

Documentation for `rac2dm`

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1 Guideline

The `rac2dm` package is designed to model a static disk structure and its thermochemistry. It is derived from the `rac2d` package. The letter “m” in the name stands for “modularized”. For the ease of implementation, the interface to each module is implemented with files.

Using script languages such as `Python`, different modules can be chained and iterated to achieve a specific goal.

2 Description of modules

2.1 Radiative transfer and bulk radiation field solution:

rtrf

2.1.1 Overall configuration

```
1 &conf_nml_rt
2   conf_rt%struct_file = "struct.dat"
3   conf_rt%opti_files = "sil.dat" "gra.dat"
4   conf_rt%rad_source_files = "s1.dat" "s2.dat"
5   conf_rt%star_mass_Msun = 0.6D0
6   conf_rt%nphoton = 1000000
7   conf_rt%nmax_encounter = 1999999999
8   conf_rt%escaped_photons_file = 'escaped_ph.dat'
9   conf_rt%collect_photon = .true.
10  conf_rt%collect_lam_min_AA = 1D0
11  conf_rt%collect_lam_max_AA = 1D8
12  conf_rt%collect_nmu = 3
13  conf_rt%collect_ang_mins_deg = 0D0 4D0 80D0
14  conf_rt%collect_ang_maxs_deg = 3D0 10D0 90D0
15  conf_rt%nlen_lut = 2048
16  conf_rt%Tdust_min_K = 1D0
17  conf_rt%Tdust_max_K = 2D3
18  conf_rt%dist_to_Earth_pc = 51.0D0
19 /
```

2.1.2 Specification of matter distribution

```
1 format ! Default to 1
2 number_of_species number_of_data_points
3 r1 z1 n11 n21 n31 ...
4 r2 z2 n12 n22 n32 ...
5 ...
```

Here n_{ij} means the density of the i th dust species at the j th location.

2.1.3 Specification of optical properties

```
1 number_of_data_points
2 angstrom1 Q_abs1 Q_sca1 g1
3 angstrom2 Q_abs2 Q_sca2 g2
4 ...
```

2.1.4 Specification of radiation sources

Multiple radiation sources are allowed. Each source can have a different type. At present the following types are supported:

- **point**: point source
- **sphere**: photons are emitted from the surface of a sphere
- **isotropic**: photons are emitted from an isotropic background
- **directional**: photons are emitted from a single direction

The following are example radiation source files:

```
1 point
2 x_au y_au z_au
3 number_of_data_points_of_spectrum total_luminosity_cgs
4 angstrom1 F1_cgs
5 angstrom2 F2_cgs
6 ...
```

```
1 sphere
2 x_cen_au y_cen_au z_cen_au r_sphere_Rsun
3 number_of_data_points_of_spectrum total_luminosity_cgs
4 angstrom1 F1_cgs
5 angstrom2 F2_cgs
6 ...
```

```
1 isotropic
2 ! Empty line
3 number_of_data_points_of_spectrum energy_density_cgs
4 angstrom1 F1_cgs
5 angstrom2 F2_cgs
6 ...
```

```
1 directional
2 v_x v_y v_z
3 number_of_data_points_of_spectrum flux_density_cgs
4 angstrom1 F1_cgs
5 angstrom2 F2_cgs
6 ...
```

2.1.5 Output

- Temperature distribution
- Internal radiation field
- Escaped radiation field

2.2 Disk structure adjustment: structadj

2.2.1 Overall configuration

```
1 &conf_nml_struct_adj
2   conf_struct_adj%density_file = "dens.dat"
3   conf_struct_adj%density_file_new = "dens_new.dat"
4   conf_struct_adj%temperature_file = "temperature.dat"
5 /
```

2.2.2 Output

- New density distribution

2.3 Thermochemical calculation: thermochem

Here we assume the thermochemical calculation is only done for a single cell. To calculate for the whole disk just call the module in a loop or use the `map` function in Python.

2.3.1 Overall configuration

```
1 &conf_nml_thermochem
2   conf_thermochem%network_file = "rate12.dat"
3   conf_thermochem%init_file = "init.dat"
4   conf_thermochem%enthalpy_file = "enthalpy.dat"
5   conf_thermochem%Tgas = 20D0
6   conf_thermochem%Tdust = 20D0
7   conf_thermochem%sigdust_cm2 = 20D0
8   conf_thermochem%G0 = 1D0
9   conf_thermochem%Xray = 0D0
10  conf_thermochem%zeta_cosmicray_H2 = 1.36D-17
11  conf_thermochem%PAH_abundance = 1.6D-9
12  conf_thermochem%Av = 10D0
13  conf_thermochem%filename_CII = ""
14  conf_thermochem%filename_OI = ""
15  conf_thermochem%filename_NII = ""
16  conf_thermochem%filename_SiII = ""
17  conf_thermochem%filename_FeII = ""
18  conf_thermochem%dt_first_step = 1D-8
19  conf_thermochem%t_max = 1D6
20  conf_thermochem%ratio_tstep = 1.1D0
21  conf_thermochem%max_runtime_allowed = 60D0
22  conf_thermochem%RTOL = 1D-6
23  conf_thermochem%ATOL = 1D-30
```

2.3.2 Output

- For a single location, output the gas temperature and chemical abundances as a function of time.
- Optional output

2.4 Make images: makeimage

3 Examples