

ELT - MICADO

SpecCADO User Manual

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Micado
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1 Scope

This document is a brief manual to get users started with SpecCADO, the spectroscopic simulator for MICADO. SpecCADO is a temporary package, intended to be merged into SimCADO, the general instrument data simulator for MICADO in the near future.

SpecCADO depends on SimCADO. For an in-depth description of SimCADO, we refer to the PDR documentation [1].

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2 Installation

Both SimCADO and SpecCADO use Python 3 (preferably 3.5 or higher, although 3.4 should work, too). If you are unsure which version of Python you have, try

```
python --version
```

If this returns "Python 2.7.9" or another version below 3, please replace the command python with python3 and pip with pip3 in the following.

2.1 Install SimCADO

SpecCADO relies on a prior installation of SimCADO. This can be achieved easily with

```
pip install [--user] [--upgrade] simcado
```

Use option --user to install into a directory that belongs to you (without this option you may need root permissions). Use option --upgrade to upgrade an existing SimCADO installation.

SimCADO and SpecCADO require a number of files that describe the telescope and instrument setup. The easiest way to obtain these is to download them using SimCADO itself. Start an interactive python (or preferably ipython session) and type:

```
In 1:    import simcado
    simcado.get_extras()
```

2.2 Install SpecCADO

The SpecCADO source package can be downloaded from https://homepage.univie.ac.at/oliver.czoske/speccado-0.1.4.tar.gz. Unpack the tar file and change to the source directory:

```
tar xvf speccado-0.1.4.tar.gz cd speccado-0.1.4/
```

It is also possible to obtain SpecCADO from the github repository at https://github.com/oczoske/SpecCADO. Note that the latest snapshots from the repository may not always be perfectly functional.

Install the package by doing

```
pip install .
```

or

```
pip install --user .
```

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To test the installation, start a python or ipython session and do

```
In 2:
         import speccado
In 3:
         speccado.__version__
Out3:
         0.1.4
In 4:
         speccado.bug_report()
         Python:
         3.6.0 (default, Feb 6 2017, 16:31:05)
         [GCC 4.9.2]
         speccado : 0.1.4dev1
         simcado : 0.6dev0
         astropy : 3.1.2
         synphot: 0.1.3
        numpy: 1.16.2
         scipy : 1.2.1
         yaml: 5.1
        poppy : 0.8.0
         wget : 3.2
         Operating system: Linux
                 Release: 4.9.0-0.bpo.8-amd64
                 Version: #1 SMP Debian 4.9.110-3+deb9u5~deb8u1 (2018-10-03)
                 Machine: x86_64
```

The last command checks that the dependencies are fulfilled. Please include the output of this command whenever you report a problem or a possible bug.

The necessary dependencies to run speccado are simcado, astropy, synphot, numpy, scipy and yaml. wget is only required for simcado.get_extras() (Sect. 2.1), poppy is not needed.

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3 Example scripts

The subdirectory example/ in the SpecCADO source distribution includes two example scripts that demonstrate how SpecCADO works:

- simulate_example.py simulates spectral traces on the MICADO detectors. The slit contains two stars, both of which have the same spectrum, viz. that of GW Ori, dimmed by 9 magnitudes to put it into the usual target range of MICADO.¹ Atmospheric emission is included as a background spectrum that fills the entire slit. The spectrum provided in atmo_emission.fits was computed by skycalc.²
- rectify_example.py takes a simulated detector frame as input and recifies it into two-dimensional $\xi \lambda$ images (ξ is the coordinate along the slit) from which one-dimensional spectra can be extracted in the usual way (e.g. using apextract in Iraf). Note that this is a perfect rectification that reverses the transformation applied by simulate_example.py. It does not include the uncertainties associated to tracing the spectra in a proper data reduction recipe.

The example/ directory includes three configuration files set up for MICADO spectroscopy using the IzJ, J and HK order sorting filters, respectively. If you run your own simulations, start from one of these configuration files and modify it according to your needs.

The interface for simulate_example.py is³

```
python simulate_example.py [-c chipno] [-s /to/data/dir/] configfile
```

configfile is the name of the configuration file; this is required.

The option -c chipno (or --chip=chipno) can be used to simulate a single detector instead of the full MI-CADO focal-plane array. Replace chipno with the chip number as defined in Fig. 1 (Sect. 6).

The option -s (or --simdatadir=) can be used to point to a directory where the data used by SimCADO are stored. This should not be necessary if you ran simcado.get_extras() as described in Sect. 2.1,

As an example, to simulate an HK spectrum of the sources as described above, type the following at the shell prompt:

```
python simulate_example.py spectro_HK.config
```

It may take about 20 minutes to simulate the nine detectors of the MICADO focal-plane array. SpecCADO writes a stream of diagnostics to the screen – if all goes well these should not be of interest to anyone except the developer.

The output file is a multi-extension FITS file (one extension per detector), named detector-<dateTtime>.fits with the time stamp of the end of the simulation, for example

```
detector-2019-02-14T17-57-20.fits
```

¹No claim of scientific realism is made for this example...

²https://www.eso.org/observing/etc/skycalc/

³If simulate_example.py is executable, then python is not needed.

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To rectify the spectra run

python rectify_example.py detector-2019-02-14T17-57-20.fits

This currently creates a separate FITS file for each order that is defined over the entire MICADO wavelength range. Due to the order sorting filter employed in the simulation, this means that most of the output files will not contain any signal. The file names are ORDER-xx_yy.fits with the numbers identifying the orders and cross-orders.

The output of rectify_example.py will be improved.

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4 Simulation of 2D spectra

In this section, the simulation of detector images will be described in more detail. This should enable the user to specify sources of their own.

4.1 Simulation parameters

Any simulation with SpecCADO or SimCADO starts by loading the configuration file:

```
In 5:     import speccado as sc
     import simcado as sim
     cmds = sim.UserCommands("spectro_HK.config")
```

If your SimCADO data are not stored in the default directory, you need to set the parameter SIM_DATA_DIR in the config file or provide the correct path as a second argument to sim.UserCommands().

It is recommended to edit the configuration file to set up the parameters for your simulation. However, it is also possible to set individual parameters separately, such as the exposure time:

```
In 6: cmds['OBS_EXPTIME'] = 60
```

This may be useful if you investigate the effect of changing a single parameter on the performance of the instrument.

For the simulation of science cases, it is often helpful to turn off detector saturation:

```
In 7: cmds['FPA_LINEARITY_CURVE'] = 'none'
```

as this makes it possible to simulate long effective exposure times without the need to break the integration up into several sub-exposures to avoid saturation.

4.2 Setting up sources and backgrounds

Currently, SpecCADO only simulates point sources (represented by the PSF) and background sources (that fill the slit homogeneously).

For point sources, two lists need to be defined, one giving the files holding the 1D spectra and one giving the positions of the sources in the slit. The following example assumes two stars are positioned on the centre line of the slit, both have the same spectrum:

```
In 8: specfiles = ['GW_Ori+9mag.fits', 'GW_Ori+9mag.fits']
sourcepos = [[-1, 0], [1, 0]]
```

The stars are positioned 1 arcsec on either side of the centre of the 3 arcsec slit ($\xi = \pm 1$, $\eta = 0$, as defined in Sect. 6.3).

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Another list gives the background spectra. Here, only one spectrum for atmospheric emission is provided:

```
In 9: bgfiles = ['atmo_emission.fits']
```

4.3 Running the simulation

The simulation can now be run with

```
In 10: outfile = sc.simulate(cmds, specfiles, sourcepos, bgfiles)
```

to simulate the entire focal-plane array or

```
In 11: outfile = sc.simulate(cmds, specfiles, sourcepos, bgfiles, chipno)
```

to simulate a single chip as numbered in Fig. 1. Both commands write their results to disk as a FITS file. The filename is returned and can be captured in a variable (here outfile) to be processed further in a script.

The following steps are performed by sc.simulate() and are described in more detail to gain an understanding of how a simulation is performed.

A SpectralSource object is created by

```
In 12: srcobj = sc.SpectralSource(cmds, specfiles, sourcepos, bgfiles)
```

The simulation further requires objects for the PSF and the detector array:

```
In 13: psfobj = sc.prepare_psf(cmds['SCOPE_PSF_FILE'])
    detector = sim.Detector(cmds, small_fov=False)
```

For the detector, we use the Detector class from SimCADO. The PSF is a slightly enhanced form of the SimCADO PSF class, created through a SpecCADO function.

The transmission of the optical system (telescope plus instrument) is extracted from SimCADO's OpticalTrain class and stored as an interpolation object that can be evaluated at any wavelength required for the simulation:⁴

The optical layout of the spectral traces in the detector focal plane is described in a FITS file that holds a number of table extensions, each describing one spectral order.⁵. It is loaded by

⁴The transmission curve tc_source describes the transmission as seen by the astronomical source, including atmosphere, telescope, instrument mirrors and filters. If your goal is to simulate calibration spectra obtained from a calibration source within the instrumen, a decent approximation to the transmission is give by tc_mirror.

⁵The most up to date description is specorders-180629.fits, based on data provided by Frank Grupp.

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```
In 15: tracelist = sc.layout.read_spec_order(cmds['SPEC_ORDER_LAYOUT'])
```

The simulation of the full detector array is then run by

The result is written to disk as the multi-extension FITS file detector-YYYY-MM-DDThh-mm-ss.fits. The file name is returned by sc.do_all_chips; you can use this in scripts to rename files according to your needs.

It is also possible to simulate just a single detector from the MICADO array:

This simulates chip number 4 of the MICADO array (as python arrays are zero-offset). The output file in this case is called chip-YYYY-MM-DDThh-mm-ss.fits and the name is again returned by the function.

⁶This differs from the interface to the convenience script simulate_example.py described in Sect. 3.

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5 Input file format

SpecCADO needs to be fully informed about the contents of the input data. It is therefore necessary for a successful simulation that input files conform to the following requirements.

5.1 Spectra

Input spectra are expected to be provided as one-dimensional FITS images (NAXIS = 1).⁷ For the transformation from pixels to wavelengths, a WCS is required in a format that can be read by the astropy.wcs module; this includes most formats defined by Greisen et al. (2006) [2]. To keep things simple, spectra sampled on a linear wavelength grid (CTYPE1='WAVE') are recommended. However, frequency or wave number grids, linear or non-linear, should be possible if the WCS is constructed correctly.

As an example, the WCS of the file GW_Ori+9mag.fits used in Sect. 4 is

```
NAXIS
                              1 / number of array dimensions
NAXIS1
                          24750
                              1 / Number of coordinate axes
WCSAXES =
CRPIX1
                            1.0 / Pixel coordinate of reference point
                         6E-11 / [m] Coordinate increment at reference point
CDELT1
CUNIT1
                                / Units of coordinate increment and value
        = 'm'
                                / Vacuum wavelength (linear)
CTYPE1
        = 'WAVE'
                    9.9402E-07 / [m] Coordinate value at reference point
CRVAL1
```

The pixel values have to be in physical units and the units have to be provided explicitly in the BUNIT keyword. SpecCADO uses the very flexible astropy.units module to convert units:

Input spectra for point sources can be given as photon or energy fluxes; internally, the units are converted to photons/s/m²/ μ m. Permitted values for BUNIT include

```
• 'erg / (Angstrom cm2 s)'
```

- 'ph / (um m2 s)'
- '1 / (nm cm2 s)'

Input spectra for background sources can be given as photon or energy flux densities; internally, the units are converted to photons/s/m²/ μ m/arcsec². Permitted values for BUNIT include

- 'ph / (s m2 micron arcsec2)'
- 'erg / (s cm2 Angstrom arcmin2)'

⁷This is admittedly rather restrictive. We hope to make SpecCADO a bit more flexible in the future.

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5.2 PSFs

The PSF used to place point sources on the slit is provided as a FITS image in a format that can be read by the SimCADO psf module.

The FITS header needs to specify the (effective) wavelength at which the PSF applies in keyword WAVEO or WAVELENG. SimCADO requires the units to be μm .

In addition, the size of the pixels in the PSF image is required. This can be given by the keyword PIXSCALE or the WCS keywords CDELT1 or CD1_1 (for the PSF a WCS with CTYPEi='LINEAR' is adequate). The units are arcsec.

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6 Coordinate systems

6.1 Detector coordinates

Detector pixels are characterised by coordinates (c, i, j), where c is the number of the chip in the MICADO focal plane array (c = 1, ..., 9); see Fig. 1) and i and j are the column and row number of a pixel (i, j = 1, ..., 4096).

6.2 Focal plane coordinates

A position in the focal plane of MICADO is characterised by the coordinate pair (x, y), given in millimeters. The origin x = 0, y = 0 is taken to coincide with the centre of detector 5, i.e. pixel coordinates i = 2048.5, j = 2048.5. The x coordinate increases along rows of the chips (increasing i), the y coordinate increases along columns of the chips (increasing j). Small rotations of the detectors with respect to the focal plane coordinates are allowed.

The locations of the detector chips within the focal plane are specified in a focal-plane array definition file that specifies for each chip the identification number id; the position of the chip centre x_cen , y_cen ; its half width xhw, yhw; the number of pixels in each direction x_len , y_len ; the physical pixel size pixsize; a rotation angle of the detector rows with respect to the x-axis, angle; and the detector gain. Listing 1 shows the focal-plane array definition file for MICADO.

The transformation between pixel coordinates and focal-plane coordinates is a linear transformation. SimCADO

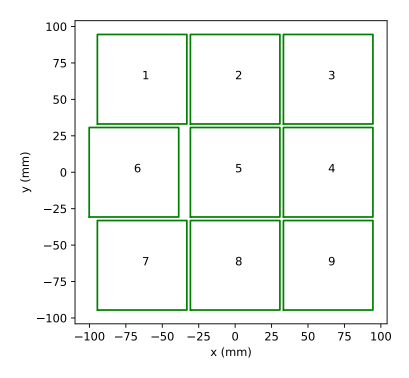


Figure 1: Numbering convention of the detectors in the MICADO focal plane.

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											-
##	MIC	CADO H4R	G-15 FI	PA							
	id	x_cen	y_cen	xhw	yhw	x_len	y_len	pixsize	angle	gain	
#		mm	mm	mm	mm	pix	pix	mm	deg	e-/ADU	
	1	-63.84	63.84	30.72	30.72	4096	4096	0.015	0.0	1.0	
	2	0.00	63.84	30.72	30.72	4096	4096	0.015	0.0	1.0	
	3	63.84	63.84	30.72	30.72	4096	4096	0.015	0.0	1.0	
	4	+63.84	0.00	30.72	30.72	4096	4096	0.015	0.0	1.0	
	5	0.00	0.00	30.72	30.72	4096	4096	0.015	0.0	1.0	
	6	-79.50	0.00	30.72	30.72	4096	4096	0.015	0.0	1.0	
	7	-63.84	-63.84	30.72	30.72	4096	4096	0.015	0.0	1.0	
	8	0.00	-63.84	30.72	30.72	4096	4096	0.015	0.0	1.0	
	9	63.84	-63.84	30.72	30.72	4096	4096	0.015	0.0	1.0	

Listing 1: Focal plane array definition file for MICADO with nine HAWAII4RG chips with 15 µm pixels

constructs a world coordinate system (CTYPEiA = 'LINEAR') from the FPA definition file and uses this internally to perform the transformations. The WCS is written to the SpecCADO output files with alternative axis descriptor A and name WCSNAMEA = 'PIX2FP'.

6.3 Spectral cube coordinates

The spectral source is described as a spectroscopic cube with spatial coordinates ξ and η along and across the slit, respectively (Fig. 2), and wavelength λ . Both ξ and η are measured in arcsec, based on the fixed imaging scale of MICADO. The along-slit coordinate ξ always starts at -1.5 arcsec. The length of the slit if set by the keyword SPEC_SLIT_LENGTH in the configuration file, meaning that the maximum value of ξ is at SPEC_SLIT_LENGTH -1.5 arcsec. Thus, for the short slit of MICADO (length 3 arcsec), ξ runs from -1.5 arcsec to 1.5 arcsec, whereas for the long slit (15 arcsec), ξ runs from -1.5 arcsec to 13.5 arcsec. The slit width is set by SPEC_SLIT_WIDTH, the centre of the slit always being at $\eta=0$.

The mapping between the centre line of the slit $(\xi, \eta = 0, \lambda)$ and focal-plane coordinates (x, y) for a given spectral order s is characterised by order-definition files provided by Frank Grupp. These files were produced using ray-tracing with Zemax and list matching cube and focal-plane coordinates for a number of points. SpecCADO models the transformations as fourth-order polynomials fitted to the order definition files:

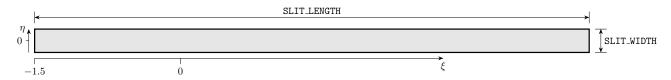


Figure 2: Slit coordinates ξ , η along and across the slit, respectively, along with the slit dimensions SPEC_SLIT_LENGTH and SPEC_SLIT_WIDTH.

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$$x(\xi,\lambda) = \sum_{i,j=0}^{4} A_{s,ij} \xi^{i} \lambda^{j}$$
 (1)

$$y(\xi,\lambda) = \sum_{i,j=0}^{4} B_{s,ij} \xi^{i} \lambda^{j}$$
 (2)

$$\xi(x,y) = \sum_{i,j=0}^{4} C_{s,ij} x^{i} y^{j}$$
 (3)

$$\lambda(x,y) = \sum_{i,j=0}^{4} D_{s,ij} x^i y^j \tag{4}$$

The across-slit coordinate η is integrated out before the spectral cube is mapped to the focal plane as illustrated in Fig. 3. A monochromatic slice $\lambda = \lambda_0$ of the spectral cube is mapped to a rectangle in the focal plane. Given a unique mapping of focal-plane coordinates to wavelength (the wavelength calibration) this means that, say, the top of the slit will be at an offset wavelength, $\lambda_0 + \Delta \lambda$. If the slit width (given by the SimCADO keyword SPEC_SLIT_WIDTH) is b and the plate scale is p (in mas μ m⁻¹), then the wavelength shift is

$$\Delta \lambda = \frac{\partial \lambda}{\partial y} \Delta y = \frac{b}{2} \frac{1}{p} \left[\frac{\partial y}{\partial \lambda} (\xi, \lambda) \right]^{-1}$$
 (5)

Each plane ($\eta = \text{const.}$) of the spectral cube is shifted by the appropriate amount in the wavelength direction. The resulting sheared cube is then summed in the η direction to give a two-dimensional spectrum with the

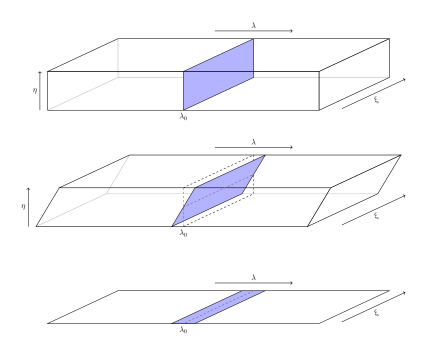


Figure 3: The spectral cube (ξ, η, λ) , top, is sheared in the wavelength direction to take into account the slit profile, middle, and then summed in the η direction to form the rectified two-dimensional spectrum (ξ, λ) , bottom, which is subsequently mapped onto the focal plane.

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spectral lines automatically broadened by the slit profile. The transformations (1) and (2) are then applied to map the 2D spectrum into the focal plane and onto the detectors.

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7 Spectral layout

The panels in Fig. 4 show the spectral traces for the 3 arcsec slit and the three order sorting filters, IzJ, J and HK. Fig. 5 identifies the orders as a function of wavelength.

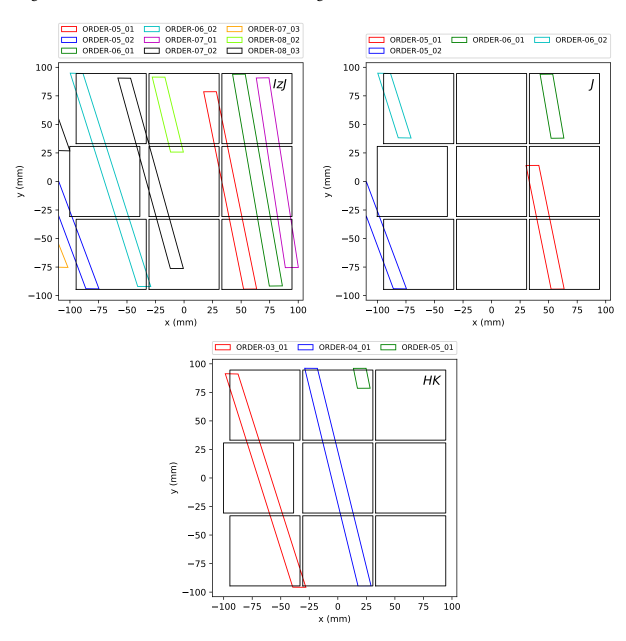


Figure 4: Spectral layout based on the PDR design. The slit has a width of 3 arcsec on the sky.

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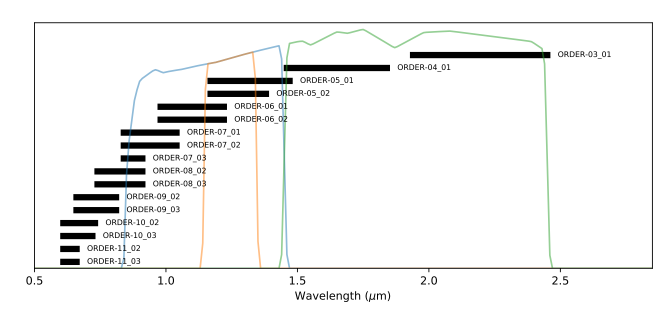


Figure 5: Wavelength ranges covered by the spectral orders. The blue, orange and green curves give the relative system transmissivities for the IzJ, J and HK order sorting filters, respectively.

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- [2] Greisen, E. W., Calabretta, M. R., Valdes, F. G., and Allen, S. L., "Representations of spectral coordinates in fits," *Astronomy & Astrophysics* **446**, 747–771 (2006). **13**