

HSIM Manual

v211

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Revision History

Version	Date	Comments
1.0	2018-09-27	HSIM v204 release
1.1	—	HSIM v210 release
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1. Quick Start Guide

1.1 Introduction and Installation

HSIM is a dedicated pipeline for simulating observations with HARMONI on the Extremely Large Telescope. HSIM takes high spectral and spatial resolution input data cubes, encoding physical descriptions of astrophysical sources, and generates mock observed data cubes. The simulations incorporate detailed models of the sky, telescope, instrument, and detectors to produce realistic mock data (?).

HSIM is programmed in Python and the source code can be found at <https://github.com/HARMONI-ELT/HSIM>. It does not require any installation, just download it from and unzip. HSIM depends on the following Python packages to work:

- astropy 2.0.4
- numpy 1.14.1
- scipy 1.0.0
- matplotlib 2.1.2
- wxPython > 3.0

The code has been tested with the indicated package version, although more recent releases of these packages are likely to work as well.

1.2 Running Simulations

1.2.1 Preparing the Input Datacube

Before running HSIM, you will need an input model of your astronomical object stored as a datacube in a FITS file. The recommended sizes for the spatial pixels (spaxels) are 1, 2, or 3 mas depending on the selected spaxel scale for HARMONI (see Table 1.1). If your datacube has a different spaxel scale, HSIM will automatically interpolate/rebin the input datacube to the recommended values.

Similarly, it is recommended to oversample the spectral dimension of the input cube by a factor of 4 with respect to the nominal resolving power (R) of the selected grating (e.g., the spectral sampling for the K-grating $R=7100$ at $2.2\mu\text{m}$ should be 0.078 nm). If your input cube has a different spectral sampling, HSIM will interpolate/rebin it. If possible, it is recommend to use input cubes with a spectral resolution a factor of 2 better than the R of the selected grating.

The information on the spatial and spectral sampling of the input cube are passed through the FITS file header to HSIM (Table 1.2 shows a summary of all the header keywords used by HSIM). In particular, the CDELT1 and CDELT2 header values are used to

Table 1.1: Recommended input pixel size

HARMONI Spaxel scale (mas)	Recommended input scale (mas)
4×4	1
10×10	2
20×20	3
30×60	3

get the spatial scale, and CDEL3 to obtain the spectral sampling. The spectral resolution of the input cube is indicated by SPECRES in wavelength units similar to CDEL3. The units of these values are those indicated by the CUNIT1, CUNIT2, and CUNIT3 header values. Currently, for the spatial units (CUNIT1 and CUNIT2), the accepted values are mas and arcsec. For the spectral units, CUNIT3 and SPECRES, HSIM recognizes the following values: angstrom, nanometers, nm, micron, and meter. The size of the input cube is obtained from NAXIS1, NAXIS2, and NAXIS3.

The wavelength of each slice of the data cube are also calculated from the FITS header using the following relation:

$$\lambda_i = \text{CRVAL3} + \text{CDEL3} \times (i - \text{CRPIX3}) \quad (1.1)$$

where i is the slice number from 1 to NAXIS3. The wavelength range of the input cube and the selected grating should, at least, partially overlap.

Finally, the input data cube should be in surface brightness units indicated by BUNIT. The accepted units are the following: erg/s/cm²/um/arcsec², erg/s/cm²/A/arcsec², J/s/m²/um/arcsec², and J/s/m²/A/arcsec².

Table 1.2: FITS header keywords used by HSIM

Keyword	Description	Accepted values
Spatial information		
NAXIS{1,2}	Number of pixels along x-axis and y-axis	
CTYPE1	Type of the spatial x-axis	x, ra
CTYPE2	Type of the spatial y-axis	y, dec
CDEL3{1,2}	X and Y pixel size	
CUNIT{1,2}	Units of CDEL3{1,2}	mas, arcsec
Spectral information		
NAXIS3	Number of spectral pixels	
CTYPE3	Type of the spectral axis	wavelength
CDEL3	Size of the spectral pixel	
SPECRES	Spectral resolution in wavelength units	
CRPIX3	Reference pixel for the spectral axis (see Equation 1.1)	
CRVAL3	Wavelength of the reference pixel	
CUNIT3	Units of CDEL3, CRVAL3, and SPECRES	angstrom, nanometers, nm, micron, meter
Flux units		
BUNIT	Flux units of the cube	erg/s/cm ² /um/arcsec ² , erg/s/cm ² /A/arcsec ² , J/s/m ² /um/arcsec ² , J/s/m ² /A/arcsec ²

1.2.2 HSIM Gratings

The gratings that can be selected in HSIM are detailed below in Table 1.3.

Table 1.3: HSIM grating parameters			
Grating	λ_{min} (μm)	λ_{max} (μm)	R
Low resolution			
V+R	0.458	0.820	3100
Iz+J	0.811	1.369	3355
H+K	1.450	2.450	3355
Medium resolution			
Iz	0.830	1.050	7104
J	1.046	1.324	7104
H	1.435	1.815	7104
K	1.951	2.469	7104
High resolution			
z-high	0.828	0.902	17385
J-short	1.012	1.102	17385
J-long	1.098	1.189	17385
J-high	1.191	1.299	17385
H-high	1.538	1.678	17385
K-short	2.017	2.201	17385
K-long	2.199	2.400	17385

1.2.3 Detector Systematics

If chosen, HSIM can be run with more detailed detector systematics to simulate some of the features seen on infra-red detectors. When this option is selected, HSIM creates 8 simulated HAWAII 4RGs where each pixel has a unique read noise value derived from analysing the KMOS H2RG detectors. The read noise values have a distribution, with the peak at the expected read noise value for the H4RGs. An example HSIM H4RG is seen in Figure 1.1.

The simulated data cube is then sliced up and added to these detectors, in a manner similar to the effect of the image slicer. This ensures that features that appear linear on the detectors no longer appear linear in the output data cube. The simulation is then folded back into a standard data cube to maintain the ‘cube in, cube out’ nature of HSIM.

When creating the reduced cube, a separate instance of the detectors is generated and used in the reduction, so the residual features will remain in the reduced cube. A separate cube is returned containing the detectors used in the simulation.

1.2.4 Running HSIM

Once the input data cube has been prepared, the next step is to run HSIM. HSIM comes with a graphical user interface (GUI) which can be used to define the input parameters of the simulations. Alternatively, the same options can be directly accessed from the command line.

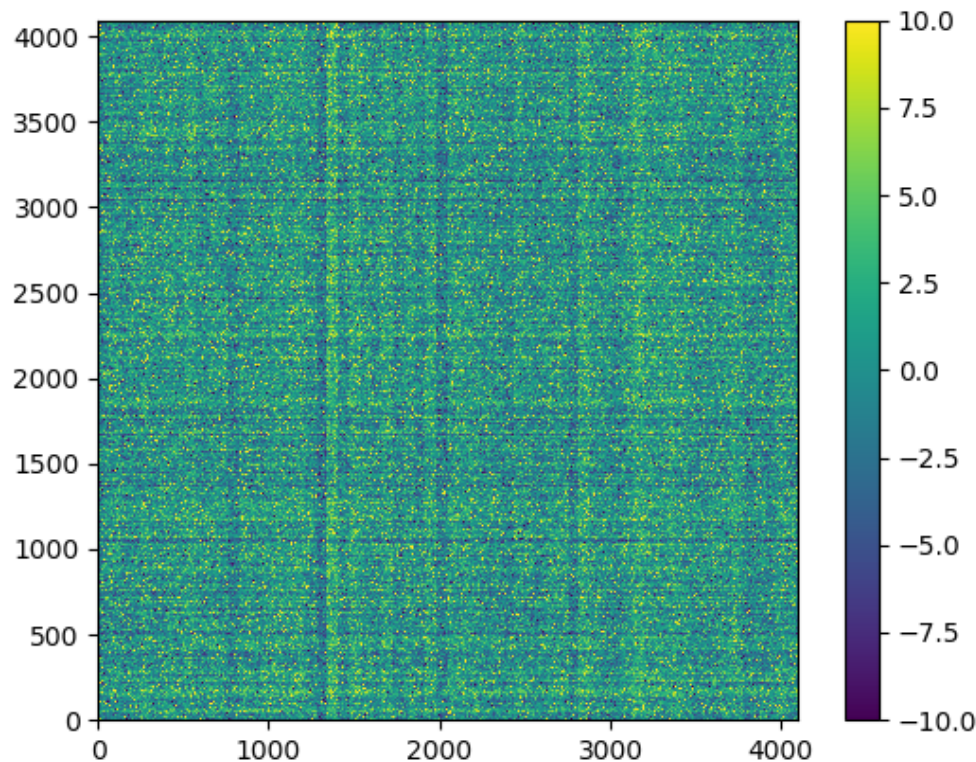


Figure 1.1: HSIM simulated detector.

GUI

The GUI provides a relatively easy way to list the available options. To launch HSIM in the GUI mode, type the following command

```
$ python hsim2.py
```

Then, a window similar to that shown in Figure 1.2 should appear. In that window, it is possible to select the input cube and the output directory and define all the parameters needed by HSIM to run the simulation (these parameters are explained in Section 1.2.5). Clicking on the **Commence simulation** button will start the simulation. You can see how the simulation advances in the terminal window from where HSIM was called.

Command-line

All the GUI options can be directly accessed from the command line mode using the following command

```
$ python hsim2.py -c arg1 arg2 ...
```

where **arg*** are the parameters of the simulation (see Section 1.2.5). All the parameters are needed to run simulation. After the **-c**, you can add **-p number**, where **number** is

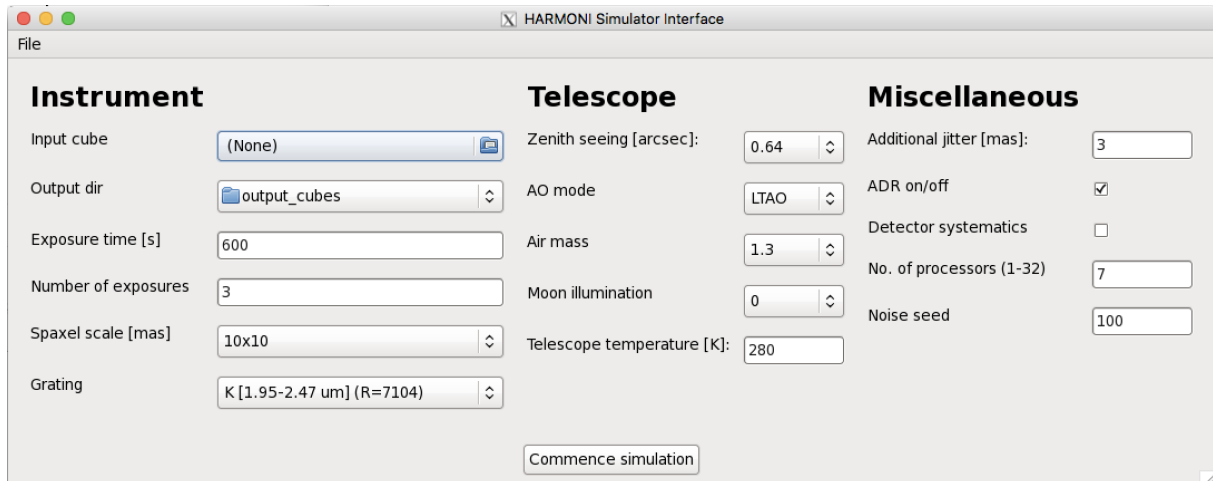


Figure 1.2: HSIM GUI.

the number of processors that will be used by HSIM. You can also add `-o directory` where `directory` is the output directory. If no output directory is specified, the default `output_cubes` will be used and results from previous simulations might be overwritten. Finally, using `-h`, will show a short help message.

1.2.5 List of Options

The available HSIM options are listed below. The number before them indicates the parameter order when used in the command line mode.

1. Input cube: Path of the input datacube (e.g., `datacube.fits`).
2. Exposure time: Integration time of each exposure in seconds (e.g., 600).
3. Number of exposures (e.g., 3).
4. Grating: Indicates the HARMONI grating. The available options are the following: V+R, Iz+J, H+K, Iz, J, H, K, z-high, J-high, H-high, K-short, K-long.
5. Spaxel Scale: HARMONI spatial scale in mas. The available values are the following: 4x4, 10x10, 20x20, 30x60.
6. Zenith seeing: Atmospheric seeing FWHM in arcsec. The available values are the following: 0.43, 0.57, 0.64, 0.72, 1.04.
7. Air mass: Average air mass during the HARMONI observation. The available values are the following: 1.1, 1.3, 1.5, 2.0, which correspond to zenith angles of 25°, 40°, 48°, and 60°, respectively.
8. Moon illumination: Fraction of the Moon that is illuminated at the time of the observation. A 30° separation between target and Moon is assumed. The available values are the following: 0, 0.5, 1.0.
9. Jitter: Additional telescope PSF blur in mas. Typical values are 2 and 3 for SCAO and LTAO, respectively.
10. Telescope temperature: Temperature of the site/telescope in K. This is used to calculate the telescope and part of the instrument background emission. Typical value is 280.

11. ADR on/off: Indicates if the atmospheric differential refraction is simulated. When simulated, the differential refraction is corrected in the output cubes in a way similar to what is expected from the pipeline. Available values are **True**, **False**.
12. Noise seed: Seed used for the random number generator during the simulation. It only affects the observed outputs (see Section 1.2.6).
13. AO mode: Adaptive optics mode of the observations. The available values are the following: **LTAO**, **SCAO**, **noAO**, **Airy**. **LTAO** corresponds to the Laser Tomography Adaptive Optics mode, **SCAO** to the Single Conjugate Adaptive Optics mode, **noAO** indicates that no adaptive optics corrections are applied, and **Airy** uses the diffraction limit of the ELT.
14. Detector systematics: Indicates if the detector systematics are simulated. It only affects the observed outputs (see Section 1.2.6). Available values are **True**, **False**.

1.2.6 Output Files

HSIM stores the results of the simulation in the specified output directory (the default output directory is `output_cubes`). The name of the output files begins with the name of the input data cube and it is followed by a suffix indicating the content of the file. The input parameters of the simulation are stored in the header of the FITS files and also in the log file.

1. `_noiseless_obj_plus_back.fits`. Data cube containing the expected number of electrons per pixel detected by HARMONI due to the target and the background (sky, telescope, and instrument) emissions. Cross-talk is also applied to the noiseless outputs to match the LSF of the observed and reduced outputs cubes.
2. `_noiseless_obj.fits`. Expected number of electrons per pixel solely due to the target.
3. `_noiseless_back.fits`. Expected number of electrons per pixel from the background sources (sky, telescope, and instrument).
4. `_observed_obj_plus_back.fits`. Simulated observed data cube taking into account the photon noise, read noise, dark current, and cross-talk.
5. `_observed_back.fits`. Simulated observed sky background taking into account the photon noise, read noise, dark current, and cross-talk.
6. `_reduced.fits`. Simulated reduced data cube created by subtracting a simulated sky observation to the `_observed_obj_plus_back.fits` data cube.
7. `_reduced.SNR.fits`. Signal to noise ratio of the reduced data cube per pixel.
8. `_dets.fits`. If detector systematics are used, a fits file containing the detectors used in the simulation.
9. `_std.fits`. Noise standard deviation. Includes object and sky noise (Poisson) and dark current and read noise (Gaussian).
10. `_PSF.fits`. Point spread function at the mean wavelength of the input data cube. The PSF is oversampled by a factor of 4 to 10 with respect to the other output FITS data cubes.
11. `_total.em.pdf`. Plot of the background emission broken up into the individual modeled components. The average HARMONI to telescope+sky background ratio is indicated.

12. `_total_tr.pdf`. Plot of system transmission broken up into the individual modeled components.
13. `_total_tr.txt`. Total transmission stored as a text file.
14. `.log`. Detailed log of the simulation. The last line of this file shows if any problem was found during the simulation.