OpTool User Guide

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Introduction

This tool allows to produce complex dust opacities right from the command line. It is derived from Michiel Min's OpacityTool for the DIANA program, and committed to its simplicity. For a more wide-ranging toolset to produce opacities, check out SIGMA, the Simple Icy Grain Model for Aggregates¹, by Lefévre et al. (2020), also derived from OpacityTool. Optool uses its effective medium mixing tool set, but is otherwise an independant development.

Capabilities

- fully command line driven, no input files need to be edited
- combining materials through mixing into a complex grain
- built-in materials, but can also be used also external data
- DHS to model shape effects and low-porosity aggregates
- full scattering matrix output in a variety of formats and normalizations.

Terms of use

The code doing the real work of optool was adopted and modified from the publicly available versions of the OpacityTool and SIGMA. I focused on the interface, on simplicity, and a clean set of conventions. If you are using this tool, you should give credit to the papers describing the methods used to compute the opacities, as well as to the original laboratory papers that published the refractive index measurements.

- DHS model for irregular grains: Min et al. 2005, A&A, 432, 909
- DIANA standard Opacities: Woitke, Min et al. 2016
- SIGMA: Lefévre, Min et al. 2020, A&A (submitted)
- Third party software: Toon et al. 1981, Applied Optics 20, 3657
- References to refractive index data used in your particular application.

¹SIGMA supports non-powerlaw size distributions, size-dependent porosities, and adding independent grain types in a single run to produce combined opacities. It can reproduce and modify many classical dust models in a computationally efficient way.

Compiling optool

On most systems, you can download and compile optool with these simple steps, using the freely available GNU FORTRAN compiler gfortran.

```
git clone https://github.com/cdominik/optool.git
cd optool
make
```

The executable is called optool, and you should put it on your execution path. If you prefer to use the Intel fortran compiler, if you have the cfitsio library installed and would like to write the opacities into FITS files instead of ASCII, or if you do not have working OpenMP installation and need to turn off parallel processing, then use one or more of these parameters during compilation:

Examples

A simple grain made only of the default pyroxene

```
optool -c pyr
```

Reproduce the DIANA standard dust model, using the pyroxene with 70% magnesium, and 13% (by mass) carbon, and 25% porosity.

```
optool -c pyr-mg70 0.87 -c c-z 0.13 -p 0.25
```

List the built-in materials

```
optool -c ?
```

Add an ice mantle that is 20% of the core mass

```
optool -c pyr-mg70 0.87 -c c-z 0.13 -m ice-w 0.2 -p 0.25
```

Like the previous example, but use ice refractive index data from a separate file.

```
optool -c pyr-mg70 0.87 -c c-z 0.13 -m ~/lnk/ice_hudgins_2.dat 0.2 -p 0.25
```

Pure ice grains in a narrow size distribution from 1 to 3 microns, with 15 sample sizes following an $f(a) \propto a^{-2.5}$ powerlaw size distribution. Also, restrict the wavelength range to $10\text{-}100\mu\text{m}$.

```
optool -c ice-w 1.0 -a 1 3 2.5 15 -l 10 100
```

Represent the default dust model (DIANA) in 42 grain sizes, and produce input files for RADMC-3D, one for each grain size, with full scattering matrix.

```
optool -na 42 -d -s -radmc
```

Command line options

-h, -help

Show a compact help message about command line options.

Grain composition and geometry

-c ?

List available built-in materials (the keys for the **-c** and **-m** options).

-c KEY-or-FILE [Mfrac]

Specify a material to include into the grain. KEYorFILE can be the key for a builtin material, or the path to the correct lnk file. Mfrac is the mass fraction of the material. You can give up to 10 -c switches to add the different contributions. All materials will be mixed together using the Bruggeman rule, and vacuum can be included through the porosity.

-m KEY-or-FILE [Mfrac]

Like -c, but use this material as a mantle material that will be mixed in using the Maxwell-Garnett rule instead of the Bruggeman rule. Only one -m switch is allowed, and it needs to come after all -c specifications.

-p POROSITY [P_MANTLE]

Porosity, the *volume* fraction of vacuum, a number smaller than 1. The default is 0. A single value will apply to both core and mantle, but a second value will be specific for the mantle (and may be 0).

-fmax VHMAX

Maximum *volume* fraction of the inner hole for the DHS approach. The default is 0.8. Zero means to use solid spheres, i.e. perfectly regular grains.

Grain size distribution

-a AMIN AMAX [NA [APOW]]

Specify minimum and maximum grain radius, and optionally the number of size bins and the size distribution powerlaw. You can also use options to set individual values with **-amin**, **-amax**, **-na**, **-apow**. The defaults are 0.05 μ m, 3000 μ m, 50, and 3.5, respectively.

Wavelength grid

-1 LMIN LMAX [NLAM]

Minimum and maximum wavelength and optionally the number of wavelengths points for the construction of the wavelength grid. The default values are 0.05 μ m, 10000 μ m, and 300, respectively. Use the options -lmin, -lmax, and -nlam (or -nl) to set individual values.

-1 FILE

Read the wavelength grid from the first column of FILE. The first number in the file must be the number of grid points. So this could be an lnk file.

Controlling the output

The standard output is the file dustkappa.dat, with the opacities and the asymmetry parameter g. The following options control and extend output. See the section Output files for details on the files that can be produced.

-o [DIR]

Put the output files in directory DIR instead of the current working directory. If you use -o without specifying a directory, ./output will be used.

-s [NANG]

Include the full scattering matrix in the output. NANG can optionally specify the number of equally-spaced steps to cover the range of angles between 0 and 180 degrees. The default for NANG is 180.

-d [NSUB]

Divide the computation up into NA parts to produce a file for each grain size. Each size will actually be an average over a small range of NSUB grains around the real size, to smear out resonances. The default for NSUB is 5.

-fits

Write dustkappa.fits with the absorption cross sections and scattering matrix elements, instead of ASCII output. With the -d switch, NA such files will be written, one for each grain size.

-radmc [LABEL]

The structure of the ASCII output of optool is already compatible with RADMC-3D. However, RADMC-3D uses a different normalization for the scattering matrix, see the appendix for more information. With this switch, the extension of the files will be changed to .inp, the normalization will be modified, and if you specify LABEL, it will be used in the file name(s).

-t [TMIN [TMAX [NT]]]

Compute mean opacities per g of dust mass, κ_{Planck} and $\kappa_{\text{Rosseland}}$, in the given temperature interval, in nt logarithmic steps, with output to dustkapmean.dat. The parameters default to 10K, 10000K, and 200, respectively.

Material properties

optool needs refractive index data to work. For your convenience, a useful list of materials is compiled into optool, but you can also find and use other data. No matter where the data is from, you should *always* cite the original laboratory papers.

Built-in materials

To access one of the built-in materials, specify the corresponding key string like pyr-mg70 instead of the path to an lnk file. In each material class I have selected a useful default, accessible with an even simpler generic key. The built-in list might be enough for opacity computation. If your goal is detailed mineralogical fits, you will have to use specialized refractive index data.

- c Key generic	- c Key full key	Material	State	$ ho$ [g/cm 3]	$\lambda_{min} \ [\mum]$	Reference
generic	pyr-mg100	MgSiO ₃	amorphous	2.71	0.2	Dorschner+1995
	pyr-mg95	$Mg_{0.95}Fe_{0.05}SiO_3$	amorphous	2.74	0.2	Dorschner+1995
	pyr-mg80	$Mg_{0.8}Fe_{0.2}SiO_3$	amorphous	2.9	0.2	Dorschner+1995
pyr	pyr-mg70	$Mg_{0.7}Fe_{0.3}SiO_3$	amorphous	3.01	0.2	Dorschner+1995
	pyr-mg60	$Mg_{0.6}Fe_{0.4}SiO_3$	amorphous	3.1	0.2	Dorschner+1995
	pyr-mg50	$Mg_{0.5}Fe_{0.5}SiO_3$	amorphous	3.2	0.2	Dorschner+1995
	pyr-mg40	$Mg_{0.4}Fe_{0.6}SiO_3$	amorphous	3.3	0.2	Dorschner+1995
ens	pyr-c-mg96	$Mg_{0.96}Fe_{0.04}SiO3$	crystalline	2.8	2.0	Jäger+1998
ol	ol-mg50	MgFeSiO ₄	amorphous	3.71	0.2	Dorschner+1995
	ol-mg40	$Mg_{0.8} Fe_{1.2} SiO_4$	amorphous	3.71	0.2	Dorschner+1995
for	ol-c-mg100	Mg ₂ SiO ₄	crystalline	3.33	3.0	Steyer+1974
С	C-Z	С	amorphous?	1.8	0.05	Zubko+1996
	c-p	C	amorphous	1.8	0.11	Preibisch+1993
ice	ice-w	Water ice	crystalline	0.92	0.04	Warren+2008
iron	fe-c	Fe	metallic	7.87	0.1	Henning+1996
cor	cor-c	Al_2O_3	crystalline	4.0	0.5	Koike+1995

External refractory index files (lnk files)

optool can use external refractive index data in files with the following format:

- The file may start with several comment lines (lines starting with !, #, or *).
- The next line contains two numbers, the number of wavelengths N_{λ} and the specific weight ρ of the material in g/cm³
- Then follow three columns of data: $\lambda[\mu m]$, and the real and imaginary parts of the refractive index, n and k.

You can use any of the refractory index files provided by Charléne Lefévre's SIGMA package out of the box, or find other files, for example by using the Jena database. Don't forget to add the line with N_{λ} and $\rho!$ If for some reason it is not convenient to add that line to the file, optool will count the lines for you and you can specify the density after the mass fraction, like this: optool -c path/to/file.lnk 0.7 3.42

Output files

dustkappa.dat

This is an ASCII file containing the basic opacity results. It starts with a comment section describing the dust model, followed by the format number (3, currently), followed by the number of wavelengths in the grid, both on lines by themselves. Then follows a block with these columns:

- 1. wavelength λ [micron]
- 2. mass absorption cross section $\kappa_{\rm abs} \ [{\rm cm}^2/{\rm g}]$
- 3. mass scattering cross section $\kappa_{\rm sca}$ [cm²/g]
- 4. asymmetry parameter g

dustkapscatmat.dat

ASCII file with cross sections and full scattering matrix. The comment section at the start of the file explains the structure. See the appendix for information about the normalization of the scattering matrix. And see the <code>-radmc</code> switch which will modify the output to make sure it can be used as an input file for RADMC-3D.

dustkappa.fits

The fits-file (ending in '.fits') is written instead of the ASCII output when using the -fits switch. It has two HDU blocks. The first block contains the cross sections per unit mass. This is a $N_{\lambda} \times 5$ matrix with these columns:

- 1. wavelengths in $[\mu m]$
- 2. mass extinction cross section $\kappa_{\rm ext}$ in [cm²/g]
- 3. mass absorption cross section $\kappa_{\rm abs}$ in [cm²/g]
- 4. mass scattering cross section $\kappa_{\rm sca}$ in [cm²/g]
- 5. asymmetry factor g

The second HDU block contains the scattering matrix elements. This is a $N_{\lambda} \times 6 \times N_{\rm ang}$ matrix. These are the 6 elements of the scattering matrix for 180 equidistant scattering angles from forward scattering (element 0) to backward scattering (element 179) for each wavelength value. The 6 elements stored are F_{11} , F_{12} , F_{22} , F_{33} , F_{34} , F_{44} respectively.

dustkapmean.dat

This file will only be written with the -t switch. It contains 3 columns: (1) T [K], (2) κ_{Planck} , (3) κ_{Ross} , both in cm² per gram of *dust*. Note that dust evaporation is not considered, and that a wide wavelengths coverage is needed for good results.

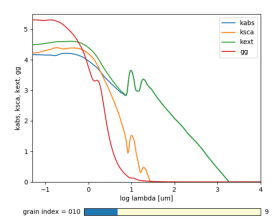
Inspecting the computed optical properties

To try out optool you could use one of these commands

The commands will run optool with the standard DIANA material properties, and then use the python script optool.py to plot the computed opacities. You will get two plots.

- a plot showing the opacities κ_{abs} , κ_{sca} , and κ_{ext} as a function of wavelength, along with the asymmetry parameter g. Note that the red g curve does not have its own scale, imagine the g axis going from 0 to 1 for g.
- a plot showing the scattering matrix elements as a function of scattering angle, and with sliders to go through grain sizes and wavelengths. The y axis of the plot is actually $\log_{10}(|F_{ij}|)$, so the downward peaks are actually places where the matrix element goes through zero.

If you want to run the plotter yourself, ipython -i path/to/optool.py is the correct command to do so, from the directory where the output files are located.



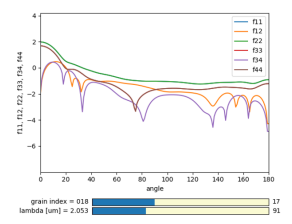


Figure 1: Screenshot of the plots created by running ipython -i optool.py. Note that we plot the logarithm of the absolute value of the scattering matrix, in order to deal with range and sign issues.

Acknowledgments

- Michiel Min for the DIANA OpacityTool and all the incredible work that went into it. optool is a direct derivative of that tool and resuses almost all of its code.
- Charléne Lefévre for SIGMA. Optool uses the generalized Bruggeman solver from SIGMA's code base.
- Kees Dullemond for his python plotting routine viewarr (available on github), and code for computing Planck and Rosseland means opacities.
- Jeroen Bouwman for pointers to refractive index data.

Appendix

Units

Due to conventions in our field, the input and output of optool uses the following units

- microns for grain sizes and wavelengths
- g/cm^3 for mass densities of materials
- cm² g⁻¹ for opacities κ_{abs} , κ_{sca} , and κ_{ext}
- sr^{-1} or cm^2 g^{-1} sr^{-1} for the scattering matrix elements, see below.
- Kelvin for temperatures

Scattering phase function normalization

A number of different normalizations for the scattering matrix are being used in the literature and in computational tools. The differences are significant, and it is important to be aware of the choice. For optool we are using a convention in which the average over all directions of the 1-1 element of the scattering matrix equals unity, i.e.

$$\oint_{(4\pi)} F_{11}(\lambda, \Theta) d\Omega = 4\pi \tag{1}$$

See Hovenier (2004) for a discussion of this normalization. optool can also produce output for RADMC-3D which uses a different normalization, namely

$$\oint_{(4\pi)} Z_{11}(\lambda, \Theta) d\Omega = \kappa_{\rm sca}(\lambda) \tag{2}$$

Note that the output in the fits files contains 180 angles for the scattering matrix, from 0-179 degrees, because this is what Michiel Min's code produces. For the ASCII files, we add a 180 degree point, because RADMC-3D requires it.

How to ingest another material

Additional refractive index data tables can be compiled into the code. Here is how:

- 1. Give your lnk file a name exactly like pyr-mg70-Dorschner1995.lnk, where pyr-mg70 is the key to access the material and Dorschner1995 is the reference.
- 2. Put this file into the lnk_data directory.
- 3. Optionally edit lnk_data/lnk-help.txt, so that optool -c? will list the new material. Note that, in order to define generic keys, optool looks for pairs like genkey -> fullkey in this file.
- 4. Run make ingest to update ref_ind.f90, now including your new material.
- 5. Recompile and install the code.

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