Manual for

NIRCam Coronagraphy Simulations

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Limitations

- In its current version, the pipeline can only be used to simulate programs consisting of one or multiple **roll 1**, **roll 2**, **reference observation sequences**. Also, all observations will be simulated at the same observation date.
- Target Acquisition and Astrometric Confirmation images are currently not simulated.
- **Standard subpixel dithering** is not supported and **small grid dithering** is only supported for reference observations, but not for science observations.
- The pipeline has been tested with the **GTO programs 1194** (HR 8799), **1411** (beta Pic), and **1412** (51 Eri). The corresponding configuration files are provided as hr8799.yaml, betapic.yaml, and 51eri.yaml and the corresponding APT files are provided in the HR8799, betaPic, and 51Eri folders. For testing the pipeline with these example programs, copy and rename the desired configuration file to config.yaml.

History

- Version 0.3.0
 - The JWST stage 3 pipeline processing was improved by fixing multiple wrong header keywords.
 - The issue with the wrongly oriented small grid dithers was fixed by inverting the RA sign of the dither offset.
 - The issue with the axis flip between pyNRC and MIRAGE data was fixed for both channels.
 - The issue with the axis flip of the coronagraphic masks was fixed for both channels.
- Version 0.2.0
 - The issue with the wrongly oriented scenes was fixed by injecting the companions with an inverted RA sign in run_pynrc.py.
 - The issue with the horizontal axis flip between pyNRC and MIRAGE data was fixed for the LW channel by flipping the CRDS reference files read by pyNRC.
 - The issue with the horizontal axis flip of the coronagraphic masks was fixed for the LW channel by flipping the masks in pyNRC.
 - The issue with the large amount of bad pixels after the JWST stage 1 pipeline processing was fixed by skipping the dark subtraction step, i.e., setting result1.dark_current.skip = True in run_jwst_s1s2.py.
 - The issue with the slightly misaligned JWST stage 3 psfalign pipeline product was fixed by explicitly specifying the bad pixel types in the PSF alignment step, i.e., setting result3.align_refs.bad_bits = 'HOT, UNRELIABLE_BIAS' in run_jwst_s3.py.
- Version 0.1.0
 - o First commit.

Installation instructions

The pipeline is separated into multiple Python scripts which need to be run one after another. The reason for this approach is that a different Python environment should be used for each of the software packages that are used by the pipeline. In the following, installation instructions for each of the different Python environments are given.

species environment

- 1) Create a new species Python environment.
- 2) Use git clone https://github.com/tomasstolker/species.git followed by git reset --hard 165554cde2ad65abc8501c11c188258264d7abfb to clone version 0.4.0 of species into a directory of your choice. Specify this directory under species_dir in the configuration file.
- 3) Install the required dependencies into the species Python environment.
- 4) Note that an installation of species itself, e.g., by running setup.py, is not recommended. Instead, just clone the repository and try running the run_species.py script. Python will crash and complain about the missing dependencies which you can then install one after another before trying it again.

pyNRC environment

- 1) Create a new pyNRC Python environment.
- 2) Install the dependencies and data files required by pyNRC following a-g below. Further information can be found at https://github.com/kammerje/pynrc/blob/kammerje-patch-1/docs/install_clean.rst. Note that steps d and e are not mentioned in this documentation, but are required to run my modified version of pyNRC.
 - a. Install Pysynphot, download the Pysynphot data files, untar them, and set the environment variable PYSYN_CDBS in your .bashrc file to point to them (export PYSYN_CDBS='\$HOME/data/cdbs/').
 - b. Install jwxml and WebbPSF, download the WebbPSF data files, untar them, and set the environment variable WEBBPSF_PATH in your .bashrc file to point to them (export WEBBPSF_PATH='\$HOME/data/webbpsf-data/').
 - c. Install JWST Backgrounds and its dependencies.
 - d. Install WebbPSF_ext using git clone https://github.com/JarronL/webbpsf_ext.git followed by git reset – -hard 19df5301f0062a3fd56a38b05ad325cdb791fafe to clone the correct version of WebbPSF_ext into a directory of your choice. Specify this directory under webbpsf_ext_dir in the configuration file. Run setup.py to complete the integration of WebbPSF_ext into WebbPSF.
 - e. Set the environment variable WEBBPSF_EXT_PATH in your .bashrc file to point to the WebbPSF data files (export WEBBPSF_EXT_PATH='\$HOME/data/webbpsf-data/').
 - f. Install pyNRC using git clone https://github.com/kammerje/pynrc.git
 --branch kammerje-patch-1
 --single-branch to clone my modified version of pyNRC into a directory of your choice. Specify this directory under pynrc_dir in the configuration file.
 - g. Download the pyNRC data files, untar them, and set the environment variable PYNRC_PATH in your bashrc file to point to them (export PYNRC_PATH='\$HOME/data/pynrc_data/').

- 3) Use git clone https://github.com/semaphoreP/whereistheplanet.git to clone the latest version of whereistheplanet into a directory of your choice. Specify this directory under whereistheplanet_dir in the configuration file.
- 4) Install the required dependencies into the pyNRC Python environment.

MIRAGE environment

- 1) Create a new MIRAGE Python environment.
- 2) Install **version 2.1.0** of MIRAGE from Pypi following https://mirage-data-simulator.readthedocs.io/en/latest/install.html#install-from-pypi.
- 3) If you encounter issues while running the MIRAGE scripts, make sure to have version 1.15.0 of healpy, version 2.1.0 of mirage, version 1.32 of grismconf, version 1.58 of nircam-gsim, and version 1.3.3 of jwst installed.
- 4) The required reference files will be downloaded using the ref_mirage.py Python script later.

JWST environment

- 1) Create a new JWST Python environment.
- 2) Install **version 1.3.3** of the JWST data reduction pipeline from Pypi following https://jwst-pipeline.readthedocs.io/en/latest/index.html.

Preparations

Before the pipeline can be run, several preparations need to be conducted. These involve modifying the APT file, modifying the configuration file, and downloading the MIRAGE reference files (the latter needs to be done only once before the pipeline is run for the first time).

APT file

- 1) Open the program with the NIRCam coronagraphy observations for which data shall be simulated in APT.
- 2) From the Observations folder of that program, remove all non-NIRCam non-coronagraphic observations.
- 3) Re-run the Visit Planner.
- 4) Under Reports → Visit X:X → Total Roll Analysis For Visit, find the preferred observation date and the corresponding roll angle constraints. These need to be specified under date, pa_roll1, pa_roll2, and pa_ref in the configuration file.
- 5) Save the xml and pointing files of the modified program using File \rightarrow Export... \rightarrow xml file & pointing file \rightarrow OK.

Configuration file

- 1) The pipeline always reads the parameters saved in the config.yaml file!
- 2) In the paths section, once the directories of species, whereistheplanet, WebbPSF_ext, and pyNRC have been set correctly, only the wdir needs to be changed if simulations for a new program shall be made.
- 3) In the apt section, the paths of the xml and pointing files (relative to the wdir) need to be specified.
- 4) In the observation section, the observation date, roll angles, wavefront drifts, and oversampling need to be specified. Furthermore, the observing sequences need to be specified as X,Y,Z, where X/Y/Z are the observation indexes (starting with 1) of the roll 1/roll 2/reference observations in the APT file (these are not necessarily the observation numbers used in the APT file!).
- 5) In the sources section, the science and reference source names and properties need to be specified. The names must match the source names ("Name in the Proposal") used in the APT file.
- 6) In the companions section, an arbitrary number of companions c1-cX can be added. For each companion, besides a mass and a specific entropy at formation, a name_witp (name in whereistheplanet) needs to be specified. A list of available companions can be found at https://github.com/semaphoreP/whereistheplanet/blob/master/whereistheplanet/ whereistheplanet.py. If a companion is not available in whereistheplanet, an arbitrary identifier (that does NOT match any of the available companions) needs to be used for name_witp and ra_off and de_off at the observation date need to be specified. Furthermore, a name_spec (name in species) needs to be specified. A list of available companions found can https://github.com/tomasstolker/species/blob/master/species/data/companions.py . If a companion is not available in species, name_spec can be left blank and the companion magnitudes need to be computed in a custom way and saved as arrays of shape (1,) under name_filter.npy (where name = name_witp and filter = FXXXW/FXXXM/FXXXN) into the pmdir specified in the configuration file.

7) In the pipeline section, the model used by species to fit the observed companion photometry and its effective temperature range can be specified. A list of available models can be found at https://github.com/tomasstolker/species/blob/1dada33c9547f5bb2721687caf3c7d5 d3d171856/species/data/database.py#L332. Furthermore, the make_plots parameter can be used to control whether plots shall be generated and saved into the pynrc_figs_dir.

MIRAGE reference files

- 1) Run the Python script ref_mirage.py in the MIRAGE Python environment, which will download the MIRAGE reference files into the mirage_refs_dir specified in the configuration file.
- 2) By default, this will only download a single linearized dark for each detector (total size of reference files ~95 GB). While it is recommended to download all linearized darks for better performance, this is irrelevant here since the ramp images in the MIRAGE data will be replaced with those from pyNRC eventually.
- 3) If you already have the MIRAGE reference files on your machine, you can simply skip this step. Just specify the mirage_refs_dir in the configuration file so that MIRAGE will be able to find them.

Running the pipeline

The pipeline is separated into multiple Python scripts which need to be run one after another. In summary, the pipeline computes the companion magnitudes in the relevant JWST bands using species, computes the companion locations at the specified observation date using whereistheplanet, simulates coronagraphic observations for the specified APT file using pyNRC, simulates the corresponding clear pupil observations using MIRAGE, replaces the ramp images in the MIRAGE data with those from the pyNRC data, and finally runs the simulated data through the JWST data reduction pipeline.

Running species

- 1) Run the Python script run_species.py in the species Python environment, which will compute the companion magnitudes in the relevant JWST filters and save them into the pmdir specified in the configuration file.
- 2) By default, the Exo-REM models with an effective temperature range of 1000-2000 K are used to fit the companion magnitudes from the literature. This can be changed using the model_spec and teff_range parameters in the configuration file. A list of available models can be found at https://github.com/tomasstolker/species/blob/1dada33c9547f5bb2721687caf3c7d5 d3d171856/species/data/database.py#L332.
- 3) If desired (or if a companion is not available in species), the companion magnitudes can also be computed in a custom way and running run_species.py can be skipped. In that case, the name_spec parameter in the configuration file can be left blank and the companion magnitudes need to be saved as arrays of shape (1,) under name_filter.npy (where name = name_witp and filter = FXXXW/FXXXM/FXXXN) into the pmdir specified in the configuration file.

Running pyNRC

- 1) Run the Python script run_pynrc.py in the pyNRC Python environment, which will simulate coronagraphic observations (ramps and noiseless slopes) for the specified APT file and save them into the pynrc_data_dir specified in the configuration file.
- 2) The make_plots parameter in the configuration file can be used to control whether plots shall be generated and saved into the pynrc_figs_dir specified in the configuration file.
- 3) The companion locations at the specified observation date are computed using whereistheplanet, but if a companion is not available in whereistheplanet, it is also possible to manually specify its ra_off and de_off in the configuration file. In that case, a name_witp still needs to be provided since this name is used as an identifier for the companion within the Python script.

Running MIRAGE

- 1) Run the Python script run_mirage.py in the MIRAGE Python environment, which will simulate clear pupil observations for the specified APT file and save them into the mirage_data_dir specified in the configuration file.
- 2) A random source at RA = DEC = 0 with a brightness of 20 mag in all JWST filters is used in this step. This should be irrelevant since the ramp images in the MIRAGE data will be replaced with those from pyNRC in the next step.

Replacing MIRAGE with pyNRC data

1) Run the Python script run_pynrc_into_mirage.py in the MIRAGE Python environment, which will replace the ramp images in the MIRAGE data with those from the pyNRC data and modify the relevant header keywords.

Running JWST data reduction pipeline

- 1) Run the Python script run_jwst_s1s2.py in the JWST Python environment, which will run the simulated ramp images through the JWST Stage 1 and 2 data reduction pipelines and save the output files into the jwst_s1s2_data_dir specified in the configuration file.
- 2) Run the Python script run_jwst_s3.py in the JWST Python environment, which will create the required ASN files, run the Stage 2 reduced data through the JWST Stage 3 data reduction pipeline and save the output files into the jwst_s3_data_dir_specified in the configuration file.
- 3) Note that the PSF alignment step can require a lot of memory allocation and processing time if the number of science integrations multiplied by the number of reference integrations is of the order of 1000 or larger.