

- SpectralModel: a high-resolution framework for petitRADTRANS 3
- Doriann Blain  $lacktrian^{1\P}$ , Paul Mollière  $lacktrian^{1}$ , and Evert Nasedkin  $lacktrian^{1}$
- 4 1 Max Planck Institut für Astronomie, DE ¶ Corresponding author

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# Summary

Atmospheric characterisation from spectroscopic data is a key to understand planetary formation. Two types of observations can be performed for this kind of analysis. Space-based observations (e.g., using the James Webb Space Telescope, JWST), are not impeded by the Earth's atmosphere, but are currently limited to low resolving powers (<3000), which can lead to ambiguities in some species detections. Ground-based observations (e.g., using the Very Large Telescope, VLT), on the other hand, can benefit from large resolving powers ( $\approx10^5$ ), allowing for unambiguous species detection, but are impacted by telluric spectral lines. petitradtrans (pRT) is a radiative transfer package used for computing emission or transmission spectra of planetary atmospheres (Mollière et al., 2019). The package has a non-negligible user base, the original article being cited in 264 refereed works at the time of writing. pRT is already relatively easy to use on space-based, low-resolution observations. However, while the package technically has the capacity to analyse high-resolution spectra, thanks to its ability to incorporate high-resolution ( $\mathcal{R}=10^6$ ) line lists, ground-based observations analysis is a complex and challenging task. The new SpectralModel object provides a powerful and flexible framework that streamlines the setup necessary to model and retrieve high-resolution spectra.

# Statement of need

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Calculating a spectrum using pRT's core object Radtrans is a two-step process in which the user first instantiates the object, giving parameters that control the loading of opacities. The second step is for the user to call one of the Radtrans function, giving "spectral" parameters such as the temperatures or the mass fractions of the atmosphere, that will be used in combination with the loaded opacities to generate the spectrum.

However, these two steps are by themselves often insufficient to build a spectrum in a real-life scenario. The spectral parameters may individually rely on arbitrarily complex models requiring their own parameters, and may depend on each other. For example, getting mass fractions from equilibrium chemistry requires knowing the temperature profile, and the mean molar mass requires knowing the mass fractions (see e.g. the built-in pRT functions). Common operations such as convolving the spectrum, scaling it to stellar flux, or more specifically for high-resolution spectra, Doppler-shiftting the spectrum and including the transit effect, must be done by post-processing the Radtrans-generated spectrum. Finally, using a retrieval requires to code a "retrieval model" including all the steps described above. This induces, especially for first-time users, a significant setup cost. The alternative is to use one of pRT's built-in models, but this lacks flexibility.

The SpectralModel object extends the base capabilities of the petitRADTRANS package by providing a standardized but flexible framework for spectral calculations. It has been especially designed to effectively erase the setup cost of modelling the spectral Doppler-shift, the transit effect, and of implementing the preparation step necessary for ground-based high-



- resolution observations analysis. SpectralModel is also interfaced with pRT's retrieval
- module (Nasedkin et al., 2024), and as such is an easy-to-use tool to perform both high- and
- 44 low-resolution atmospheric retrievals.
- 45 The combination of ease-of-use and flexibility offered by SpectralModel makes it a powerful tool
- 46 for high-resolution (but also low-resolution) atmospheric characterisation. With the upcoming
- 47 first light of a new generation of ground based telescopes, such as the Extremely Large
- Telescope, SpectralModel makes petitRADTRANS ready for the new scientific discoveries
- that will be unveiled in the next era of high-resolution observations.

# 55 The SpectralModel object

## 51 Main features

## 52 Spectral parameter calculation framework

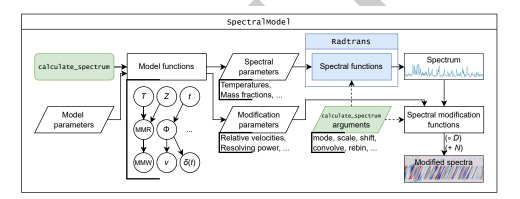


Figure 1: Flowchart of SpectralModel.calculate\_spectrum function. The annotation below the model functions represents an example of execution order of these function after topological sorting, involving the temperature (T), the metallicity (Z), the time (t), the mass fractions (MMR), the mean molar masses (MMW), the orbital phases  $(\phi)$ , the relative velocities (v), and the transit effect  $(\delta)$ .

SpectralModel provides a framework to automatise the calculation of the spectral parameters. Each spectral parameter is linked to a function, called here "model function", which calculates its value. This feature can be extended to the parameters required for these functions, and so on. Before calculating spectra, the function's execution order is automatically determined through a topological sorting algorithm<sup>1</sup> (Kahn, 1962). SpectralModel comes with built-in functions (Blain et al., 2024) for all the spectral parameters, so that the object can be used "out-of-the box". Parameters that ultimately do not depend on any function are called "model parameters", and must be given during instantiation.

- In addition, SpectralModel provides built-in functions (Blain et al., 2024) to scale, convolve,
- Doppler-shift, rebin, include planet transit effect, and prepare a spectrum after it has been
- 63 calculated. Similarly to model functions, these "spectral modification functions" must be given,
- if used, their own model parameters during instantiation.
- The spectral calculation is done within the calculate\_spectrum function (see Figure 1). The spectral mode (emission or transmission), as well as which of the spectral modification to
- activate (i.e. only scaling, or both convolving and rebinning, etc.), are controlled through the
- function's arguments ("spectral modification parameters").

<sup>&</sup>lt;sup>1</sup>Cyclic dependencies are not supported.



## 69 Automatic optimal wavelength range calculation

High-resolution spectra require high-resolution opacities, which, when loaded, can take a lot of Random-Access Memory (RAM). For example, loading pRT's 1H2-160\_P0KAZATEL line-by-line line list ( $\mathcal{R}=10^6$ ) between 1 and 2  $\mu$ m takes 804 MB² of RAM. Moreover, it is not unusual for a model to incorporate multiple species opacities. Fast retrievals in pRT are also performed in parallel, using multiple processes on distributed memory. The total RAM usage of the opacities is thus the amount of bytes taken by one species on the required wavelength range, times the number of species, time the number of processes. Using too many processes can therefore overload hardware RAM. To avoid out-of-memory errors, the number of processes must be limited, decreasing retrieval speed.

A way to slightly reduce this memory usage is to load exactly the wavelength range required for an analysis, instead of relying on manual inputs. This task is complicated in high-resolution retrievals due to parameters influencing the Doppler-shift (that is, the radial velocity semi-amplitude  $K_p$ , the rest frame velocity shift Vrest, and the mid transit time offset  $T_0$ ) being retrieved. SpectralModel comes with a class method with\_velocity\_range, which takes into account the (uniform) prior range of these parameters to automatically calculate the optimal wavelength range to load.

## 86 Interface with pRT's retrieval module

The Retrieval object has been extended to support spectra with up to 3 dimensions, intended to be spectral order, exposure (time), and spectral pixel (wavelength). It now also has a class method that instantiates a Retrieval object from Data objects<sup>3</sup>. The Data object has also been extended in two ways: it now allows taking directly data as arrays, instead of requiring a file, and allows taking directly Radtrans (or by extension SpectralModel) objects, instead of generating a new one during a Retrieval instantiation.

In addition, SpectralModel's model parameters and spectral modification functions can be advantageously used to simplify the retrieval setup compared to Radtrans'. This removes the need for several steps:

- building the RetrievalConfig object, as this has been automated,
- declaring the fixed parameters, as all model parameters that are not retrieved parameters are de facto fixed parameters,
- writing the retrieval model function, as it is given by the SpectralModel itself.

Ground-based high-resolution spectra contain telluric and stellar lines that must be removed. This is usually done with a "preparing" pipeline (also called "detrending" or "pre-processing" pipeline). To this end, a new retrieval.preparing sub-module has been implemented, containing the "Polyfit" pipeline (Blain et al., 2024) and the "SysRem" pipeline (Tamuz et al., 2005). To perform a retrieval when the data are prepared with "Polyfit", the forward model must be prepared in the same way (Blain et al., 2024). This forward model preparation step can be activated when calculating a spectrum with SpectralModel.

# Ground-based data simulation

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Data (F) taken from ground telescopes can be expressed as  $F=M_\Theta\circ D+N$  (Blain et al., 2024), where  $M_\Theta$  is an exact model with true parameters  $\Theta$ , D ("deformation matrix") represents the combination of telluric lines, stellar lines, and instrumental deformations (pseudocontinuum, blaze function, ...), and N is the noise. The operator " $\circ$ " represents the element-wise product. Telluric lines, noise, and other deformations can be included in a SpectralModel object. A time-varying airmass can be added as model parameter to better model the telluric lines. Finally, a command-line interface (CLI) with ESO's SKYCALC sky model calculator has been implemented, adapting the CLI provided on the ESO's website.

<sup>&</sup>lt;sup>2</sup>According to numpy.ndarray.nbytes.

<sup>&</sup>lt;sup>3</sup>pRT's retrieval module allows for the retrieval of a combination of datasets.



#### 116 Workflows

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Examples for these workflows are available in the pRT's documentation.

#### Spectra calculation

119 Calculating spectra with SpectralModel is done in two steps:

- 1. Instantiation: similarly to Radtrans, this step is done to load the opacities, and thus requires the same parameter as a Radtrans instantiation. In addition, the user can provide model parameters, that will give the spectral parameters and the modification parameters. Finally, a custom dict can be given if the user desires to use different functions than the built-in ones.
- Calculation: spectral calculation is done with a unique function. The spectrum type (emission or transmission), as well as modification flags (for scaling, Doppler-shifting, etc.) are given as arguments.

#### 128 Retrievals

Retrieving spectra with SpectralModel is done in seven steps:

- 1. Loading the data,
- 2. For high-resolution ground-based data: preparing the data,
- 3. Setting the retrieved parameters, this is done by filling a dict,
- 4. Setting the forward model, by instantiating a SpectralModel object,
- 5. Instantiating a Data object with the SpectralModel dedicated function,
- 6. Instantiating a Retrieval object from the previously built Data object(s),
- 7. Running the retrieval.
- In addition, a new corner plot function, based on the corner package (Foreman-Mackey, 2016), has been implemented to ease the representation of the retrieval results with this framework.

# The petitRADTRANS 3 update

Test	pRT 2.7.7 time (s)	pRT 3.1.0 time (s)	pRT 2.7.7 RAM (MB)	pRT 3.1.0 RAM (MB)
Opacity loading, 'c-k'	3.2	0.9	_	_
Opacity loading, 'lbl'	6.3	0.4	_	_
Emission, 'c-k'	6.4	5.2	2428	1472
Emission, 'lbl'	7.8	4.4	3929	2643
Transmission, 'c-k'	1.2	0.6	992	757
Transmission, 'lbl'	6.6	3.1	3929	2230

- Times are measured using the cProfile standard library, from the average of 7 runs.
- "RAM": peak RAM usage as reported by the tracemalloc standard library.
- 'c-k': using correlated-k opacities (CH<sub>4</sub> and H<sub>2</sub>O), from 0.3 to 28  $\mu$ m.
- 'lbl': using line-by-line opacities (CO and  $H_2O$ ), from 0.9 to 1.2  $\mu$ m.
- All spectra calculations are done using 100 pressure levels. Emission scattering is activated in 'c-k' mode.
- Results obtained on Debian 12.5 (WSL2), CPU: AMD Ryzen 9 3950X @ 3.50 GHz.

Along with SpectralModel, major changes have been made to pRT. The changes focus on optimisations (both for speed and RAM usage) for high-resolution spectra computing, but this also impacts the correlated-k (low-resolution) part of the code (see Table 1). To speed-up "input data" (opacities, pre-calculated equilibrium chemistry table, star spectra table) loading



times, pRT's loading system has been overhauled and the loaded files have been converted from a mix of ASCII, Fortran unformatted and HDF5 files to HDF5-only. Opacities now also follow an extended ExoMol database naming and structure convention. The package's installation process has been made compatible with Python  $\geq 3.12^4$ . The package's code has also been rationalised, clarified, and refactored. Finally, several quality-of-life features (e.g., missing requested opacities can be automatically downloaded from the project's Keeper library, or the Planet object) have been implemented.

# 151 Acknowledgements

We thank the pRT users, who greatly helped improving the package by sharing their suggestions and reporting their issues.

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<sup>&</sup>lt;sup>4</sup>pRT 2 used the numpy.distutils module to compile its Fortran extensions. This module is deprecated and is removed for Python 3.12. pRT 3 uses the Meson build system instead, with almost unnoticeable changes for users.