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# **RAPOC Documentation**

***Release 1.0.4***

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**May 05, 2021**



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Developed by Lorenzo V. Mugnai and Darius Modirrousta-Galian, the RAPOC code is the product of a collaboration between Sapienza Università di Roma, Università degli Studi di Palermo and INAF - Osservatorio Astronomico di Palermo.





*RAPOC* uses molecular absorption measurements (i.e. wavelength-dependent opacities) to calculate Rosseland and Planck mean opacities that are commonly used in atmospheric modelling.

*RAPOC* is designed to be simple, straightforward, and easily incorporated into other codes. It is completely written in Python and documented with docstrings.

## INSTALLATION & UPDATES

### 1.1 Installing from Pypi

*RAPOC* can be installed from the Pypi repository with the following script:

```
pip install rapoc
```

### 1.2 Installing from git

*RAPOC* may also be cloned from the main git repository:

```
git clone https://github.com/ExObsSim/Rapoc-public.git
```

The next step is to move into the *RAPOC* folder:

```
cd /your_path/Rapoc
```

Then:

```
pip install .
```

To check if one has the correct setup:

```
python -c "import rapoc"
```

### 1.3 Uninstall Rapoc

*RAPOC* is installed in your system as a standard python package: to uninstall it completely:

```
pip uninstall rapoc
```

## 1.4 Update Rapoc

### 1.4.1 Update from Pypi

To update *RAPOC* to the latest stable version:

```
pip install rapoc --upgrade
```

### 1.4.2 Update from source

One can download or pull a newer version of *RAPOC* over the old one, replacing all modified data.

Then one has to place themselves inside the installation directory within the console:

```
cd /your_path/Rapoc
```

*RAPOC* can now be updated:

```
pip install . --upgrade
```

or simply:

```
pip install .
```



## USER GUIDE

This guide is an instruction manual on the *RAPOC* (Rosseland And Planck Opacity Converter) tool and how it can be included in one's own code.

For a quickstart, please refer to the python notebook inside the *examples* directory. If *RAPOC* has been installed from Pypi, an example notebook can be found on the [GitHub](#) repository.

### 2.1 Input

Everything inside *RAPOC* is managed by the *Model* class. The only input required by *RAPOC* is the opacity data. Opacity data can be directly downloaded from dedicated repositories such as ExoMol or, instead, by a custom made Python dictionary. In this guide both approaches will be explored:

#### 2.1.1 Using ExoMol (easier)

This is the most straightforward approach. Note that *RAPOC* only accepts data with the TauRex format. Once the opacities have been downloaded from [ExoMol](#), *RAPOC* will load them using the *ExoMolFileLoader*. For practice, we use the TauRex formatted opacities for [water](#). However, if other molecules are required then they can be downloaded [here](#).

#### 2.1.2 Python dictionary (harder)

To load data from a Python dictionary, the code uses the *DictLoader* class. The user may also refer to its documentation. To summarise, the python dictionary must have the layout described in the [Input data layout](#).

Table 1: Input data layout

Information	data format	supported keys names
molecular name	string	mol
pressure grid in [ <i>Pa</i> ]	list or np.array	p, P, pressure, pressure_grid
temperature grid [ <i>K</i> ]	list or np.array	t, T, temperature, temperature_grid
wavenumber grid [1/ <i>cm</i> ]	list or np.array	wn, wavenumber, wavenumbers, wavenumbers_grid, wavenumber_grid
opacities [ <i>m</i> <sup>2</sup> / <i>kg</i> ]	list or np.array	opacities

Every value in the dictionary can be assigned units by using the *units* `astropy.units.Quantity`.

## 2.2 The model

In this section we will assume that the data has been downloaded from [ExoMol](#) and that the `input_file` variable is a string pointing to that file.

Once the data is ready, it may loaded into the `Model` class. After this, two classes can be produced, one for Rosseland and the other for Planck mean opacities: `Rosseland`, `Planck`. These models can be initialised as such:

```
from rapoc import Rosseland, Planck

ross = Rosseland(input_data=input_file)
plan = Planck(input_data=input_file)
```

### 2.2.1 Calculating the mean opacities

For purely illustrative purposes, a temperature (T) of 1000 K with a pressure (P) of 1000 Pa in the wavelength range of 1-10 micron will be used. Therefore, the first step is to define the temperature, pressure and wavelength range:

```
import astropy.units as u

P = 1000.0 * u.Pa
T = 1000.0 * u.K
wl = (1 * u.um, 10 * u.um)
```

Due to the units provided by the `astropy.units.Quantity` module, *RAPOC* can handle the conversions automatically. To perform the opacity calculations the following function is used: `estimate()`

```
r_estimate = ross.estimate(P_input = P, T_input=T, band=wl, mode='closest')

0.00418 m2 / kg
```

```
p_estimate = plan.estimate(P_input = P, T_input=T, band=wl, mode='closest')

17.9466 m2 / kg
```

The investigated wavelength range may also be expressed as a wavenumbers range or frequencies range. This is achieved by attaching the corresponding units.

There are two estimation modes available in *RAPOC*:

- *closest*: the code estimates the mean opacity for the closest pressure and temperature values found within the input data grid.
- *linear* or *loglinear*: the code estimates the mean opacity by performing an interpolation that makes use of the `scipy.interpolate.griddata()`.

In the second case, *RAPOC* needs to first build a `map()` of the mean opacities in the input pressure and temperature data grid. This may be slow. To make this process faster, *RAPOC* will continue to reuse this map until the user asks for estimates in another wavelength range. Only then would *RAPOC* produce a new map.

*RAPOC* can also perform estimates of multiple temperatures and pressures at once (e.g. lists and arrays):

```
P = [1, 10] * u.bar
T = [500.0, 1000.0, 2000.0] * u.K
wl = (1 * u.um, 10 * u.um)
```

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```
r_estimate = ross.estimate(P_input = P, T_input=T, band=wl, mode='closest')

[[0.00094728 0.03303942 0.17086334]
 [0.00484263 0.08600765 0.21673218]] m2 / kg
```

```
p_estimate = plan.estimate(P_input = P, T_input=T, band=wl, mode='closest')

[[27.18478963 17.36108261 8.57782242]
 [27.17086583 17.78923892 8.8249388 ]] m2 / kg
```

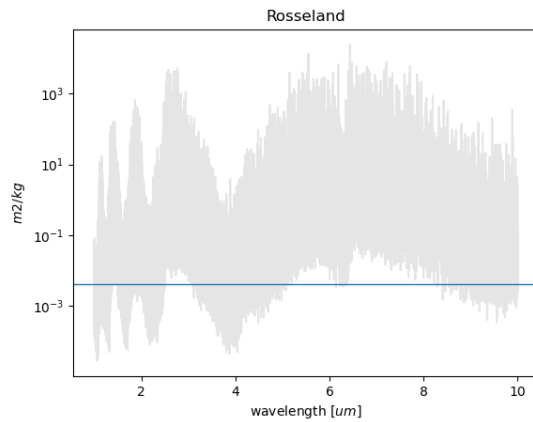
## 2.2.2 Build some plots

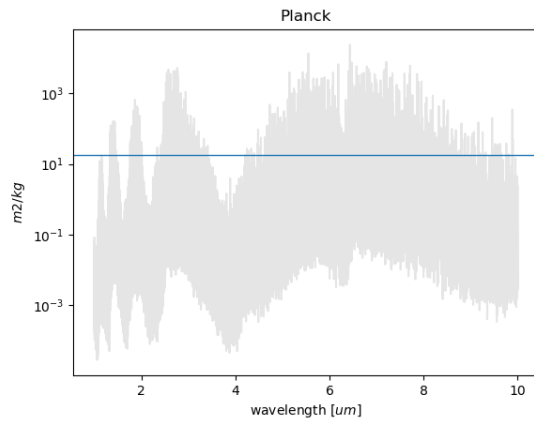
RAPOC can build plots of the calculated Rosseland and Planck mean opacities by using `estimate_plot()`.

If only a single value has been produced, then to produce plots one can use the following script:

```
P = 1000.0 *u.Pa
T = 1000.0 *u.K
wl = (1 * u.um, 10 * u.um)

fig0, ax0 = ross.estimate_plot(P_input = P, T_input=T, band=wl, mode='closest
↪')
fig1, ax1 = plan.estimate_plot(P_input = P, T_input=T, band=wl, mode='closest
↪')
```

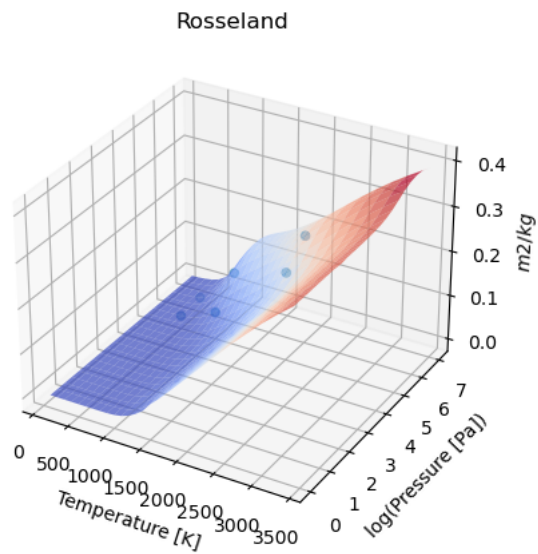


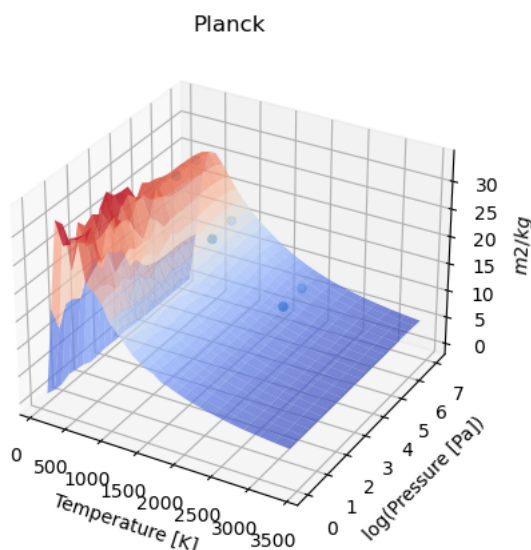


If instead multiple opacities have been calculated, then then the script should be:

```
P = [1, 10 ] *u.bar
T = [500.0, 1000.0, 2000.0] *u.K
wl = (1 * u.um, 10 * u.um)

fig0, ax0 = ross.estimate_plot(P_input = P, T_input=T, band=wl, mode='closest
↪')
fig1, ax1 = plan.estimate_plot(P_input = P, T_input=T, band=wl, mode='closest
↪')
```





Notwithstanding, to force the production of a `map_plot()` with a single value, the following should be used `force_map=True`

## 2.3 Include Rayleigh scattering

### 2.3.1 Initialising Rayleigh data

*RAPOC* can estimate the Rayleigh scattering absorption for a list of atoms, using the tables and equations in Chapter 10 of David R. Lide, ed., CRC Handbook of Chemistry and Physics, Internet Version 2005, [hbcponline](#), CRC Press, Boca Raton, FL, 2005.

To compute the Rayleigh scattering mean opacities, the first step is to produce the Rayleigh data set. This can be done initialising the *Rayleigh* class. This class is similar to a *FileLoader* class: it can be injected into a *Model* as input data, and therefore used as any other input data to estimate the mean opacities.

To initialise the *Rayleigh* it is needed the atom and the wavenumber grid to use to sample the Rayleigh scattering. In the following example we estimate the scattering data for hydrogen in a simple wavenumber grid: 100000, 10000, 1000 cm<sup>-1</sup>

```
from rapoc import Rayleigh
rayleigh = Rayleigh('H', wavenumber_grid=[100000, 10000, 1000])
```

Another way to initialise the *Rayleigh* is to use an already initialised *Model*. This can be useful is the Rayleigh scattering is to be used along with other absorptions. Let's assume that a *Planck* class has been initialised already. That model can be used to produce a Rayleigh scattering dataset sampled at the same wavenumbers grid.

```
plan = Planck(input_data=input_file)
rayleigh = Rayleigh('H', model=plan)
```

Once the wavenumber grid is produced, either using the *wavenumber\_grid* or the *model* keyword, the class produces the opacities data using the `compute_opacities()`.

## 2.3.2 Estimating Rayleigh mean opacities

The initialised `Rayleigh` can be now used as input data for *Rosseland* and *Planck*

```
ross = Rosseland(input_data=rayleigh)
plan = Planck(input_data=rayleigh)
```

Because the Rayleigh scattering doesn't depend on temperature and pressure, is not sampled in a temperatures and pressures grid as the molecular opacities. Therefore, to estimate its mean opacities the pressure must not be indicated. The temperature, on the contrary, must be indicated because of the Rosseland and Planck equation involving a black body. For the same reason, the estimation mode is forced to *closest*, so should not be indicated by the user. Here is an example

```
T = 1000.0 * u.K
wl = (1 * u.um, 10 * u.um)

r_estimate = ross.estimate(T_input=T, band=wl)

6.725572872420127e-08 m2 / kg
```

```
p_estimate = plan.estimate(T_input=T, band=wl)

6.518714389415313e-09 m2 / kg
```

Because, again, the Rayleigh scattering doesn't depend on temperature and pressure, the `map()` is disabled for this kind of data

## 3.1 Planck Model

**class** rapoc.**Planck** (*input\_data*)

Planck model class. This class implements the Planck Opacity model:

$$k = \frac{\int_{\nu_1}^{\nu_2} k_{\nu} B_{\nu}(T) d\nu}{\int_{\nu_1}^{\nu_2} B_{\nu}(T) d\nu}$$

where  $\nu_1$  and  $\nu_2$  are the investigated frequency band edges,  $k_{\nu}$  is the input data opacity at a given frequency, and  $B_{\nu}(T)$  is the black body radiation energy distribution computed at a certain temperature:

$$B_{\nu}(T) = \frac{2h\nu^3}{c^2} \frac{1}{e^{\frac{h\nu}{k_B T}} - 1}$$

### Variables

- **input\_data** (*str* or *dict*) – data file name or data dictionary
- **model\_name** (*str*) – model name
- **band** (*tuple*) – investigated band. Is initialised only after the use of `map()`
- **selected\_grid** (*astropy.units.Quantity*) – investigated grid. Is initialised only after the use of `map()`
- **mol** (*str*) – molecule name
- **mol\_mass** (*astropy.units.Quantity*) – molecular mass
- **pressure\_grid** (*astropy.units.Quantity*) – data pressure grid in *si* units
- **temperature\_grid** (*astropy.units.Quantity*) – data temperature grid in *si* units
- **wavenumber\_grid** (*astropy.units.Quantity*) – data wavenumber grid
- **opacities** (*astropy.units.Quantity*) – data opacities grid in *si* units
- **frequency\_grid** (*astropy.units.Quantity*) – data frequency grid
- **wavelength\_grid** (*astropy.units.Quantity*) – data wavelength grid

**Parameters** **input\_data** (*str* or *dict*) – data file name or `input_data` dictionary. If the input is a *str*, the correspondent loader is used ([ExoMolFileLoader](#)) if the file format is supported. If is *dict*, then [DictLoader](#) is used.

**Raises** **IOError** – if data format is not supported:

## Examples

First we prepare a data dictionary:

```
>>> import numpy as np
>>> data_dict = {'mol': 'H2O',
>>>               'pressure': np.array([1.00000000e+00, 1.00000000e+01, 1.
↪ 00000000e+02, 1.00000000e+03,
>>>                                     1.00000000e+04, 1.00000000e+05, 1.
↪ 00000000e+06, 1.00000000e+07]),
>>>               'temperature': np.array([500, 1000, 1500, 2000, 3000]),
>>>               'wavenumber': np.array([100000, 1000])}
>>> ktab = np.ones((data_dict['pressure'].size, data_dict['temperature'].size, data_
↪ dict['wavenumber'].size,))
>>> data_dict['opacities'] = ktab
```

Let's build the Planck method using an Exomol file as input

```
>>> from rapoc import Planck
>>> dict_data = 'example.TauREx.h5'
>>> model = Planck(input_data=dict_data)
```

Now the model is ready to be used

**opacity\_model** (*opacities, nu, T\_input*)

This function computes the Planck Opacity model:

$$k = \frac{\int_{\nu_1}^{\nu_2} k_{\nu} B_{\nu}(T) d\nu}{\int_{\nu_1}^{\nu_2} B_{\nu}(T) d\nu}$$

where  $\nu_1$  and  $\nu_2$  are the investigated frequency band edges,  $k_{\nu}$  is the input data opacity at a given frequency, and  $B_{\nu}(T)$  is the black body radiation energy distribution computed at a certain temperature:

$$B_{\nu}(T) = \frac{2h\nu^3}{c^2} \frac{1}{e^{\frac{h\nu}{k_B T}} - 1}$$

### Parameters

- **opacities** (*np.array*) – opacity array. Has the same dimension of the frequency grid *nu*
- **nu** (*np.array*) – frequency grid
- **T\_input** (*float*) – temperature

**Returns** mean opacity computed from the model

**Return type** float

## 3.2 Rosseland Model

**class** rapoc.Rosseland (*input\_data*)

Rosseland model class. This class implements the Rosseland Opacity model:

$$\frac{1}{k} = \frac{\int_{\nu_1}^{\nu_2} k_{\nu}^{-1} u(\nu, T) d\nu}{\int_{\nu_1}^{\nu_2} u(\nu, T) d\nu}$$



where  $\nu_1$  and  $\nu_2$  are the investigated frequency band edges,  $k_\nu$  is the input data opacity at a given frequency, and  $u(\nu, T)$  is the black body temperature derivative:

$$u(\nu, T) = \frac{\partial B(\nu, T)}{\partial T} = 2 \frac{h^2 \nu^4}{c^2 k_b} \frac{1}{T^2} \frac{e^{\frac{h\nu}{k_b T}}}{(e^{\frac{h\nu}{k_b T}} - 1)^2}$$

### Variables

- **input\_data** (*str* or *dict*) – data file name or data dictionary
- **model\_name** (*str*) – model name
- **band** (*tuple*) – investigated band. Is initialised only after the use of `map()`
- **selected\_grid** (*astropy.units.Quantity*) – investigated grid. Is initialised only after the use of `map()`
- **mol** (*str*) – molecule name
- **mol\_mass** (*astropy.units.Quantity*) – molecular mass
- **pressure\_grid** (*astropy.units.Quantity*) – data pressure grid in *si* units
- **temperature\_grid** (*astropy.units.Quantity*) – data temperature grid in *si* units
- **wavenumber\_grid** (*astropy.units.Quantity*) – data wavenumber grid
- **opacities** (*astropy.units.Quantity*) – data opacities grid in *si* units
- **frequency\_grid** (*astropy.units.Quantity*) – data frequency grid
- **wavelength\_grid** (*astropy.units.Quantity*) – data wavelength grid

**Parameters** **input\_data** (*str* or *dict*) – data file name or `input_data` dictionary. If the input is a *str*, the correspondent loader is used (`ExoMolFileLoader`) if the file format is supported. If is *dict*, then `DictLoader` is used.

**Raises** **IOError** – if data format is not supported:

### Examples

Let's build the Rosseland method using an Exomol file as input

```
>>> from rapoc import Rosseland
>>> exomolFile = 'example.TauREx.h5'
>>> model = Rosseland(input_data=exomolFile)
```

Now the model is ready to be used

**opacity\_model** (*opacities, nu, T\_input*)

This function computes the Rosseland Opacity model:

$$\frac{1}{k} = \frac{\int_{\nu_1}^{\nu_2} k_\nu^{-1} u(\nu, T) d\nu}{\int_{\nu_1}^{\nu_2} u(\nu, T) d\nu}$$

where  $\nu_1$  and  $\nu_2$  are the investigated frequency band edges,  $k_\nu$  is the input data opacity at a given frequency, and  $u(\nu, T)$  is the black body temperature derivative:

$$u(\nu, T) = \frac{\partial B(\nu, T)}{\partial T} = 2 \frac{h^2 \nu^4}{c^2 k_b} \frac{1}{T^2} \frac{e^{\frac{h\nu}{k_b T}}}{(e^{\frac{h\nu}{k_b T}} - 1)^2}$$

**Parameters**

- **opacities** (*np.array*) – opacity array. Has the same dimension of the frequency grid *nu*
- **nu** (*np.array*) – frequency grid
- **T\_input** (*float*) – temperature

**Returns** mean opacity computed from the model

**Return type** float

## 3.3 Rayleigh Model

**class** `rapoc.Rayleigh` (*atom*, *wavenumber\_grid=None*, *model=None*)

The Rayleigh class estimates the Rayleigh scattering for the indicated atom using the equations from Chapter 10 of David R. Lide, ed., CRC Handbook of Chemistry and Physics, Internet Version 2005, [hbcponline](#), CRC Press, Boca Raton, FL, 2005. These values won't depend on pressure or temperature and therefore only a wavenumbers grid is needed to produce them. The wavenumbers grid can be given by the user or loaded from an already initialised `Model` class (as `Rosseland` or `Planck`). The latter solution might be of help in using the Rayleigh scattering along with other opacities.

**Parameters**

- **atom** (*str*) – name of the considered atom
- **wavenumber\_grid** (*list or numpy.array or astropy.units.Quantity*) – data wavenumber grid
- **model** (*Model*) – built model to use to load the wavenumbers grid.

### Examples

First we prepare a data wavenumbers grid, then we can produce the Rayleigh data:

```
>>> rayleigh = Rayleigh('H', wavenumber_grid=[100000, 10000, 1000])
```

if we already have a molecular model loaded, as example built from a datafile, we can use it to initialise the Rayleigh class:

```
>>> input_data = Rosseland(input_data=exomol_file)
>>> rayleigh = Rayleigh(atom='Na', model=input_data)
```

Now the Rayleigh data are ready to be used as input data for any RAPOC `Model` class:

```
>>> pl = Planck(input_data=rayleigh)
>>> rs = Rosseland(input_data=rayleigh)
```

**compute\_opacities()**

It computes the opacities for Rayleigh scattering as described in David R. Lide, ed., CRC Handbook of Chemistry and Physics, Internet Version 2005, [hbcponline](#), CRC Press, Boca Raton, FL, 2005. chapter 10 table 1:

$$\alpha(\lambda) = \frac{128\pi^5}{3\lambda^4} \frac{a^2}{m}$$

where  $a$  is depending on the atom polarizability listed in table 2 chapter 10 of CRC handbook, and  $m$  is the atom mass.

**Returns** returns the estimated opacity sampled at the indicated wavenumbers.

**Return type** `astropy.units.Quantity`

**read\_content** ()

Reads the class content and returns the needed values for the opacity models.

**Returns**

- *str* – molecule name
- `astropy.units.Quantity` – molecular mass
- `astropy.units.Quantity` – data pressure grid in si units
- `astropy.units.Quantity` – data temperature grid in si units
- `astropy.units.Quantity` – data wavenumber grid
- `astropy.units.Quantity` – data opacities grid in si units

## 3.4 Base Model

**class** `rapoc.models.model.Model` (*input\_data*)

Base model class.

**Variables**

- **input\_data** (*str* or *dict*) – data file name or data dictionary
- **model\_name** (*str*) – model name
- **band** (*tuple*) – investigated band. Is initialised only after the use of `map` ()
- **selected\_grid** (`astropy.units.Quantity`) – investigated grid. Is initialised only after the use of `map` ()
- **mol** (*str*) – molecule name
- **mol\_mass** (`astropy.units.Quantity`) – molecular mass
- **pressure\_grid** (`astropy.units.Quantity`) – data pressure grid in *si* units
- **temperature\_grid** (`astropy.units.Quantity`) – data temperature grid in *si* units
- **wavenumber\_grid** (`astropy.units.Quantity`) – data wavenumber grid
- **opacities** (`astropy.units.Quantity`) – data opacities grid in *si* units
- **frequency\_grid** (`astropy.units.Quantity`) – data frequency grid
- **wavelength\_grid** (`astropy.units.Quantity`) – data wavelength grid

**Parameters** **input\_data** (*str* or *dict*) – data file name or *input\_data* dictionary. If the input is a *str*, the correspondent loader is used (`ExoMolFileLoader`) if the file format is supported. If is *dict*, then `DictLoader` is used.

**Raises** `IOError` – if data format is not supported:

## Examples

For the sake of simplicity let's assume we use an ExoMol file as input

```
>>> from rapoc.models import Model
>>> exomolFile = 'example.TauREx.h5'
>>> model = Model(input_data=exomolFile)
```

**opacity\_model** (*opacities, nu, T\_input*)

Computes the mean opacity in the investigated range.

### Parameters

- **opacities** (*np.array*) – opacity array. Has the same dimension of the frequency grid *nu*
- **nu** (*np.array*) – frequency grid
- **T\_input** (*float*) – temperature

**Returns** mean opacity computed from the model

**Return type** float

**extract\_opacities** (*T\_input, band, P\_input=None, units='si'*)

Returns the input\_data opacities for given pressure, temperature and band.

### Parameters

- **T\_input** (*float or np.array*) – temperature to use for the estimation. This should be a *astropy.units.Quantity* with specified units, if not *K* are assumed as units. Can either be a single value or an array.
- **band** (*tuple*) – this should be a tuple of *astropy.units.Quantity* indicating the edges of the band to use in the estimation. The units used reveal the intention to work in wavelengths, wavenumbers or frequencies.
- **P\_input** (*float or np.array*) – pressure to use for the estimation. This should be a *astropy.units.Quantity* with specified units, if not *Pa* are assumed as units. Can either be a single value or an array. It can be *None* in the case of data not depending on pressure, as Rayleigh scattering. Default is *None*.
- **units** (*str or astropy.units, optional*) – indicates the output units system. If *si* the opacity is returned as  $m^2/kg$ , if *cgs* is returned as  $cm^2/g$ . Instead of a string, you can also indicate the units using *astropy.units*. Units is *si* by default.

### Returns

- *astropy.units.Quantity* – returns the estimated opacity for the pressures and temperatures indicated. If only one pressure and temperature are indicated, it returns a single *Quantity*. If *n* pressures and *m* temperatures are indicated, it return a *Quantity* array of dimensions (n,m)
- *list* – list of wavenumber (or wavelength or frequency) index relative to the investigated band

**estimate** (*T\_input, band, P\_input=None, mode='closest', units='si'*)

It estimates the model opacity in the indicated pressure and temperature grid for the indicated band, using the *opacity\_model()*.

### Parameters

- **T\_input** (*float or np.array*) – temperature to use for the estimation. This should be a *astropy.units.Quantity* with specified units, if not *K* are assumed as units. Can either be a single value or an array.
- **band** (*tuple*) – this should be a tuple of *astropy.units.Quantity* indicating the edges of the band to use in the estimation. The units used reveal the intention to work in wavelengths, wavenumbers or frequencies.
- **P\_input** (*float or np.array*) – pressure to use for the estimation. This should be a *astropy.units.Quantity* with specified units, if not *Pa* are assumed as units. Can either be a single value or an array. It can be *None* in the case of data not depending on pressure, as Rayleigh scattering. Default is *None*.
- **mode** (*str, optional*) – indicates the estimation mode desired. If *closest* the output will be the opacity computed from the data at the nearest pressure on the data grid to the indicated one, and at the nearest temperature on the data grid to the indicated one. If {'linear', 'loglinear'} it's used the *scipy.interpolate.griddata()* with the indicated *kind*. To interpolate a map of opacities over the data grid is computed first using *map()*. Mode is *closest* by default.
- **units** (*str or astropy.units, optional*) – indicates the output units system. If *si* the opacity is returned as  $m^2/kg$ , if *cgs* is returned as  $cm^2/g$ . Instead of a string, you can also indicate the units using *astropy.units*. Units is *si* by default.

**Returns** returns the estimated opacity for the pressures and temperatures indicated. If only one pressure and temperature are indicated, it returns a single *Quantity*. If *n* pressures and *m* temperatures are indicated, it return a *Quantity* array of dimensions (n,m)

**Return type** *astropy.units.Quantity*

**Raises**

- **NotImplementedError**: – if the indicated mode is not supported
- **astropy.units.UnitConversionError**: – if the input units cannot be converted to *si*
- **AttributeError**: – if the band cannot be interpreted as wavelength, wavenumber or frequency

**estimate\_plot** (*T\_input, band, P\_input=None, mode='closest', force\_map=False, fig=None, ax=None, yscale='log', xscale='linear', grid='wavelength'*)

Produces a plot of the estimates (produced with *estimate()*), comparing the raw data with the result. If a single estimate is to be plotted, this method produces a plot of raw opacities vs wavelength from the opacities (in grey) and the mean opacity estimated. If multiple estimates are to be plotted, it produces a 3D plot, with the surface of mean opacity vs temperature and pressure from the data grid (using *map\_plot()*) and the interpolated data superimposed.

**Parameters**

- **T\_input** (*float or np.array*) – temperature to use for the estimation. This should be a *astropy.units.Quantity* with specified units, if not *K* are assumed as units. Can either be a single value or an array.
- **band** (*tuple*) – this should be a tuple of *astropy.units.Quantity* indicating the edges of the band to use in the estimation. The units used reveal the intention to work in wavelengths, wavenumbers or frequencies.
- **P\_input** (*float or np.array*) – pressure to use for the estimation. This should be a *astropy.units.Quantity* with specified units, if not *Pa* are assumed as units. Can either be a single value or an array. It can be *None* in the case of data not depending on pressure, as Rayleigh scattering. Default is *None*.

- **mode** (*str*, *optional*) – indicates the estimation mode desired. If *closest* the output will be the opacity computed from the data at the nearest pressure on the data grid to the indicated one, and at the nearest temperature on the data grid to the indicated one. If {'linear', 'loglinear'} it's used the `scipy.interpolate.griddata()` with the indicated *kind*. Mode is *closest* by default.
- **force\_map** (*bool*) – If True a 3D map plot is generate even for a single estimate. Default is *False*.
- **fig** (`matplotlib.pyplot.figure`, *optional*) – indicates the figure to use. If *None* a new figure is produced. Default is *None*
- **ax** (`matplotlib.pyplot.axis`, *optional*) – indicates the axis of the figure to use. If *None* a new axis is produced. Default is *None*
- **yscale** (*str*) – y-axis scale to use. Default is *log*
- **xscale** (*str*) – x-axis scale to use. Default is *linear*
- **grid** (*str*) – x-axis grid format. {*wavelength*, *frequency*, *wavenumber*} are supported. Default is *wavelength*.

**Returns**

- `matplotlib.pyplot.figure`, *optional* – figure containing the plot
- `matplotlib.pyplot.axis`, *optional* – axis containing the plot

**Raises** **KeyError**: – if the indicated grid is not supported

**map** (*band*)

Returns the mean opacity map for the indicated model.

**Parameters** **band** (*tuple*) – this should be a tuple of *astropy.units.Quantity* indicating the edges of the band to use in the estimation. The units used reveal the intention to work in wavelengths, wavenumbers or frequencies.

**Returns** mean opacity map in *si* units

**Return type** `np.array`

**Notes**

If the map has been already built for the indicated band, then the method returns the previous map. This speeds up the use of the code in external functions.

**map\_plot** (*band*, *fig=None*, *ax=None*)

Produces a 3D-plot of the model mean opacity map built with `map()`.

**Parameters**

- **band** (*tuple*) – this should be a tuple of *astropy.units.Quantity* indicating the edges of the band to use in the estimation. The units used reveal the intention to work in wavelengths, wavenumbers or frequencies.
- **fig** (`matplotlib.pyplot.figure`, *optional*) – indicates the figure to use. If *None* a new figure is produced. Default is *None*
- **ax** (`matplotlib.pyplot.axis`, *optional*) – indicates the axis of the figure to use. If *None* a new axis is produced. Default is *None*

**Returns**

- `matplotlib.pyplot.figure`, *optional* – figure containing the plot

- `matplotlib.pyplot.axis, optinal` – axis containing the plot

## 3.5 ExoMol file loader

**class** `rapoc.loaders.ExoMolFileLoader` (*filename*)

File loader for ExoMol file. It works for opacities file in the *TauREx.h5* format. These file can be downloaded from [ExoMol website](#).

### Notes

This class is implicitly called by the model when initialised if the matched input is used.

### Examples

Let's build the Rosseland method using an Exomol file as input

```
>>> from rapoc import Rosseland
>>> exomolFile = 'example.TauREx.h5'
>>> model = Rosseland(input_data=exomolFile)
```

Now the model is ready to be used

**Parameters** `filename` (*str*) – data file name

## 3.6 Base file loader

**class** `rapoc.loaders.DictLoader` (*input\_data*)

Dict loader class. The dictionary should contain the following information under certain keys:

information	default units	supported keys
molecular name		mol
pressure grid	<i>Pa</i>	p, P, pressure, pressure_grid
temperature grid	<i>K</i>	t, T, temperature, temperature_grid
wavenumber grid	<i>1/cm</i>	wn, wavenumber, wavenumbers, wavenumbers_grid, wavenumber_grid
opacities	<i>m<sup>2</sup>/kg</i>	opacities

If the input data has no units attached, the defaults units are assume. If they have units, a simple conversion is performed by the code to match the default values.

## Notes

This class is implicitly called by the model when initialised if the matched input is used.

## Examples

First we prepare a data dictionary. Please be aware that this are not representative of any molecule.

```
>>> import numpy as np
>>> data_dict = {'mol': 'None',
>>>              'mol_mass': 42,
>>>              'pressure': np.array([1.00000000e+00, 1.00000000e+01, 1.
↪ 00000000e+02, 1.00000000e+03,
>>>                                  1.00000000e+04, 1.00000000e+05, 1.
↪ 00000000e+06, 1.00000000e+07]),
>>>              'temperature': np.array([500, 1000, 1500, 2000, 3000]),
>>>              'wavenumber': np.array([100000, 1000])}
>>> opac = np.ones((data_dict['pressure'].size, data_dict['temperature'].size, data_
↪ dict['wavenumber'].size,))
>>> data_dict['opacities'] = opac
```

Let's build the Planck method using the dictionary as input

```
>>> from rapoc import Planck
>>> model = Planck(input_data=data_dict)
```

Now the model is ready to be used

**Parameters** `input_data` (*dict*) – input\_data dictionary

**read\_content** ()

Reads the dict content and returns the needed valued for the opacity models.

### Returns

- *str* – molecule name
- *astropy.units.Quantity* – molecular mass
- *astropy.units.Quantity* – data pressure grid in si units
- *astropy.units.Quantity* – data temperature grid in si units
- *astropy.units.Quantity* – data wavenumber grid
- *astropy.units.Quantity* – data opacities grid in si units

## 3.7 Base file loader

**class** `rapoc.loaders.loader.FileLoader` (*filename*)

Base file loaders class.

**Parameters** `filename` (*str*) – data file name

**read\_content** ()

Reads the file content and returns the needed valued for the opacity models.

### Returns

- *str* – molecule name



- *astropy.units.Quantity* – molecular mass
- *astropy.units.Quantity* – data pressure grid in si units
- *astropy.units.Quantity* – data temperature grid in si units
- *astropy.units.Quantity* – data wavenumber grid
- *astropy.units.Quantity* – data opacities grid in si units



CITE

If you use this code or its results, please cite *RAPOC: the Rosseland and Planck opacity converter* by Mugnai L. V. and Modirrousta-Galian D. (submitted).



## MEET THE AUTHORS

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