
engine Documentation

Release

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Tools to help the community with planning exoplanet observations.

PandExo is both an online tool and a python package for generating instrument simulations of JWST's NIRSpec, NIRCam, NIRISS and NIRCам and HST WFC3. It uses throughput calculations from STScI's Exposure Time Calculator, Pandeia: Pandeia + Exoplanets = PandExo. This documentation contains information on how to download, install and analyze PandExo output.

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GETTING STARTED

Tools to help the community with planning exoplanet observations.

PandExo is both an online tool and a python package for generating instrument simulations of JWST's NIRSpec, NIRCam, NIRISS and NIRCams and HST WFC3. It uses throughput calculations from STScI's Exposure Time Calculator, Pandeia: Pandeia + Exoplanets = PandExo. This documentation contains information on how to download, install and analyze PandExo output.

1.1 Should I install PandExo or use the online interface?

Install if...

- I will be using PandExo for more than 10 runs
- I want to submit bash runs
- I want to use the online GUI but don't want to be subject to any slow downs because of high user frequency
- I want to be able to use plotting functions to analyze my output

Install EVEN if...

- I am scared of Python

Do not install if...

- I will only be using PandExo fewer than 10 times
- I have a student who can install it for me

1.2 Requires

- Python >2.7, that's it

INSTALLATION

Warning: Before reading further, if you do not wish to install PandExo, there is an online version of the code here at [PSU Science](#) or here at NASA GSFC.

2.1 Installation Procedure

1. Easiest way to Install:

```
pip install pandexo.engine
```

OR Download PandExo's repository:

```
git clone --recursive https://github.com/natashabatalha/pandexo
cd pandexo
python setup.py install
```

2. Download the Phoenix Stellar Atlas [FROM THIS LINK](#) in order to easily pull Stellar SED's from the phoenix database. Then type the following commands. It might be helpful to add the export command to your ~/.bashrc file.

```
mkdir pysynphot_data
mv synphot5.tar.gz pysynphot_data
tar -xvf synphot5.tar.gz
export PYSYN_CDBS=USRDIR/pysynphot_data
```

Once you do this your untarred file should automatically have this structure. If you use wget you might get something different. So double check this.

```
ls pysynphot_data/grid/phoenix/
.DS_Store      catalog.fits  phoenix05/   phoenix15/   phoenix25/   phoenix35/  ↵
↪phoenixp03/
AA_README      phoenixm00/  phoenixm10/  phoenixm20/  phoenixm30/  phoenixm40/  ↵
↪phoenixp05/
```

3. Download the JWST Reference Data [FROM HERE](#) . This is a big file (6 gigs) so think carefully about where you want to store it. Don't accidentally download it on your Mac Air then wonder why you can't save a 32 Kb doc file.

Then make sure you untar and point to the file so PandExo knows where it is. Like above, it might be helpful to put this in your ~/.bashrc file.

```
tar xf pandeia_data-1.0.tar.gz
export pandeia_refdata=USRDIR/pandeia_data
```

Your Pandeia data reference file should look like this:

```
ls pandeia_data/
.DS_Store      README.md      background/    extinction/    jwst/          sed/           ↵
↪ strategy/
.git/          VERSION_PSF    devtools/     hst/          normalization/ source/         ↵
↪ wfirst/
```

2.2 Troubleshooting

2.2.1 Problems with PYFFTW?

Many users experience issues when downloading Pandeia because of its dependency on *pyfftw*. If you experience this problem try these steps:

- If you do not have a non-LLVM based GCC installation on your system, you can obtain one from here but gcc 5.1 does not produce a usable FFTW installation so make sure you download **gcc 4.9 or below**
- STScI created the following script to successfully install *pyfftw*

```
mv $(which gcc) $(which gcc).orig
curl -O https://bitbucket.org/api/2.0/snippets/jhunkeler/R7gy5/
↪3265aea27175817087ab4a39c21157d926f8afc3/files/build_fftw.sh
chmod +x build_fftw.sh
./build_fftw
```

2.2.2 Can't find Pandeia Reference Data

This usually looks like NoneType errors.

- Make sure PandExo knows where the Pandeia reference data is:

```
export pandeia_refdata=USRDIR/pandeia_data
```

JWST TUTORIAL

This tutorial can be downloaded as a iPython notebook on the PandExo Github.

```
import pandexo.engine.justdoit as jdi # THIS IS THE HOLY GRAIL OF PANDEXO
```

3.1 Editting Input Dictionaries

3.1.1 Step 1) Load in a blank exoplanet dictionary

To start, load in a blank exoplanet dictionary with empty keys. You will fill these out for yourself in the next step.

```
exo_dict = jdi.load_exo_dict()
```

Edit exoplanet observation inputs

Editting each keys are annoying. But, do this carefully or it could result in nonsense runs

```
exo_dict['observation']['sat_level'] = 80      #saturation level in percent of full well
exo_dict['observation']['noccultations'] = 2   #number of transits
exo_dict['observation']['R'] = None           #fixed binning. I usually suggest ZERO
↳binning.. you can always bin later

exo_dict['observation']['fraction'] = 1.0      #without having to redo the calcualtion
↳= in/out                                     #fraction of time in transit versus out

exo_dict['observation']['noise_floor'] = 0     #this can be a fixed level or it can be
↳a filepath                                  #to a wavelength dependent noise floor
↳solution (units are ppm)
```

Edit exoplanet star inputs

Note... If you select 'phoenix' you **do not** have to provide a starpath, w_unit or f_unit, but you **do** have to provide a temp, metal andlogg. If you select 'user' you **do not** need to provide a temp, metal andlogg, but you **do** need to provide units and starpath.

```
exo_dict['star']['type'] = 'phoenix'           #phoenix or user (if you have your own)
exo_dict['star']['mag'] = 8.0                  #magnitude of the system
exo_dict['star']['ref_wave'] = 1.25            #For J mag = 1.25, H = 1.6, K =2.22.. etc
↳(all in micron)
```

```
exo_dict['star']['temp'] = 5500           #in K
exo_dict['star']['metal'] = 0.0          # as log Fe/H
exo_dict['star']['logg'] = 4.0           #log surface gravity cgs
```

Edit exoplanet planet inputs

```
exo_dict['planet']['exopath'] = '/Users/nbatalh1/Desktop/Simulations/wasp12b.txt'
exo_dict['planet']['w_unit'] = 'cm'      #other options include "um",
↳ "Angs", "secs" (for phase curves)
exo_dict['planet']['f_unit'] = 'rp^2/r^2' #other options are 'fp/f*'
exo_dict['planet']['transit_duration'] = 2.0*60.0*60.0 #transit duration in seconds
```

3.1.2 Step 2) Load in instrument dictionary (optional)

Step 2 is optional because PandExo has templates for the most common modes. Within those templates the subarrays selected are the ones with the lowest frame times, and the read modes selected are the ones with one frame per 1 group (standard). If this suits you see Option 1 below. There is additional fine tuning that can be done within each of these observing modes. As a first pass, I'd suggest skipping this for now. Then come back and fine tune after your first run.

- NIRCam F444W
- NIRSpec Prism
- NIRSpec G395M
- NIRSpec G395H
- NIRSpec G235H
- NIRSpec G235M
- NIRCam F322W2
- NIRSpec G140M
- NIRSpec G140H
- MIRI LRS
- NIRISS SOSS

```
jdi.print_instruments()
```

Choose **from the** following:

```
['NIRCam F444W', 'NIRSpec Prism', 'NIRSpec G395M', 'NIRCam F322W2', 'NIRSpec G395H',
↳ 'NIRSpec G235H', 'NIRSpec G235M', 'NIRSpec G140M', 'NIRSpec G140H', 'MIRI LRS',
↳ 'NIRISS SOSS', 'WFC3 G141']
```

```
inst_dict = jdi.load_mode_dict('NIRSpec Prism')
```

Change subarray:

```
inst_dict["configuration"]["detector"]["subarray"] = 'sub512'
```

Change the read mode

```
inst_dict["configuration"]["detector"]["readmode"] = 'nrs'
```

The last thing to note is that PandExo (by Default) optimizes the amount of groups that can fit into an integration. If you want to set your own number of groups you can change that too.

..code:: python

```
inst_dict["configuration"]["detector"]["ngroup"] = 5
```

3.2 Running PandExo Command Line

You have **four options** for running PandExo. All of them are accessed through attribute **jdi.run_pandexo**. See examples below.

```
jdi.run_pandexo(exo, inst, param_space = 0, param_range = 0, save_file = True,
output_path=os.getcwd(), output_file = '')
```

3.2.1 Option 1- Run single instrument mode, single planet

If you forget which instruments are available run **jdi.print_instruments()** and pick one

```
jdi.print_instruments()
```

Choose **from the** following:

```
['NIRCam F444W', 'NIRSpec Prism', 'NIRSpec G395M', 'NIRCam F322W2', 'NIRSpec G395H',
↪ 'NIRSpec G235H', 'NIRSpec G235M', 'NIRSpec G140M', 'NIRSpec G140H', 'MIRI LRS',
↪ 'NIRISS SOSS_Or1', 'NIRISS SOSS_Or2', 'WFC3 G141']
```

```
result = jdi.run_pandexo(exo_dict, ['NIRCam F322W2'])
```

```
Running Single Case for: NIRCam F322W2
Computing Duty Cycle
Finished Duty Cycle Calc
Starting Out of Transit Simulation
End out of Transit
Starting In Transit Simulation
End In Transit
```

3.2.2 Option 2- Run single instrument mode (with user dict), single planet

This is the same thing as option 1 but instead of feeding it a list of keys, you can feed it a instrument dictionary (this is for users who wanted to simulate something NOT pre defined within pandexo)

```
inst_dict = jdi.load_mode_dict('NIRSpec G140H')
#personalize subarray
inst_dict["configuration"]["detector"]["subarray"] = 'sub2048'
result = jdi.run_pandexo(exo_dict, inst_dict)
```

```
Running Single Case w/ User Instrument Dict
Computing Duty Cycle
Finished Duty Cycle Calc
Starting Out of Transit Simulation
```

```
End out of Transit
Starting In Transit Simulation
End In Transit
```

3.2.3 Option 3- Run several modes, single planet

Use several modes from `print_instruments()` options.

```
#choose select
result = jdi.run_pandexo(exo_dict, ['NIRCam F444W', 'NIRCam F322W2', 'MIRI LRS'],
                          output_path = '/Users/nbatalh1/Desktop/JWSTFUN')

#run all
result = jdi.run_pandexo(exo_dict, ['RUN ALL'], save_file = False)
```

3.2.4 Option 4- Run single mode, several planet cases

Use a single modes from `print_instruments()` options. But explore parameter space with respect to **any** parameter in the exo dict. The example below shows how to loop over several planet models

You can loop through anything in the exoplanet dictionary. It will be planet, star or observation followed by whatever you want to loop through in that set.

i.e. planet+exopath, star+temp, star+metal, star+logg, observation+sat_level.. etc

```
#looping over different exoplanet models
jdi.run_pandexo(exo_dict, ['NIRCam F444W'], param_space = 'planet+exopath',
                param_range = os.listdir('/Users/nbatalha1/all_my_models_here'),
                output_path = '/Users/nbatalh1/Desktop/JWSTFUN')

#looping over different stellar temperatures
jdi.run_pandexo(exo_dict, ['NIRCam F444W'], param_space = 'star+temp',
                param_range = np.linspace(5000,8000,2),
                output_path = '/Users/nbatalh1/Desktop/JWSTFUN')

#looping over different saturation levels
jdi.run_pandexo(exo_dict, ['NIRCam F444W'], param_space = 'observation+sat_level',
                param_range = np.linspace(.5,1,5),
                output_path = '/Users/nbatalh1/Desktop/JWSTFUN')
```

3.3 Running PandExo GUI

The same interface that is available online is also available for use on your machine. Using the GUI is very simple and good alternative if editing the input dictionaries is confusing.

```
import pandexo.engine.run_online as ro
ro.main()
```

Then open up your favorite internet browser and go to: <http://localhost:1111>

Note: Some WebApp functions may not be available. For example, the precomputed noise simulations and transmission and emission modeling are only available online.

3.4 Analyzing Output

There are pre computed functions for analyzing most common outputs. You can also explore the dictionary structure yourself.

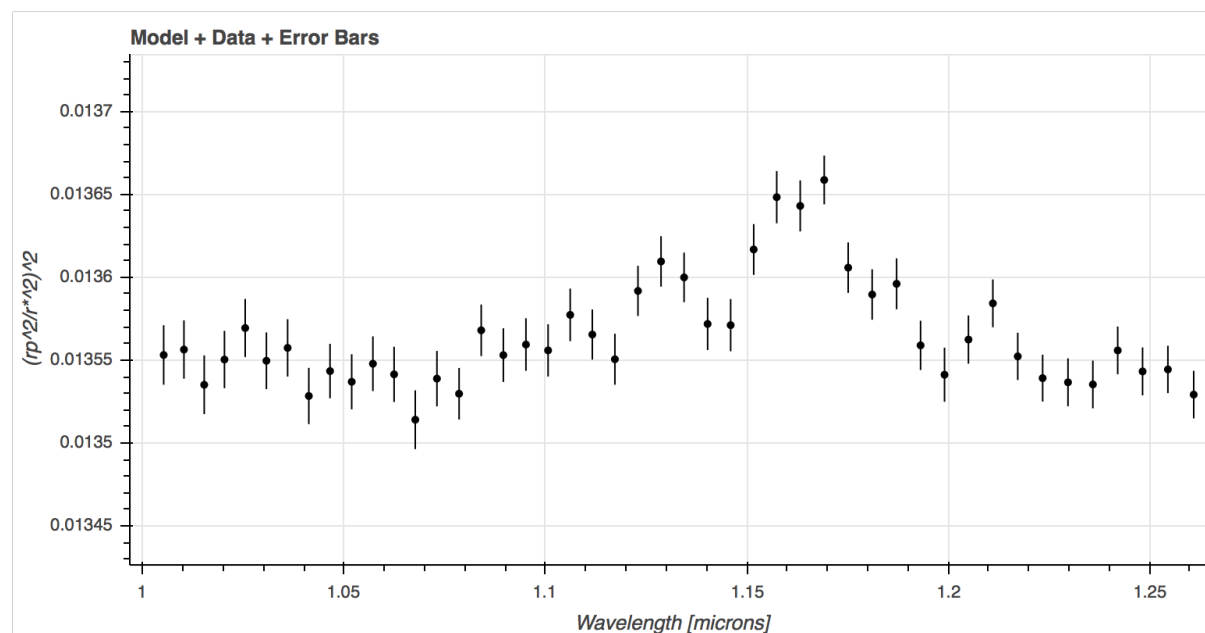
```
import pandexo.engine.justplotit as jpi
import pickle as pk
```

3.4.1 Plot 1D Data with Errorbars

Multiple plotting options exist within *jwst_1d_spec*

1. Plot a single run

```
#load in output from run
out = pk.load(open('singlerun.p', 'r'))
#for a single run
x,y, e = jpi.jwst_1d_spec(out, R=100, num_tran=10, model=False, x_range=[.8,1.28])
```

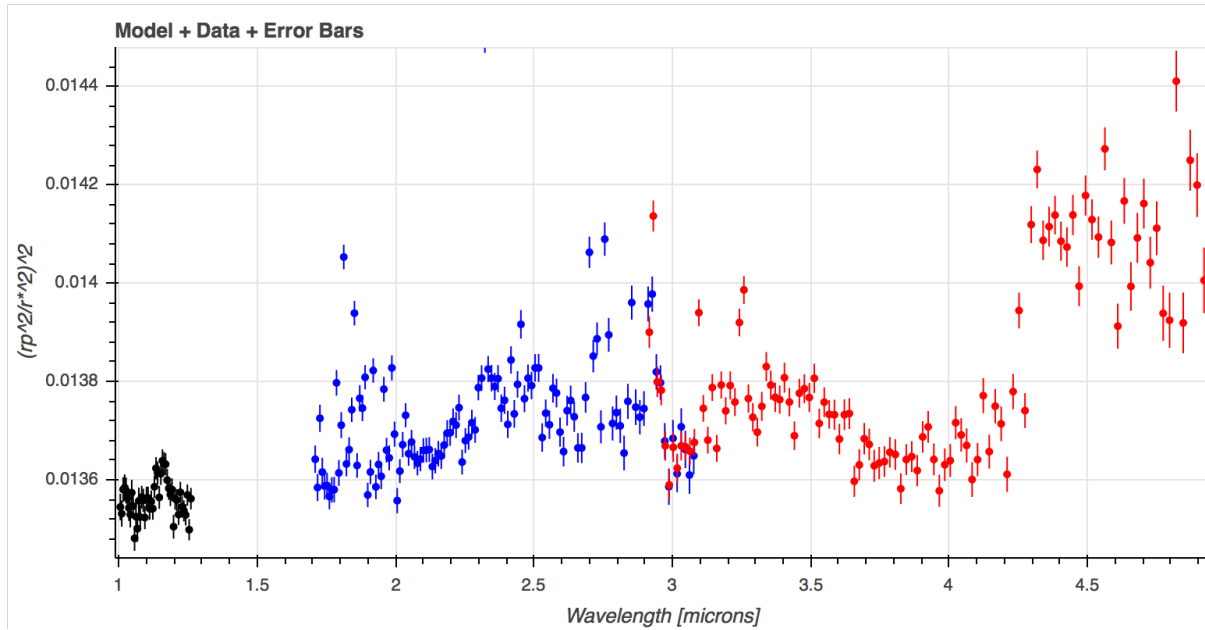


2. Plot several runs from parameters space run

```
#load in output from multiple runs
multi = pk.load(open('three_nirspec_modes.p', 'r'))

#get into list format
list_multi = [multi[0]['NIRSpec G140M'], multi[1]['NIRSpec G235M'], multi[2]['NIRSpec_
↪G395M']]

x,y,e = jpi.jwst_1d_spec(list_multi, R=100, model=False, x_range=[1,5])
```

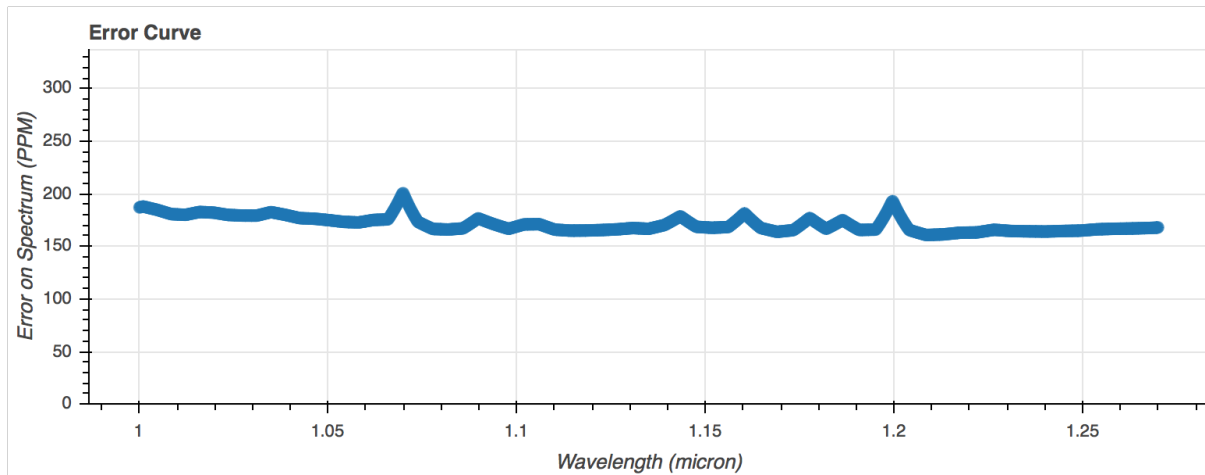


3.4.2 Plot Noise & More

Several functions exist to plot various outputs.

See also `jwst_1d_bkg`, `jwst_1d_snr`, `jwst_1d_flux`,

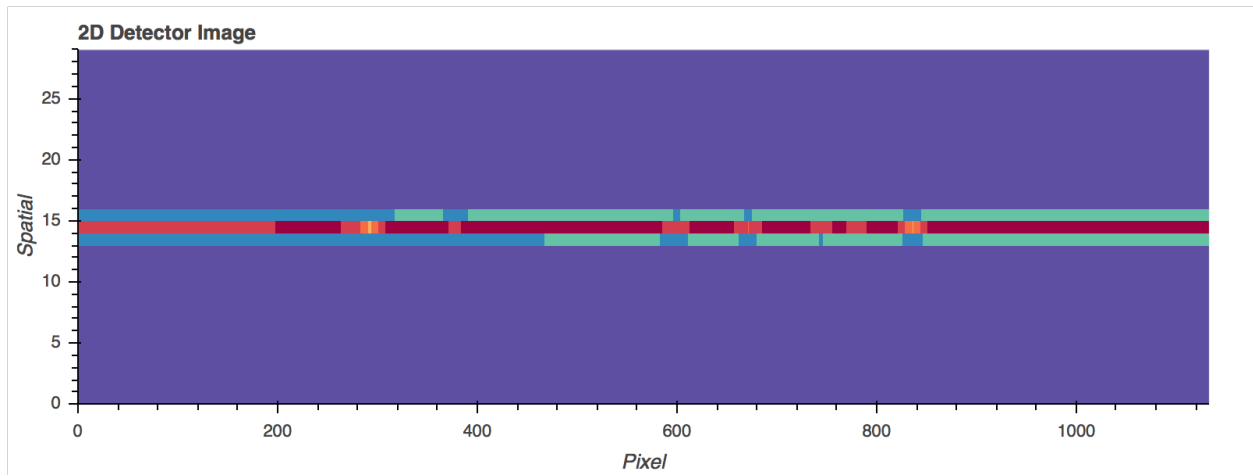
```
x, y = jpi.jwst_noise(out)
```



3.4.3 Plot 2D Detector Profile

See also `jwst_2d_sat` to plot saturation profile

```
data = jpi.jwst_2d_det(out)
```

POSSIBLE INSTRUMENT INPUT PARAMS

Below are all the possible instrument input parameters, which match the JWST APT ([downloadable here](#)).

4.1 NIRSpec

4.1.1 Grisms/Filter combinations

4.1.2 Subarrays

4.2 NIRISS

4.3 NIRCам

4.4 MIRI

PY DICT STRUCTURE OF JWST OUTPUT

The PandExo output is organized into [Python Dictionaries](#). See [example notebooks](#) for an explanation of how to manipulate these and plot the most common outputs. Below is a breakdown of everything contained in the PandExo output dictionary.

5.1 FinalSpectrum (contains 4 keys)

- **spectrum_w_rand**: Planet spectrum with random noise added in units of $(Rp/Rs)^2$ or (Fp/Fs)
- **spectrum**: Planet spectrum with no random noise
- **error_w_floor**: Error with user defined noise floor. If no floor was specified, there is no floor.
- **wave**: wavelength (microns)

```
print dict['FinalSpectrum']['spectrum_w_rand']
```

5.2 OriginalInput (contains 2 keys)

- **model_wave**: original wavelength input by user
- **model_spec**: original spectrum input by user

```
print dict['OriginalInput']['model_wave']
```

5.3 Warning (contains 6 keys)

- **Num Groups Reset?**: Before PandExo simulates in and out of transit observations, it computes a single integration with 2 groups in order to figure out how many additional groups it can add before hitting saturation. If it computes a number less than 2, it resets the number of groups to 2.
- **Group Number Too Low?**: Prints out warning if number of groups is less than 5 and the saturation level less than 60%.
- **Group Number Too High?**: Prints out warning if number of groups per integration exceeds 65536
- **Saturated?**: This is an output directly taken from Pandeia's "hard saturation" flag. If there are any saturated pixels, it will alert you here. You can also see the saturation profile.
- **Non linear?**: This is an output directly taken from Pandeia's "soft saturation" flag.

- **% full well high?**: If you've set the saturation level over 80%, it will warn you.

```
print dict['Warning']['Num Groups Reset?']
```

5.4 PandeiaOutTrans (contains 9 keys)

This is the raw output of Pandeia's simulation of the out of transit observation. For a complete breakdown of these outputs go to [STScI's Pandeia Documentation](#).

- **sub_reports**
- **information**
- **warnings**
- **transform**
- **2d**
- **scalar**
- **1d**
- **input**
- **3d**

```
print dict['PandeiaOutTrans']['information']
```

5.5 RawData (contains 6 keys)

- **var_in**: The variance of only the in transit data
- **wave**: wavelength vector
- **flux_in**: Flux of the in transit data in units of e-/s
- **flux_out**: Flux of the out of transit data in units of e-/s
- **error_no_floor**: Error without any noise floor
- **var_out**: The variance of only the out of transit data

```
print dict['RawData']['var_in']
```

5.6 Timing (contains 10 keys)

- **Seconds per Frame**
- **Number of Transits**
- **Observing Efficiency (%)** = $(\text{num groups} - 1) / (\text{num groups} + 1)$
- **Num Integrations Out of Transit**
- **On Source Time**
- **Exposure Time Per Integration (secs)**

- **Reset time Plus TA time (hrs):** Target acquisition time is assumed to be 30 minutes
- **Num Integrations In Transit**
- **Num Groups per Integration**
- **Num Integrations per Occultation**

```
print dict['Timing']['Seconds per Frame']
```

5.7 Input (contains 10 keys)

- **Target Mag**
- **Readmode**
- **Disperser**
- **Filter**
- **Instrument**
- **Mode**
- **Saturation Level (electons)**
- **Aperture**
- **Subarray**
- **Primary/Secondary**

```
print dict['Input']['Target Mag']
```

5.8 Timing, Warning, and Input Divs (all contain html script)

Here are the html scripts for the three tables that are rendered on the website output page.

HST TUTORIAL

This file demonstrates how to use TExoNS to predict the: 1. Transmission/emission spectrum S/N ratio 2. Observation start window for any system observed with WFC3/IR.

```
import pandexo.engine.justdoit as jdi
```

6.1 Editing Input Dictionaries

6.1.1 Step 1) Load in a blank exoplanet dictionary

```
exo_dict = jdi.load_exo_dict()
```

Edit stellar and planet inputs

```
#WASP-43
exo_dict['star']['mag']      = 9.397                # H magnitude of the system
#WASP-43b
exo_dict['planet']['type']   = 'user'                # user specified inputs
exo_dict['planet']['exopath'] = 'WASP43b-Eclipse_Spectrum.txt' # filename for model_
↪ spectrum
exo_dict['planet']['w_unit']  = 'um'                 # wavelength unit
exo_dict['planet']['f_unit']  = 'fp/f*'              # flux ratio unit (can also put
↪ "rp^2/r*^2")
exo_dict['planet']['depth']   = 4.0e-3               # flux ratio
exo_dict['planet']['i']       = 82.6                 # Orbital inclination in degrees
exo_dict['planet']['ars']     = 5.13                 # Semi-major axis / stellar_
↪ radius
exo_dict['planet']['period']  = 0.8135               # Orbital period in days
exo_dict['planet']['transit_duration'] = 4170.0/60/60/24 # (optional if given above_
↪ info) transit duration in days
exo_dict['planet']['w']       = 90                   # (optional) longitude of_
↪ periastron. Default is 90
exo_dict['planet']['ecc']     = 0                     # (optional) eccentricity._
↪ Default is 0
```

6.1.2 Step 2) Load in instrument dictionary

- WFC3 G141

- WFC3 G102

```
inst_dict = jdi.load_mode_dict('WFC3 G141')
```

Edit HST/WFC3 detector and observation inputs

```
exo_dict['observation']['noccultations'] = 5 # Number of_
↳transits/eclipses
inst_dict['configuration']['detector']['subarray'] = 'GRISM256' # GRISM256 or_
↳GRISM512
inst_dict['configuration']['detector']['nsamp'] = 10 # WFC3 N_SAMP,_
↳1..15
inst_dict['configuration']['detector']['samp_seq'] = 'SPARS5' # WFC3 SAMP_SEQ,
↳ SPARS5 or SPARS10
inst_dict['strategy']['norbits'] = 4 # Number of HST_
↳orbits
inst_dict['strategy']['nchan'] = 15 # Number of_
↳spectrophotometric channels
inst_dict['strategy']['scanDirection'] = 'Forward' # Spatial scan_
↳direction, Forward or Round Trip
inst_dict['strategy']['schedulability'] = 30 # 30 for small/
↳medium program, 100 for large program
inst_dict['strategy']['windowSize'] = 20 # (optional)_
↳Observation start window size in minutes. Default is 20 minutes.
```

6.2 Run PandExo Command Line

```
jdi.run_pandexo(exo, inst, param_space = 0, param_range = 0, save_file = True,
output_path=os.getcwd(), output_file = '')
```

See wiki Attributes for more thorough explanation fo inputs

```
foo = jdi.run_pandexo(exo_dict, inst_dict, output_file='wasp43b.p')
Running Single Case w/ User Instrument Dict
***WARNING: Observing plan may incur mid-orbit buffer dumps. Check with APT.
```

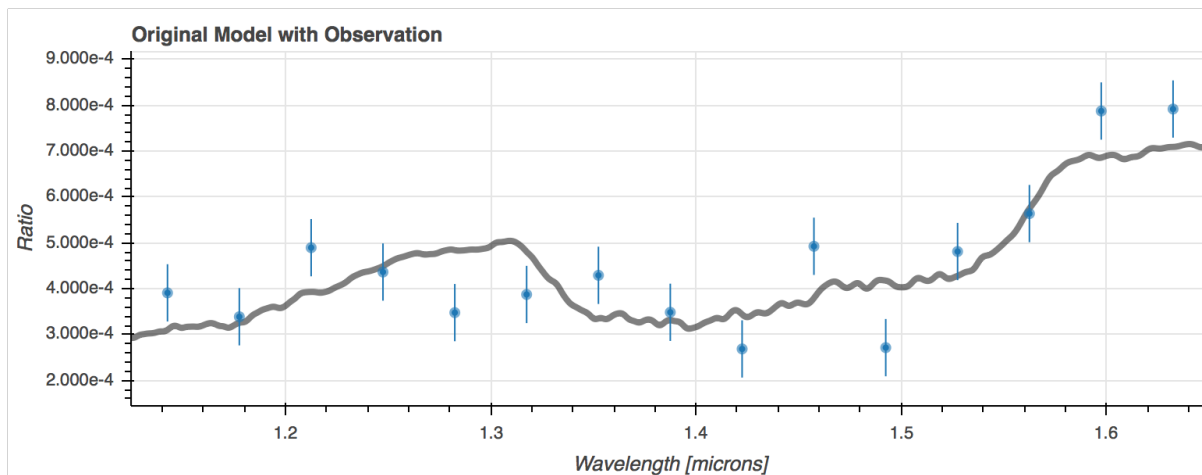
```
inst_dict['configuration']['detector']['nsamp'] = None
inst_dict['configuration']['detector']['samp_seq'] = None
bar = jdi.run_pandexo(exo_dict, inst_dict, output_file='wasp43b.p')
Running Single Case w/ User Instrument Dict
```

```
exo_dict['observation']['scanDirection'] = 'Round Trip'
hst = jdi.run_pandexo(exo_dict, inst_dict, output_file='wasp43b.p')
Running Single Case w/ User Instrument Dict
```

6.3 Plot Results

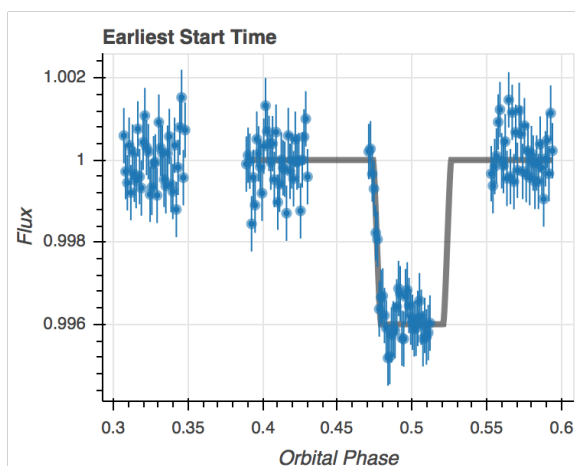
6.3.1 Plot simulated spectrum using specified file

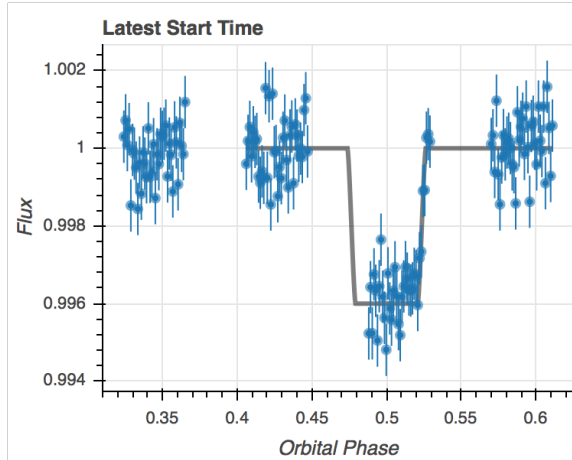
```
import pandexo.engine.justplotit as jpi
#using foo from above
#other keys include model=True/False
datawave, dataspec, dataerror, modelwave, modelspec = jpi.hst_spec(foo)
```



6.3.2 Compute earliest and latest start times for given start window size

```
#using foo from above
obsphase1, obstr1, obsphase2, obstr2, rms = jpi.hst_time(foo)
```





Print important info for observation

```
foo['wfc3_TExoNS']['info']
{'Estimated duty cycle (outside of Earth occultation)': 24.991166666666668,
 'Maximum pixel fluence (electrons)': 30266.370139081948,
 'Number of HST orbits': 4,
 'Number of Transits': 5,
 'Number of channels': 15,
 'Recommended scan rate (arcsec/s)': 0.07599999999999998,
 'Scan height (pixels)': 13.454512396694216,
 'Start observations between orbital phases': '0.307075052926-0.324148057092',
 'Transit depth uncertainty(ppm)': 62.433045276228441,
 'WFC3 parameters: NSAMP': 10,
 'WFC3 parameters: SAMP_SEQ': 'SPARS5'}
```

THE CODE

7.1 engine.justdoit

`engine.justdoit.get_thruput (inst)`

Returns complete instrument photon to electron conversion efficiency Pulls complete instrument photon to electron conversion efficiency (PCE) based on instrument key input

Parameters `inst (str)` – One of the instrument keys in *print_instruments*

Returns Dictionary with wave solution and PCE

Return type dict

Example

```
>>> thru_dict = get_thruput('NIRISS SOSS_Or1')
```

`engine.justdoit.load_exo_dict ()`

Loads in empty exoplanet dictionary for pandexo input

Loads in empty exoplanet dictionary so that the user can manually edit different planet parameters. Must be done for every bash run. User must edit each keys within the dictionary.

Returns Empty dictionary to be filled out by the user before running PandExo

Return type dict

Example

```
>>> exo_dict = load_exo_dict()
>>> exo_dict['planet']['transit_duration'] = 2*60*60 #2 hours
```

`engine.justdoit.load_mode_dict (inst)`

Function to pull in correct instrument dictionary

This is the instrument counterpart to `load_exo_dict`. It loads in a template instrument dictionary for a specific instrument mode.

Parameters `inst (str)` – One of the allowable instrument keys. To see which instrument modes are available use *print_instruments()*

Returns Filled out template of instrument dictionary, which can be edited before running to PandExo (not required).

Return type dict

Example

```
>>> inst_dict = load_mode_dict('MIRI LRS')
>>> inst_dict['configuration']['instrument']['aperture'] = 'lrsslit'
```

`engine.justdoit.print_instruments()`

Prints a list of the possible instrument templates to load

`engine.justdoit.run_inst_space(inst, exo)`

Changes inst dictionary and submits run

This function is used to reset the instrument dictionary.

Parameters

- **exo** (*dict*) – Exoplanet dictionary which can be loaded in and edited through `load_exo_dict`
- **inst** (*str*) – Key which indicates with instrument

Returns Dictionary with output of pandexo. Key is the value of the parameter that was looped through.

Return type dict

`engine.justdoit.run_pandexo(exo, inst, param_space=0, param_range=0, save_file=True, output_path='/Users/nbatalh1/Desktop/JWST/pandexo/docs', output_file='')`

Submits multiple runs of pandexo in parallel.

Functionality: program contains functionality for running single or multiple runs of PandExo

Parameters

- **exo** (*dict*) – exoplanet input dictionary
- **inst** (*dict or str or list of str*) – instrument input dictionary OR LIST of keys (for allowable keys see `print_instruments()`)
- **param_space** (*str or 0*) – (Optional) Default is 0 = no exoplanet parameter space. To run through a parameter specify which one need to specify two keys from exo dict with + in between. i.e. observation+fraction star+temp planet+exopath
- **param_range** (*list of str or list of float*) – (Optional) Default = 0 An array or list over which to run the parameters space. i.e. array of temperatures if running through stellar temp or array of files if running through planet models. Must specify param_space if using this.
- **save_file** (*bool*) – (Optional) Default = True saves file, False does not
- **output_path** (*str*) – (Optional) Defaults to current working directory
- **output_file** (*str*) – (Optional) Default is “singlerun.p” for single runs, “param_space.p” for exo parameter runs or “instrument_run.p” for instrument parameter space runs.

Returns For single run output will just be a single PandExo output dictionary <https://github.com/natashabatalha/PandExo/wiki/PandExo-Output> For multiple runs the output will be organized into a list with each a dictionary named by whatever you are looping through i.e. [{ ‘First temp’: PandExoDict}, { ‘Second temp’: PandExoDict}, etc..]

Return type dict

Example

For single run:

```
>>> a = run_pandexo(exo_dict, ['MIRI LRS'])
```

For multiple instruments:

```
>>> a = run_pandexo(exo_dict, ['MIRI LRS', 'NIRSpec G395H'])
```

Loop through a exoplanet parameter (stellar magnitude):

```
>>> a = run_pandexo(exo_dict, ['NIRSpec G395M'],
                    param_space='star+mag', param_range=np.linspace(6,10,5))
```

`engine.justdoit.run_param_space(i, exo, inst, param_space)`

Changes exo dictionary and submits run

This function is used to reset the exo dictionary based on what parameter is being looped over and submits run to *wrapper* so that all the jobs are run in parallel

Parameters

- **i** (*str or float*) – Can be either a str or float based on what you are looping through (str for filenames, float for stellar temps, float for magnitudes, etc)
- **exo** (*dict*) – Exoplanet dictionary which can be loaded in and edited through *load_exo_dict*
- **inst** (*str*) – Key which indicates with instrument
- **param_space** (*str*) – Set of keys within exo_dict to indicate which parameter to loop through. Should be in the format of “first level of dict”+”second level of dict”. For example, for stellar temp *param_space* would be “star+temp”

Returns Dictionary with output of pandexo. Key is the value of the parameter that was looped through.

Return type dict

7.2 engine.justplotit

`engine.justplotit.bin_wave_to_R(w, R)`

Creates new wavelength axis at specified resolution

Parameters

- **w** (*list of float or numpy array of float*) – Wavelength axis to be rebinned
- **R** (*float or int*) – Resolution to bin axis to

Returns New wavelength axis at specified resolution

Return type list of float

Examples

```
>>> newwave = bin_wave_to_R(np.linspace(1,2,1000), 10)
>>> print len(newwave)
11
```

`engine.justplotit.hst_spec(result_dict, plot=True, output_file='hstspec.html', model=True)`
 Plot 1d spec with error bars for hst

Parameters

- **result_dict** (*dict*) – Dictionary from pandexo output.
- **plot** (*bool*) – (Optional) True renders plot, False does not. Default=True
- **model** (*bool*) – (Optional) Plot model under data. Default=True
- **output_file** (*str*) – (Optional) Default = 'hstspec.html'

Returns

- **x** (*numpy array*) – micron
- **y** (*numpy array*) – 1D spec fp/f^* or rp^2/r^{*2}
- **e** (*numpy array*) – 1D rms noise
- **modelx** (*numpy array*) – micron
- **modely** (*numpy array*) – 1D spec fp/f^* or rp^2/r^{*2}

See also:

[`hst_time\(\)`](#)

`engine.justplotit.hst_time(result_dict, plot=True, output_file='hsttime.html', model=True)`
 Plot earliest and latest start times for hst observation

Parameters

- **result_dict** (*dict*) – Dictionary from pandexo output.
- **plot** (*bool*) – (Optional) True renders plot, False does not. Default=True
- **model** (*bool*) – (Optional) Plot model under data. Default=True
- **output_file** (*str*) – (Optional) Default = 'hsttime.html'

Returns

- **obsphase1** (*numpy array*) – earliest start time
- **obstr1** (*numpy array*) – white light curve
- **obsphase2** (*numpy array*) – latest start time
- **obstr2** (*numpy array*) – white light curve
- **rms** (*numpy array*) – 1D rms noise

See also:

[`hst_spec\(\)`](#)

`engine.justplotit.jwst_1d_bkg(result_dict, plot=True, output_file='bkg.html')`
 Plot background

Parameters

- **result_dict** (*dict*) – Dictionary from pandexo output. If parameter space was run in `run_pandexo` make sure to restructure the input as a list of dictionaries without they key words that `run_pandexo` assigns.
- **plot** (*bool*) – (Optional) True renders plot, Flase does not. Default=True
- **output_file** (*str*) – (Optional) Default = `bkt.html`

Returns

- **x** (*numpy array*) – micron
- **y** (*numpy array*) – 1D bakground e/s

See also:

`jwst_1d_spec()`, `jwst_noise()`, `jwst_1d_flux()`, `jwst_1d_snr()`, `jwst_2d_det()`, `jwst_2d_sat()`

`engine.justplotit.jwst_1d_flux(result_dict, plot=True, output_file='flux.html')`

Plot flux rate in e/s

Parameters

- **result_dict** (*dict*) – Dictionary from pandexo output. If parameter space was run in `run_pandexo` make sure to restructure the input as a list of dictionaries without they key words that `run_pandexo` assigns.
- **plot** (*bool*) – (Optional) True renders plot, Flase does not. Default=True
- **output_file** (*str*) – (Optional) Default = `'flux.html'`

Returns

- **x** (*numpy array*) – micron
- **y** (*numpy array*) – 1D flux rate in electrons/s

See also:

`jwst_1d_spec()`, `jwst_1d_bkg()`, `jwst_noise()`, `jwst_1d_snr()`, `jwst_2d_det()`, `jwst_2d_sat()`

`engine.justplotit.jwst_1d_snr(result_dict, plot=True, output_file='snr.html')`

Plot SNR

Parameters

- **result_dict** (*dict*) – Dictionary from pandexo output. If parameter space was run in `run_pandexo` make sure to restructure the input as a list of dictionaries without they key words that `run_pandexo` assigns.
- **plot** (*bool*) – (Optional) True renders plot, Flase does not. Default=True
- **output_file** (*str*) – (Optional) Default = `'snr.html'`

Returns

- **x** (*numpy array*) – micron
- **y** (*numpy array*) – 1D SNR

See also:

`jwst_1d_bkg()`, `jwst_noise()`, `jwst_1d_flux()`, `jwst_1d_spec()`, `jwst_2d_det()`, `jwst_2d_sat()`

```
engine.justplotit.jwst_1d_spec(result_dict, model=True, title='Model + Data + Error
                             Bars', output_file='data.html', legend=False, R=False,
                             num_tran=False, plot_width=800, plot_height=400,
                             x_range=[1, 10])
```

Plots 1d simulated spectrum and rebin or rescale for more transits

Plots 1d data points with model in the background (if wanted). Designed to read in exact output of run_pandexo.

Parameters

- **result_dict** (*dict or list of dict*) – Dictionary from pandexo output. If parameter space was run in run_pandexo make sure to restructure the input as a list of dictionaries without they key words that run_pandexo assigns.
- **model** (*bool*) – (Optional) True is default. True plots model, False does not plot model
- **title** (*str*) – (Optional) Title of plot. Default is “Model + Data + Error Bars”.
- **output_file** (*str*) – (Optional) name of html file for you bokeh plot. After bokeh plot is rendered you will have the option to save as png.
- **legend** (*bool*) – (Optional) Default is False. True, plots legend.
- **R** (*float*) – (Optional) Rebin data from native instrument resolution to specified resolution. Dafult is False, no binning.
- **num_tran** (*float*) – (Optional) Scales data by number of transits to improve error by $\sqrt{\text{num_trans}}$
- **plot_width** (*int*) – (Optional) Sets the width of the plot. Default = 800
- **plot_height** (*int*) – (Optional) Sets the height of the plot. Default = 400
- **x_range** (*list of int*) – (Optional) Sets x range of plot. Default = [1,10]

Returns **x,y,e** – Returns wave axis, spectrum and associated error in list format. x[0] will be correspond to the first dictionary input, x[1] to the second, etc.

Return type list of arrays

Examples

```
>>> jwst_1d_spec(result_dict, num_tran = 3, R = 35) #for a single plot
```

If you wanted to save each of the axis that were being plotted:

```
>>> x,y,e = jwst_1d_data([result_dict1, result_dict2], model=False, num_tran = 5,
↳ R = 100) #for multiple
```

See also:

```
jwst_noise(), jwst_1d_bkg(), jwst_1d_flux(), jwst_1d_snr(), jwst_2d_det(),
jwst_2d_sat()
```

```
engine.justplotit.jwst_2d_det(result_dict, plot=True, output_file='det2d.html')
```

Plot 2d detector image

Parameters

- **result_dict** (*dict*) – Dictionary from pandexo output. If parameter space was run in run_pandexo make sure to restructure the input as a list of dictionaries without they key words that run_pandexo assigns.

- **plot** (*bool*) – (Optional) True renders plot, Flase does not. Default=True
- **output_file** (*str*) – (Optional) Default = 'det2d.html'

Returns 2D array of out of transit detector simulation

Return type numpy array

See also:

`jwst_1d_spec()`, `jwst_1d_bkg()`, `jwst_1d_flux()`, `jwst_1d_snr()`, `jwst_noise()`,
`jwst_2d_sat()`

`engine.justplotit.jwst_2d_sat(result_dict, plot=True, output_file='sat2d.html')`

Plot 2d saturation profile

Parameters

- **result_dict** (*dict*) – Dictionary from pandexo output. If parameter space was run in `run_pandexo` make sure to restructure the input as a list of dictionaries without they key words that `run_pandexo` assigns.
- **plot** (*bool*) – (Optional) True renders plot, Flase does not. Default=True
- **output_file** (*str*) – (Optional) Default = 'sat2d.html'

Returns 2D array of out of transit detector simulation

Return type numpy array

See also:

`jwst_1d_spec()`, `jwst_1d_bkg()`, `jwst_1d_flux()`, `jwst_1d_snr()`, `jwst_2d_det()`,
`jwst_noise()`

`engine.justplotit.jwst_noise(result_dict, plot=True, output_file='noise.html')`

Plot background

Parameters

- **result_dict** (*dict*) – Dictionary from pandexo output. If parameter space was run in `run_pandexo` make sure to restructure the input as a list of dictionaries without they key words that `run_pandexo` assigns.
- **plot** (*bool*) – (Optional) True renders plot, Flase does not. Default=True
- **output_file** (*str*) – (Optional) Default = 'noise.html'

Returns

- **x** (*numpy array*) – micron
- **y** (*numpy array*) – 1D noise (ppm)

See also:

`jwst_1d_spec()`, `jwst_1d_bkg()`, `jwst_1d_flux()`, `jwst_1d_snr()`, `jwst_2d_det()`,
`jwst_2d_sat()`

`engine.justplotit.uniform_tophat_mean(xnew, x, y)`

Adapted from Mike R. Line to rebin spectra

Takes average of group of points in bin

Parameters

- **xnew** (*list of float or numpy array of float*) – New wavelength grid to rebin to

- **x**(*list of float or numpy array of float*) – Old wavelength grid to get rid of
- **y**(*list of float or numpy array of float*) – New rebinned y axis

Returns new y axis

Return type array of floats

Examples

```
>>> oldgrid = np.linspace(1,3,100)
>>> y = np.zeros(100)+10.0
>>> newy = uniform_tophat_sum(np.linspace(2,3,3), oldgrid, y)
>>> newy
array([ 240.,  250.,  130.] )
```

engine.justplotit.**uniform_tophat_sum**(*xnew*, *x*, *y*)

Adapted from Mike R. Line to rebin spectra

Takes sum of group of points in bin of wave points :param *xnew*: New wavelength grid to rebin to :type *xnew*: list of float or numpy array of float :param *x*: Old wavelength grid to get rid of :type *x*: list of float or numpy array of float :param *y*: New rebinned y axis :type *y*: list of float or numpy array of float

Returns new y axis

Return type array of floats

Examples

```
>>> oldgrid = np.linspace(1,3,100)
>>> y = np.zeros(100)+10.0
>>> newy = uniform_tophat_sum(np.linspace(2,3,3), oldgrid, y)
>>> newy
array([ 240.,  250.,  130.] )
```

7.3 engine.run_online

class engine.run_online.**AboutHandler**(*application*, *request*, ****kwargs**)

Bases: [engine.run_online.BaseHandler](#)

get ()

Render about PandExo Page

class engine.run_online.**Application**

Bases: [tornado.web.Application](#)

Gobal settings of the server This defines the global settings of the server. This parses out the handlers, and includes settings for if ever we want to tie this to a database.

class engine.run_online.**BaseHandler**(*application*, *request*, ****kwargs**)

Bases: [tornado.web.RequestHandler](#)

Logic to handle user information and database access might go here.

buffer = [OrderedDict](#)()

executor = <concurrent.futures.process.ProcessPoolExecutor object>

write_error (*status_code*, ***kwargs*)

This renders a customized error page

class `engine.run_online.CalculationDownloadHandler` (*application*, *request*, ***kwargs*)

Bases: `engine.run_online.BaseHandler`

Handlers returning the downloaded data of a particular calculation task. Handlers returning the status of a particular calculation task.

get (*id*)

class `engine.run_online.CalculationDownloadPandInHandler` (*application*, *request*, ***kwargs*)

Bases: `engine.run_online.BaseHandler`

Handlers returning the downloaded data of a particular calculation task. Handlers returning the status of a particular calculation task.

get (*id*)

class `engine.run_online.CalculationDownloadSpecHandler` (*application*, *request*, ***kwargs*)

Bases: `engine.run_online.BaseHandler`

Handlers returning the downloaded data of a particular calculation task. Handlers returning the status of a particular calculation task.

get (*id*)

class `engine.run_online.CalculationNewHSTHandler` (*application*, *request*, ***kwargs*)

Bases: `engine.run_online.BaseHandler`

This request handler deals with processing the form data and submitting a new HST calculation task to the parallelized workers.

get ()

post ()

The post method contains the returned data from the form data (accessed by using `self.get_argument(...)` for specific arguments, or `self.request.body` to grab the entire returned object.

class `engine.run_online.CalculationNewHandler` (*application*, *request*, ***kwargs*)

Bases: `engine.run_online.BaseHandler`

This request handler deals with processing the form data and submitting a new calculation task to the parallelized workers.

get ()

post ()

The post method contains the returned data from the form data (accessed by using `self.get_argument(...)` for specific arguments, or `self.request.body` to grab the entire returned object.

class `engine.run_online.CalculationNewSpecHandler` (*application*, *request*, ***kwargs*)

Bases: `engine.run_online.BaseHandler`

This request handler deals with processing the form data and submitting a new calculation task to the parallelized workers.

get ()

post ()

The post method contains the returned data from the form data (accessed by using `self.get_argument(...)` for specific arguments, or `self.request.body` to grab the entire returned object.

```
class engine.run_online.CalculationStatusHSTHandler (application, request, **kwargs)
    Bases: engine.run_online.BaseHandler

    Handlers returning the status of a particular HST calculation task.

    get (id)

class engine.run_online.CalculationStatusHandler (application, request, **kwargs)
    Bases: engine.run_online.BaseHandler

    Handlers returning the status of a particular JWST calculation task.

    get (id)

class engine.run_online.CalculationStatusSpecHandler (application, request, **kwargs)
    Bases: engine.run_online.BaseHandler

    Handlers returning the status of a particular Spec calculation task.

    get (id)

class engine.run_online.CalculationTask (id, name, task, cookie, count)
    Bases: tuple

    cookie
        Alias for field number 3

    count
        Alias for field number 4

    id
        Alias for field number 0

    name
        Alias for field number 1

    task
        Alias for field number 2

class engine.run_online.CalculationViewHSTHandler (application, request, **kwargs)
    Bases: engine.run_online.BaseHandler

    This handler deals with passing the results from Pandeia to the create_component_hst function which generates
    the Bokeh interactive plots.

    get (id)

class engine.run_online.CalculationViewHandler (application, request, **kwargs)
    Bases: engine.run_online.BaseHandler

    This handler deals with passing the results from Pandeia to the create_component_jwst function which generates
    the Bokeh interactive plots.

    get (id)

class engine.run_online.CalculationViewSpecHandler (application, request, **kwargs)
    Bases: engine.run_online.BaseHandler

    This handler deals with passing the results from Pandeia to the create_component_spec function which generates
    the Bokeh interactive plots.

    get (id)

class engine.run_online.DashboardHSTHandler (application, request, **kwargs)
    Bases: engine.run_online.BaseHandler
```

Request handler for the dashboard page. This will retrieve and render the html template, along with the list of current task objects.

`get()`

class `engine.run_online.DashboardHandler` (*application, request, **kwargs*)

Bases: `engine.run_online.BaseHandler`

Request handler for the dashboard page. This will retrieve and render the html template, along with the list of current task objects.

`get()`

class `engine.run_online.DashboardSpecHandler` (*application, request, **kwargs*)

Bases: `engine.run_online.BaseHandler`

Request handler for the dashboard page. This will retrieve and render the html template, along with the list of current task objects.

`get()`

class `engine.run_online.HelpfulPlotsHandler` (*application, request, **kwargs*)

Bases: `engine.run_online.BaseHandler`

`get()`

Renders helpful bokeh plots

class `engine.run_online.HomeHandler` (*application, request, **kwargs*)

Bases: `engine.run_online.BaseHandler`

`get()`

This sets an **unsecured** cookie. If user accounts gets implemented, this must be changed to a secure cookie.

class `engine.run_online.TablesHandler` (*application, request, **kwargs*)

Bases: `engine.run_online.BaseHandler`

`get()`

Render tables with confirmed candidates

`engine.run_online.main()`

7.4 engine.compute_noise

class `engine.compute_noise.ExtractSpec` (*inn, out, rn, extraction_area, timing*)

Different methods for computing noise.

PandExo has several different methods of computing noise. MULTIACCUM (slope method) assumes that each frame is fit for up the ramp and that the final noise includes correlated noise from fitting each frame up the ramp. First Minus Last assumes that intermediate frames are not used and final noise is just the first frame minus the last frame. 2d extract uses Pandeia's 2d simulations to extract the spectrum from the 2d extraction box. It extracts the entire postage stamp so it might be an overestimate of flux (in contrast pandeia requires background extraction region to be at least equal to the flux extraction region)

Noise Components Included:

- Shot
- Background and Dark Current
- Read noise

Parameters

- **inn** (*dict*) – In transit dictionary computed from PandExo
- **out** (*dict*) – Out of transit dictionary computed from PandExo
- **rn** (*float*) – Read noise electrons
- **extraction_area** (*float*) – Number of extracted pixels (square pixels)
- **timing** (*dict*) – Dictionary of computed JWST timing products

loopingL ()

extracts pixels from center to bottom

loopingU ()

extracts pixels from center to top

sum_spatial ()

sums pixels in optimal extraction region

extract_region ()

determines optimal extraction region

run_2d_extract ()

top level to extract spec from 2d pandeia output

run_slope_method ()

computes noise using multiaccum formulation

run_f_minus_1 ()

computes noise using first minus last

run_phase_spec ()

computes noise for phase curve observations

extract_region ()

Determine extraction Region

Contains functionality to determine extraction region from Pandeia 2d noise simulations. Calls *self.loopingL* and *self.loopingU*.

Returns bounding regions, photon and noise to be summed

Return type dict

loopingL (*cen, signal_col, noise_col, bkg_col*)

Finds bottom of the optimal extraction region.

Find location where SNR is the highest and loop from highest value downward

Parameters

- **cen** (*float or int*) – Pixel where SNR is the highest
- **signal_col** (*array of float*) – Array of fluxes to be extracted in a single column on the detector
- **noise_col** (*array of float*) – Array of noise to be extracted in a single column on the detector
- **bkg_col** (*array of float*) – Array of background fluxes to be extracted in a single column on the detector

Returns Bottom most pixel to be extracted

Return type int

loopingU (*cen, signal_col, noise_col, bkg_col*)

Finds top of the optimal extraction region.

Find location where SNR is the highest and loop from highest value upward

Parameters

- **cen** (*float or int*) – Pixel where SNR is the highest
- **signal_col** (*array of float*) – Array of fluxes to be extracted in a single column on the detector
- **noise_col** (*array of float*) – Array of noise to be extracted in a single column on the detector
- **bkg_col** (*array of float*) – Array of background fluxes to be extracted in a single column on the detector

Returns Top most pixel to be extracted

Return type int

run_2d_extract ()

Extract noise from 2d detector image

Contains functionality to extract noise from 2d detector image

Returns all optimally extracted 1d products

Return type dict

run_f_minus_1 ()

Compute noise using first minus last formula

Uses 1d extracted products from pandeia to compute noise without multiaccum noise formula from Rauscher 07. Includes readnoise background.

Returns all optimally extracted 1d products

Return type dict

run_phase_spec ()

Computes time dependent noise for phase curves.

Computes noise for phase curve analysis instead of spectroscopy. Using MULTIACCUM formula here but this could be changed in the future. Does not return a spectra for each time element... On your own for that.

Returns all optimally extracted 1d products

Return type dict

run_slope_method ()

Compute noise using Pandeia 1d noise

Contains functionality to compute noise using Pandeia 1d noise output products (uses MULTIACCUM) noise formula

Returns all optimally extracted 1d products

Return type dict

sum_spatial (*extract_info*)

Sum pixel in the spatial direction

Takes extraction info from *extract_region* and sums pixels in that region taking into account integrations and number of transits

Parameters `extract_info` (*dict*) – Dictionary with information on extraction box, flux and noise products

Returns Dictionary with all extracted 1d products including noise, background, fluxes

Return type dict

7.5 engine.create_input

`engine.create_input.binning` (*x*, *y*, *R*=2000.0)

Bins spectra at fixed dlambda

Takes 2 arrays *x* and *y* and bins them into groups of blength. Uses uniform top hat to bin. This is necessary because really high resolution spectra cannot go through pandeia without crashing it. User is advised to bin spectra before running it through pandexo.

Parameters

- **x** (*list or numpy array of float*) – *x* (micron)
- **y** (*list or numpy array of float*) – (any unit)
- **R** (*int or float*) – (optional) Default is 2000.0 at micron which is approximately max resolution pandeia can handle.

Returns

- **xout** (*numpy array of float*) – 1D numpy arrays at new resolution
- **yout** (*numpy array of float*) – 1D numpy arrays at new resolution

`engine.create_input.bothTrans` (*out_trans*, *planet*)

Calculates in transit flux

Takes output from *outTrans*, which is the normalized stellar flux, and creates either a transit transmission spectrum, phase curve or emission spectrum. Magnitude

Parameters

- **out_trans** (*dict*) – includes dictionary from *outTrans* output.
- **planet** (*dict*) – dictionary with direction to planet spectra, wavelength and flux units

Returns dictionary with out of transit flux, in transit flux, original model and corresponding wavelengths

Return type dict

`engine.create_input.outTrans` (*input*)

Compute out of transit spectra

Computes the out of transit spectra by normalizing flux to specified magnitude and convert to specified Pandeia units of milliJy and microns.

Parameters **input** (*dict*) – stellar scene which includes parameters to extract phoenix database or a filename which points to a stellar spectrum

Returns contains wave and flux_out_trans

Return type dict

7.6 engine.hst

`engine.hst.calc_start_window(eventType, rms, ptsOrbit, numOrbits, depth, inc, aRs, period, windowSize, ecc=0, w=90.0, duration=None, offset=0.0)`

Calculate earliest and latest start times

Plot earliest and latest possible spectroscopic light curves for given start window size

Parameters

- **eventType** (*str*) – ‘transit’ or ‘eclipse’
- **rms** (*float*) – light curve root-mean-square
- **ptsOrbit** (*float*) – number of frames per HST orbit
- **numOrbits** (*float*) – number of HST orbits per visit
- **depth** (*float*) – transit/eclipse depth
- **inc** (*float*) – orbital inclination in degrees
- **aRs** (*float*) – Semi-major axis in units of stellar radii (a/R^*)
- **period** (*float*) – orbital period in days
- **windowSize** (*float*) – observation start window size in minutes
- **ecc** (*float*) – (Optional) eccentricity
- **w** (*float*) – (Optional) longitude of periastron
- **duration** (*float*) – (Optional) full transit/eclipse duration in days
- **offset** (*float*) – (Optional) manual offset in observation start time, in minutes

Returns

- *float* – minphase–earliest observation start phase
- *float* – maxphase–latest observation start phase

`engine.hst.compute_sim_hst(dictinput)`

Sets up HST simulations

Function to set up exoplanet observations for HST only and compute simulated spectrum.

Parameters **dictinput** (*dict*) – instrument and pandexo dictionaries in format {“pandea_input”:dict1, “pandexo_input”:dict2}

Returns All hst output info needed to plot simulated data, light curves timing info

Return type dict

`engine.hst.create_out_div(input_dict, minphase, maxphase)`

Function to render input dicts in html format for web front end

Parameters **input_dict** (*dict*) – any input dictionary

Returns html rendered table

Return type div

`engine.hst.planet_spec(specfile, w_unit, disperser, deptherr, nchan, smooth=None)`

Plot exoplanet transmission/emission spectrum

Parameters

- **specfile** (*str*) – filename for model spectrum [wavelength, flux]
 - **w_unit** (*str*) – wavelength unit (um or nm)
 - **disperser** – grism (g102 or g141)
 - **deptherr** (*float*) – simulated transit/eclipse depth uncertainty
 - **nchan** (*float*) – number of spectrophotometric channels
 - **smooth** (*float*) – (Optional)length of smoothing kernel
- returns** contains following keys {‘model_wave’,‘model_spec’,‘binwave’,‘binspec’,
‘error’,‘wmin’,‘wmax’}
- rtype** dict

`engine.hst.wfc3_GuessNOrbits` (*trdur*)

Predict number of HST orbits

Predict number of HST orbits for transit observation when not provided by the user.

Parameters *trdur* (*float*) – transit duration in days

Returns number of requested orbits per transit (including discarded thermal-settling orbit)

Return type float

`engine.hst.wfc3_GuessParams` (*hmag*, *disperser*, *scanDirection*, *subarray*, *obsTime*, *maxExptime=150.0*)

Predict nsamp and samp_seq when values not provided by the user.

Parameters

- **hmag** (*float*) – H-band magnitude
- **disperser** (*str*) – grism (‘G141’ or ‘G102’)
- **scanDirection** (*str*) – spatial scan direction (‘Forward’ or ‘Round Trip’)
- **subarray** (*str*) – Subarray aperture (‘grism256’ or ‘grism512’)
- **obsTime** (*float*) – Available observing time per HST orbit in seconds
- **maxExptime** (*float*) – (Optional) default=150.0, maximum exposure time in seconds

Returns

- *float* – nsamp–number of up-the-ramp samples (1..15)
- *str* – samp_seq–time between non-destructive reads

`engine.hst.wfc3_TExoNS` (*dictinput*)

Compute Transit depth uncertainty

Compute the transit depth uncertainty for a defined system and number of spectrophotometric channels. Written by Kevin Stevenson October 2016

Parameters *dictinput* (*dict*) – dictionary containing instrument parameters and exoplanet specific parameters. {“pandea_input”:dict1, “pandexo_input”:dict1}

Returns

- *float* – deptherr–transit depth uncertainty per spectrophotometric channel
- *float* – chanrms–light curve root mean squarermes
- *float* – ptsOrbit–number of HST orbits per visit

`engine.hst.wfc3_obs(hmag, disperser, scanDirection, subarray, nsamp, samp_seq)`

Determine the recommended exposure time, scan rate, scan height, and overheads.

Parameters

- **hmag** (*float*) – H-band magnitude
- **disperser** (*str*) – Grism ('G141' or 'G102')
- **scanDirection** (*str*) – spatial scan direction ('Forward' or 'Round Trip')
- **subarray** (*str*) – Subarray aperture ('grism256' or 'grism512')
- **nsamp** (*float*) – Number of up-the-ramp samples (1..15)
- **samp_seq** (*str*) – Time between non-destructive reads ('SPARS5', 'SPARS10', or 'SPARS25')

Returns

- *float* – exptime–exposure time in seconds
- *float* – tottime–total frame time including overheads in seconds
- *float* – scanRate–recommended scan rate in arcsec/s
- *float* – scanHeight–scan height in pixels
- *float* – fluence–maximum pixel fluence in electrons

7.7 engine.jwst

`engine.jwst.add_noise_floor(noise_floor, wave_bin, error_spec)`

Add in noise floor

This adds in a user specified noise floor. Does not add the noise floor in quadrature instead it sets `error[error<noise_floor] = noise_floor`. If a wavelength dependent noise floor is given and the wavelength ranges are off, it interpolates the out of range noise floor.

Parameters

- **noise_floor** (*str or int*) – file with two column [wavelength, noise(ppm)] or single number with constant noise floor in ppm
- **wave_bin** (*array of float*) – final binned wavelength grid from simulation
- **error_spec** (*array of float*) – final computed error on the planet spectrum in units of rp^2/r^2 or fp/f^*

Returns `error_spec`– new error

Return type array of float

Examples

```
>>> import numpy as np
>>> wave = np.linspace(1,2.7,10)
>>> error = np.zeros(10)+1e-6
>>> newerror = add_noise_floor(20, wave, error)
>>> print newerror
[ 2.00000000e-05  2.00000000e-05  2.00000000e-05  2.00000000e-05
```

2.00000000e-05	2.00000000e-05	2.00000000e-05	2.00000000e-05
2.00000000e-05	2.00000000e-05]		

`engine.jwst.add_warnings(pand_dict, timing, sat_level, flags, instrument)`

Add warnings for front end

Adds in necessary warning flags for a JWST observation usually associated with too few or too many groups or saturation. Alerts user if saturation level is higher than 80 percent and if the number of groups is less than 5. Or, if the full well is greater than 80. These warnings are currently very arbitrary. Will be updated as better JWST recommendations are made.

Parameters

- **pand_dict** – output from pandeia run
- **timing** (*dict*) – output from **compute_timing**
- **sat_level** (*int or float*) – user specified saturation level in electrons
- **flags** (*dict*) – warning flags taken from output of **compute_timing**
- **instrument** (*str*) – Only allowable strings are: “nirspec”, “niriss”, “nircam”, “miri”

Returns all warnings

Return type dict

Notes

These are warnings are just suggestions and are not yet required.

`engine.jwst.as_dict(out, both_spec, binned, timing, mag, sat_level, warnings, punit, unbinned, calculation)`

Format dictionary for output data

Takes all output from jwst run and converts it to simple dictionary

Parameters

- **out** (*dict*) – output dictionary from **compute_out**
- **both_spec** (*dict*) – output dictionary from **createInput.bothTrans**
- **binned** (*dict*) – dictionary from **wrapper**
- **timing** (*dict*) – dictionary from **compute_timing**
- **mag** (*dict*) – magnitude of system
- **sat_level** (*float or int*) – saturation level in electrons
- **warnings** (*dict*) – warning dictionary from **add_warnings**
- **punit** (*"fp/f*" or "rp^2/r*^2"*) – unit of supplied spectra options are: only options are fp/f* or rp^2/r*^2
- **unbinned** (*dict*) – unbinned raw data from **wrapper**
- **calculation** (*str*) – noise calculation type

Returns compressed dictionary

Return type dict

`engine.jwst.bin_wave_to_R(w, R)`

Creates new wavelength axis at specified resolution

Rebins wavelength bin to a certain R

Parameters

- **w** (*list of float or numpy array of float*) – Wavelength axis to be rebinned
- **R** (*float or int*) – Resolution to bin axis to

Returns New wavelength axis at specified resolution

Return type list of float

Examples

```
>>> newwave = bin_wave_to_R(np.linspace(1,2,1000), 10)
>>> print len(newwave)
11
```

`engine.jwst.compute_full_sim(dictinput)`

Top level function to set up exoplanet obs. for JW

Function to set up exoplanet observations for JWST only and compute simulated spectrum. It uses STScI's Pandeia to compute instrument throughputs and WebbPSF to compute PSFs.

Parameters `dictinput` (*dict*) – dictionary containing instrument parameters and exoplanet specific parameters. {"pandeia_input":dict1, "pandexo_input":dict1}

Returns large dictionary with 1d, 2d simulations, timing info, instrument info, warnings

Return type dict

Examples

```
>>> from pandexo.engine.jwst import compute_full_sim
>>> from pandexo.engine.justplotit import jwst_1d_spec
>>> a = compute_full_sim({"pandeia_input": pandeiadict, "pandexo_input":exodict})
>>> jwst_1d_spec(a)
.. image:: 1d_spec.png
```

Notes

It is much easier to run simulations through either **run_online** or **justdoit**. **justdoit** contains functions to create input dictionaries and **run_online** contains web forms to create input dictionaries.

See also:

pandexo.engine.justdoit.run_pandexo() Best function for running pandexo runs

pandexo.engine.run_online() Allows running functions through online interface

`engine.jwst.compute_maxexptime_per_int(pandeia_input, sat_level)`

Computes optimal maximum exposure time per integration

Function to simulate 2d jwst image with 2 groups, 1 integration, 1 exposure and return the maximum time for one integration before saturation occurs. If saturation has already occurred, returns `maxexptime_per_int` as `np.nan`. This then tells Pandexo to set min number of groups (`ngroups=2`). This avoids error if saturation occurs. This routine assumes that min `ngroups` is 2.

Parameters

- **pandeia_input** (*dict*) – pandeia dictionary input
- **sat_level** (*int or float*) – user defined saturation level in units of electrons

Returns Maximum exposure time per integration before specified saturation level

Return type float

Examples

```
>>> max_time = compute_maxexptime_per_int(pandeia_input, 50000.0)
>>> print max_time
12.0
```

`engine.jwst.compute_timing(m, transit_duration, expfact_out, noccultations)`

Computes all timing info for observation

Computes all JWST specific timing info for observation including. Some pertinent JWST terminology:

- **frame**: The result of sequentially clocking and digitizing all pixels in a rectangular area of an SCA. **Full-frame readout** means to digitize all pixels in an SCA, including reference pixels. **Frame** also applies to the result of clocking and digitizing a subarray on an SCA.
- **group**: One or more consecutively read frames. There are no intervening resets. Frames may be averaged to form a group but for exoplanets the read out scheme is always 1 frame = 1 group
- **integration**: The end result of resetting the detector and then non-destructively sampling it one or more times over a finite period of time before resetting the detector again. This is a unit of data for which signal is proportional to intensity, and it consists of one or more GROUPS.
- **exposure**: The end result of one or more INTEGRATIONS over a finite period of time. EXPOSURE defines the contents of a single FITS file.

Parameters

- **m** (*dict*) – Dictionary output from `compute_maxexptime_per_int`
- **transit_duration** (*float or int*) – transit duration in seconds
- **expfact_out** (*float or int*) – fraction of time spent in transit versus out of transit
- **noccultations** (*int*) – number of transits

Returns

- **timing** (*dict*) – All timing info
- **warningflag** (*dict*) – Warning flags

Examples

```
>>> timing, flags = compute_timing(m, 2*60.0*60.0, 1.0, 1.0)
>>> print timing.keys()
['Number of Transits', 'Num Integrations Out of Transit', 'Num Integrations In_
↳Transit',
'APT: Num Groups per Integration', 'Seconds per Frame', 'Observing Efficiency (%)
↳', 'On Source Time(sec)',
'Exposure Time Per Integration (secs)', 'Reset time Plus 30 min TA time (hrs)',
'APT: Num Integrations per Occultation', 'Transit Duration']
```

`engine.jwst.perform_in(pandeia_input, pandexo_input, timing, both_spec, out, calculation)`

Computes in transit data

Runs Pandeia for the in transit data or computes the in transit simulation from the out of transit pandeia run

Parameters

- **pandeia_input** (*dict*) – pandeia specific input info
- **pandexo_input** (*dict*) – exoplanet specific observation info
- **timing** (*dict*) – timing dictionary from **compute_timing**
- **both_spec** (*dict*) – dictionary transit spectra computed from **createInput.bothTrans**
- **out** (*dict*) – out of transit dictionary from **perform_in**
- **calculation** (*str*) – key which specifies the kind of noise calculation (2d extract, slope method, fml, phase_spec). Recommended for transit transmission spectra = fml

Returns pandeia output dictionary

Return type dict

`engine.jwst.perform_out(pandeia_input, pandexo_input, timing, both_spec)`

Runs pandeia for the out of transit data

Parameters

- **pandeia_input** (*dict*) – pandeia specific input info
- **pandexo_input** (*dict*) – exoplanet specific observation info
- **timing** (*dict*) – timing dictionary from **compute_timing**
- **both_spec** (*dict*) – dictionary transit spectra computed from **createInput.bothTrans**

Returns pandeia output dictionary for out of transit data

Return type dict

`engine.jwst.uniform_tophat_sum(xnew, x, y)`

Adapted from Mike R. Line to rebin spectra

Sums groups of points in certain wave bin

Parameters

- **xnew** (*list of float or numpy array of float*) – New wavelength grid to rebin to
- **x** (*list of float or numpy array of float*) – Old wavelength grid to get rid of
- **y** (*list of float or numpy array of float*) – New rebinned y axis

Returns new wavelength grid

Return type array of floats

Examples

```
>>> from pandexo.engine.jwst import uniform_tophat_sum
>>> oldgrid = np.linspace(1,3,100)
>>> y = np.zeros(100)+10.0
>>> newy = uniform_tophat_sum(np.linspace(2,3,3), oldgrid, y)
>>> newy
array([ 240.,  250.,  130.] )
```

7.8 engine.load_modes

class engine.load_modes.**SetDefaultModes** (*inst*)

Class to load default instrument mode dicts

This class contains functionality for loading observing modes for exoplanet observations This is NOT a complete set of ALL possible observing modes. Instead, it offers a starting point for choosing one instrument specification. There is one function for each instrument. For example, It sets slitless mode for MIRI LRS. If users are interested in other specific observation modes they should load in the dictionary and then edit individual keys.

Included modes are:

- “MIRI LRS”
- “NIRISS SOSS”
- “NIRSpec G140M”
- “NIRSpec G140H”
- “NIRSpec G235M”
- “NIRSpec G235H”
- “NIRSpec G395M”
- “NIRSpec G395H”
- “NIRSpec Prism”
- “NIRCam F322W2”
- “NIRCam F444W”
- “WFC3 G102”
- “WFC3 G141”

Parameters **inst** (*str*) – Allowable strings listed above

miri ()

Handles MIRI template

nircam ()

Handles NIRCcam template

niriss()
Handles NIRISS template

nirspec()
Handles NIRSpec template

pick()
Points to specific instrument based on key choice

wfc3()
Handles WFC3 template

7.9 engine.pandexo

`engine.pandexo.wrapper(dictinput)`

Top level function to call either jwst, hst, wfirst

Top level function which calls either jwst, hst or wfirst noise simulation.

Parameters `dictinput` – dictionary containing instrument parameters and exoplanet specific parameters. {"pandea_input":dict1, "pandexo_input":dict1}

Returns output specific to observatory requested

Return type dict

Notes

You should not run simulations through this. It is much easier to run simulations through either **run_online** or **justdoit**. **justdoit** contains functions to create input dictionaries and **run_online** contains web forms to create input dictionaries.

See also:

`pandexo.engine.justdoit()` gives ability to simply submit runs

`pandexo.engine.run_online()` submit runs through user interface

7.10 engine.ComputeZ

`engine.ComputeZ.addBackground(molDict)`

`engine.ComputeZ.computeAlpha(molDict, plot=False)`

`engine.ComputeZ.computeJac(molDict)`

Function to construct Jacobian of equation alpha <http://arxiv.org/pdf/1511.09443v2.pdf> eqn. 4

Inputs: T, mu, g, beta: float xsec: dict of cross sections squig: dict of mixing ratios

`engine.ComputeZ.computeMu(squig)`

Compute mean molecular weight of an atmosphere

Input: Dict of molecules with mixing ratios i.e. {"H2O": 0.98, "H2":0.2}

Output: mean molecular weight, float

`engine.ComputeZ.computeZSum(xsec, squig, waves, plot2, plot)`

`engine.ComputeZ.find_nearest` (*array, value*)

`engine.ComputeZ.readXSecs` (*molDict*)

7.11 engine.elements

Properties of the chemical elements.

Each chemical element is represented as an object instance. Physicochemical and descriptive properties of the elements are stored as instance attributes.

Author Christoph Gohlke

Version 2015.01.29

7.11.1 Requirements

- CPython 2.7 or 3.4

References

1. <http://physics.nist.gov/PhysRefData/Compositions/>
2. <http://physics.nist.gov/PhysRefData/IonEnergy/tblNew.html>
3. [http://en.wikipedia.org/wiki/%\(element.name\)s](http://en.wikipedia.org/wiki/%(element.name)s)
4. <http://www.miranda.org/~jkominek/elements/elements.db>

Examples

```
>>> from elements import ELEMENTS
>>> len(ELEMENTS)
109
>>> str(ELEMENTS[109])
'Meitnerium'
>>> ele = ELEMENTS['C']
>>> ele.number, ele.symbol, ele.name, ele.eleconfig
(6, 'C', 'Carbon', '[He] 2s2 2p2')
>>> ele.eleconfig_dict
{(1, 's'): 2, (2, 'p'): 2, (2, 's'): 2}
>>> sum(ele.mass for ele in ELEMENTS)
14659.111599
>>> for ele in ELEMENTS:
...     ele.validate()
...     ele = eval(repr(ele))
```

7.12 engine.hst_smooth

`engine.hst_smooth.medfilt` (*x, window_len*)

Apply a length-k median filter to a 1D array *x*. Boundaries are extended by repeating endpoints.

`engine.hst_smooth.smooth(x, window_len=10, window='hanning')`
smooth the data using a window with requested size.

This method is based on the convolution of a scaled window with the signal. The signal is prepared by introducing reflected copies of the signal (with the window size) in both ends so that transient parts are minimized in the beginning and end part of the output signal.

Parameters

- **x** (*array of floats*) – the input signal
- **window_len** (*int*) – (Optional) Default=10. The dimension of the smoothing window
- **window** (*str*) – the type of window from ‘flat’, ‘hanning’, ‘hamming’, ‘bartlett’, ‘blackman’ flat window will produce a moving average smoothing.

Returns The smoothed signal

Return type array of floats

Examples

```
>>> t=linspace(-2,2,0.1)
>>> x=sin(t)+randn(len(t))*0.1
>>> y=smooth(x)
```

See also:

`numpy.hanning()`, `numpy.hamming()`, `numpy.bartlett()`, `numpy.blackman()`, `numpy.convolve()`, `scipy.signal.lfilter()`, `Todos()`, `-----()`, `The()`

Source () <http://www.scipy.org/Cookbook/SignalSmooth> 2009-03-13

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