_1 Namespace Reference	. 18
	3.8.1	Detailed	Description	. 18
	3.8.2	Function	Documentation	. 18
		3.8.2.1	getDefaultParams	. 18
		3.8.2.2	getDustDensity	. 19
		3.8.2.3	getModelDesc	. 19
3.9	radmc	3dPy.mode	el_spher2d_1 Namespace Reference	. 19
	3.9.1	Detailed	Description	. 19
	3.9.2	Function	Documentation	. 19

CONTENTS

		3.9.2.1	getDefaultParams	19
		3.9.2.2	getDustDensity	19
		3.9.2.3	getModelDesc	20
3.10	radmc3	BdPy.mode	el_template Namespace Reference	20
	3.10.1	Detailed	Description	20
	3.10.2	Function	Documentation	20
		3.10.2.1	getDefaultParams	20
		3.10.2.2	getDustDensity	21
		3.10.2.3	getDustTemperature	21
		3.10.2.4	getGasAbundance	21
		3.10.2.5	getGasDensity	21
		3.10.2.6	getGasTemperature	21
		3.10.2.7	getModelDesc	22
		3.10.2.8	getVelocity	22
		3.10.2.9	getVTurb	22
3.11	radmc3	BdPy.mode	el_test_scattering_1 Namespace Reference	22
	3.11.1	Detailed	Description	22
	3.11.2	Function	Documentation	22
		3.11.2.1	getDefaultParams	22
		3.11.2.2	getDustDensity	23
		3.11.2.3	getModelDesc	23
3.12	radmc3	BdPy.natco	onst Namespace Reference	23
	3.12.1	Detailed	Description	24
3.13	radmc3	BdPy.setup	Namespace Reference	24
	3.13.1	Detailed	Description	24
	3.13.2	Function	Documentation	24
		3.13.2.1	getModelDesc	24
		3.13.2.2	getModelNames	24
		3.13.2.3	getTemplateModel	25
		3.13.2.4	problemSetupDust	25
		3.13.2.5	problemSetupGas	25
		3.13.2.6	writeLinesInp	26
		3.13.2.7	writeRadmc3dInp	26
Clas	s Docu	mentation		27
4.1			'ze.radmc3dData Class Reference	
	4.1.1		Description	
	4.1.2		Function Documentation	
		4.1.2.1	getSigmaDust	
		4.1.2.2	getSigmaGas	
			goto-graduo	20

4

iv CONTENTS

		4.1.2.3	getTau	28
		4.1.2.4	getTauOneDust	28
		4.1.2.5	readDustDens	29
		4.1.2.6	readDustTemp	29
		4.1.2.7	readGasDens	29
		4.1.2.8	readGasTemp	29
		4.1.2.9	readGasVel	29
		4.1.2.10	readVTurb	30
		4.1.2.11	writeDustDens	30
		4.1.2.12	writeDustTemp	30
		4.1.2.13	writeGasDens	30
		4.1.2.14	writeGasTemp	30
		4.1.2.15	writeGasVel	31
		4.1.2.16	writeVTK	31
		4.1.2.17	writeVTurb	31
4.2	radmc	3dPy.analy	ze.radmc3dDustOpac Class Reference	31
	4.2.1	Detailed	Description	32
	4.2.2	Member	Function Documentation	32
		4.2.2.1	makeOpac	32
		4.2.2.2	mixOpac	33
		4.2.2.3	readMasterOpac	33
		4.2.2.4	readOpac	33
		4.2.2.5	runMakedust	33
		4.2.2.6	writeMasterOpac	34
4.3	radmc	3dPy.analy	ze.radmc3dGrid Class Reference	34
	4.3.1	Detailed	Description	34
	4.3.2	Member	Function Documentation	35
		4.3.2.1	getCellVolume	35
		4.3.2.2	makeSpatialGrid	35
		4.3.2.3	makeWavelengthGrid	35
		4.3.2.4	readGrid	35
		4.3.2.5	writeSpatialGrid	36
		4.3.2.6	writeWavelengthGrid	36
4.4	radmc	3dPy.image	e.radmc3dImage Class Reference	36
	4.4.1	Detailed	Description	37
	4.4.2	Member	Function Documentation	37
		4.4.2.1	getClosurePhase	37
		4.4.2.2	getMomentMap	37
		4.4.2.3	getVisibility	38
		4.4.2.4	imConv	38

CONTENTS

		4.4.2.5	plotMomentMap	38
		4.4.2.6	readImage	38
		4.4.2.7	writeFits	39
4.5	radmc	3dPy.analy	ze.radmc3dPar Class Reference	39
	4.5.1	Detailed	Description	40
	4.5.2	Member	Function Documentation	40
		4.5.2.1	loadDefaults	40
		4.5.2.2	printPar	40
		4.5.2.3	readPar	40
		4.5.2.4	setPar	41
		4.5.2.5	writeParfile	41
4.6	radmc	3dPy.analy	ze.radmc3dStars Class Reference	41
	4.6.1	Detailed	Description	42
	4.6.2	Member	Function Documentation	42
		4.6.2.1	findPeakStarspec	42
		4.6.2.2	getStellarSpectrum	42
		4.6.2.3	readStarsinp	42
		4.6.2.4	writeStarsinp	42
Index				42

# Chapter 1

# Namespace Index

# 1.1 Packages

Here are the packages with brief descriptions (if available):

radmc3dPy	5
radmc3dPy.analyze	5
radmc3dPy.crd_trans	8
radmc3dPy.image	
radmc3dPy.model_lines_nlte_lvg_1d_11	3
radmc3dPy.model_ppdisk	
radmc3dPy.model_simple_1 1	
radmc3dPy.model_spher1d_1 1	
radmc3dPy.model_spher2d_1 1	
radmc3dPy.model_template	
radmc3dPy.model_test_scattering_1	
radmc3dPy.natconst	23
radmc3dPv setup 2	۷4

2 Namespace Index

# **Chapter 2**

# **Class Index**

# 2.1 Class List

	Here are the classes,	structs,	unions	and	interfaces	with	brief	descri	ptions
--	-----------------------	----------	--------	-----	------------	------	-------	--------	--------

admc3dPy.analyze.radmc3dData	27
admc3dPy.analyze.radmc3dDustOpac	31
admc3dPy.analyze.radmc3dGrid	34
admc3dPy.image.radmc3dImage	36
admc3dPy.analyze.radmc3dPar	39
admc3dPy.analyze.radmc3dStars	41

Class Index

# **Chapter 3**

# **Namespace Documentation**

# 3.1 radmc3dPy Namespace Reference

### **Namespaces**

- · namespace analyze
- namespace crd\_trans
- namespace image
- namespace model\_lines\_nlte\_lvg\_1d\_1
- namespace model\_ppdisk
- namespace model\_simple\_1
- namespace model\_spher1d\_1
- namespace model\_spher2d\_1
- namespace model\_template
- namespace model\_test\_scattering\_1
- namespace natconst
- namespace setup

#### **Variables**

```
• string __version__ "0.25"
```

• list \_\_all\_\_ ["analyze", "setup", "image", "crd\_trans", "natconst"]

#### 3.1.1 Detailed Description

```
RADMC-3D Python module (c) Attila Juhasz, Leiden, 2011,2012,2013,2014
```

# 3.2 radmc3dPy.analyze Namespace Reference

### Classes

- class radmc3dGrid
- · class radmc3dData
- · class radmc3dStars
- · class radmc3dDustOpac
- · class radmc3dPar

#### **Functions**

- def readOpac
- · def readData
- · def readGrid
- def readParams
- def writeDefaultParfile
- def readSpectrum
- · def getDensVstruct

#### 3.2.1 Detailed Description

```
PYTHON module for RADMC3D
(c) Attila Juhasz 2011,2012,2013,2014

This sub-module contains classes and functions to read and write input/output data to/from RADMC3D

CLASSES:
-------
radmc3dData
radmc3dDustOpac
radmc3dSrid
radmc3dStars

FUNCTIONS:
-----
read_data()
readGrid()
readMasterOpac()
writeMasterOpac()
readOpac()
readOpac()
readOpac()
readOpac()
readOpac()
```

#### 3.2.2 Function Documentation

INPUT:

```
3.2.2.1 def radmc3dPy.analyze.getDensVstruct( data = None, vmean_temp = False, ispec_tgas = 0, gsize = [], idust = None, mstar = 0.)
```

Calculates the vertical hydrostatic equilibrium  ${\bf r}$ 

```
data - An instance of the radmc3DData class
  vmean_temp - If True (T(z) = T(-z) = 0.5*(T(z) + T(-z))) if False (T(z)!=T(-z))
  idust - List of dust indices whose structure must be calculated
  mstar - Stellar mass

OPTIONS:

-----

ispec_tgas - Index of dust species whose temperature is taken to be the gas temperature
  gsize - Dust grain sizes - If specified, the gas temperature is calculated as the average temperature
  of all dust grains in the grid cell weighted by the total surface area of dust grains with a
  size - NOTE: this approach assumes that all dust grains of a given size have the same bulk of
```

Returns a Numpy array with the dust density

3.2.2.2 def radmc3dPy.analyze.readData ( ddens = False, dtemp = False, gdens = False, gtemp = False, gvel = False, ispec = None, vturb = False, binary = True )

Function to read the model data (e.g. density, velocity, temperature)

```
INPUT:
    ddens - If True dust density will be read (all dust species and grain sizes)
    dtemp - If True dust temperature will be read (all dust species and grain sizes)
    gdens - If True gas density will be read (NOTE: the gas density will be number density in 1/cm^3)
    gtemp - If True gas temperature will be read (all dust species and grain sizes)
           If True the velocity field will be read
    ispec - Name of the molecule in the 'molecule_ispec.inp' filename
OUTPUT:
    Returns an instance of the radmc3dData class with the following attributes:
        rhodust - Dust density in g/cm^3
        dusttemp - Dust temperature in K
rhogas - Gas density in molecule/cm^3
        gasvel
                  - Gas velocity in cm/s
                     Gas temperature in K
        gastemp -
                  - Mictroturbulence in cm/s
        vturb
        taux
                  - Optical depth along the x (cartesian) / r (cylindrical) / r (spherical) dimension
                     Optical depth along the y (cartesian) / theta (cylindrical) / theta (spherical) dimension
        tauv
                  - Optical depth along the z (cartesian) / z (cylindrical) / phi (spherical) dimension
        sigmadust - Dust surface density in g/cm^2
        sigmagas - Gas surface density in molecule/cm^2 (or g/cm^2 depending on the dimension of rhogas)
3.2.2.3 def radmc3dPy.analyze.readGrid ( )
Function to read the spatial and frequency grid
OUTPUT
    Returns an instance of the radmc3dGrid class with the following attributes:
               - 'car'/'cyl'/'sph' coordinate system of the spatial grid
               - A three element vector the i-th element is 1 if the i-th dimension is active, otherwize the i
               - Number of grid points in the x (cartesian) / r (cylindrical) / r (spherical) dimension
               - Number of grid points in the y (cartesian) / theta (cylindrical) / theta (spherical) dimension
    nv
               - Number of grid points in the z (cartesian) / z (cylindrical) / phi (spherical) dimension
               - Number of cell interfaces in the x (cartesian) / r (cylindrical) / r (spherical) dimension - Number of cell interfaces in the y (cartesian) / theta (cylindrical) / theta (spherical) dimension
    nxi
    nyi
               - Number of cell interfaces in the z (cartesian) / z (cylindrical) / phi (spherical) dimension
               - Number of wavelengths in the wavelength grid
    nwav
    freq
               - Number of frequencies in the grid (equal to nwav)
               - Cell centered x (cartesian) / r (cylindrical) / r (spherical) grid points
               - Cell centered y (cartesian) / theta (cylindrical) / theta (spherical) \,\, grid points
    У
               - Cell centered z (cartesian) / z (cylindrical) / phi (spherical) grid points
               - Cell interfaces in the x (cartesian) / r (cylindrical) / r (spherical) dimension
    хi
               - Cell interfaces in the y (cartesian) / theta (cylindrical) / theta (spherical) dimension
    уi
    zi
               - Cell interfaces in the z (cartesian) / z (cylindrical) / phi (spherical) dimension
    wav
               - Wavelengh grid
    freq
               - Frequency grid
3.2.2.4 def radmc3dPy.analyze.readOpac ( ext = [''], idust = None )
Function to read the dust opacity files
This function is an interface to radmc3dDustOpac.readOpac()
INPUT:
    ext : file name extension (file names should look like 'dustkappa_ext.inp')
    idust: index of the dust species in the master opacity file (dustopac.inp')
OUTPUT:
    Returns an instance of the radmc3dDustOpac class with the following attributes:
    waw
            - wavelength grid
    freq
            - frequency grid
```

```
- number of wavelengths
    kabs
            - absorption coefficient per unit mass
    ksca
          - scattering coefficient per unit mass
    phase_g - phase function
            - if set it contains the file name extension of the duskappa_ext.Kappa file
    therm - if False the dust grains are quantum-heated (default: True)
    idust — index of the dust species in the dust density distribution array
3.2.2.5 def radmc3dPy.analyze.readParams ( )
Function to read the problem_params.inp file (interface function to radmc3dPar.readPar())
OUTPUT:
    Returns an instance of the radmc3dPar class with the following attributes:
           : Dictionary containing parameter values with parameter names as keys
    pdesc : Disctionary containing parameter description (comments in the parameter file) with parameter name
    pblock : Dictionary containing the block names in the parameter file and parameter names as values
    pvalstr: Dictionary containing parameter values as strings with parameter names as keys
3.2.2.6 def radmc3dPy.analyze.readSpectrum ( fname = ' ')
Function to read the spectrum / SED
OPTIONS:
   fname - Name of the file to be read
OUTPUT:
    Returns a two dimensional Numpy array with [Nwavelength, 2] dimensions
    [Nwavelength,0] is the wavelength / velocity and
    [Nwavelength,1] is the flux density
3.2.2.7 def radmc3dPy.analyze.writeDefaultParfile ( model = ' ' , fname = ' ' )
Function to write a parameter file (problem_params.inp) with default parameters for a given model
INPUT:
   model - Name of the model whose parameter should be written to the file
OPTIONS:
   fname - Name of the parameter file to be written (if omitted problem_params.inp will be used)
```

# 3.3 radmc3dPy.crd\_trans Namespace Reference

#### **Functions**

- def ctrans\_sph2cyl
- · def ctrans\_sph2cart
- · def vtrans sph2cart
- · def csrot
- def vrot

#### 3.3.1 Detailed Description

```
PYTHON module for RADMC3D
(c) Attila Juhasz 2011,2012,2013

This sub-module contains functions for coordinate transformations (e.g. rotation)

3.3.2 Function Documentation

3.3.2.1 def radmc3dPy.crd_trans.csrot ( crd = None, ang = None, xang = 0.0, yang = 0.0, zang = 0.0, deg = False )

Function to make coordinate system rotation
```

Rotation matrices :

```
| 1 0 0 |
| 0 cos(alpha) -sin(alpha) |
| 0 sin(alpha) cos(alpha) |
```

Y-axis

X-axis

INPUT:

Z-axis

```
| cos(gamma) -sin(gamma) 0 |
| sin(gamma) cos(gamma) 0 |
| 0 0 1
```

### 3.3.2.2 def radmc3dPy.crd\_trans.ctrans\_sph2cart ( crd = [0, reverse = False )

```
Function to transform coordinates between spherical to cartesian systems
```

```
3.3.2.3 def radmc3dPy.crd_trans.ctrans_sph2cyl ( crd = None, theta = None, reverse = False )
Function to transform coordinates between spherical to cylindrical systems
INPUT :
       r,phi,theta: numpy arrays containing the spherical coordinates
OPTIONS :
_____
       reverse=False : Calculates the inverse trasnformation
                   (cartesian -> spherical). In this case crd should be [r,phi,theta]
OUTPUT :
        result : a numpy array of [Nr, Nphi, Ntheta, 3] dimensions containing the cylindrical
                   coordinates [rcyl, z, phi]
3.3.2.4 def radmc3dPy.crd_trans.vrot ( crd = None, v = None, ang = None )
Function to rotate a vector in spherical coordinate system
First transform the vector to cartesian coordinate system do the rotation then make the
inverse transformation
TNPUT :
       crd : three element vector containing the coordinates of a
              given point in a cartesian system
       {\bf v} : three element array, angles of rotation around the {\bf x}, {\bf y}, {\bf z} axes
       ang : angle around the x, y, z, axes with which the vector should be rotated
3.3.2.5 def radmc3dPy.crd_trans.vtrans_sph2cart ( crd = [0, v = [0, reverse = False )
Function to transform velocities between spherical to cartesian systems
INPUT :
        ard
                : Three element array containing the input
                   coordinates [x,y,z] or [r,phi,theta] by default
                   the coordinates assumed to be in the cartesian system
                 : Three element array containing the input
                   velocities in the same coordinate system as crd
OPTIONS :
        reverse=False : Calculates the inverse trasnformation (cartesian -> spherical)
OUTPUT :
       result : A three element array containg the output
                   velocities [vr, vphi, vtheta] or [vx, vy, vz]
NOTE!!!!! The velocities in the spherical system are not angular velocities!!!!
v[1] = dphi/dt * r
v[2] = dtheta/dt * r
```

# 3.4 radmc3dPy.image Namespace Reference

#### **Classes**

· class radmc3dlmage

#### **Functions**

- · def getPSF
- def readImage
- · def plotImage
- · def makelmage
- · def cmask

#### 3.4.1 Detailed Description

```
PYTHON module for RADMC3D
(c) Attila Juhasz 2011,2012,2013,2014
This sub-module contains classes/functions to create and read images with radmc3d and to calculate
interferometric visibilities and write fits files
For help on the syntax or functionality of each function see the help of the individual functions
CLASSES:
radmc3dImage - RADMC3D image class
radmc3dVisibility - Class of interferometric visibilities
FUNCTIONS:
getPSF() - Calculates a Gaussian PSF/beam
getVisibility() - Calculates interferometric visiblities
makeImage() - Runs RADMC3D to calculate images/channel maps
plotImage() - Plots the image
readImage() - Reads RADMC3D image(s)
3.4.2 Function Documentation
3.4.2.1 def radmc3dPy.image.cmask ( im = None, rad = 0.0, au = False, arcsec = False, dpc = None )
Function to simulate a coronographic mask by
```

res

```
image center
INPUT:
          : a radmc3dImage class
          : radius of the mask
   rad
          : if true the radius is taken to have a unit of AU
    arcsec : if true the radius is taken to have a unit of arcsec (dpc
              should also be set)
           : distance of the source (required if arcsec = True)
    NOTE: if arcsec=False and au=False rad is taken to have a unit of pixel
OUTPUT:
```

: a radmc3dImage class containing the masked image

setting the image values to zero within circle of a given radius around the

#### 3.4.2.2 def radmc3dPy.image.getPSF ( nx = None, ny = None, fwhm = None, pa = None, pscale = None)

Function to generate a two dimensional Gaussian PSF

```
INPUT:
             : image size in the first dimension
              : image size in the second dimension
     nv
              : full width at half maximum of the psf in each dimension [fwhm_x, fwhm_y]
     fwhm
              : position angle of the gaussian if the gaussian is not symmetric
      pscale : pixelscale of the image, if set fwhm should be in the same unit, if not set unit of fwhm is pixelscale.
OUTPUT:
      result : dictionary containing the following keys
      'psf' : two dimensional numpy array containing the normalized psf
      ' x'
             : first coordinate axis of the psf
              : seonc coordinate axis of the psf
3.4.2.3 def radmc3dPy.image.makelmage ( npix = None, incl = None, wav = None, sizeau = None, phi = None,
      posang = None, pointau = None, fluxcons = True, nostar = False, noscat = False, widthkms = None,
      linenlam = None, vkms = None, iline = None, lambdarange = None, nlam = None )
Function to call RADMC3D to calculate a rectangular image
SYNTAX:
      makeImage(npix=100, incl=60.0, wav=10.0, sizeau=300., phi=0., posang=15.,
                 pointau=[0., 0.,0.], fluxcons=True, nostar=False, noscat=False)
INPUT:
                   : number of pixels on the rectangular images
       npix
                   : diameter of the image in au
       sizeau
                  : inclination angle of the source
                   : distance of the source in parsec
       dpc
                   : azimuthal rotation angle of the source in the model space
       phi
                  : position angle of the source in the image plane
                  : three elements list of the cartesian coordinates of the image center
       pointau
       widthkms
                   : width of the frequency axis of the channel maps
       linenlam : number of wavelengths to calculate images at
              : a single velocity value at which a channel map should be calculated
                   : line transition index
       lambdarange: two element list with the wavelenght boundaries between which
                    multiwavelength images should be calculated
                   : number of wavelengths to be calculated in lambdarange
KEYWORDS:
       fluxcons : this should not even be a keyword argument, it ensures flux conservation
       (adaptive subpixeling) in the rectangular images
       \hbox{nostar} \quad \hbox{: if True the calculated images will not contain stellar emission}
                : if True, scattered emission will be neglected in the source function, however,
       noscat
                      extinction will contain scattering if kappa_scat is not zero.
3.4.2.4 def radmc3dPy.image.plotImage ( image = None, arcsec = False, au = False, log = False, dpc = None,
      maxlog = None, saturate = None, bunit = None, ifreq = 0, cmask_rad = None, interpolation = 'nearest',
      cmap = cm.gist_gray, stokes = 'I', kwargs )
Function to plot a radmc3d image
SYNTAX:
      result = plotImage(image='image.out', arcsec=True, au=False, log=True, dpc=140, maxlog=-6.,
                         saturate=0.1, bunit='Jy')
INPUT:
              : A radmc3dImage class returned by readimage
     arcsec : If True image axis will have the unit arcsec (NOTE: dpc keyword should also be set!)
               : If True image axis will have the unit AU
```

```
: If True image scale will be logarithmic, otherwise linear
                                       : Distance to the source in parsec (This keywords should be set if arcsec=True, or bunit!=None)
                                    : Logarithm of the lowest pixel value to be plotted, lower pixel values will be clippde
                saturate : Highest pixel values to be plotted in terms of the peak value, higher pixel values will be cl
                                   : Unit of the image, (None - Inu/max(Inu), 'inu' - Inu, fnu - Jy/pixel)
                                       : If the image file/array consists of multiple frequencies/wavelengths ifreq denotes the index
                                            of the frequency/wavelength in the image array to be plotted
                \verb|cmask_rad|: Simulates coronographyic mask|: sets the image values to zero within this radius of the image values to zero within this radius of the image values to zero within this radius of the image values to zero within this radius of the image values to zero within this radius of the image values to zero within this radius of the image values to zero within this radius of the image values to zero within this radius of the image values to zero within this radius of the image values to zero within this radius of the image values to zero within this radius of the image values to zero within this radius of the image values to zero within this radius of the image values to zero within this radius of the image values to zero within this radius of the image values to zero within this radius of the image values to zero within this radius of the image values are the image values of the image value of the image
                                               The unit is the same as the image axis (au, arcsec, cm)
                                               NOTE: this works only on the plot, the image array is not changed (for that used the cmask()
                                       : matplotlib color map
                interpolation: interpolation keyword for imshow (e.g. 'nearest', 'bilinear', 'bicubic')
3.4.2.5 def radmc3dPy.image.readImage ( fname = None, binary = False )
Function to read an image calculated by RADMC3D
INPUT:
                            : file name of the radmc3d output image (if omitted 'image.out' is used)
          binary : False - the image format is formatted ASCII if True - C-compliant binary
```

### 3.5 radmc3dPy.model\_lines\_nlte\_lvg\_1d\_1 Namespace Reference

#### **Functions**

- · def getModelDesc
- def getDefaultParams
- def getGasTemperature
- def getDustTemperature
- def getGasAbundance
- · def getGasDensity
- def getDustDensity
- def getVTurb
- def getVelocity

### 3.5.1 Detailed Description

```
PYTHON module for RADMC3D (c) Attila Juhasz, Kees Dullemond 2011,2012,2013,2014
Original IDL model by Kees Dullemond, Python translation by Attila Juhasz
```

#### 3.5.2 Function Documentation

#### 3.5.2.1 def radmc3dPy.model\_lines\_nlte\_lvg\_1d\_1.getDefaultParams ( )

```
Function to provide default parameter values
```

OUTPUT:

Returns a list whose elements are also lists with three elements:
1) parameter name, 2) parameter value, 3) parameter description
All three elements should be strings. The string of the parameter
value will be directly written out to the parameter file if requested,
and the value of the string expression will be evaluated and be put
to radmc3dData.ppar. The third element contains the description of the
parameter which will be written in the comment field of the line when
a parameter file is written.

# 3.5.2.2 def radmc3dPy.model\_lines\_nlte\_lvg\_1d\_1.getDustDensity ( grid = None, ppar = None ) Function to create the dust density distribution INPUT: grid - An instance of the radmc3dGrid class containing the spatial and wavelength grid ppar - Dictionary containing all parameters of the model OUTPUT: returns the volume density in q/cm^3 3.5.2.3 def radmc3dPy.model\_lines\_nlte\_lvg\_1d\_1.getDustTemperature ( grid = None, ppar = None ) Function to calcualte/set the dust temperature INPUT: $\operatorname{grid}$ - An instance of the radmc3dGrid class containing the spatial and wavelength $\operatorname{grid}$ ppar - Dictionary containing all parameters of the model OUTPUT: returns the dust temperature in K 3.5.2.4 def radmc3dPy.model\_lines\_nlte\_lvg\_1d\_1.getGasAbundance ( grid = None, ppar = None, ispec = '') Function to create the conversion factor from volume density to number density of molecule ispec. The number density of a molecule is rhogas $\star$ abun INPUT: grid - An instance of the radmc3dGrid class containing the spatial and wavelength grid $\ensuremath{\mathsf{ppar}}\xspace$ - Dictionary containing all parameters of the model ispec - The name of the gas species whose abundance should be calculated OUTPUT: returns the abundance as a Numpy array 3.5.2.5 def radmc3dPy.model\_lines\_nlte\_lvg\_1d\_1.getGasDensity ( grid = None, ppar = None ) Function to create the total gas density distribution INPUT: grid - An instance of the radmc3dGrid class containing the spatial and wavelength grid ppar - Dictionary containing all parameters of the model OUTPUT: returns the volume density in $g/cm^3$

#### 3.5.2.6 def radmc3dPy.model\_lines\_nlte\_lvg\_1d\_1.getGasTemperature ( grid = None, ppar = None)

Function to calcualte/set the gas temperature  $\ensuremath{\mathsf{Set}}$ 

```
INPUT:
```

-----

grid - An instance of the radmc3dGrid class containing the spatial and wavelength grid ppar - Dictionary containing all parameters of the model

```
OUTPUT:
    returns the gas temperature in K
3.5.2.7 def radmc3dPy.model_lines_nlte_lvg_1d_1.getModelDesc ( )
Function to provide a brief description of the model
3.5.2.8 def radmc3dPy.model_lines_nlte_lvg_1d_1.getVelocity ( grid = None, ppar = None )
Function to create the turbulent velocity field
INPUT:
    grid - An instance of the radmc3dGrid class containing the spatial and wavelength grid
    ppar - Dictionary containing all parameters of the model
OUTPUT:
    returns the turbulent velocity in cm/s
3.5.2.9 def radmc3dPy.model_lines_nlte_lvg_1d_1.getVTurb ( grid = None, ppar = None )
Function to create the turbulent velocity field
INPUT:
    grid - An instance of the radmc3dGrid class containing the spatial and wavelength grid
    ppar - Dictionary containing all parameters of the model
OUTPUT:
   returns the turbulent velocity in cm/s
```

# 3.6 radmc3dPy.model\_ppdisk Namespace Reference

#### **Functions**

- · def getModelDesc
- · def getDefaultParams
- · def getDustDensity

WARNING!!!!! At the moment I assume that the multiple dust population differ from each other only in grain size but not in bulk density thus when I calculate the abundances / mass fractions they are independent of the grains bulk density since abundances/mass fractions are normalized to the total mass.

- def getGasDensity
- def getGasAbundance
- def getVTurb
- · def getVelocity

#### 3.6.1 Detailed Description

```
PYTHON module for RADMC3D (c) Attila Juhasz 2011,2012,2013,2014

Generic protoplanetary disk model with a simple chemistry
```

The density is given by

The molecular abundance function takes into account dissociation and freeze-out of the molecules For photodissociation only the continuum (dust) shielding is taken into account in a way that whenever the continuum optical depth radially drops below a threshold value the molecular abundance is dropped to zero. For freeze-out the molecular abundance below a threshold temperature is decreased by a given fractor.

#### 3.6.2 Function Documentation

#### 3.6.2.1 def radmc3dPy.model\_ppdisk.getDefaultParams ( )

Function to provide default parameter values

OUTPUT:

Returns a list whose elements are also lists with three elements:
1) parameter name, 2) parameter value, 3) parameter description
All three elements should be strings. The string of the parameter
value will be directly written out to the parameter file if requested,
and the value of the string expression will be evaluated and be put
to radmc3dData.ppar. The third element contains the description of the
parameter which will be written in the comment field of the line when
a parameter file is written.

3.6.2.2 def radmc3dPy.model\_ppdisk.getDustDensity ( rcyl = None, phi = None, z = None, z0 = None, hp = None, sigma = None, grid = None, ppar = None )

WARNING!!!!!! At the moment I assume that the multiple dust population differ from each other only in grain size but not in bulk density thus when I calculate the abundances / mass fractions they are independent of the grains bulk density since abundances/mass fractions are normalized to the total mass.

```
Function to create the density distribution in a protoplanetary disk $\operatorname{\textsc{OUTPUT}}:$
```

returns the volume density in  $g/cm^3$ , whether the density is that of the gas or dust or both depends on what is specified in the surface density/mass

Thus I use 1g/cm<sup>^3</sup> for all grain sizes. TODO: Add the possibility to handle multiple dust species with different bulk densities and with multiple grain sizes.

 $gdens = zeros(ngs, dtype=float) + 1.0 \ gs = ppar['gsmin'] * (ppar['gsmax']/ppar['gsmin']) ** (arange(ppar['ngs'], dtype=float64) / (float(ppar['ngs'])-1.)) \ gmass = 4./3.*np.pi*gs**3. * gdens gsfact = gmass * gs**(ppar['gsdist_-powex']+1) \ gsfact = gsfact / gsfact.sum()$ 

3.6.2.3 def radmc3dPy.model\_ppdisk.getGasAbundance ( grid = None, ppar = None, ispec = '')

Function to create the conversion factor from volume density to number density of molecule ispec. The number density of a molecule is rhogas  $\star$  abun

INPUT:

```
grid - An instance of the radmc3dGrid class containing the spatial and wavelength grid
    ppar - Dictionary containing all parameters of the model
    ispec - The name of the gas species whose abundance should be calculated
OUTPUT:
    returns the abundance as a Numpy array
3.6.2.4 def radmc3dPy.model_ppdisk.getGasDensity ( rcyl = None, phi = None, z = None, z0 = None, hp = None,
       sigma = None, grid = None, ppar = None )
Function to create the density distribution in a protoplanetary disk
OUTPUT:
   returns the volume density in g/cm^3, whether the density is that of the gas
    or dust or both depends on what is specified in the surface density/mass
3.6.2.5 def radmc3dPy.model_ppdisk.getModelDesc ( )
Function to provide a brief description of the model
3.6.2.6 def radmc3dPy.model_ppdisk.getVelocity ( rcyl = None, phi = None, z = None, z0 = None, grid = None, ppar
       = None )
Function to create the velocity field in a protoplanetary disk
3.6.2.7 def radmc3dPy.model_ppdisk.getVTurb ( grid = None, ppar = None )
Function to create the turbulent velocity field
INPUT:
    grid - An instance of the radmc3dGrid class containing the spatial and wavelength grid
    ppar - Dictionary containing all parameters of the model
OUTPUT:
   returns the turbulent velocity in cm/s
```

# 3.7 radmc3dPy.model\_simple\_1 Namespace Reference

#### **Functions**

- def getModelDesc
- def getDefaultParams
- · def getDustDensity

### 3.7.1 Detailed Description

```
PYTHON module for RADMC3D (c) Attila Juhasz, Kees Dullemond 2011,2012,2013
Original IDL model by Kees Dullemond, Python translation by Attila Juhasz
```

#### 3.7.2 Function Documentation

#### 3.7.2.1 def radmc3dPy.model\_simple\_1.getDefaultParams ( )

Function to provide default parameter values

OUTPUT:

Returns a list whose elements are also lists with three elements: 1) parameter name, 2) parameter value, 3) parameter description All three elements should be strings. The string of the parameter value will be directly written out to the parameter file if requested, and the value of the string expression will be evaluated and be put to radmc3dData.ppar. The third element contains the description of the parameter which will be written in the comment field of the line when a parameter file is written.

#### 3.7.2.2 def radmc3dPy.model\_simple\_1.getDustDensity ( grid = None, ppar = None )

Function to create the dust density distribution

OUTPUT:

returns the volume density in g/cm^3

#### 3.7.2.3 def radmc3dPy.model\_simple\_1.getModelDesc ( )

Function to provide a brief description of the model

#### radmc3dPy.model\_spher1d\_1 Namespace Reference 3.8

#### **Functions**

- def getModelDesc
- · def getDefaultParams
- · def getDustDensity

#### 3.8.1 Detailed Description

```
PYTHON module for RADMC3D
(c) Attila Juhasz, Kees Dullemond 2011, 2012, 2013, 2014
Original IDL model by Kees Dullemond, Python translation by Attila Juhasz
```

### 3.8.2 Function Documentation

### 3.8.2.1 def radmc3dPy.model\_spher1d\_1.getDefaultParams ( )

Function to provide default parameter values

OUTPUT:

Returns a list whose elements are also lists with three elements: 1) parameter name, 2) parameter value, 3) parameter description All three elements should be strings. The string of the parameter value will be directly written out to the parameter file if requested, and the value of the string expression will be evaluated and be put to radmc3dData.ppar. The third element contains the description of the parameter which will be written in the comment field of the line when a parameter file is written.

#### 3.8.2.2 def radmc3dPy.model\_spher1d\_1.getDustDensity ( grid = None, ppar = None )

Function to create the dust density distribution OUTPUT:
----returns the volume density in q/cm^3

#### 3.8.2.3 def radmc3dPy.model\_spher1d\_1.getModelDesc ( )

Function to provide a brief description of the model

# 3.9 radmc3dPy.model\_spher2d\_1 Namespace Reference

#### **Functions**

- · def getModelDesc
- · def getDefaultParams
- def getDustDensity

#### 3.9.1 Detailed Description

```
PYTHON module for RADMC3D (c) Attila Juhasz, Kees Dullemond 2011,2012,2013,2014
Original IDL model by Kees Dullemond, Python translation by Attila Juhasz
```

#### 3.9.2 Function Documentation

#### 3.9.2.1 def radmc3dPy.model\_spher2d\_1.getDefaultParams ( )

Function to provide default parameter values OUTPUT:

Returns a list whose elements are also lists with three elements:
1) parameter name, 2) parameter value, 3) parameter description
All three elements should be strings. The string of the parameter
value will be directly written out to the parameter file if requested,
and the value of the string expression will be evaluated and be put
to radmc3dData.ppar. The third element contains the description of the
parameter which will be written in the comment field of the line when
a parameter file is written.

# 3.9.2.2 def radmc3dPy.model\_spher2d\_1.getDustDensity ( grid = None, ppar = None )

Function to create the dust density distribution  $\mbox{\tt OUTPUT:}$   $\mbox{\tt -----}$ 

returns the volume density in  $g/cm^3$ 

#### 3.9.2.3 def radmc3dPy.model\_spher2d\_1.getModelDesc ( )

Function to provide a brief description of the model

# 3.10 radmc3dPy.model\_template Namespace Reference

#### **Functions**

- · def getModelDesc
- def getDefaultParams
- · def getGasTemperature
- def getDustTemperature
- · def getGasAbundance
- · def getGasDensity
- · def getDustDensity
- def getVTurb
- · def getVelocity

### 3.10.1 Detailed Description

```
This is a radmc3dPy model template
A radmc3dPy model file can contain any / all of the functions below

getDefaultParams()
getModelDesc()
getDustDensity()
getDustTemperature()
getGasAbundance()
getGasDensity()
getGasTemperature()
getGasTemperature()
getVelocity()
```

The description of the individual functions can be found in the docstrings below the function name. If a model does not provide a variable or the variable should be calculated by RADMC-3D (e.g. dust temperature) the corresponding function (e.g. get\_dust\_temperature) should be removed from or commented out in the model file.

NOTE: When using this template it is strongly advised to renme the template model (to e.g. model\_mydisk.py) as the get\_model\_names() function in the setup module removes the name 'template' from the list of available models.

#### 3.10.2 Function Documentation

#### 3.10.2.1 def radmc3dPy.model\_template.getDefaultParams ( )

```
Function to provide default parameter values
```

OUTPUT:

Returns a list whose elements are also lists with three elements:
1) parameter name, 2) parameter value, 3) parameter description
All three elements should be strings. The string of the parameter
value will be directly written out to the parameter file if requested,
and the value of the string expression will be evaluated and be put
to radmc3dData.ppar. The third element contains the description of the
parameter which will be written in the comment field of the line when
a parameter file is written.

# 3.10.2.2 def radmc3dPy.model\_template.getDustDensity ( grid = None, ppar = None ) Function to create the dust density distribution INPUT: grid - An instance of the radmc3dGrid class containing the spatial and wavelength grid ppar - Dictionary containing all parameters of the model OUTPUT: returns the volume density in q/cm^3 3.10.2.3 def radmc3dPy.model\_template.getDustTemperature ( grid = None, ppar = None ) Function to calcualte/set the dust temperature INPUT: $\operatorname{grid}$ - An instance of the radmc3dGrid class containing the spatial and wavelength $\operatorname{grid}$ ppar - Dictionary containing all parameters of the model OUTPUT: returns the dust temperature in K 3.10.2.4 def radmc3dPy.model\_template.getGasAbundance( grid = None, ppar = None, ispec = '') Function to create the conversion factor from volume density to number density of molecule ispec. The number density of a molecule is rhogas $\star$ abun INPUT: grid - An instance of the radmc3dGrid class containing the spatial and wavelength grid ppar - Dictionary containing all parameters of the model ispec - The name of the gas species whose abundance should be calculated OUTPUT: returns the abundance as a Numpy array 3.10.2.5 def radmc3dPy.model\_template.getGasDensity ( grid = None, ppar = None ) Function to create the total gas density distribution INPUT: grid - An instance of the radmc3dGrid class containing the spatial and wavelength grid ppar - Dictionary containing all parameters of the model OUTPUT: returns the volume density in $g/cm^3$ 3.10.2.6 def radmc3dPy.model\_template.getGasTemperature ( grid = None, ppar = None ) Function to calcualte/set the gas temperature INPUT:

 $\verb|grid - An instance of the radmc3dGrid class containing the spatial and wavelength grid$ 

ppar - Dictionary containing all parameters of the model

```
OUTPUT:
    returns the gas temperature in K
3.10.2.7 def radmc3dPy.model_template.getModelDesc ( )
Function to provide a brief description of the model
3.10.2.8 def radmc3dPy.model_template.getVelocity ( grid = None, ppar = None )
Function to create the turbulent velocity field
INPUT:
    grid - An instance of the radmc3dGrid class containing the spatial and wavelength grid
    ppar - Dictionary containing all parameters of the model
OUTPUT:
   returns the turbulent velocity in cm/s
3.10.2.9 def radmc3dPy.model_template.getVTurb ( grid = None, ppar = None )
Function to create the turbulent velocity field
INPUT:
    \operatorname{grid} - An instance of the radmc3dGrid class containing the spatial and wavelength \operatorname{grid}
    ppar - Dictionary containing all parameters of the model
OUTPUT:
   returns the turbulent velocity in cm/s
```

# 3.11 radmc3dPy.model\_test\_scattering\_1 Namespace Reference

### **Functions**

- def getModelDesc
- def getDefaultParams
- def getDustDensity

### 3.11.1 Detailed Description

```
PYTHON module for RADMC3D (c) Attila Juhasz, Kees Dullemond 2011,2012,2013
Original IDL model by Kees Dullemond, Python translation by Attila Juhasz
```

#### 3.11.2 Function Documentation

#### 3.11.2.1 def radmc3dPy.model\_test\_scattering\_1.getDefaultParams ( )

Function to provide default parameter values

```
OUTPUT:
```

Returns a list whose elements are also lists with three elements:
1) parameter name, 2) parameter value, 3) parameter description
All three elements should be strings. The string of the parameter
value will be directly written out to the parameter file if requested,
and the value of the string expression will be evaluated and be put
to radmc3dData.ppar. The third element contains the description of the
parameter which will be written in the comment field of the line when
a parameter file is written.

#### 3.11.2.2 def radmc3dPy.model\_test\_scattering\_1.getDustDensity ( grid = None, ppar = None )

```
Function to create the dust density distribution

OUTPUT:
----
returns the volume density in g/cm^3
```

#### 3.11.2.3 def radmc3dPy.model\_test\_scattering\_1.getModelDesc ( )

Function to provide a brief description of the model

# 3.12 radmc3dPy.natconst Namespace Reference

#### **Variables**

- float gg 6.672e-8
- float mp 1.6726e-24
- float me 9.1095e-28
- float kk 1.3807e-16
- float **hh** 6.6262e-27
- float ee 4.8032e-10
- float cc 2.99792458e10
- float st 6.6524e-25
- float ss 5.6703e-5
- float aa 7.5657e-15
- float muh2 2.3000e0
- float ev 1.6022e-12
- float kev 1.6022e-9
- int micr 1
- int **km** 1
- int angs 1
- float Is 3.8525e33
- float rs 6.96e10
- float ms 1.99e33
- float ts 5.780e3
- float au 1.496e13
- float pc 3.08572e18
- float mea 5.9736e27
- float rea 6.375e8
- float mmo 7.347e25
- float rmo 1.738e8
- float dmo 3.844e10

- float mju 1.899e30
- float rju 7.1492e9
- float dju 7.78412e13
- float year 3.1536e7
- float hour 3.6000e3float day 8.6400e4

### 3.12.1 Detailed Description

```
PYTHON module for RADMC3D
(c) Attila Juhasz 2011,2012,2013,2014

This sub-module contains natural constants in CGS units
(Translated from RADMC's IDL function problem_natconst.pro)
```

# 3.13 radmc3dPy.setup Namespace Reference

#### **Functions**

- · def getTemplateModel
- · def getModelNames
- def getModelDesc
- · def problemSetupDust
- · def problemSetupGas
- · def writeRadmc3dInp
- def writeLinesInp

#### 3.13.1 Detailed Description

PYTHON module for RADMC3D

```
(c) Attila Juhasz 2011,2012,2013,2014

This sub-module functions to set up a RADMC3D model for dust and/or line simulations

For help on the syntax or functionality of each function see the help of the individual functions

FUNCTIONS:

------

get_model_desc() - Returns the brief description of a model (if the model file contains a get_desc() for get_model_names() - Returns the list of available models

get_template_model() - Copy the template model file from the library directory (radmc3dPy) to the current problem_setup_dust() - Function to set up a dust model

problem_setup_gas() - Function to set up a line simulation

writeLinesInp() - Writes the lines.inp master command file for line simulations

writeRadmc3dInp() - Writes the radmc3d.inp master command file required for all RADMC3D runs
```

#### 3.13.2 Function Documentation

#### 3.13.2.1 def radmc3dPy.setup.getModelDesc ( model = ' ' )

Returns the brief description of the selected model

### 3.13.2.2 def radmc3dPy.setup.getModelNames ( )

Returns the name of the available models

#### 3.13.2.3 def radmc3dPy.setup.getTemplateModel ( )

Create a copy of the template model file in the current working directory. The PYTHONPATH environment variable is checked for the installation path of radmc3dPy and the template file is copied from the first hit in the path list.

#### 3.13.2.4 def radmc3dPy.setup.problemSetupDust ( model = ' ', binary = True, writeDustTemp = False, kwargs )

Function to set up a dust model for RADMC3D

#### TNPHT.

model : Name of the model that should be used to create the density structure. The file should be in a directory from where it can directly be imported (i.e. the directory should be in the PYTHON\_PATH environment variable or it should be in the current working directory)  $% \frac{1}{2}\left( \frac{1}{2}\right) =\frac{1}{2}\left( \frac{1}{2}\right) +\frac{1}{2}\left( \frac{1}{2}\right) +$ and the file name should be 'model\_xxx.py', where xxx stands for the string that should be specified in this variable

binary: If True input files will be written in binary format, if False input files are written as formatted ascii text.

writeDustTemp: If True a separate dust\_temperature.inp/dust\_tempearture.binp file will be written under the condition that the model contains a function getDustTemperature()

#### OPTIONS:

Any varible name in problem\_params.inp can be used as a keyword argument. At first all variables are read from problem\_params.in to a dictionary called ppar. Then if there is any keyword argument set in the call of problem\_setup\_dust the ppar dictionary is searched for this key. If found the value belonging to that key in the ppar dictionary is changed to the value of the keyword argument. If no such key is found then the dictionary is simply extended by the keyword argument. Finally the problem\_params.inp file is updated with the new parameter values.

#### FILES WRITTEN DURING THE SETUP:

dustopac.inp dust opacity master file wavelength\_micron.inp - wavelength grid - spatial grid amr\_grid.inp

stars.inp - input radiation field

dust\_density.inp - dust density distribution - parameters for RADMC3D (e.g. Nr of photons to be used, scattering type, etc)

#### STEPS OF THE SETUP:

- 1) Create the spatial and frequency grid
- 2) Create the master opacity file and calculate opacities with the Mie-code if necessary
- 3) Set up the input radiation field (generate stars.inp)
- 4) Calculate the dust density
- 5) If specified; calculatest the dust temperature (e.g. for gas simulations, or if it is taken from an external input (e.g. from another model))
- 6) Write all output files

### 3.13.2.5 def radmc3dPy.setup.problemSetupGas ( model = '', fullsetup = False, binary = True, writeGasTemp = False, kwargs)

Function to set up a gas model for RADMC3D

#### INPUT:

model: Name of the model that should be used to create the density structure the file should be in a directory from where it can directly be imported (i.e. the directory should be in the PYTHON\_PATH environment variable, or it should be the current working directory) and the file name should be 'model\_xxx.py', where xxx stands for the string that should be specified in this variable

fullsetup : if False only the files related to the gas simulation is written out
 (i.e. no grid, stellar parameter file and radmc3d master command file is written)
 if True the spatial and wavelength grid as well as the input radiation field
 and the radmc3d master command file will be (over)written.

binary : If True input files will be written in binary format, if False input files are written as formatted ascii text.

writeGasTemp: If True a separate gas\_temperature.inp/gas\_tempearture.binp file will be written under the condition that the model contains a function get\_gas\_temperature()

#### OPTIONS:

-----

Any varible name in problem\_params.inp can be used as a keyword argument. At first all variables are read from problem\_params.in to a dictionary called ppar. Then if there is any keyword argument set in the call of problem\_setup\_gas the ppar dictionary is searched for such key. If found the value belonging to that key in the ppar dictionary is changed to the value of the keyword argument. If no such key is found then the dictionary is simply extended by the keyword argument. Finally the problem\_params.inp file is updated with the new parameter values.

#### FILES WRITTEN DURING THE SETUP:

\_\_\_\_\_

fullsetup = True

amr\_grid.inp - spatial grid
wavelength\_micron.inp - wavelength grid

stars.inp - input radiation field

radmc3d.inp - parameters for RADMC3D (e.g. Nr of photons to be used, scattering type, etc)

lines.inp - line mode master command file

numberdens\_xxx.inp - number density of molecule/atomic species 'xxx'

gas\_velocity.inp - Gas velocity

microturbulence.inp - The standard deviation of the Gaussian line profile caused by turbulent broadening (doublecheck if it is really the standard deviation or a factor

of sqrt(2) less than that!)

gas\_temperature.inp - Gas temperature (which may be different from the dust temperature). If

tgas\_eq\_tdust is set to zero in radmc3d.inp the gas temperature in this

file will be used instead of the dust temperature.

fullsetup = False

lines.inp - line mode master command file

numberdens\_xxx.inp - number density of molecule/atomic species 'xxx'

gas\_velocity.inp - Gas velocity

microturbulence.inp - The standard deviation of the Gaussian line profile caused by turbulent

broadening (doublecheck if it is really the standard deviation or a factor

of sqrt(2) less than that!)

gas\_temperature.inp - Gas temperature (which may be different from the dust temperature). If

tgas\_eq\_tdust is set to zero in radmc3d.inp the gas temperature in this

file will be used instead of the dust temperature.

#### 3.13.2.6 def radmc3dPy.setup.writeLinesInp ( ppar = None )

Function to write the lines.inp master command file for line simulation in RADMC3D

### INPUT:

\_\_\_\_\_

ppar : dictionary containing all parameters of a RADMC3D run

#### 3.13.2.7 def radmc3dPy.setup.writeRadmc3dInp ( modpar = None )

Function to write the radmc3d.inp master command file for RADMC3D

#### INPUT:

\_\_\_\_

 $\ensuremath{\mathsf{ppar}}$  : dictionary containing all parameters of a RADMC3D run

# **Chapter 4**

# **Class Documentation**

# 4.1 radmc3dPy.analyze.radmc3dData Class Reference

### **Public Member Functions**

- def \_\_init\_\_
- def getTauOneDust
- def getTau
- def readDustDens
- def readDustTemp
- def readGasVel
- def readVTurb
- def readGasDens
- def readGasTemp
- def writeDustDens
- def writeDustTemp
- def writeGasDens
- def writeGasTemp
- def writeGasVeldef writeVTurb
- def writeVTK
- def getSigmaDust
- def getSigmaGas

### **Public Attributes**

- grid
- rhodust
- dusttemp
- rhogas
- · ndens\_mol
- · ndens\_cp
- gasvel
- · gastemp
- vturb
- taux
- tauy
- tauz
- · sigmadust
- · sigmagas

28 **Class Documentation** 

```
4.1.1 Detailed Description
RADMC3D data class
    reading and writing dust density/temperature, gas density/temperature/velocity,
    generating a legacy vtk file for visualization
ATTRIBUTES:
    rhodust - Dust density in g/cm^3
dusttemp - Dust temperature in K
              - Gas density in g/cm^3
    ndens_mol - Number density of the molecule [molecule/cm^3]
    ndens_cp - Number density of the collisional partner [molecule/cm^3]
               - Gas velocity in cm/s
              - Gas temperature in K
    gastemp
               - Mictroturbulence in cm/s
    vturb
              - Optical depth along the x (cartesian) / r (cylindrical) / r (spherical) dimension
    taux
              - Optical depth along the y (cartesian) / theta (cylindrical) / theta (spherical) dimension - Optical depth along the z (cartesian) / z (cylindrical) / phi (spherical) dimension
    tauy
    sigmadust - Dust surface density in q/cm^2
    sigmagas - Gas surface density in molecule/cm^2 (or g/cm^2 depending on the dimension of rhogas)
4.1.2 Member Function Documentation
4.1.2.1 def radmc3dPy.analyze.radmc3dData.getSigmaDust ( self, idust = 0 )
Function to calculate dust surface density
OPTIONS:
    idust - index of the dust species for which the surface density should be calculated
    if omitted the calculated surface density will be the sum over all dust species
4.1.2.2 def radmc3dPy.analyze.radmc3dData.getSigmaGas ( self )
```

```
Function to calculate gas surface density
This function uses radmc3dData.rhogas to calculate the surface density, thus the
unit of surface density depends on the unit of radmc3dData.rhogas (g/cm^2) or molecule/cm<sup>2</sup>)
```

#### 4.1.2.3 def radmc3dPy.analyze.radmc3dData.getTau ( self, idust = [], axis = 'xy', wav = 0., kappa = None )

Function to calculate the optical depth along any given combination of the axes

```
INPUT:
```

```
idust : List of dust component indices whose optical depth should be calculated
If multiple indices are set the total optical depth is calculated summing
over all dust species in idust
axis - Name of the axis/axes along which the optical depth should be calculated
(e.g. '{\rm x}' for the first dimension or '{\rm xyz}' for all three dimensions)
wav : Wavelength at which the optical depth should be calculated
kappa : If set it should be a list of mass extinction coefficients at the desired wavelength
The number of elements in the list should be equal to that in the idust keyword
```

#### 4.1.2.4 def radmc3dPy.analyze.radmc3dData.getTauOneDust ( self, idust = 0, axis = ' ', kappa = 0 . )

Function to calculate the optical depth of a single dust species along any given combination of the axes

# INPUT:

```
idust - Index of the dust species whose optical depth should be calculated
axis - Name of the axis/axes along which the optical depth should be calculated
```

```
(e.g. 'x' for the first dimension or 'xyz' for all three dimensions)
    kappa - Mass extinction coefficients of the dust species at the desired wavelength
   Returns a dictionary with the following keys;
    \hbox{\tt taux - optical depth along the first dimension}\\
    tauy - optical depth along the second dimension
    (tauz is not yet implemented)
4.1.2.5 def radmc3dPy.analyze.radmc3dData.readDustDens ( self, fname = '', binary = True )
Function to read the dust density
OPTIONS:
    fname - Name of the file that contains the dust density. If omitted 'dust_density.inp' is used
    (or if binary=True the 'dust_density.binp' is used).
    binary - If true the data will be read in binary format, otherwise the file format is ascii
4.1.2.6 def radmc3dPy.analyze.radmc3dData.readDustTemp ( self, fname = '', binary = True )
Function to read the dust temperature
OPTIONS:
    fname - Name of the file that contains the dust temperature.
    If omitted 'dust_temperature.dat' (if binary=True 'dust_temperature.bdat') is used.
    binary - If true the data will be read in binary format, otherwise the file format is ascii
4.1.2.7 def radmc3dPy.analyze.radmc3dData.readGasDens ( self, ispec = '', binary = True )
Function to read the gas density
INPUT:
   ispec - File name extension of the 'numberdens_ispec.inp' (or if binary=True 'numberdens_ispec.binp') file
OPTIONS:
   binary - If true the data will be read in binary format, otherwise the file format is ascii
4.1.2.8 def radmc3dPy.analyze.radmc3dData.readGasTemp ( self, fname = '', binary = True )
Function to read the gas temperature
OPTIONS:
   fname - Name of the file that contains the gas temperature. If omitted 'gas_temperature.inp'
    (or if binary=True 'gas_tempearture.binp') is used.
    binary - If true the data will be read in binary format, otherwise the file format is ascii
4.1.2.9 def radmc3dPy.analyze.radmc3dData.readGasVel ( self, fname = ' ', binary = True )
Function to read the gas velocity.
OPTIONS:
```

```
fname - Name of the file that contains the gas velocity
    If omitted 'gas_velocity.inp' (if binary=True 'gas_velocity.binp')is used.
    binary - If true the data will be read in binary format, otherwise the file format is ascii
4.1.2.10 def radmc3dPy.analyze.radmc3dData.readVTurb ( self, fname = ' ', binary = True )
Function to read the turbulent velocity field.
OPTIONS:
   fname - Name of the file that contains the turbulent velocity field
    If omitted 'microturbulence.inp' (if binary=True 'microturbulence.binp') is used.
    binary - If true the data will be read in binary format, otherwise the file format is ascii
4.1.2.11 def radmc3dPy.analyze.radmc3dData.writeDustDens ( self, fname = ' ', binary = True )
Function to write the dust density
OPTIONS:
    fname - Name of the file into which the dust density should be written. If omitted 'dust_density.inp' is a
    binary - If true the data will be written in binary format, otherwise the file format is ascii
4.1.2.12 def radmc3dPy.analyze.radmc3dData.writeDustTemp ( self, fname = ' ', binary = True )
Function to write the dust density
OPTIONS:
    fname - Name of the file into which the dust density should be written. If omitted 'dust_density.inp' is u
    binary - If true the data will be written in binary format, otherwise the file format is ascii
4.1.2.13 def radmc3dPy.analyze.radmc3dData.writeGasDens ( self, fname = '', ispec = '', binary = True )
Function to write the gas density
INPUT:
fname - Name of the file into which the data will be written. If omitted "numberdens_xxx.inp" and
 "numberdens_xxx.binp" will be used for ascii and binary format, respectively (xxx is the name of the molecule
ispec - File name extension of the 'numberdens_ispec.inp' (if binary=True 'numberdens_ispec.binp')
file into which the gas density should be written
binary - If true the data will be written in binary format, otherwise the file format is ascii
4.1.2.14 def radmc3dPy.analyze.radmc3dData.writeGasTemp ( self, fname = ' ', binary = True )
Function to write the gas temperature
OPTIONS:
    fname - Name of the file into which the gas temperature should be written. If omitted
    'gas_temperature.inp' (if binary=True 'gas_tempearture.binp') is used.
    binary - If true the data will be written in binary format, otherwise the file format is ascii
```

# 4.1.2.15 def radmc3dPy.analyze.radmc3dData.writeGasVel ( self, fname = ' ' , binary = True ) Function to write the gas velocity OPTIONS: fname - Name of the file into which the gas temperature should be written. If omitted 'gas\_velocity.inp' (if binary=True 'gas\_velocity.binp') is used. binary - If true the data will be written in binary format, otherwise the file format is ascii 4.1.2.16 def radmc3dPy.analyze.radmc3dData.writeVTK( self, vtk\_fname = '', ddens = False, dtemp = False, idust = [0], gdens = False, gvel = False, gtemp = False) Function to dump all physical variables to a legacy vtk file INPUT: vtk\_fname : name of the file to be written, if not specified 'radmc3d\_data.vtk' will be used ddens $\hspace{0.1in}:\hspace{0.1in}$ if set to True the dust density will be written to the vtk file : if set to True the dust temperature will be written to the vtk file : a list of indices that specifies which dust component should be written idust. if not set then the first dust species (zero index) will be used gdens : if set to True the gas density will be written to the vtk file : if set to True the gas temperature will be written to the vtk file gtemp gvel : if set to True the gas velocity will be written to the vtk file 4.1.2.17 def radmc3dPy.analyze.radmc3dData.writeVTurb ( self, fname = ' ' , binary = True ) Function to write the microturbulence file OPTIONS: fname - Name of the file into which the turubulent velocity field should be written. If omitted 'microturbulence.inp' (if binary=True 'microturbulence.binp') is used. binary - If true the data will be written in binary format, otherwise the file format is ascii

# 4.2 radmc3dPy.analyze.radmc3dDustOpac Class Reference

## **Public Member Functions**

- def \_\_init\_\_
- def readOpac
- def makeOpac
- def mixOpac
- def readMasterOpac
- def writeMasterOpac
- · def runMakedust

#### **Public Attributes**

- · freq
- nwav
- nfreq
- · kabs
- ksca
- phase\_g
- ext
- · idust
- therm

#### **Static Public Attributes**

- tuple grid radmc3dGrid()
- wav grid.wav
- list **ext** []
- tuple rfile open(ppar['lnk fname'][idust], 'r')
- list w []
- list **n** []
- list k []
- tuple dum rfile.readline()
- tuple w array(w, dtype=float)
- tuple **n** array(n, dtype=float)
- tuple k array(k, dtype=float)
- tuple wfile open('opt const.dat', 'w')
- string Ink\_fname 'opt\_const.dat'
- list **mixnames** ['dustkappa\_igsize\_'+str(igs+1)+'.inp']
- list mixspecs [['dustkappa\_idust\_'+str(idust+1)+'\_igsize\_'+str(igs+1)+'.inp' for idust in range(len(ppar['lnk\_-fname']))]]
- list therm [True for i in range(len(ext))]

## 4.2.1 Detailed Description

```
Dust opacity class
ATTRIBUTES:
           - wavelength grid
    waw
    freq
           - frequency grid
          - number of wavelengths
    nwav
    kabs
           - absorption coefficient per unit mass
    ksca
           - scattering coefficient per unit mass
    phase_g - phase function
          - if set it contains the file name extension of the duskappa_ext.Kappa file
           - if False the dust grains are quantum-heated (default: True)
    therm
           - index of the dust species in the dust density distribution array
    NOTE: Each attribute is a list with each element containing the corresponding data for
          a given dust species
METHODS:
                      - Read the dust opacity files
    readOpac()
    readMasterOpac() - Read the master opacity file
    writeMasterOpac() - Write the master opacity file
                       - Calculates opacities with the Mie-code that comes with RADMC-3D (using the runMakedu
    makeOpac()
                     - Runs the Mie-code to calculate dust opacities
    runMakedust()
```

#### 4.2.2 Member Function Documentation

```
4.2.2.1 def radmc3dPy.analyze.radmc3dDustOpac.makeOpac ( self, ppar = None, wav = None )
```

```
Function to create dust opacities for RADMC3D using MIE calculation

INPUT:
-----
ppar - dictionary containing all parameter of the simulation

OPTIONS:
------
wav - numpy.ndarray containing the wavelength grid on which the mass absorption coefficients should be calculation
```

```
4.2.2.2 def radmc3dPy.analyze.radmc3dDustOpac.mixOpac ( self, ppar = None, mixnames = [], mixspecs = [],
       mixabun = [], writefile = True )
Function to mix opacities
INPUT:
             - A dictionary containing all parameters of the actual model setup
       If any keyword is set besides ppar, the value of the separate keyword
        will be taken instead of that in ppar. If mixname, mixspecs, and mixabun are all set
        ppar is completely omitted and not necessary to set when mixOpac is called.
    mixnames - Names of the files into which the mixed dust opacities will be written (not needed if writefil
    mixspecs - Names of the files from which the dust opacities are read (not needed if readfile=False)
    mixabun - Abundances of different dust species
    writefile - If False the mixed opacities will not be written out to files given in mixnames.
4.2.2.3 def radmc3dPy.analyze.radmc3dDustOpac.readMasterOpac ( self )
Function to read the master opacity file 'dustopac.inp'
it reads the dustkappa filename extensions (dustkappa_ext.inp) corresponding to dust species indices
OUTPUT:
Returns a dictionary with the following keys:
           - list of dustkappa file name extensions
    ^{\prime}therm^{\prime} - a list of integers specifying whether the dust grain is thermal or quantum heated
      (0 - thermal, 1 - quantum heated)
4.2.2.4 def radmc3dPy.analyze.radmc3dDustOpac.readOpac ( self, ext = [''], idust = None )
Function to read the dust opacity files
INPUT:
    ext : file name extension (file names should look like 'dustkappa_ext.inp')
    idust: index of the dust species in the master opacity file (dustopac.inp') - starts at 0
4.2.2.5 def radmc3dPy.analyze.radmc3dDustOpac.runMakedust ( self, freq = None, gmin = None, gmax = None, ngs =
       None, Ink_fname = None, gdens = None)
Interface function to the F77 code makedust to calculate mass absorption
coefficients from the optical constants using Mie-theory
INPUT:
   freq
              - numpy.ndarray containing the frequency grid on which the opacities should be calculated
              - minimum grain size
    gmin
               - maximum grain size
    amax
              - number of grain sizes
              - density of the dust grain in g/cm^3
    lnk\_faname - name of the file in which the optical constants are stored
OUTPUT:
   result
                   - numpy.ndarray[nfreq,ngs] containing the resulting opacities
FILE OUTPUT:
    dustopac_i.inp - Contains the dust opacities in radmc3d format
    dustopac.inp - Master dust opacity file
```

# **4.2.2.6** def radmc3dPy.analyze.radmc3dDustOpac.writeMasterOpac ( self, ext = None, therm = None, scattering\_mode\_max = 1 )

```
Function to write the master opacity file 'dustopac.inp'

INPUT:
-----
ext : list of dustkappa file name extensions
therm : list of integers specifying whether the dust grain is thermal or quantum heated
(0-thermal, 1-quantum)
```

# 4.3 radmc3dPy.analyze.radmc3dGrid Class Reference

#### **Public Member Functions**

- · def \_\_init\_\_
- · def makeWavelengthGrid
- · def writeWavelengthGrid
- · def makeSpatialGrid
- · def writeSpatialGrid
- def readGrid
- · def getCellVolume

#### **Public Attributes**

- · crd\_sys
- · act dim
- nx
- ny
- nz
- nxi
- nyi
- nzinwav
- nfreq
- x
- у
- z
- xi

This has to be done properly if ppar.has\_key('xres\_nlev'):  $ri_ext = array([self.xi[0], self.xi[0], self.xi[ppar['xres_nspan']]])$  for i in  $range(ppar['xres_nlev'])$ :  $dum_ri = ri_ext[0] + (ri_ext[1]-ri_ext[0]) * arange(ppar['xres_nstep']+1, dtype=float64) / float(ppar['xres_nstep'])$  print  $ri_ext[0:2]/au$  print  $dum_ri/au$   $ri_ext_old = array(ri_ext)$   $ri_ext = array(dum_ri)$   $ri_ext = append(ri_ext_old[2:])$  print  $ri_ext_old[2:])$  print  $ri_ext_old[2:])$  print  $ri_ext_old[2:])$  print  $ri_ext_old[2:])$ 

- yi
- zi
- wav
- freq

### 4.3.1 Detailed Description

Class for the spatial and frequency grid used by RADMC3D

```
ATTRIBUTES:
```

```
crd_sys - 'car'/'cyl'/'sph' coordinate system of the spatial grid
act_dim - A three element vector the i-th element is 1 if the i-th dimension is active, otherwize the i
```

```
- Number of grid points in the x (cartesian) / r (cylindrical) / r (spherical) dimension
   nx
              - Number of grid points in the y (cartesian) / theta (cylindrical) / theta (spherical) dimension
   nv
              - Number of grid points in the z (cartesian) / z (cylindrical) / phi (spherical) dimension
              - Number of cell interfaces in the x (cartesian) / r (cylindrical) / r (spherical) dimension
   nxi
              - Number of cell interfaces in the y (cartesian) / theta (cylindrical) / theta (spherical) dime
   nyi
              - Number of cell interfaces in the z (cartesian) / z (cylindrical) / phi (spherical) dimension
   nwav
              - Number of wavelengths in the wavelength grid
    freq
              - Number of frequencies in the grid (equal to nwav)
              - Cell centered x (cartesian) / r (cylindrical) / r (spherical) grid points
              - Cell centered y (cartesian) / theta (cylindrical) / theta (spherical) \,\, grid points
    У
              - Cell centered z (cartesian) / z (cylindrical) / phi (spherical) grid points
    хi
              - Cell interfaces in the x (cartesian) / r (cylindrical) / r (spherical) dimension
              - Cell interfaces in the y (cartesian) / theta (cylindrical) / theta (spherical) dimension
    уi
    zi
              - Cell interfaces in the z (cartesian) / z (cylindrical) / phi (spherical) dimension
              - Wavelengh grid
    wav
   freq
              - Frequency grid
METHODS:
```

#### 4.3.2 Member Function Documentation

# 4.3.2.1 def radmc3dPy.analyze.radmc3dGrid.getCellVolume ( self )

Function to calculate the volume of grid cells

# 4.3.2.2 def radmc3dPy.analyze.radmc3dGrid.makeSpatialGrid ( self, crd\_sys = None, xbound = None, ybound = None, vbound = None, nxi = None,

Function to create the spatial grid

```
TNPHT.
                - 'car'/'sph' Coordinate system of the spatial grid
    crd svs
                - List (with at least two elements) of boundaries for the grid along the first dimension
                - List (with at least two elements) of boundaries for the grid along the second dimension
                - List (with at least two elements) of boundaries for the grid along the third dimension
    zbound
                 - Number of grid points along the first dimension. List with len(xbound)-1 elements with
    nxi
            \mathtt{nxi}[\mathtt{i}] being the number of grid points between \mathtt{xbound}[\mathtt{i}] and \mathtt{xbound}[\mathtt{i+1}]
                - Same as nxi but for the second dimension
    nvi
                - Same as nxi but for the third dimension
    nzi
OPTIONS:
                - Dictionary containing all input parameters of the model (from the problem_params.inp file)
   ppar
          if ppar is set all keyword arguments that are not set will be taken from this dictionary
```

## 4.3.2.3 def radmc3dPy.analyze.radmc3dGrid.makeWavelengthGrid ( self, wbound = None, nw = None, ppar = None )

```
Function to create a wavelength/frequency grid

INPUT:
-----
    wbound: list of at least two elements containing the wavelength boundaries of the wavelength grid
    nw : list of len(wbound)-1 elements containing the number of wavelengths between the bounds
    set by wbound

OPTIONS:
------
    ppar : parameter dictionary
```

#### 4.3.2.4 def radmc3dPy.analyze.radmc3dGrid.readGrid ( self, fname = ' ' )

Function to read the spatial (amr\_grid.inp) and frequency grid (wavelength\_micron.inp).

```
OPTIONS:

fname - File name from which the spatial grid should be read. If omitted 'amr_grid.inp' will be used.

4.3.2.5 def radmc3dPy.analyze.radmc3dGrid.writeSpatialGrid( self, fname = '')

Function to write the wavelength grid to a file (e.g. amr_grid.inp)

OPTIONS:

fname - File name into which the spatial grid should be written. If omitted 'amr_grid.inp' will be used.

4.3.2.6 def radmc3dPy.analyze.radmc3dGrid.writeWavelengthGrid( self, fname = '')

Function to write the wavelength grid to a file (e.g. wavelength_micron.inp)

OPTIONS:

fname - File name into which the wavelength grid should be written. If omitted 'wavelength_micron.inp' will be used.
```

# 4.4 radmc3dPy.image.radmc3dImage Class Reference

#### **Public Member Functions**

- def \_\_init\_\_
- · def getClosurePhase
- def getVisibility
- def writeFits
- def plotMomentMap
- def getMomentMap
- def readImage
- def imConv

# **Public Attributes**

- · image
- imageJyppix
- x
- у
- nx
- ny
- sizepix\_x
- sizepix\_y
- nfreq
- freq
- nwav
- wav
- stokes
- · psf
- fwhm
- pa
- dpc

```
4.4.1 Detailed Description
RADMC3D image class
ATTRIBUTES:
              - The image as calculated by radmc3d (the values are intensities in erg/s/cm^2/Hz/ster)
    imageJyppix - The image with pixel units of Jy/pixel
               - x coordinate of the image [cm]
               - y coordinate of the image [cm]
               - Number of pixels in the horizontal direction
   nx
   ny
               - Number of pixels in the vertical direction
               - Pixel size in the horizontal direction [cm]
   sizepix_x
              - Pixel size in the vertical direction [cm]
   sizepix_y
   nfreq
               - Number of frequencies in the image cube
   freq
               - Frequency grid in the image cube
               - Number of wavelengths in the image cube (same as nfreq)
   nwav
   wav
               - Wavelength grid in the image cube
4.4.2 Member Function Documentation
4.4.2.1 def radmc3dPy.image.radmc3dImage.getClosurePhase ( self, bl = None, pa = None, dpc = None)
Function to calculate clusure phases for a given model image for any arbitrary baseline triplet
INPUT:
   bl
            - a list or Numpy array containing the length of projected baselines in meter(!)
            - a list or Numpy array containing the position angles of projected baselines in degree(!)
            - distance of the source in parsec
```

NOTE, bl and pa should either be an array with dimension [N,3] or if they are lists each element of the list should be a list of length 3, since closure phases are calculated only for closed triangles

# OUTPUT:

```
returns a dictionary with the following keys:
b1
      - projected baseline in meter
      - position angle of the projected baseline in degree
ра
      - number of baselines
nbl
      - spatial frequency along the x axis of the image
      - spatial frequency along the v axis of the image
      - complex visibility at points (u,v)
vis
      - correlation amplitude
phase - Fourier phase
ср
      - closure phase
wav
      - wavelength
```

# 4.4.2.2 def radmc3dPy.image.radmc3dImage.getMomentMap ( self, moment = 0, nu0 = 0, wav0 = 0 )

nwav - number of wavelengths

### 4.4.2.3 def radmc3dPy.image.radmc3dImage.getVisibility ( self, bl = None, pa = None, dpc = None)

```
Function to calculate visibilities for a given set of projected baselines and position angles
with the Discrete Fourier Transform
   bl
             - a list or Numpy array containing the length of projected baselines in meter(!)
             - a list or Numpy array containing the position angles of projected baselines in degree(!)
             - distance of the source in parsec
OUTPUT:
   returns a dictionary with the following keys:
bl
      - projected baseline in meter
      - position angle of the projected baseline in degree
ра
      - number of baselines
nhl
       - spatial frequency along the x axis of the image
       - spatial frequency along the v axis of the image
vis
       - complex visibility at points (u,v)
       - correlation amplitude
amp
phase - phase
wav
      - wavelength
      - number of wavelengths
nwav
4.4.2.4 def radmc3dPy.image.radmc3dImage.imConv ( self, fwhm = None, pa = None, dpc = 1.)
Function to convolve a radmc3d image with a two dimensional Gaussian psf
INPUT:
      fwhm
             : A list of two numbers; the FWHM of the two dimensional psf along the two principal axes
           The unit is assumed to be arcsec
            : Position angle of the psf ellipse (counts from North counterclockwise)
              : Distance of the source in pc
      dpc
OUTPUT:
      result : same
      'cimage': The convolved image with the psf (unit is erg/s/cm/cm/Hz/ster)
      'image' : The original unconvolved image (unit is erg/s/cm/Hz/ster)
            : Two dimensional psf
             : first coordinate axis of the psf/image
      ′ x′
              : second coordinate axis of the psf/image
4.4.2.5 def radmc3dPy.image.radmc3dImage.plotMomentMap ( self, moment = 0, nu0 = 0, wav0 = 0, dpc = 1., au = 0
      False, arcsec = False, cmap = None, vclip = None)
Function to plot moment maps
INPUT:
   moment : moment of the channel maps to be calculated
    nu0
          : rest frequency of the line in Hz
    wav0
           : rest wavelength of the line in micron
           : distance of the source in pc
    dpc
```

#### 4.4.2.6 def radmc3dPy.image.radmc3dImage.readImage ( self, fname = None, binary = False )

: If true displays the image with AU as the spatial axis unit

arcsec : If true displays the image with arcsec as the spatial axis unit (dpc should also be set!)

vclip : two element list / Numpy array containin the lower and upper limits for the values in the moment

Function to read an image calculated by RADMC3D  $\,$ 

: matplotlib colormap

map to be displayed

```
INPUT:
         : file name of the radmc3d output image (if omitted 'image.out' is used)
binary : False - the image format is formatted ASCII if True - C-compliant binary
4.4.2.7 def radmc3dPy.image.radmc3dImage.writeFits ( self, fname = ' ', dpc = 1 ., coord =
      '03h10m05s -10d05m30s', bandwidthmhz = 2000.0, casa = False, nu0 = 0., wav0 = 0., stokes =
      ' I', fitsheadkeys = [ ] )
Function to write out a RADMC3D image data in fits format (CASA compatible)
fname
             : File name of the radmc3d output image (if omitted 'image.fits' is used)
             : Image center coordinates
bandwidthmhz : Bandwidth of the image in MHz (equivalent of the CDELT keyword in the fits header)
         : If set to True a CASA compatible four dimensional image cube will be written
 nu0
              : Rest frequency of the line (for channel maps)
 wav0
              : Rest wavelength of the line (for channel maps)
 stokes
              : Stokes parameter to be written if the image contains Stokes IQUV (possible
       choices: 'I', 'Q', 'U', 'Y', 'PI' -Latter being the polarized intensity)
 fitsheadkeys: Dictionary containing all (extra) keywords to be added to the fits header. If
        the keyword is already in the fits header (e.g. CDELT1) it will be updated/changed
        to the value in fitsheadkeys, if the keyword is not present the keyword is added to
        the fits header.
```

# 4.5 radmc3dPy.analyze.radmc3dPar Class Reference

#### **Public Member Functions**

- def init
- · def readPar
- · def setPar
- def loadDefaults
- def printPar
- · def writeParfile

### **Public Attributes**

- ppar
- pdesc
- pblock
- pvalstr
- pvarstr

#### **Static Public Attributes**

- list dumlist []
- string dumline '-'
- tuple dumline rfile.readline()
- · comment False
- list varlist []
- int iline 0
- list ind dumlist[iline]
- list blockname dumlist[iline]

- list vlist dumlist[iline]
- list **lbind** vlist[1]
- list cind vlist[1]
- · inBrokenLine False
- list expr vlist[1]
- · list com vlist[1]
- · string com ' '
- int iline2 0
- list dummy dumlist[iline + iline2]
- tuple cind2 dummy.find('#')
- tuple Ibind2 dummy.find('\\')
- iline iline+iline2
- tuple glob globals()
- tuple loc locals()
- tuple val eval(varlist[i][1], glob)

## 4.5.1 Detailed Description

```
Class for parameters in a RADMC-3D model
ATTRIBUTES:
           : Dictionary containing parameter values with parameter names as keys
    pdesc : Disctionary containing parameter description (comments in the parameter file) with parameter name
    pblock : Dictionary containing the block names in the parameter file and parameter names as values
    pvalstr: Dictionary containing parameter values as strings with parameter names as keys
4.5.2 Member Function Documentation
4.5.2.1 def radmc3dPy.analyze.radmc3dPar.loadDefaults ( self, model = ' ' , ppar = {}, reset = True )
Function to fill up the classs attributes with default values
OPTIONS:
   model - Model name whose paraemters should also be loaded
    ppar - Dictionary containing parameter values as string and parameter names as keys
   Default values will be re-set to the values in this dictionary
    reset - If True the all class attributes will be re-initialized before
    the default values would be loaded. I.e. it will remove all entries
    from the dictionary that does not conain default values either in this
    function or in the optional ppar keyword argument
```

#### 4.5.2.2 def radmc3dPy.analyze.radmc3dPar.printPar ( self )

Print the parameters of the current model

## 4.5.2.3 def radmc3dPy.analyze.radmc3dPar.readPar ( self, fname = ' ' )

```
Function to read a parameter file
The parameters in the files should follow the python syntax
INPUT:
```

fname : file name to be read (if omitted problem\_params.inp is used)

```
OUTPUT:

Returns a dictionary with the parameter names as keys

4.5.2.4 defradmc3dPy.analyze.radmc3dPar.setPar( self, parlist = [] )

Function to add parameter to the radmc3DPar parameter class
If the parameter is already defined its value will be modified

INPUT:

parlist - If the parameter is already defined parlist should be a two element list 1) parameter name, 2) parameter expression/value as a string

If the parameter is not yet defined parlist should be a four element list 1) parameter name, 2) parameter expression/value as a string

3) Parameter description (= comment field in the parameter file)

4.5.2.5 defradmc3dPy.analyze.radmc3dPar.writeParfile( self, fname = '' )

Function to write a parameter file

INPUT:

-----
fname : File name to be read (if omitted problem_params.inp is used)
```

# 4.6 radmc3dPy.analyze.radmc3dStars Class Reference

# **Public Member Functions**

- def \_\_init\_\_
- · def findPeakStarspec
- · def readStarsinp
- · def writeStarsinp
- def getStellarSpectrum

## **Public Attributes**

- · mstar
- tstar
- rstar
- Istar
- nstar
- pstar
- wav
- freq
- fnu
- nwav
- · nfreq

# 4.6.1 Detailed Description

```
Class of the radiation sources (currently only stars)
ATTRIBUTES:
   mstar - List of stellar masses
    tstar - List of stellar effective temperatures
    rstar - List of stellar radii
    lstar - List of stellar luminosities
    nstar - Number of stars
    pstar - Locations (coordinates) of the stars
    wav - Wavelength for the stellar spectrum
    freq - Frequency for the stellar spectrum
          - Stellar spectrum (flux@1pc)
    nwav - Number of wavelenghts in the stellar spectrum
    nfreq - Number of frequencies in the stellar spectrum
4.6.2 Member Function Documentation
4.6.2.1 def radmc3dPy.analyze.radmc3dStars.findPeakStarspec ( self )
Function to calculate the peak wavelength of the stellar spectrum
OUTPUT:
  Returns the peak wavelength of the stellar spectrum in nu*Fnu for all
stars as a list
4.6.2.2 def radmc3dPy.analyze.radmc3dStars.getStellarSpectrum ( self, tstar = None, rstar = None, lstar = None, nu =
       None, wav = None )
Function to calculate a blackbody stellar spectrum
INPUT:
   tstar : Effective temperature of the star in [K]
    rstar : Radius of the star in [cm]
    lstar : Bolometric luminosity of the star [erg/s] (either rstar or lstar should be given)
         : frequency grid on which the spectrum should be calculated [Hz]
         : wavelength grid on which the spectrum should be calculated [micron]
4.6.2.3 def radmc3dPy.analyze.radmc3dStars.readStarsinp ( self, fname = ' ' )
Function to read the stellar data from the stars.inp file
OPTIONS:
    fname - File name of the file that should be read (if omitted stars.inp will be used)
4.6.2.4 def radmc3dPy.analyze.radmc3dStars.writeStarsinp ( self, wav = None, freq = None, pstar = None, tstar =
       None )
Writes the stars.inp file
INPUT:
         - Wavelength grid for the stellar spectrum
    freq - Frequency grid for the stellar spectrum (either freq or wav should be set)
    pstar - List of the cartesian coordinates of the stars (each element of pstar should be a list of three el
    with the [x,y,z] coordinate of the individual stars)
    tstar - List containing the effective temperature of the stars
```

# Index

add_par	get_gas_density
radmc3dPy::analyze::radmc3dPar, 40	radmc3dPy::model_lines_nlte_lvg_1d_1, 14
	radmc3dPy::model_ppdisk, 16
cmask	radmc3dPy::model_template, 21
radmc3dPy::image, 11	get_gas_temperature
csrot	radmc3dPy::model_lines_nlte_lvg_1d_1, 14
radmc3dPy::crd_trans, 8	radmc3dPy::model_template, 21
ctrans_sph2cart	get_model_desc
radmc3dPy::crd_trans, 9	radmc3dPy::setup, 24
ctrans_sph2cyl	get_model_names
radmc3dPy::crd_trans, 9	radmc3dPy::setup, 24
, _ ,	get momentmap
find_peak_starspec	radmc3dPy::image::radmc3dImage, 37
radmc3dPy::analyze::radmc3dStars, 42	get_psf
	radmc3dPy::image, 11
get_cell_volume	get_sigmadust
radmc3dPy::analyze::radmc3dGrid, 35	radmc3dPy::analyze::radmc3dData, 28
get closure phase	get_sigmagas
radmc3dPy::image::radmc3dImage, 37	
get_default_params	radmc3dPy::analyze::radmc3dData, 28
radmc3dPy::model_lines_nlte_lvg_1d_1, 13	get_stellar_spectrum
radmc3dPy::model ppdisk, 15	radmc3dPy::analyze::radmc3dStars, 42
radmc3dPy::model_simple_1, 17	get_tau
radmc3dPy::model_spher1d_1, 18	radmc3dPy::analyze::radmc3dData, 28
radmc3dPy::model_spher2d_1, 18	get_tau_1dust
radmc3dPy::model_template, 20	radmc3dPy::analyze::radmc3dData, 28
radmc3dPy::model_test_scattering_1, 22	get_template_model
get_desc	radmc3dPy::setup, 24
radmc3dPy::model_lines_nlte_lvg_1d_1, 13	get_velocity
radmc3dPy::model_ppdisk, 16	radmc3dPy::model_lines_nlte_lvg_1d_1, 14
radmc3dPy::model_simple_1, 17	radmc3dPy::model_ppdisk, 16
	radmc3dPy::model_template, 21
radmc3dPy::model_spher1d_1, 18	get_visibility
radmc3dPy::model_spher2d_1, 19	radmc3dPy::image::radmc3dImage, 37
radmc3dPy::model_template, 20	get_vturb
radmc3dPy::model_test_scattering_1, 22	radmc3dPy::model_lines_nlte_lvg_1d_1, 15
get_dust_density	radmc3dPy::model_template, 21
radmc3dPy::model_lines_nlte_lvg_1d_1, 13	
radmc3dPy::model_ppdisk, 16	imconv
radmc3dPy::model_simple_1, 17	radmc3dPy::image::radmc3dImage, 38
radmc3dPy::model_spher1d_1, 18	
radmc3dPy::model_spher2d_1, 19	load_defaults
radmc3dPy::model_template, 20	radmc3dPy::analyze::radmc3dPar, 40
radmc3dPy::model_test_scattering_1, 22	
get_dust_temperature	make_spatial_grid
radmc3dPy::model_lines_nlte_lvg_1d_1, 13	radmc3dPy::analyze::radmc3dGrid, 35
radmc3dPy::model_template, 20	make_wav_grid
get_gas_abundance	radmc3dPy::analyze::radmc3dGrid, 35
radmc3dPy::model_lines_nlte_lvg_1d_1, 14	makeimage
radmc3dPy::model_ppdisk, 16	radmc3dPy::image, 11
radmc3dPv::model_template, 20	makeopac

44 INDEX

radmc3dPy::analyze::radmc3dDustOpac, 32	mixopac, 32
mixopac	read_masteropac, 33
radmc3dPy::analyze::radmc3dDustOpac, 32	readopac, 33
	run_makedust, 33
plot_momentmap	write_masteropac, 33
radmc3dPy::image::radmc3dImage, 38	radmc3dPy::analyze::radmc3dGrid
plotimage	get_cell_volume, 35
radmc3dPy::image, 12	make_spatial_grid, 35
problem_setup_dust	make_wav_grid, 35
radmc3dPy::setup, 24	read_grid, 35
problem_setup_gas	write_spatial_grid, 36
radmc3dPy::setup, 25	write_wav_grid, 36
	radmc3dPy::analyze::radmc3dPar
radmc3dPy, 5	add_par, 40
radmc3dPy.analyze, 5	load_defaults, 40
radmc3dPy.analyze.radmc3dData, 27	readparams, 40
radmc3dPy.analyze.radmc3dDustOpac, 31	write_parfile, 41
radmc3dPy.analyze.radmc3dGrid, 34	radmc3dPy::analyze::radmc3dStars
radmc3dPy.analyze.radmc3dPar, 39	find_peak_starspec, 42
radmc3dPy.analyze.radmc3dStars, 41	get_stellar_spectrum, 42
radmc3dPy.crd_trans, 8	read_starsinp, 42
radmc3dPy.image, 10	write_starsinp, 42
radmc3dPy.image.radmc3dImage, 36	radmc3dPy::crd_trans
radmc3dPy.model_lines_nlte_lvg_1d_1, 13	csrot, 8
radmc3dPy.model_ppdisk, 15	ctrans_sph2cart, 9
radmc3dPy.model_simple_1, 17	ctrans_sph2cult, 9
radmc3dPy.model_spher1d_1, 17	— · ·
radmc3dPy.model_spher2d_1, 18	vrot, 9
radmc3dPy.model_template, 19	vtrans_sph2cart, 10
radmc3dPy.model_test_scattering_1, 22	radmc3dPy::image
radmc3dPy.natconst, 22	cmask, 11
radmc3dPy.setup, 23	get_psf, 11
radmc3dPy::analyze	makeimage, 11
read_data, 6	plotimage, 12
read_grid, 6	readimage, 12
read_spectrum, 7	radmc3dPy::image::radmc3dImage
readopac, 7	get_closure_phase, 37
readparams, 8	get_momentmap, 37
write_default_parfile, 8	get_visibility, 37
radmc3dPy::analyze::radmc3dData	imconv, 38
get_sigmadust, 28	plot_momentmap, 38
get_sigmagas, 28	readimage, 38
get_tau, 28	writefits, 39
get_tau_1dust, 28	radmc3dPy::model_lines_nlte_lvg_1d_1
read_dustdens, 29	get_default_params, 13
read_dusttemp, 29	get_desc, 13
read_gasdens, 29	get_dust_density, 13
read_gastemp, 29	get_dust_temperature, 13
read_gasvel, 29	get_gas_abundance, 14
read_vturb, 30	get_gas_density, 14
write_dustdens, 30	get_gas_temperature, 14
write_dusttemp, 30	get_velocity, 14
write_gasdens, 30	get_vturb, 15
write_gastemp, 30	radmc3dPy::model_ppdisk
write_gasvel, 30	get_default_params, 15
write_vtk, 31	get_desc, 16
write_vturb, 31	get_dust_density, 16
radmc3dPy::analyze::radmc3dDustOpac	get_gas_abundance, 16
makeopac, 32	get_gas_density, 16
/ -	gg · · · · · · · · · · ·

INDEX 45

get_velocity, 16	readimage
radmc3dPy::model_simple_1	radmc3dPy::image, 12
get_default_params, 17	radmc3dPy::image::radmc3dImage, 38
get_desc, 17	readopac
get_dust_density, 17	radmc3dPy::analyze, 7
radmc3dPy::model_spher1d_1	radmc3dPy::analyze::radmc3dDustOpac, 33
get_default_params, 18	readparams
get_desc, 18	radmc3dPy::analyze, 8
get_dust_density, 18	radmc3dPy::analyze::radmc3dPar, 40
radmc3dPy::model_spher2d_1	run_makedust
get_default_params, 18	radmc3dPy::analyze::radmc3dDustOpac, 33
get_desc, 19	
get_dust_density, 19	vrot
radmc3dPy::model_template	radmc3dPy::crd_trans, 9
get_default_params, 20	vtrans_sph2cart
get_desc, 20	radmc3dPy::crd_trans, 10
get_dust_density, 20	
get_dust_temperature, 20	write_default_parfile
get_gas_abundance, 20	radmc3dPy::analyze, 8
get_gas_density, 21	write_dustdens
get_gas_temperature, 21	radmc3dPy::analyze::radmc3dData, 30
	write_dusttemp
get_velocity, 21	radmc3dPy::analyze::radmc3dData, 30
get_vturb, 21	write_gasdens
radmc3dPy::model_test_scattering_1	radmc3dPy::analyze::radmc3dData, 30
get_default_params, 22	write_gastemp
get_desc, 22	radmc3dPy::analyze::radmc3dData, 30
get_dust_density, 22	write_gasvel
radmc3dPy::setup	radmc3dPy::analyze::radmc3dData, 30
get_model_desc, 24	write_lines_inp
get_model_names, 24	radmc3dPy::setup, 26
get_template_model, 24	write_masteropac
problem_setup_dust, 24	radmc3dPy::analyze::radmc3dDustOpac, 33
problem_setup_gas, 25	write_parfile
write_lines_inp, 26	radmc3dPy::analyze::radmc3dPar, 41
write_radmc3d_inp, 26	write_radmc3d_inp
read_data	radmc3dPy::setup, 26
radmc3dPy::analyze, 6	write_spatial_grid
read_dustdens	radmc3dPy::analyze::radmc3dGrid, 36
radmc3dPy::analyze::radmc3dData, 29	write starsinp
read_dusttemp	radmc3dPy::analyze::radmc3dStars, 42
radmc3dPy::analyze::radmc3dData, 29	write_vtk
read_gasdens	radmc3dPy::analyze::radmc3dData, 31
radmc3dPy::analyze::radmc3dData, 29	write_vturb
read gastemp	radmc3dPy::analyze::radmc3dData, 31
radmc3dPy::analyze::radmc3dData, 29	write wav grid
read_gasvel	radmc3dPy::analyze::radmc3dGrid, 36
radmc3dPy::analyze::radmc3dData, 29	writefits
read grid	radmc3dPy::image::radmc3dImage, 39
radmc3dPy::analyze, 6	radificoar ymageradificoaffiage, 00
radmc3dPy::analyze::radmc3dGrid, 35	
read_masteropac	
radmc3dPy::analyze::radmc3dDustOpac, 33	
read_spectrum	
radmc3dPy::analyze, 7	
read_starsinp	
radmc3dPy::analyze::radmc3dStars, 42	
read_vturb	
radmc3dPy::analyze::radmc3dData, 30	