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# **Codex Africanus Documentation**

***Release 0.3.2***

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**CHAPTER  
ONE**

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**CODEX AFRICANUS**

Radio Astronomy Building Blocks

## **1.1 Documentation**

<https://codex-africanus.readthedocs.io>.



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## CHAPTER TWO

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# INSTALLATION

## 2.1 Stable release

To install Codex Africanus, run this command in your terminal:

```
$ pip install codex-africanus
```

This is the preferred method to install Codex Africanus, as it will always install the most recent stable release.

If you don't have `pip` installed, this [Python installation guide](#) can guide you through the process.

By default, Codex Africanus will install with a minimal set of dependencies, `numpy` and `numba`.

Further functionality can be enabled by installing extra requirements as follows:

```
$ pip install codex-africanus[dask]
$ pip install codex-africanus[scipy]
$ pip install codex-africanus[astropy]
$ pip install codex-africanus[python-casacore]
```

To install the complete set of dependencies for the CPU:

```
$ pip install codex-africanus[complete]
```

To install the complete set of dependencies including CUDA:

```
$ pip install codex-africanus[complete-cuda]
```

## 2.2 From sources

The sources for Codex Africanus can be downloaded from the [Github repo](#).

You can either clone the public repository:

```
$ git clone git://github.com/ska-sa/codex-africanus
```

Or download the [tarball](#):

```
$ curl -OL https://github.com/ska-sa/codex-africanus/tarball/master
```

Once you have a copy of the source, you can install it with:

```
$ python setup.py install
```

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**CHAPTER  
THREE**

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**USAGE**

To use Codex Africanus in a project:

```
import africanus
```



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CHAPTER  
**FOUR**

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## **COMMAND LINE UTILITIES**

The following command line utilities are installed. Run each utility's help for further information.

```
$ utility --help
```

### **4.1 plot-filter**

Plots convolution filters.

### **4.2 plot-taper**

Plots tapers associated with convolution filters.



## 5.1 Radio Interferometer Measurement Equation

Functions used to compute the terms of the Radio Interferometer Measurement Equation (RIME). It describes the response of an interferometer to a sky model.

$$V_{pq} = G_p \left( \sum_s E_{ps} L_p K_{ps} B_s K_{qs}^H L_q^H E_{qs}^H \right) G_q^H$$

where for antenna  $p$  and  $q$ , and source  $s$ :

- $G_p$  represents direction-independent effects.
- $E_{ps}$  represents direction-dependent effects.
- $L_p$  represents the feed rotation.
- $K_{ps}$  represents the phase delay term.
- $B_s$  represents the brightness matrix.

The RIME is more formally described in the following four papers:

- I. A full-sky Jones formalism
- II. Calibration and direction-dependent effects
- III. Addressing direction-dependent effects in 21cm WSRT observations of 3C147
- IV. A generalized tensor formalism

### 5.1.1 Numpy

<code>predict_vis(time_index, antenna1, antenna2)</code>	Multiply Jones terms together to form model visibilities according to the following formula:
<code>phase_delay(lm, uvw, frequency[, convention])</code>	Computes the phase delay (K) term:
<code>parallactic_angles(times, antenna_positions, ...)</code>	Computes parallactic angles per timestep for the given reference antenna position and field centre.
<code>feed_rotation(parallactic_angles[, feed_type])</code>	Computes the 2x2 feed rotation (L) matrix from the <code>parallactic_angles</code> .
<code>transform_sources(lm, parallactic_angles, ...)</code>	Creates beam sampling coordinates suitable for use in <code>beam_cube_dde()</code> by:

continues on next page

Table 1 – continued from previous page

<code>beam_cube_dde(beam, beam_lm_extents, ...)</code>	Evaluates Direction Dependent Effects along a source's path by interpolating the values of a complex beam cube at the source location.
<code>zernike_dde(coords, coeffs, noll_index, ...)</code>	Computes Direction Dependent Effects by evaluating Zernicke Polynomials defined by coefficients <code>coeffs</code> and noll indexes <code>noll_index</code> at the specified coordinates <code>coords</code> .
<code>wsclean_predict(uvw, lm, source_type, flux, ...)</code>	Predict visibilities from a WSClean sky model.

`africanus.rime.predict_vis(time_index, antenna1, antenna2, dde1_jones=None, source_coh=None, dde2_jones=None, die1_jones=None, base_vis=None, die2_jones=None)`

Multiply Jones terms together to form model visibilities according to the following formula:

$$V_{pq} = G_p \left( B_{pq} + \sum_s E_{ps} X_{pqs} E_{qs}^H \right) G_q^H$$

where for antenna  $p$  and  $q$ , and source  $s$ :

- $B_{pq}$  represent base coherencies.
- $E_{ps}$  represents Direction-Dependent Jones terms.
- $X_{pqs}$  represents a coherency matrix (per-source).
- $G_p$  represents Direction-Independent Jones terms.

Generally,  $E_{ps}$ ,  $G_p$ ,  $X_{pqs}$  should be formed by using the [RIME API](#) functions and combining them together with `einsum()`.

## Please read the Notes

### Parameters

**time\_index** [numpy.ndarray] Time index used to look up the antenna Jones index for a particular baseline with shape (row,). Obtainable via `np.unique(time, return_inverse=True)[1]`.

**antenna1** [numpy.ndarray] Antenna 1 index used to look up the antenna Jones for a particular baseline. with shape (row,).

**antenna2** [numpy.ndarray] Antenna 2 index used to look up the antenna Jones for a particular baseline. with shape (row,).

**dde1\_jones** [numpy.ndarray, optional]  $E_{ps}$  Direction-Dependent Jones terms for the first antenna. shape (source, time, ant, chan, corr\_1, corr\_2)

**source\_coh** [numpy.ndarray, optional]  $X_{pqs}$  Direction-Dependent Coherency matrix for the baseline. with shape (source, row, chan, corr\_1, corr\_2)

**dde2\_jones** [numpy.ndarray, optional]  $E_{qs}$  Direction-Dependent Jones terms for the second antenna. This is usually the same array as `dde1_jones` as this preserves the symmetry of the RIME. `predict_vis` will perform the conjugate transpose internally. shape (source, time, ant, chan, corr\_1, corr\_2)

**die1\_jones** [numpy.ndarray, optional]  $G_p$  Direction-Independent Jones terms for the first antenna of the baseline. with shape (time, ant, chan, corr\_1, corr\_2)

**base\_vis** [numpy.ndarray, optional]  $B_{pq}$  base coherencies, added to source coherency summation before multiplication with `die1_jones` and `die2_jones`. shape (row, chan, corr\_1, corr\_2).

**die2\_jones** [numpy.ndarray, optional]  $G_{ps}$  Direction-Independent Jones terms for the second antenna of the baseline. This is usually the same array as `die1_jones` as this preserves the symmetry of the RIME. `predict_vis` will perform the conjugate transpose internally.  
shape (time, ant, chan, corr\_1, corr\_2)

#### Returns

**visibilities** [numpy.ndarray] Model visibilities of shape (row, chan, corr\_1, corr\_2)

#### Notes

- Direction-Dependent terms (`dde{1,2}_jones`) and Independent (`die{1,2}_jones`) are optional, but if one is present, the other must be present.
- The inputs to this function involve `row`, `time` and `ant` (antenna) dimensions.
- Each `row` is associated with a pair of antenna Jones matrices at a particular timestep via the `time_index`, `antenna1` and `antenna2` inputs.
- The `row` dimension must be an increasing partial order in time.

`africanus.rime.phase_delay(lm, uvw, frequency, convention='fourier')`

Computes the phase delay (K) term:

$$e^{-2\pi i(ul+vm+w(n-1))}$$

where  $n = \sqrt{1 - l^2 - m^2}$

#### Parameters

**lm** [numpy.ndarray] LM coordinates of shape (source, 2) with L and M components in the last dimension.

**uvw** [numpy.ndarray] UVW coordinates of shape (row, 3) with U, V and W components in the last dimension.

**frequency** [numpy.ndarray] frequencies of shape (chan, )

**convention** [{‘fourier’, ‘casa’}] Uses the  $e^{-2\pi i}$  sign convention if `fourier` and  $e^{2\pi i}$  if `casa`.

#### Returns

**complex\_phase** [numpy.ndarray] complex of shape (source, row, chan)

#### Notes

Corresponds to the complex exponential of the Van Cittert-Zernike Theorem.

`MeqTrees` uses the CASA sign convention.

`africanus.rime.parallactic_angles(times, antenna_positions, field_centre, backend='casa')`

Computes parallactic angles per timestep for the given reference antenna position and field centre.

#### Parameters

**times** [numpy.ndarray] Array of Mean Julian Date times in *seconds* with shape (time,),

**antenna\_positions** [numpy.ndarray] Antenna positions of shape (ant, 3) in *metres* in the *ITRF* frame.

**field\_centre** [numpy.ndarray] Field centre of shape (2,) in *radians*

**backend** [{‘casa’, ‘test’}, optional] Backend to use for calculating the parallactic angles.

- `casa` defers to an implementation depending on `python-casacore`. This backend should be used by default.
- `test` creates parallactic angles by multiplying the `times` and `antenna_position` arrays. It exists solely for testing.

**Returns**

`parallactic_angles` [numpy.ndarray] Parallactic angles of shape (time, ant)

`africanus.rime.feed_rotation(parallactic_angles, feed_type='linear')`

Computes the 2x2 feed rotation (L) matrix from the `parallactic_angles`.

$$\text{linear} \begin{bmatrix} \cos(pa) & \sin(pa) \\ -\sin(pa) & \cos(pa) \end{bmatrix} \quad \text{circular} \begin{bmatrix} e^{-ipa} & 0 \\ 0 & e^{ipa} \end{bmatrix}$$

**Parameters**

`parallactic_angles` [numpy.ndarray] floating point parallactic angles. Of shape (pa0, pa1, ..., pan).

`feed_type` [{‘linear’, ‘circular’}] The type of feed

**Returns**

`feed_matrix` [numpy.ndarray] Feed rotation matrix of shape (pa0, pa1, ..., pan, 2, 2)

`africanus.rime.transform_sources(lm, parallactic_angles, pointing_errors, antenna_scaling, frequency, dtype=None)`

Creates beam sampling coordinates suitable for use in `beam_cube_dde()` by:

1. Rotating `lm` coordinates by the `parallactic_angles`
2. Adding `pointing_errors`
3. Scaling by `antenna_scaling`

**Parameters**

`lm` [numpy.ndarray] LM coordinates of shape (src, 2) in radians offset from the phase centre.

`parallactic_angles` [numpy.ndarray] parallactic angles of shape (time, antenna) in radians.

`pointing_errors` [numpy.ndarray] LM pointing errors for each antenna at each timestep in radians. Has shape (time, antenna, 2)

`antenna_scaling` [numpy.ndarray] antenna scaling factor for each channel and each antenna. Has shape (antenna, chan)

`frequency` [numpy.ndarray] frequencies for each channel. Has shape (chan,)

`dtype` [numpy.dtype, optional] Numpy dtype of result array. Should be float32 or float64. Defaults to float64

**Returns**

`coords` [numpy.ndarray] coordinates of shape (3, src, time, antenna, chan) where each coordinate component represents `l`, `m` and `frequency`, respectively.

`africanus.rime.beam_cube_dde(beam, beam_lm_extents, beam_freq_map, lm, parallactic_angles, point_errors, antenna_scaling, frequency)`

Evaluates Direction Dependent Effects along a source’s path by interpolating the values of a complex beam cube at the source location.

## Parameters

**beam** [numpy.ndarray] Complex beam cube of shape (beam\_lw, beam\_mh, beam\_nud, corr, corr). *beam\_lw*, *beam\_mh* and *beam\_nud* define the size of the cube in the l, m and frequency dimensions, respectively.

**beam\_lm\_extents** [numpy.ndarray] lm extents of the beam cube of shape (2, 2). [[lower\_l, upper\_l], [lower\_m, upper\_m]].

**beam\_freq\_map** [numpy.ndarray] Beam frequency map of shape (beam\_nud, ). This array is used to define interpolation along the (chan,) dimension.

**lm** [numpy.ndarray] Source lm coordinates of shape (source, 2). These coordinates are:

1. Scaled if the associated frequency lies outside the beam cube.
2. Offset by pointing errors: **point\_errors**
3. Rotated by parallactic angles: **parallactic\_angles**.
4. Scaled by antenna scaling factors: **antenna\_scaling**.

**parallactic\_angles** [numpy.ndarray] Parallactic angles of shape (time, ant).

**point\_errors** [numpy.ndarray] Pointing errors of shape (time, ant, chan, 2).

**antenna\_scaling** [numpy.ndarray] Antenna scaling factors of shape (ant, chan, 2)

**frequency** [numpy.ndarray] Frequencies of shape (chan, ).

## Returns

**ddes** [numpy.ndarray] Direction Dependent Effects of shape (source, time, ant, chan, corr, corr)

## Notes

1. Sources are clamped to the provided *beam\_lm\_extents*.
2. Frequencies outside the cube (i.e. outside *beam\_freq\_map*) introduce linear scaling to the lm coordinates of a source.

africanus.rime.zernike\_dde(coords, coeffs, noll\_index, parallactic\_angles, frequency\_scaling, antenna\_scaling, pointing\_errors)

Computes Direction Dependent Effects by evaluating Zernicke Polynomials defined by coefficients **coeffs** and noll indexes **noll\_index** at the specified coordinates **coords**.

Decomposition of a voxel beam cube into Zernicke polynomial coefficients can be achieved through the use of the **eidos** package.

## Parameters

**coords** [numpy.ndarray] Float coordinates at which to evaluate the zernike polynomials. Has shape (3, source, time, ant, chan). The three components in the first dimension represent l, m and frequency coordinates, respectively.

**coeffs** [numpy.ndarray] complex Zernicke polynomial coefficients. Has shape (ant, chan, corr\_1, ..., corr\_n, poly) where poly is the number of polynomial coefficients and corr\_1, ..., corr\_n are a variable number of correlation dimensions.

**noll\_index** [numpy.ndarray] Noll index associated with each polynomial coefficient. Has shape (ant, chan, corr\_1, ..., corr\_n, poly). correlation dimensions.

**parallactic\_angles** [numpy.ndarray] Parallactic angle rotation. Has shape (time, ant).

**frequency\_scaling** [numpy.ndarray] The scaling of frequency of the beam. Has shape (chan, ).

**antenna\_scaling** [numpy.ndarray] The antenna scaling. Has shape (ant, chan, 2).

**pointing\_errors** [numpy.ndarray] The pointing error. Has shape (time, ant, chan, 2).

#### Returns

**dde** [numpy.ndarray] complex values with shape (source, time, ant, chan, corr\_1, ..., corr\_n)

africanus.rime.wsclean\_predict(*uvw*, *lm*, *source\_type*, *flux*, *coeffs*, *log\_poly*, *ref\_freq*, *gauss\_shape*, *frequency*)

Predict visibilities from a WSClean sky model.

#### Parameters

**uvw** [numpy.ndarray] UVW coordinates of shape (row, 3)

**lm** [numpy.ndarray] Source LM coordinates of shape (source, 2), in radians. Derived from the Ra and Dec fields.

**source\_type** [numpy.ndarray] Strings defining the source type of shape (source,). Should be either "POINT" or "GAUSSIAN". Contains the Type field.

**flux** [numpy.ndarray] Source flux of shape (source,). Contains the I field.

**coeffs** [numpy.ndarray] Source Polynomial coefficients of shape (source, coeffs). Contains the SpectralIndex field.

**log\_poly** [numpy.ndarray] Source polynomial type of shape (source,). If True, logarithmic polynomials are used. If False, standard polynomials are used. Contains the LogarithmicSI field.

**ref\_freq** [numpy.ndarray] Source Reference frequency of shape (source,). Contains the ReferenceFrequency field.

**gauss\_shape** [numpy.ndarray] Gaussian shape parameters of shape (source, 3) used when the corresponding source\_type is "GAUSSIAN". The 3 components should contain the MajorAxis, MinorAxis and Orientation fields in radians, respectively.

**frequency** [numpy.ndarray] Frequency of shape (chan,).

#### Returns

**visibilities** [numpy.ndarray] Complex visibilities of shape (row, chan, 1)

## 5.1.2 Cuda

<code><i>predict_vis</i>(time_index, antenna1, antenna2)</code>	Multiply Jones terms together to form model visibilities according to the following formula:
<code><i>phase_delay</i>(lm, uvw, frequency)</code>	Computes the phase delay (K) term:
<code><i>feed_rotation</i>(parallactic_angles[, feed_type])</code>	Computes the 2x2 feed rotation (L) matrix from the parallactic_angles.
<code><i>beam_cube_dde</i>(beam, beam_lm_ext, ...)</code>	Evaluates Direction Dependent Effects along a source's path by interpolating the values of a complex beam cube at the source location.

---

```
africanus.rime.cuda.predict_vis(time_index, antenna1, antenna2, dde1_jones=None, source_coh=None,
                                 dde2_jones=None, die1_jones=None, base_vis=None, die2_jones=None)
```

Multiply Jones terms together to form model visibilities according to the following formula:

$$V_{pq} = G_p \left( B_{pq} + \sum_s E_{ps} X_{pqs} E_{qs}^H \right) G_q^H$$

where for antenna  $p$  and  $q$ , and source  $s$ :

- $B_{pq}$  represent base coherencies.
- $E_{ps}$  represents Direction-Dependent Jones terms.
- $X_{pqs}$  represents a coherency matrix (per-source).
- $G_p$  represents Direction-Independent Jones terms.

Generally,  $E_{ps}$ ,  $G_p$ ,  $X_{pqs}$  should be formed by using the [RIME API](#) functions and combining them together with `einsum()`.

## Please read the Notes

### Parameters

**time\_index** [cupy.ndarray] Time index used to look up the antenna Jones index for a particular baseline with shape (row,). Obtainable via `cp.unique(time, return_inverse=True)[1]`.

**antenna1** [cupy.ndarray] Antenna 1 index used to look up the antenna Jones for a particular baseline. with shape (row,).

**antenna2** [cupy.ndarray] Antenna 2 index used to look up the antenna Jones for a particular baseline. with shape (row,).

**dde1\_jones** [cupy.ndarray, optional]  $E_{ps}$  Direction-Dependent Jones terms for the first antenna. shape (source, time, ant, chan, corr\_1, corr\_2)

**source\_coh** [cupy.ndarray, optional]  $X_{pqs}$  Direction-Dependent Coherency matrix for the baseline. with shape (source, row, chan, corr\_1, corr\_2)

**dde2\_jones** [cupy.ndarray, optional]  $E_{qs}$  Direction-Dependent Jones terms for the second antenna. This is usually the same array as `dde1_jones` as this preserves the symmetry of the RIME. `predict_vis` will perform the conjugate transpose internally. shape (source, time, ant, chan, corr\_1, corr\_2)

**die1\_jones** [cupy.ndarray, optional]  $G_{ps}$  Direction-Independent Jones terms for the first antenna of the baseline. with shape (time, ant, chan, corr\_1, corr\_2)

**base\_vis** [cupy.ndarray, optional]  $B_{pq}$  base coherencies, added to source coherency summation before multiplication with `die1_jones` and `die2_jones`. shape (row, chan, corr\_1, corr\_2).

**die2\_jones** [cupy.ndarray, optional]  $G_{qs}$  Direction-Independent Jones terms for the second antenna of the baseline. This is usually the same array as `die1_jones` as this preserves the symmetry of the RIME. `predict_vis` will perform the conjugate transpose internally. shape (time, ant, chan, corr\_1, corr\_2)

### Returns

**visibilities** [cupy.ndarray] Model visibilities of shape (row, chan, corr\_1, corr\_2)

## Notes

- Direction-Dependent terms (`dde{1,2}_jones`) and Independent (`die{1,2}_jones`) are optional, but if one is present, the other must be present.
- The inputs to this function involve `row`, `time` and `ant` (antenna) dimensions.
- Each `row` is associated with a pair of antenna Jones matrices at a particular timestep via the `time_index`, `antenna1` and `antenna2` inputs.
- The `row` dimension must be an increasing partial order in time.

`africanus.rime.cuda.phase_delay(lm, uvw, frequency)`

Computes the phase delay (K) term:

$$e^{-2\pi i(ul+vm+w(n-1))}$$

where  $n = \sqrt{1 - l^2 - m^2}$

## Parameters

**lm** [cupy.ndarray] LM coordinates of shape (`source`, 2) with L and M components in the last dimension.

**uvw** [cupy.ndarray] UVW coordinates of shape (`row`, 3) with U, V and W components in the last dimension.

**frequency** [cupy.ndarray] frequencies of shape (`chan`,

**convention** [{‘fourier’, ‘casa’}] Uses the  $e^{-2\pi i}$  sign convention if `fourier` and  $e^{2\pi i}$  if `casa`.

## Returns

**complex\_phase** [cupy.ndarray] complex of shape (`source`, `row`, `chan`)

## Notes

Corresponds to the complex exponential of the Van Cittert-Zernike Theorem.

MeqTrees uses the CASA sign convention.

`africanus.rime.cuda.feed_rotation(parallactic_angles, feed_type='linear')`

Computes the 2x2 feed rotation (L) matrix from the `parallactic_angles`.

$$\text{linear} \begin{bmatrix} \cos(pa) & \sin(pa) \\ -\sin(pa) & \cos(pa) \end{bmatrix} \quad \text{circular} \begin{bmatrix} e^{-ipa} & 0 \\ 0 & e^{ipa} \end{bmatrix}$$

## Parameters

**parallactic\_angles** [cupy.ndarray] floating point parallactic angles. Of shape (`pa0`, `pa1`, ..., `pan`).

**feed\_type** [{‘linear’, ‘circular’}] The type of feed

## Returns

**feed\_matrix** [cupy.ndarray] Feed rotation matrix of shape (`pa0`, `pa1`, ..., `pan`, 2, 2)

`africanus.rime.cuda.beam_cube_dde(beam, beam_lm_ext, beam_freq_map, lm, parangles, pointing_errors, antenna_scaling, frequencies)`

Evaluates Direction Dependent Effects along a source’s path by interpolating the values of a complex beam cube at the source location.

## Parameters

**beam** [cupy.ndarray] Complex beam cube of shape (beam\_lw, beam\_mh, beam\_nud, corr, corr). *beam\_lw*, *beam\_mh* and *beam\_nud* define the size of the cube in the l, m and frequency dimensions, respectively.

**beam\_lm\_extents** [cupy.ndarray] lm extents of the beam cube of shape (2, 2). [[lower\_l, upper\_l], [lower\_m, upper\_m]].

**beam\_freq\_map** [cupy.ndarray] Beam frequency map of shape (beam\_nud, ). This array is used to define interpolation along the (chan,) dimension.

**lm** [cupy.ndarray] Source lm coordinates of shape (source, 2). These coordinates are:

1. Scaled if the associated frequency lies outside the beam cube.
2. Offset by pointing errors: **point\_errors**
3. Rotated by parallactic angles: **parallactic\_angles**.
4. Scaled by antenna scaling factors: **antenna\_scaling**.

**parallactic\_angles** [cupy.ndarray] Parallactic angles of shape (time, ant).

**point\_errors** [cupy.ndarray] Pointing errors of shape (time, ant, chan, 2).

**antenna\_scaling** [cupy.ndarray] Antenna scaling factors of shape (ant, chan, 2)

**frequency** [cupy.ndarray] Frequencies of shape (chan, ).

## Returns

**ddes** [cupy.ndarray] Direction Dependent Effects of shape (source, time, ant, chan, corr, corr)

## Notes

1. Sources are clamped to the provided *beam\_lm\_extents*.
2. Frequencies outside the cube (i.e. outside *beam\_freq\_map*) introduce linear scaling to the lm coordinates of a source.

### 5.1.3 Dask

<code>predict_vis</code> (time_index, antenna1, antenna2)	Multiply Jones terms together to form model visibilities according to the following formula:
<code>phase_delay</code> (lm, uvw, frequency[, convention])	Computes the phase delay (K) term:
<code>parallactic_angles</code> (times, antenna_positions, ...)	Computes parallactic angles per timestep for the given reference antenna position and field centre.
<code>feed_rotation</code> (parallactic_angles, feed_type)	Computes the 2x2 feed rotation (L) matrix from the <code>parallactic_angles</code> .
<code>transform_sources</code> (lm, parallactic_angles, ...)	Creates beam sampling coordinates suitable for use in <code>beam_cube_dde()</code> by:
<code>beam_cube_dde</code> (beam, beam_lm_extents, ...)	Evaluates Direction Dependent Effects along a source's path by interpolating the values of a complex beam cube at the source location.

continues on next page

Table 3 – continued from previous page

<code>zernike_dde(coords, coeffs, noll_index, ...)</code>	Computes Direction Dependent Effects by evaluating Zernicke Polynomials defined by coefficients <code>coeffs</code> and noll indexes <code>noll_index</code> at the specified coordinates <code>coords</code> .
<code>wsclean_predict(uvw, lm, source_type, flux, ...)</code>	Predict visibilities from a WSClean sky model.

---

```
africanus.rime.dask.predict_vis(time_index, antenna1, antenna2, dde1_jones=None, source_coh=None,
                                  dde2_jones=None, die1_jones=None, base_vis=None, die2_jones=None,
                                  streams=None)
```

Multiply Jones terms together to form model visibilities according to the following formula:

$$V_{pq} = G_p \left( B_{pq} + \sum_s E_{ps} X_{pqs} E_{qs}^H \right) G_q^H$$

where for antenna  $p$  and  $q$ , and source  $s$ :

- $B_{pq}$  represent base coherencies.
- $E_{ps}$  represents Direction-Dependent Jones terms.
- $X_{pqs}$  represents a coherency matrix (per-source).
- $G_p$  represents Direction-Independent Jones terms.

Generally,  $E_{ps}$ ,  $G_p$ ,  $X_{pqs}$  should be formed by using the [RIME API](#) functions and combining them together with `einsum()`.

## Please read the Notes

### Parameters

**time\_index** [dask.array.Array] Time index used to look up the antenna Jones index for a particular baseline with shape (row,). Obtainable via `time.map_blocks(lambda a: np.unique(a, return_inverse=True)[1])`.

**antenna1** [dask.array.Array] Antenna 1 index used to look up the antenna Jones for a particular baseline. with shape (row,).

**antenna2** [dask.array.Array] Antenna 2 index used to look up the antenna Jones for a particular baseline. with shape (row,).

**dde1\_jones** [dask.array.Array, optional]  $E_{ps}$  Direction-Dependent Jones terms for the first antenna. shape (source, time, ant, chan, corr\_1, corr\_2)

**source\_coh** [dask.array.Array, optional]  $X_{pqs}$  Direction-Dependent Coherency matrix for the baseline. with shape (source, row, chan, corr\_1, corr\_2)

**dde2\_jones** [dask.array.Array, optional]  $E_{qs}$  Direction-Dependent Jones terms for the second antenna. This is usually the same array as `dde1_jones` as this preserves the symmetry of the RIME. `predict_vis` will perform the conjugate transpose internally. shape (source, time, ant, chan, corr\_1, corr\_2)

**die1\_jones** [dask.array.Array, optional]  $G_{ps}$  Direction-Independent Jones terms for the first antenna of the baseline. with shape (time, ant, chan, corr\_1, corr\_2)

**base\_vis** [dask.array.Array, optional]  $B_{pq}$  base coherencies, added to source coherency summation before multiplication with `die1_jones` and `die2_jones`. shape (row, chan, corr\_1, corr\_2).

**die2\_jones** [`dask.array.Array`, optional]  $G_{ps}$  Direction-Independent Jones terms for the second antenna of the baseline. This is usually the same array as `die1_jones` as this preserves the symmetry of the RIME. `predict_vis` will perform the conjugate transpose internally.  
`shape (time, ant, chan, corr_1, corr_2)`

**streams** [{False, True}] If True the coherencies are serially summed in a linear chain. If False, `dask` uses a tree style reduction algorithm.

### Returns

**visibilities** [`dask.array.Array`] Model visibilities of shape `(row, chan, corr_1, corr_2)`

### Notes

- Direction-Dependent terms (`dde{1,2}_jones`) and Independent (`die{1,2}_jones`) are optional, but if one is present, the other must be present.
- The inputs to this function involve `row`, `time` and `ant` (antenna) dimensions.
- Each `row` is associated with a pair of antenna Jones matrices at a particular timestep via the `time_index`, `antenna1` and `antenna2` inputs.
- The `row` dimension must be an increasing partial order in time.
- The `ant` dimension should only contain a single chunk equal to the number of antenna. Since each `row` can contain any antenna, random access must be preserved along this dimension.
- The chunks in the `row` and `time` dimension **must** align. This subtle point **must be understood otherwise invalid results will be produced** by the chunking scheme. In the example below we have four unique time indices [0, 1, 2, 3], and four unique antenna [0, 1, 2, 3] indexing 10 rows.

```
# Row indices into the time/antenna indexed arrays
time_idx = np.asarray([0, 0, 1, 1, 2, 2, 2, 2, 3, 3])
ant1 = np.asarray([0, 0, 0, 1, 1, 1, 2, 2, 3])
ant2 = np.asarray([0, 1, 2, 3, 1, 2, 3, 2, 3, 3])
```

A reasonable chunking scheme for the `row` and `time` dimension would be (4, 4, 2) and (2, 1, 1) respectively. Another way of explaining this is that the first four rows contain two unique timesteps, the second four rows contain one unique timestep and the last two rows contain one unique timestep.

Some rules of thumb:

1. The number chunks in `row` and `time` must match although the individual chunk sizes need not.
2. Unique timesteps should not be split across row chunks.
3. For a Measurement Set whose rows are ordered on the TIME column, the following is a good way of obtaining the row chunking strategy:

```
import numpy as np
import pyrap.tables as pt

ms = pt.table("data.ms")
times = ms.getcol("TIME")
unique_times, chunks = np.unique(times, return_counts=True)
```

4. Use `aggregate_chunks()` to aggregate multiple `row` and `time` chunks into chunks large enough such that functions operating on the resulting data can drop the GIL and spend time processing the data. Expanding the previous example:

```
# Aggregate row
utimes = unique_times.size
# Single chunk for each unique time
time_chunks = (1,)*utimes
# Aggregate row chunks into chunks <= 10000
aggregate_chunks((chunks, time_chunks), (10000, utimes))
```

`africanus.rime.dask.phase_delay(lm, uvw, frequency, convention='fourier')`

Computes the phase delay (K) term:

$$e^{-2\pi i(ul+vm+w(n-1))}$$

where  $n = \sqrt{1 - l^2 - m^2}$

#### Parameters

**lm** [dask.array.Array] LM coordinates of shape (source, 2) with L and M components in the last dimension.

**uvw** [dask.array.Array] UVW coordinates of shape (row, 3) with U, V and W components in the last dimension.

**frequency** [dask.array.Array] frequencies of shape (chan,)

**convention** [{‘fourier’, ‘casa’}] Uses the  $e^{-2\pi i}$  sign convention if `fourier` and  $e^{2\pi i}$  if `casa`.

#### Returns

**complex\_phase** [dask.array.Array] complex of shape (source, row, chan)

#### Notes

Corresponds to the complex exponential of the Van Cittert-Zernike Theorem.

MeqTrees uses the CASA sign convention.

`africanus.rime.dask.parallactic_angles(times, antenna_positions, field_centre, **kwargs)`

Computes parallactic angles per timestep for the given reference antenna position and field centre.

#### Parameters

**times** [dask.array.Array] Array of Mean Julian Date times in *seconds* with shape (time,),

**antenna\_positions** [dask.array.Array] Antenna positions of shape (ant, 3) in *metres* in the *ITRF* frame.

**field\_centre** [dask.array.Array] Field centre of shape (2,) in *radians*

**backend** [{‘casa’, ‘test’}, optional] Backend to use for calculating the parallactic angles.

- `casa` defers to an implementation depending on `python-casacore`. This backend should be used by default.
- `test` creates parallactic angles by multiplying the `times` and `antenna_position` arrays. It exists solely for testing.

#### Returns

**parallactic\_angles** [dask.array.Array] Parallactic angles of shape (time, ant)

`africanus.rime.dask.feed_rotation(parallactic_angles, feed_type)`

Computes the 2x2 feed rotation (L) matrix from the `parallactic_angles`.

$$\text{linear} \begin{bmatrix} \cos(pa) & \sin(pa) \\ -\sin(pa) & \cos(pa) \end{bmatrix} \quad \text{circular} \begin{bmatrix} e^{-ipa} & 0 \\ 0 & e^{ipa} \end{bmatrix}$$

#### Parameters

`parallactic_angles` [numpy.ndarray] floating point parallactic angles. Of shape (pa0, pa1, ..., pan).

`feed_type` [{‘linear’, ‘circular’}] The type of feed

#### Returns

`feed_matrix` [numpy.ndarray] Feed rotation matrix of shape (pa0, pa1, ..., pan, 2, 2)

`africanus.rime.dask.transform_sources(lm, parallactic_angles, pointing_errors, antenna_scaling, frequency, dtype=None)`

Creates beam sampling coordinates suitable for use in `beam_cube_dde()` by:

1. Rotating `lm` coordinates by the `parallactic_angles`
2. Adding `pointing_errors`
3. Scaling by `antenna_scaling`

#### Parameters

`lm` [dask.array.Array] LM coordinates of shape (src, 2) in radians offset from the phase centre.

`parallactic_angles` [dask.array.Array] parallactic angles of shape (time, antenna) in radians.

`pointing_errors` [dask.array.Array] LM pointing errors for each antenna at each timestep in radians. Has shape (time, antenna, 2)

`antenna_scaling` [dask.array.Array] antenna scaling factor for each channel and each antenna. Has shape (antenna, chan)

`frequency` [dask.array.Array] frequencies for each channel. Has shape (chan,)

`dtype` [numpy.dtype, optional] Numpy dtype of result array. Should be float32 or float64. Defaults to float64

#### Returns

`coords` [dask.array.Array] coordinates of shape (3, src, time, antenna, chan) where each coordinate component represents `l`, `m` and `frequency`, respectively.

`africanus.rime.dask.beam_cube_dde(beam, beam_lm_extents, beam_freq_map, lm, parallactic_angles, point_errors, antenna_scaling, frequencies)`

Evaluates Direction Dependent Effects along a source’s path by interpolating the values of a complex beam cube at the source location.

#### Parameters

`beam` [dask.array.Array] Complex beam cube of shape (beam\_lw, beam\_mh, beam\_nud, corr, corr). `beam_lw`, `beam_mh` and `beam_nud` define the size of the cube in the l, m and frequency dimensions, respectively.

`beam_lm_extents` [dask.array.Array] lm extents of the beam cube of shape (2, 2). [[lower\_l, upper\_l], [lower\_m, upper\_m]].

**beam\_freq\_map** [dask.array.Array] Beam frequency map of shape (beam\_nud,). This array is used to define interpolation along the (chan,) dimension.

**lm** [dask.array.Array] Source lm coordinates of shape (source, 2). These coordinates are:

1. Scaled if the associated frequency lies outside the beam cube.
2. Offset by pointing errors: `point_errors`
3. Rotated by parallactic angles: `parallactic_angles`.
4. Scaled by antenna scaling factors: `antenna_scaling`.

**parallactic\_angles** [dask.array.Array] Parallactic angles of shape (time, ant).

**point\_errors** [dask.array.Array] Pointing errors of shape (time, ant, chan, 2).

**antenna\_scaling** [dask.array.Array] Antenna scaling factors of shape (ant, chan, 2)

**frequency** [dask.array.Array] Frequencies of shape (chan,).

#### Returns

**ddes** [dask.array.Array] Direction Dependent Effects of shape (source, time, ant, chan, corr, corr)

#### Notes

1. Sources are clamped to the provided `beam_lm_extents`.
2. Frequencies outside the cube (i.e. outside `beam_freq_map`) introduce linear scaling to the lm coordinates of a source.

`africanus.rime.dask.zernike_dde(coords, coeffs, noll_index, parallactic_angle, frequency_scaling, antenna_scaling, pointing_errors)`

Computes Direction Dependent Effects by evaluating Zernicke Polynomials defined by coefficients `coeffs` and noll indexes `noll_index` at the specified coordinates `coords`.

Decomposition of a voxel beam cube into Zernicke polynomial coefficients can be achieved through the use of the `eidos` package.

#### Parameters

**coords** [dask.array.Array] Float coordinates at which to evaluate the zernike polynomials.  
Has shape (3, source, time, ant, chan). The three components in the first dimension represent l, m and frequency coordinates, respectively.

**coeffs** [dask.array.Array] complex Zernicke polynomial coefficients. Has shape (ant, chan, corr\_1, ..., corr\_n, poly) where poly is the number of polynomial coefficients and corr\_1, ..., corr\_n are a variable number of correlation dimensions.

**noll\_index** [dask.array.Array] Noll index associated with each polynomial coefficient. Has shape (ant, chan, corr\_1, ..., corr\_n, poly). correlation dimensions.

**parallactic\_angles** [dask.array.Array] Parallactic angle rotation. Has shape (time, ant).

**frequency\_scaling** [dask.array.Array] The scaling of frequency of the beam. Has shape (chan, ).

**antenna\_scaling** [dask.array.Array] The antenna scaling. Has shape (ant, chan, 2).

**pointing\_errors** [dask.array.Array] The pointing error. Has shape (time, ant, chan, 2).

**Returns**

**dde** [dask.array.Array] complex values with shape (source, time, ant, chan, corr\_1, ..., corr\_n)

africanus.rime.dask.wsclean\_predict(uvw, lm, source\_type, flux, coeffs, log\_poly, ref\_freq, gauss\_shape, frequency)

Predict visibilities from a WSClean sky model.

**Parameters**

**uvw** [dask.array.Array] UVW coordinates of shape (row, 3)

**lm** [dask.array.Array] Source LM coordinates of shape (source, 2), in radians. Derived from the Ra and Dec fields.

**source\_type** [dask.array.Array] Strings defining the source type of shape (source,). Should be either "POINT" or "GAUSSIAN". Contains the Type field.

**flux** [dask.array.Array] Source flux of shape (source,). Contains the I field.

**coeffs** [dask.array.Array] Source Polynomial coefficients of shape (source, coeffs). Contains the SpectralIndex field.

**log\_poly** [dask.array.Array] Source polynomial type of shape (source,). If True, logarithmic polynomials are used. If False, standard polynomials are used. Contains the LogarithmicSI field.

**ref\_freq** [dask.array.Array] Source Reference frequency of shape (source,). Contains the ReferenceFrequency field.

**gauss\_shape** [dask.array.Array] Gaussian shape parameters of shape (source, 3) used when the corresponding source\_type is "GAUSSIAN". The 3 components should contain the MajorAxis, MinorAxis and Orientation fields in radians, respectively.

**frequency** [dask.array.Array] Frequency of shape (chan,).

**Returns**

**visibilities** [dask.array.Array] Complex visibilities of shape (row, chan, 1)

## 5.2 Direct Fourier Transform

Functions used to compute the discretised direct Fourier transform (DFT) for an ideal interferometer. The DFT for an ideal interferometer is defined as

$$V(u, v, w) = \int B(l, m) e^{-2\pi i(u l + v m + w(n-1))} \frac{dl dm}{n}$$

where  $u, v, w$  are data space coordinates and where visibilities  $V$  have been obtained. The  $l, m, n$  are signal space coordinates at which we wish to reconstruct the signal  $B$ . Note that the signal corresponds to the brightness matrix and not the Stokes parameters. We adopt the convention where we absorb the fixed coordinate  $n$  in the denominator into the image. Note that the data space coordinates have an implicit dependence on frequency and time and that the image has an implicit dependence on frequency. The discretised form of the DFT can be written as

$$V(u, v, w) = \sum_s e^{-2\pi i(u l_s + v m_s + w(n_s-1))} \cdot B_s$$

where  $s$  labels the source (or pixel) location. If only a single correlation is present  $B = I$ , this can be cast into a matrix equation as follows

$$V = RI$$

where  $R$  is the operator that maps an image to visibility space. This mapping is implemented by the `im_to_vis()` function. If multiple correlations are present then each one is mapped to its corresponding visibility. An imaging algorithm also requires the adjoint denoted  $R^\dagger$  which is simply the complex conjugate transpose of  $R$ . The dirty image is obtained by applying the adjoint operator to the visibilities

$$I^D = R^\dagger V$$

This is implemented by the `vis_to_im()` function. Note that an imaging algorithm using these operators will actually reconstruct  $\frac{I}{n}$  but that it is trivial to obtain  $I$  since  $n$  is known at each location in the image.

## 5.2.1 Numpy

---

<code>im_to_vis(image, uvw, lm, frequency[, ...])</code>	Computes the discrete image to visibility mapping of an ideal interferometer:
<code>vis_to_im(vis, uvw, lm, frequency, flags[, ...])</code>	Computes visibility to image mapping of an ideal interferometer:

---

`africanus.dft.im_to_vis(image, uvw, lm, frequency, convention='fourier', dtype=None)`

Computes the discrete image to visibility mapping of an ideal interferometer:

$$\sum_s e^{-2\pi i(u l_s + v m_s + w(n_s - 1))} \cdot I_s$$

### Parameters

**image** [numpy.ndarray] image of shape (source, chan, corr) The brightness matrix in each pixel (flatten 2D array per channel and corr). Note not Stokes terms

**uvw** [numpy.ndarray] uvw coordinates of shape (row, 3) with u, v and w components in the last dimension.

**lm** [numpy.ndarray] lm coordinates of shape (source, 2) with l and m components in the last dimension.

**frequency** [numpy.ndarray] frequencies of shape (chan,)

**convention** [{‘fourier’, ‘casa’}] Uses the  $e^{-2\pi i}$  sign convention if `fourier` and  $e^{2\pi i}$  if `casa`.

**dtype** [np.dtype, optional] Datatype of result. Should be either np.complex64 or np.complex128. If None, `numpy.result_type()` is used to infer the data type from the inputs.

### Returns

**visibilities** [numpy.ndarray] complex of shape (row, chan, corr)

`africanus.dft.vis_to_im(vis, uvw, lm, frequency, flags, convention='fourier', dtype=None)`

Computes visibility to image mapping of an ideal interferometer:

$$\sum_k e^{2\pi i(u_k l + v_k m + w_k(n-1))} \cdot V_k$$

### Parameters

**vis** [numpy.ndarray] visibilities of shape (row, chan, corr) Visibilities corresponding to brightness terms. Note the dirty images produced do not necessarily correspond to Stokes terms and need to be converted.

**uvw** [numpy.ndarray] uvw coordinates of shape (row, 3) with u, v and w components in the last dimension.

**lm** [numpy.ndarray] lm coordinates of shape (source, 2) with l and m components in the last dimension.

**frequency** [numpy.ndarray] frequencies of shape (chan,)

**flags** [numpy.ndarray] Boolean array of shape (row, chan, corr) Note that if one correlation is flagged we discard all of them otherwise we end up irretrievably mixing Stokes terms.

**convention** [{‘fourier’, ‘casa’}] Uses the  $e^{-2\pi i}$  sign convention if **fourier** and  $e^{2\pi i}$  if **casa**.

**dtype** [np.dtype, optional] Datatype of result. Should be either np.float32 or np.float64. If None, `numpy.result_type()` is used to infer the data type from the inputs.

#### Returns

**image** [numpy.ndarray] float of shape (source, chan, corr)

### 5.2.2 Dask

<code>im_to_vis(image, uvw, lm, frequency[, ...])</code>	Computes the discrete image to visibility mapping of an ideal interferometer:
<code>vis_to_im(vis, uvw, lm, frequency, flags[, ...])</code>	Computes visibility to image mapping of an ideal interferometer:

`africanus.dft.dask.im_to_vis(image, uvw, lm, frequency, convention='fourier', dtype=numpy.complex128)`

Computes the discrete image to visibility mapping of an ideal interferometer:

$$\sum_s e^{-2\pi i(u l_s + v m_s + w(n_s - 1))} \cdot I_s$$

#### Parameters

**image** [dask.array.Array] image of shape (source, chan, corr) The brightness matrix in each pixel (flatten 2D array per channel and corr). Note not Stokes terms

**uvw** [dask.array.Array] uvw coordinates of shape (row, 3) with u, v and w components in the last dimension.

**lm** [dask.array.Array] lm coordinates of shape (source, 2) with l and m components in the last dimension.

**frequency** [dask.array.Array] frequencies of shape (chan,)

**convention** [{‘fourier’, ‘casa’}] Uses the  $e^{-2\pi i}$  sign convention if **fourier** and  $e^{2\pi i}$  if **casa**.

**dtype** [np.dtype, optional] Datatype of result. Should be either np.complex64 or np.complex128. If None, `numpy.result_type()` is used to infer the data type from the inputs.

#### Returns

**visibilities** [dask.array.Array] complex of shape (row, chan, corr)

`africanus.dft.dask.vis_to_im(vis, uvw, lm, frequency, flags, convention='fourier', dtype=numpy.float64)`

Computes visibility to image mapping of an ideal interferometer:

$$\sum_k e^{2\pi i(u_k l + v_k m + w_k(n - 1))} \cdot V_k$$

### Parameters

**vis** [dask.array.Array] visibilities of shape (row, chan, corr) Visibilities corresponding to brightness terms. Note the dirty images produced do not necessarily correspond to Stokes terms and need to be converted.

**uvw** [dask.array.Array] uvw coordinates of shape (row, 3) with u, v and w components in the last dimension.

**lm** [dask.array.Array] lm coordinates of shape (source, 2) with l and m components in the last dimension.

**frequency** [dask.array.Array] frequencies of shape (chan,)

**flags** [dask.array.Array] Boolean array of shape (row, chan, corr) Note that if one correlation is flagged we discard all of them otherwise we end up irretrievably mixing Stokes terms.

**convention** [{‘fourier’, ‘casa’}] Uses the  $e^{-2\pi i}$  sign convention if `fourier` and  $e^{2\pi i}$  if `casa`.

**dtype** [np.dtype, optional] Datatype of result. Should be either np.float32 or np.float64. If None, `numpy.result_type()` is used to infer the data type from the inputs.

### Returns

**image** [dask.array.Array] float of shape (source, chan, corr)

## 5.3 Gridding and Degridding

This section contains routines for

1. Gridding complex visibilities onto an image.
2. Degridding complex visibilities from an image.

### 5.3.1 Nifty

Dask wrappers around Nifty’s Gridder.

#### Dask

<code>grid_config([nx, ny, eps, cell_size_x, ...])</code>	Returns a wrapper around a NIFTY GridderConfiguration object.
<code>grid(vis, uvw, flags, weights, frequencies, ...)</code>	Grids the supplied visibilities in parallel.
<code>dirty(grid, grid_config)</code>	Computes the dirty image from gridded visibilities and the gridding configuration.
<code>degrid(grid, uvw, flags, weights, ...[, ...])</code>	Degrids the visibilities from the supplied grid in parallel.
<code>model(image, grid_config)</code>	Computes model visibilities from an image and a gridding configuration.

`africanus.gridding.nifty.dask.grid_config(nx=1024, ny=1024, eps=2e-13, cell_size_x=2.0, cell_size_y=2.0)`

Returns a wrapper around a NIFTY GridderConfiguration object.

### Parameters

**nx** [int, optional] Number of X pixels in the grid. Defaults to 1024.  
**ny** [int, optional] Number of Y pixels in the grid. Defaults to 1024.  
**cell\_size\_x** [float, optional] Cell size of the X pixel in arcseconds. Defaults to 2.0.  
**cell\_size\_y** [float, optional] Cell size of the Y pixel in arcseconds. Defaults to 2.0.  
**eps** [float] Gridder accuracy error. Defaults to 2e-13

**Returns**

**grid\_config** [GridderConfigWrapper] The NIFTY Gridder Configuration

```
africanus.gridding.nifty.dask.grid(vis, uvw, flags, weights, frequencies, grid_config, wmin=- 1e+30,
                                     wmax=1e+30, streams=None)
```

Grids the supplied visibilities in parallel. Note that a grid is created for each visibility chunk.

**Parameters**

**vis** [dask.array.Array] visibilities of shape (row, chan, corr)  
**uvw** [dask.array.Array] uvw coordinates of shape (row, 3)  
**flags** [dask.array.Array] flags of shape (row, chan, corr)  
**weights** [dask.array.Array] weights of shape (row, chan, corr).  
**frequencies** [dask.array.Array] frequencies of shape (chan,)  
**grid\_config** [GridderConfigWrapper] Gridding Configuration  
**wmin** [float] Minimum W coordinate to grid. Defaults to -1e30.  
**wmax** [float] Maximum W coordinate to grid. Default to 1e30.  
**streams** [int, optional] Number of parallel gridding operations. Default to None, in which case as many grids as visibility chunks will be created.

**Returns**

**grid** [dask.array.Array] grid of shape (ny, nx, corr)

```
africanus.gridding.nifty.dask.dirty(grid, grid_config)
```

Computes the dirty image from gridded visibilities and the gridding configuration.

**Parameters**

**grid** [dask.array.Array] Gridded visibilities of shape (nv, nu, ncorr)  
**grid\_config** [GridderConfigWrapper] Gridding configuration

**Returns**

**dirty** [dask.array.Array] dirty image of shape (ny, nx, corr)

```
africanus.gridding.nifty.dask.degrid(grid, uvw, flags, weights, frequencies, grid_config, wmin=- 1e+30,
                                         wmax=1e+30)
```

Degrids the visibilities from the supplied grid in parallel.

**Parameters**

**grid** [dask.array.Array] gridded visibilities of shape (ny, nx, corr)  
**uvw** [dask.array.Array] uvw coordinates of shape (row, 3)  
**flags** [dask.array.Array] flags of shape (row, chan, corr)

**weights** [dask.array.Array] weights of shape (row, chan, corr). Currently unsupported and ignored.

**frequencies** [dask.array.Array] frequencies of shape (chan,)

**grid\_config** [GridderConfigWrapper] Gridding Configuration

**wmin** [float] Minimum W coordinate to grid. Defaults to -1e30.

**wmax** [float] Maximum W coordinate to grid. Default to 1e30.

#### Returns

**grid** [dask.array.Array] grid of shape (ny, nx, corr)

`africanus.gridding.nifty.dask.model(image, grid_config)`

Computes model visibilities from an image and a gridding configuration.

#### Parameters

**image** [dask.array.Array] Image of shape (ny, nx, corr).

**grid\_config** [GridderConfigWrapper] nifty gridding configuration object

#### Returns

**model\_vis** [dask.array.Array] Model visibilities of shape (nu, nv, corr).

## 5.3.2 wgriddler

Wrappers around ‘ducc.wgriddler <<https://gitlab.mpcdf.mpg.de/mtr/ducc>>`\_.

### Numpy

<code>dirty</code> (uvw, freq, vis, freq_bin_idx, ...[, ...])	Compute visibility to image mapping using ducc griddler i.e.
<code>model</code> (uvw, freq, image, freq_bin_idx, ...[, ...])	Compute image to visibility mapping using ducc degriddler i.e.
<code>residual</code> (uvw, freq, image, vis, ...[, ...])	Compute residual image given a model and visibilities using ducc degriddler i.e.
<code>hessian</code> (uvw, freq, image, freq_bin_idx, ...)	Compute action of Hessian on an image using ducc

`africanus.gridding.wgriddler.dirty(uvw, freq, vis, freq_bin_idx, freq_bin_counts, nx, ny, cell, weights=None, flag=None, celly=None, epsilon=1e-05, nthreads=1, do_wstacking=True, double_accum=False)`

Compute visibility to image mapping using ducc griddler i.e.

$$I^D = R^\dagger \Sigma^{-1} V$$

where  $R^\dagger$  is an implicit gridding operator,  $V$  denotes visibilities of shape (row, chan) and  $I^D$  is the dirty image of shape (band, nx, ny).

The number of imaging bands (band) must be less than or equal to the number of channels (chan) at which the data were obtained. The mapping from (chan) to (band) is described by freq\_bin\_idx and freq\_bin\_counts as described below.

Note that, if self adjoint gridding and degridding operators are required then weights should be the square root of what is typically referred to as imaging weights and should also be passed into the degriddler. In this case, the data needs to be pre-whitened.

**Parameters**

**uvw** [numpy.ndarray] uvw coordinates at which visibilities were obtained with shape (row, 3).

**freq** [numpy.ndarray] Observational frequencies of shape (chan, ).

**vis** [numpy.ndarray] Visibilities of shape (row, chan).

**freq\_bin\_idx** [numpy.ndarray] Starting indices of frequency bins for each imaging band of shape (band, ).

**freq\_bin\_counts** [numpy.ndarray] The number of channels in each imaging band of shape (band, ).

**cell** [float] The cell size of a pixel along the  $x$  direction in radians.

**weights** [numpy.ndarray, optional] Imaging weights of shape (row, chan).

**flag:** [class:numpy.ndarray, optional] Flags of shape (row, chan). Will only process visibilities for which flag!=0

**celly** [float, optional] The cell size of a pixel along the  $y$  direction in radians. By default same as cell size along  $x$  direction.

**epsilon** [float, optional] The precision of the gridded with respect to the direct Fourier transform. By default, this is set to 1e-5 for single precision and 1e-7 for double precision.

**nthreads** [int, optional] The number of threads to use. Defaults to one. If set to zero will use all available cores.

**do\_wstacking** [bool, optional] Whether to correct for the w-term or not. Defaults to True

**double\_accum** [bool, optional] If true ducc will accumulate in double precision regardless of the input type.

**Returns**

**model** [numpy.ndarray] Dirty image corresponding to visibilities of shape (nband, nx, ny).

```
africanus.gridding.wgridded.model(uvw,freq,image,freq_bin_idx,freq_bin_counts,cell,weights=None,
                                     flag=None,celly=None,epsilon=1e-05,nthreads=1,
                                     do_wstacking=True)
```

Compute image to visibility mapping using ducc degridded i.e.

$$V = Rx$$

where  $R$  is an implicit degridding operator,  $V$  denotes visibilities of shape (row, chan) and  $x$  is the image of shape (band, nx, ny).

The number of imaging bands (band) has to be less than or equal to the number of channels (chan) at which the data were obtained. The mapping from (chan) to (band) is described by freq\_bin\_idx and freq\_bin\_counts as described below.

There is an option to provide weights during degridding to cater for self adjoint gridding and degridding operators. In this case weights should actually be the square root of what is typically referred to as imaging weights. In this case the degridded computes the whitened model visibilities i.e.

$$V = \Sigma^{-\frac{1}{2}} Rx$$

where  $\Sigma$  refers to the inverse of the weights (i.e. the data covariance matrix when using natural weighting).

**Parameters**

**uvw** [numpy.ndarray] uvw coordinates at which visibilities were obtained with shape (row, 3).

**freq** [numpy.ndarray] Observational frequencies of shape (chan, ).

**model** [numpy.ndarray] Model image to degrid of shape (nband, nx, ny).

**freq\_bin\_idx** [numpy.ndarray] Starting indices of frequency bins for each imaging band of shape (band, ).

**freq\_bin\_counts** [numpy.ndarray] The number of channels in each imaging band of shape (band, ).

**cell** [float] The cell size of a pixel along the *x* direction in radians.

**weights** [numpy.ndarray, optional] Imaging weights of shape (row, chan).

**flag:** [class:numpy.ndarray, optional] Flags of shape (row, chan). Will only process visibilities for which flag!=0

**celly** [float, optional] The cell size of a pixel along the *y* direction in radians. By default same as cell size along *x* direction.

**epsilon** [float, optional] The precision of the gridded with respect to the direct Fourier transform. By default, this is set to 1e-5 for single precision and 1e-7 for double precision.

**nthreads** [int, optional] The number of threads to use. Defaults to one. If set to zero will use all available cores.

**do\_wstacking** [bool, optional] Whether to correct for the w-term or not. Defaults to True

## Returns

**vis** [numpy.ndarray] Visibilities corresponding to model of shape (row, chan).

```
africanus.gridding.wgridded.residual(uvw, freq, image, vis, freq_bin_idx, freq_bin_counts, cell,
                                         weights=None, flag=None, celly=None, epsilon=1e-05, nthreads=1,
                                         do_wstacking=True, double_accum=False)
```

Compute residual image given a model and visibilities using ducc degridder i.e.

$$I^R = R^\dagger \Sigma^{-1} (V - Rx)$$

where  $R$  is an implicit degridding operator,  $V$  denotes visibilities of shape (row, chan) and  $x$  is the image of shape (band, nx, ny).

The number of imaging bands (band) must be less than or equal to the number of channels (chan) at which the data were obtained. The mapping from (chan) to (band) is described by freq\_bin\_idx and freq\_bin\_counts as described below.

Note that, if the gridding and degridding operators both apply the square root of the imaging weights then the visibilities that are passed in should be pre-whitened. In this case the function computes

$$I^R = R^\dagger \Sigma^{-\frac{1}{2}} (\tilde{V} - \Sigma^{-\frac{1}{2}} Rx)$$

which is identical to the above expression if  $\tilde{V} = \Sigma^{-\frac{1}{2}} V$ .

## Parameters

**uvw** [numpy.ndarray] uvw coordinates at which visibilities were obtained with shape (row, 3).

**freq** [numpy.ndarray] Observational frequencies of shape (chan, ).

**model** [numpy.ndarray] Model image to degrid of shape (band, nx, ny).

**vis** [numpy.ndarray] Visibilities of shape (row, chan).

**weights** [numpy.ndarray] Imaging weights of shape (row, chan).

**freq\_bin\_idx** [numpy.ndarray] Starting indices of frequency bins for each imaging band of shape (band, ).

**freq\_bin\_counts** [numpy.ndarray] The number of channels in each imaging band of shape (band, ).

**cell** [float] The cell size of a pixel along the *x* direction in radians.

**flag:** [class: numpy.ndarray, optional] Flags of shape (row, chan). Will only process visibilities for which flag!=0

**celly** [float, optional] The cell size of a pixel along the *y* direction in radians. By default same as cell size along *x* direction.

**nu** [int, optional] The number of pixels in the padded grid along the *x* direction. Chosen automatically by default.

**nv** [int, optional] The number of pixels in the padded grid along the *y* direction. Chosen automatically by default.

**epsilon** [float, optional] The precision of the gridded with respect to the direct Fourier transform. By default, this is set to 1e-5 for single precision and 1e-7 for double precision.

**nthreads** [int, optional] The number of threads to use. Defaults to one.

**do\_wstacking** [bool, optional] Whether to correct for the w-term or not. Defaults to True

**double\_accum** [bool, optional] If true ducc will accumulate in double precision regardless of the input type.

### Returns

**residual** [numpy.ndarray] Residual image corresponding to model of shape (band, nx, ny).

```
africanus.gridding.wgridder.hessian(uvw, freq, image, freq_bin_idx, freq_bin_counts, cell, weights=None,
                                      flag=None, celly=None, epsilon=1e-05, nthreads=1,
                                      do_wstacking=True, double_accum=False)
```

Compute action of Hessian on an image using ducc

$$R^\dagger \Sigma^{-1} R x$$

where  $R$  is an implicit degridding operator and  $x$  is the image of shape (band, nx, ny).

The number of imaging bands (band) must be less than or equal to the number of channels (chan) at which the data were obtained. The mapping from (chan) to (band) is described by freq\_bin\_idx and freq\_bin\_counts as described below.

### Parameters

**uvw** [numpy.ndarray] uvw coordinates at which visibilities were obtained with shape (row, 3).

**freq** [numpy.ndarray] Observational frequencies of shape (chan, ).

**model** [numpy.ndarray] Model image to degrid of shape (band, nx, ny).

**weights** [numpy.ndarray] Imaging weights of shape (row, chan).

**freq\_bin\_idx** [numpy.ndarray] Starting indices of frequency bins for each imaging band of shape (band, ).

**freq\_bin\_counts** [numpy.ndarray] The number of channels in each imaging band of shape (band, ).

**cell** [float] The cell size of a pixel along the  $x$  direction in radians.

**flag:** [class: numpy.ndarray, optional] Flags of shape (row, chan). Will only process visibilities for which flag!=0

**celly** [float, optional] The cell size of a pixel along the  $y$  direction in radians. By default same as cell size along  $x$  direction.

**nx** [int, optional] The number of pixels in the padded grid along the  $x$  direction. Chosen automatically by default.

**ny** [int, optional] The number of pixels in the padded grid along the  $y$  direction. Chosen automatically by default.

**epsilon** [float, optional] The precision of the gridded with respect to the direct Fourier transform. By default, this is set to 1e-5 for single precision and 1e-7 for double precision.

**nthreads** [int, optional] The number of threads to use. Defaults to one.

**do\_wstacking** [bool, optional] Whether to correct for the w-term or not. Defaults to True

**double\_accum** [bool, optional] If true ducc will accumulate in double precision regardless of the input type.

#### Returns

**residual** [numpy.ndarray] Residual image corresponding to model of shape (band, nx, ny).

## Dask

<code>dirty</code> (uvw, freq, vis, freq_bin_idx, ...[, ...])	Compute visibility to image mapping using ducc gridded i.e.
<code>model</code> (uvw, freq, image, freq_bin_idx, ...[, ...])	Compute image to visibility mapping using ducc degridded i.e.
<code>residual</code> (uvw, freq, image, vis, ...[, ...])	Compute residual image given a model and visibilities using ducc degridded i.e.
<code>hessian</code> (uvw, freq, image, freq_bin_idx, ...)	Compute action of Hessian on an image using ducc

`africanus.gridding.wgridder.dask.dirty`(uvw, freq, vis, freq\_bin\_idx, freq\_bin\_counts, nx, ny, cell, weights=None, flag=None, celly=None, epsilon=1e-05, nthreads=1, do\_wstacking=True, double\_accum=False)

Compute visibility to image mapping using ducc gridded i.e.

$$I^D = R^\dagger \Sigma^{-1} V$$

where  $R^\dagger$  is an implicit gridding operator,  $V$  denotes visibilities of shape (row, chan) and  $I^D$  is the dirty image of shape (band, nx, ny).

The number of imaging bands (band) must be less than or equal to the number of channels (chan) at which the data were obtained. The mapping from (chan) to (band) is described by freq\_bin\_idx and freq\_bin\_counts as described below.

Note that, if self adjoint gridding and degridding operators are required then weights should be the square root of what is typically referred to as imaging weights and should also be passed into the degridded. In this case, the data needs to be pre-whitened.

**Parameters**

**uvw** [dask.array.Array] uvw coordinates at which visibilities were obtained with shape (row, 3).

**freq** [dask.array.Array] Observational frequencies of shape (chan, ).

**vis** [dask.array.Array] Visibilities of shape (row, chan).

**freq\_bin\_idx** [dask.array.Array] Starting indices of frequency bins for each imaging band of shape (band, ).

**freq\_bin\_counts** [dask.array.Array] The number of channels in each imaging band of shape (band, ).

**cell** [float] The cell size of a pixel along the  $x$  direction in radians.

**weights** [dask.array.Array, optional] Imaging weights of shape (row, chan).

**flag:** [class:dask.array.Array, optional] Flags of shape (row, chan). Will only process visibilities for which flag!=0

**celly** [float, optional] The cell size of a pixel along the  $y$  direction in radians. By default same as cell size along  $x$  direction.

**epsilon** [float, optional] The precision of the gridded with respect to the direct Fourier transform. By deafult, this is set to 1e-5 for single precision and 1e-7 for double precision.

**nthreads** [int, optional] The number of threads to use. Defaults to one. If set to zero will use all available cores.

**do\_wstacking** [bool, optional] Whether to correct for the w-term or not. Defaults to True

**double\_accum** [bool, optional] If true ducc will accumulate in double precision regardless of the input type.

**Returns**

**model** [dask.array.Array] Dirty image corresponding to visibilities of shape (nband, nx, ny).

```
africanus.gridding.wgridded.dask.model(uvw,freq,image,freq_bin_idx,freq_bin_counts,cell,
                                         weights=None,flag=None,celly=None,epsilon=1e-05,
                                         nthreads=1,do_wstacking=True)
```

Compute image to visibility mapping using ducc degridded i.e.

$$V = Rx$$

where  $R$  is an implicit degridding operator,  $V$  denotes visibilities of shape (row, chan) and  $x$  is the image of shape (band, nx, ny).

The number of imaging bands (band) has to be less than or equal to the number of channels (chan) at which the data were obtained. The mapping from (chan) to (band) is described by freq\_bin\_idx and freq\_bin\_counts as described below.

There is an option to provide weights during degridding to cater for self adjoint gridding and degridding operators. In this case weights should actually be the square root of what is typically referred to as imaging weights. In this case the degridded computes the whitened model visibilities i.e.

$$V = \Sigma^{-\frac{1}{2}} Rx$$

where  $\Sigma$  refers to the inverse of the weights (i.e. the data covariance matrix when using natural weighting).

**Parameters**

**uvw** [dask.array.Array] uvw coordinates at which visibilities were obtained with shape (row, 3).

**freq** [dask.array.Array] Observational frequencies of shape (chan, ).

**model** [dask.array.Array] Model image to degrid of shape (nband, nx, ny).

**freq\_bin\_idx** [dask.array.Array] Starting indices of frequency bins for each imaging band of shape (band, ).

**freq\_bin\_counts** [dask.array.Array] The number of channels in each imaging band of shape (band, ).

**cell** [float] The cell size of a pixel along the *x* direction in radians.

**weights** [dask.array.Array, optional] Imaging weights of shape (row, chan).

**flag:** [class:dask.array.Array, optional] Flags of shape (row, chan). Will only process visibilities for which flag!=0

**celly** [float, optional] The cell size of a pixel along the *y* direction in radians. By default same as cell size along *x* direction.

**epsilon** [float, optional] The precision of the gridded with respect to the direct Fourier transform. By default, this is set to 1e-5 for single precision and 1e-7 for double precision.

**nthreads** [int, optional] The number of threads to use. Defaults to one. If set to zero will use all available cores.

**do\_wstacking** [bool, optional] Whether to correct for the w-term or not. Defaults to True

### Returns

**vis** [dask.array.Array] Visibilities corresponding to model of shape (row, chan).

africanus.gridding.wgriddler.dask.residual(uvw, freq, image, vis, freq\_bin\_idx, freq\_bin\_counts, cell, weights=None, flag=None, celly=None, epsilon=1e-05, nthreads=1, do\_wstacking=True, double\_accum=False)

Compute residual image given a model and visibilities using ducc degriddler i.e.

$$I^R = R^\dagger \Sigma^{-1} (V - Rx)$$

where  $R$  is an implicit degridding operator,  $V$  denotes visibilities of shape (row, chan) and  $x$  is the image of shape (band, nx, ny).

The number of imaging bands (band) must be less than or equal to the number of channels (chan) at which the data were obtained. The mapping from (chan) to (band) is described by freq\_bin\_idx and freq\_bin\_counts as described below.

Note that, if the gridding and degridding operators both apply the square root of the imaging weights then the visibilities that are passed in should be pre-whitened. In this case the function computes

$$I^R = R^\dagger \Sigma^{-\frac{1}{2}} (\tilde{V} - \Sigma^{-\frac{1}{2}} Rx)$$

which is identical to the above expression if  $\tilde{V} = \Sigma^{-\frac{1}{2}} V$ .

### Parameters

**uvw** [dask.array.Array] uvw coordinates at which visibilities were obtained with shape (row, 3).

**freq** [dask.array.Array] Observational frequencies of shape (chan, ).

**model** [dask.array.Array] Model image to degrid of shape (band, nx, ny).

**vis** [dask.array.Array] Visibilities of shape (row, chan).

**weights** [dask.array.Array] Imaging weights of shape (row, chan).

**freq\_bin\_idx** [dask.array.Array] Starting indices of frequency bins for each imaging band of shape (band, ).

**freq\_bin\_counts** [dask.array.Array] The number of channels in each imaging band of shape (band, ).

**cell** [float] The cell size of a pixel along the  $x$  direction in radians.

**flag:** [class:dask.array.Array, optional] Flags of shape (row, chan). Will only process visibilities for which flag!=0

**celly** [float, optional] The cell size of a pixel along the  $y$  direction in radians. By default same as cell size along  $x$  direction.

**nu** [int, optional] The number of pixels in the padded grid along the  $x$  direction. Chosen automatically by default.

**nv** [int, optional] The number of pixels in the padded grid along the  $y$  direction. Chosen automatically by default.

**epsilon** [float, optional] The precision of the gridded with respect to the direct Fourier transform. By default, this is set to 1e-5 for single precision and 1e-7 for double precision.

**nthreads** [int, optional] The number of threads to use. Defaults to one.

**do\_wstacking** [bool, optional] Whether to correct for the w-term or not. Defaults to True

**double\_accum** [bool, optional] If true ducc will accumulate in double precision regardless of the input type.

### Returns

**residual** [dask.array.Array] Residual image corresponding to model of shape (band, nx, ny).

```
africanus.gridding.wgriddler.dask.hessian(uvw, freq, image, freq_bin_idx, freq_bin_counts, cell,
                                             weights=None, flag=None, celly=None, epsilon=1e-05,
                                             nthreads=1, do_wstacking=True, double_accum=False)
```

Compute action of Hessian on an image using ducc

$$R^\dagger \Sigma^{-1} R x$$

where  $R$  is an implicit degridding operator and  $x$  is the image of shape (band, nx, ny).

The number of imaging bands (band) must be less than or equal to the number of channels (chan) at which the data were obtained. The mapping from (chan) to (band) is described by freq\_bin\_idx and freq\_bin\_counts as described below.

### Parameters

**uvw** [dask.array.Array] uvw coordinates at which visibilities were obtained with shape (row, 3).

**freq** [dask.array.Array] Observational frequencies of shape (chan, ).

**model** [dask.array.Array] Model image to degrid of shape (band, nx, ny).

**weights** [dask.array.Array] Imaging weights of shape (row, chan).

**freq\_bin\_idx** [dask.array.Array] Starting indices of frequency bins for each imaging band of shape (band, ).

**freq\_bin\_counts** [dask.array.Array] The number of channels in each imaging band of shape (band, ).

**cell** [float] The cell size of a pixel along the  $x$  direction in radians.

**flag:** [class:dask.array.Array, optional] Flags of shape (row, chan). Will only process visibilities for which flag!=0

**celly** [float, optional] The cell size of a pixel along the  $y$  direction in radians. By default same as cell size along  $x$  direction.

**nu** [int, optional] The number of pixels in the padded grid along the  $x$  direction. Chosen automatically by default.

**nv** [int, optional] The number of pixels in the padded grid along the  $y$  direction. Chosen automatically by default.

**epsilon** [float, optional] The precision of the gridded with respect to the direct Fourier transform. By default, this is set to 1e-5 for single precision and 1e-7 for double precision.

**nthreads** [int, optional] The number of threads to use. Defaults to one.

**do\_wstacking** [bool, optional] Whether to correct for the w-term or not. Defaults to True

**double\_accum** [bool, optional] If true ducc will accumulate in double precision regardless of the input type.

#### Returns

**residual** [dask.array.Array] Residual image corresponding to model of shape (band, nx, ny).

### 5.3.3 Utilities

---

<code>africanus.gridding.util.estimate_cell_size(u, v, wavelength[, ...])</code>	Estimate the cell size in arcseconds given baseline u and v coordinates, as well as the wavelengths, $\lambda$ .
--	--

---

`africanus.gridding.util.estimate_cell_size(u, v, wavelength, factor=3.0, ny=None, nx=None)`  
Estimate the cell size in arcseconds given baseline u and v coordinates, as well as the wavelengths,  $\lambda$ .

The cell size is computed as:

$$\Delta u = 1.0 / (2 \times \text{factor} \times \max(|u|) / \min(\lambda))$$
$$\Delta v = 1.0 / (2 \times \text{factor} \times \max(|v|) / \min(\lambda))$$

If ny and nx are provided the following checks are performed and exceptions are raised on failure:

$$\Delta u * ny \leq \min(\lambda) / \min(|u|)$$
$$\Delta v * nx \leq \min(\lambda) / \min(|v|)$$

#### Parameters

**u** [numpy.ndarray or float] Maximum u coordinate in metres.

**v** [numpy.ndarray or float] Maximum v coordinate in metres.

**wavelength** [numpy.ndarray or float] Wavelengths, in metres.

**factor** [float, optional] Scaling factor

**ny** [int, optional] Grid y dimension

**nx** [int, optional] Grid x dimension

**Returns**

**numpy.ndarray** Cell size of u and v in arcseconds with shape (2,)

**Raises**

**ValueError** If the cell size criteria are not matched.

## 5.4 Deconvolution Algorithms

`africanus.deconv.hogbom.hogbom_clean(dirty, psf, gamma=0.1, threshold='default', niter='default')`

Performs Hogbom Clean on the dirty image given the psf.

**Parameters**

**dirty** [np.ndarray] float64 dirty image of shape (ny, nx)

**psf** [np.ndarray] float64 Point Spread Function of shape (2\*ny, 2\*nx)

**gamma (optional) float** the gain factor (must be less than one)

**threshold (optional)** [float or str] the threshold to clean to

**niter (optional)** [integer] the maximum number of iterations allowed

**Returns**

**np.ndarray** float64 clean image of shape (ny, nx)

**np.ndarray** float64 residual image of shape (ny, nx)

## 5.5 Coordinate Transforms

### 5.5.1 Numpy

<code>radec_to_lm(radec[, phase_centre])</code>	Converts Right-Ascension/Declination coordinates in radians to a Direction Cosine lm coordinates, relative to the Phase Centre.
<code>radec_to_lmn(radec[, phase_centre])</code>	Converts Right-Ascension/Declination coordinates in radians to a Direction Cosine lmn coordinates, relative to the Phase Centre.
<code>lm_to_radec(lm[, phase_centre])</code>	Convert Direction Cosine lm coordinates to Right Ascension/Declination coordinates in radians, relative to the Phase Centre.
<code>lmn_to_radec(lmn[, phase_centre])</code>	Convert Direction Cosine lmn coordinates to Right Ascension/Declination coordinates in radians, relative to the Phase Centre.

`africanus.coordinates.radec_to_lm(radec, phase_centre=None)`

Converts Right-Ascension/Declination coordinates in radians to a Direction Cosine lm coordinates, relative to

the Phase Centre.

$$l = \cos \delta \sin \Delta\alpha \quad (5.1)$$

$$m = \sin \delta \cos \delta_0 - \cos \delta \sin \delta_0 \cos \Delta\alpha \quad (5.2)$$

$$n = \sqrt{1 - l^2 - m^2} - 1 \quad (5.3)$$

where  $\Delta\alpha = \alpha - \alpha_0$  is the difference between the Right Ascension of each coordinate and the phase centre and  $\delta_0$  is the Declination of the phase centre.

### Parameters

**radec** [numpy.ndarray] radec coordinates of shape (coord, 2) where Right-Ascension and Declination are in the last 2 components, respectively.

**phase\_centre** [numpy.ndarray, optional] radec coordinates of the Phase Centre. Shape (2,)

### Returns

**numpy.ndarray** lm Direction Cosines of shape (coord, 2)

`africanus.coordinates.radec_to_lm`(*radec*, *phase\_centre=None*)

Converts Right-Ascension/Declination coordinates in radians to a Direction Cosine lm coordinates, relative to the Phase Centre.

$$l = \cos \delta \sin \Delta\alpha \quad (5.4)$$

$$m = \sin \delta \cos \delta_0 - \cos \delta \sin \delta_0 \cos \Delta\alpha \quad (5.5)$$

$$n = \sqrt{1 - l^2 - m^2} - 1 \quad (5.6)$$

where  $\Delta\alpha = \alpha - \alpha_0$  is the difference between the Right Ascension of each coordinate and the phase centre and  $\delta_0$  is the Declination of the phase centre.

### Parameters

**radec** [numpy.ndarray] radec coordinates of shape (coord, 2) where Right-Ascension and Declination are in the last 2 components, respectively.

**phase\_centre** [numpy.ndarray, optional] radec coordinates of the Phase Centre. Shape (2,)

### Returns

**numpy.ndarray** lm Direction Cosines of shape (coord, 3)

`africanus.coordinates.lm_to_radec`(*lm*, *phase\_centre=None*)

Convert Direction Cosine lm coordinates to Right Ascension/Declination coordinates in radians, relative to the Phase Centre.

$$\delta = \arcsin(m \cos \delta_0 + n \sin \delta_0) \quad (5.7)$$

$$\alpha = \arctan\left(\frac{l}{n \cos \delta_0 - m \sin \delta_0}\right) \quad (5.8)$$

$$(5.9)$$

where  $\alpha$  is the Right Ascension of each coordinate and the phase centre and  $\delta_0$  is the Declination of the phase centre.

### Parameters

**lm** [numpy.ndarray] lm Direction Cosines of shape (coord, 2)

**phase\_centre** [numpy.ndarray, optional] radec coordinates of the Phase Centre. Shape (2,)

### Returns

`numpy.ndarray` radec coordinates of shape (coord, 2) where Right-Ascension and Declination are in the last 2 components, respectively.

`africanus.coordinates.lmn_to_radec(lmn, phase_centre=None)`

Convert Direction Cosine lm coordinates to Right Ascension/Declination coordinates in radians, relative to the Phase Centre.

$$\delta = \arcsin(m \cos \delta_0 + n \sin \delta_0) \quad (5.10)$$

$$\alpha = \arctan\left(\frac{l}{n \cos \delta_0 - m \sin \delta_0}\right) \quad (5.11)$$

$$(5.12)$$

where  $\alpha$  is the Right Ascension of each coordinate and the phase centre and  $\delta_0$  is the Declination of the phase centre.

#### Parameters

`lmn` [numpy.ndarray] lm Direction Cosines of shape (coord, 3)

`phase_centre` [numpy.ndarray, optional] radec coordinates of the Phase Centre. Shape (2, )

#### Returns

`numpy.ndarray` radec coordinates of shape (coord, 2) where Right-Ascension and Declination are in the last 2 components, respectively.

## 5.5.2 Dask

<code>radec_to_lm(radec[, phase_centre])</code>	Converts Right-Ascension/Declination coordinates in radians to a Direction Cosine lm coordinates, relative to the Phase Centre.
<code>radec_to_lmn(radec[, phase_centre])</code>	Converts Right-Ascension/Declination coordinates in radians to a Direction Cosine lm coordinates, relative to the Phase Centre.
<code>lm_to_radec(lm[, phase_centre])</code>	Convert Direction Cosine lm coordinates to Right Ascension/Declination coordinates in radians, relative to the Phase Centre.
<code>lmn_to_radec(lmn[, phase_centre])</code>	Convert Direction Cosine lm coordinates to Right Ascension/Declination coordinates in radians, relative to the Phase Centre.

`africanus.coordinates.dask.radec_to_lm(radec, phase_centre=None)`

Converts Right-Ascension/Declination coordinates in radians to a Direction Cosine lm coordinates, relative to the Phase Centre.

$$l = \cos \delta \sin \Delta\alpha \quad (5.13)$$

$$m = \sin \delta \cos \delta_0 - \cos \delta \sin \delta_0 \cos \Delta\alpha \quad (5.14)$$

$$n = \sqrt{1 - l^2 - m^2} - 1 \quad (5.15)$$

where  $\Delta\alpha = \alpha - \alpha_0$  is the difference between the Right Ascension of each coordinate and the phase centre and  $\delta_0$  is the Declination of the phase centre.

#### Parameters

`radec` [dask.array.Array] radec coordinates of shape (coord, 2) where Right-Ascension and Declination are in the last 2 components, respectively.

**phase\_centre** [`dask.array.Array`, optional] radec coordinates of the Phase Centre. Shape  $(2,)$

#### Returns

`dask.array.Array` lm Direction Cosines of shape  $(\text{coord}, 2)$

`africanus.coordinates.dask.radec_to_lmn(radec, phase_centre=None)`

Converts Right-Ascension/Declination coordinates in radians to a Direction Cosine lm coordinates, relative to the Phase Centre.

$$l = \cos \delta \sin \Delta\alpha \quad (5.16)$$

$$m = \sin \delta \cos \delta 0 - \cos \delta \sin \delta 0 \cos \Delta\alpha \quad (5.17)$$

$$n = \sqrt{1 - l^2 - m^2} - 1 \quad (5.18)$$

where  $\Delta\alpha = \alpha - \alpha 0$  is the difference between the Right Ascension of each coordinate and the phase centre and  $\delta 0$  is the Declination of the phase centre.

#### Parameters

**radec** [`dask.array.Array`] radec coordinates of shape  $(\text{coord}, 2)$  where Right-Ascension and Declination are in the last 2 components, respectively.

**phase\_centre** [`dask.array.Array`, optional] radec coordinates of the Phase Centre. Shape  $(2,)$

#### Returns

`dask.array.Array` lm Direction Cosines of shape  $(\text{coord}, 3)$

`africanus.coordinates.dask.lm_to_radec(lm, phase_centre=None)`

Convert Direction Cosine lm coordinates to Right Ascension/Declination coordinates in radians, relative to the Phase Centre.

$$\delta = \arcsin(m \cos \delta 0 + n \sin \delta 0) \quad (5.19)$$

$$\alpha = \arctan\left(\frac{l}{n \cos \delta 0 - m \sin \delta 0}\right) \quad (5.20)$$

$$(5.21)$$

where  $\alpha$  is the Right Ascension of each coordinate and the phase centre and  $\delta 0$  is the Declination of the phase centre.

#### Parameters

**lm** [`dask.array.Array`] lm Direction Cosines of shape  $(\text{coord}, 2)$

**phase\_centre** [`dask.array.Array`, optional] radec coordinates of the Phase Centre. Shape  $(2,)$

#### Returns

`dask.array.Array` radec coordinates of shape  $(\text{coord}, 2)$  where Right-Ascension and Declination are in the last 2 components, respectively.

`africanus.coordinates.dask.lmn_to_radec(lmn, phase_centre=None)`

Convert Direction Cosine lm coordinates to Right Ascension/Declination coordinates in radians, relative to the Phase Centre.

$$\delta = \arcsin(m \cos \delta 0 + n \sin \delta 0) \quad (5.22)$$

$$\alpha = \arctan\left(\frac{l}{n \cos \delta 0 - m \sin \delta 0}\right) \quad (5.23)$$

$$(5.24)$$

where  $\alpha$  is the Right Ascension of each coordinate and the phase centre and  $\delta 0$  is the Declination of the phase centre.

#### Parameters

**lmn** [dask.array.Array] lm Direction Cosines of shape (coord, 3)  
**phase\_centre** [dask.array.Array, optional] radec coordinates of the Phase Centre. Shape (2,)

#### Returns

dask.array.Array radec coordinates of shape (coord, 2) where Right-Ascension and Declination are in the last 2 components, respectively.

## 5.6 Sky Model

Functionality related to the Sky Model.

### 5.6.1 Coherency Conversion

Utilities for converting back and forth between stokes parameters and correlations

#### Numpy

---

<code>convert</code> (input, input_schema, output_schema)	This function converts forward and backward from stokes I,Q,U,V to both linear XX,XY,YX,YY and circular RR, RL, LR, LL correlations.
---	--

---

`africanus.model.coherency.convert`(*input, input\_schema, output\_schema*)

This function converts forward and backward from stokes I,Q,U,V to both linear XX,XY,YX,YY and circular RR, RL, LR, LL correlations.

For example, we can convert from stokes parameters to linear correlations:

```
stokes.shape == (10, 4, 4)
corrs = convert(stokes, ["I", "Q", "U", "V"],
                [['XX', 'XY'], ['YX', 'YY']])
assert corrs.shape == (10, 4, 2, 2)
```

Or circular correlations to stokes:

```
vis.shape == (10, 4, 2, 2)
stokes = convert(vis, [['RR', 'RL'], ['LR', 'LL']],
                  ['I', 'Q', 'U', 'V'])
assert stokes.shape == (10, 4, 4)
```

input can output can be arbitrarily nested or ordered lists, but the appropriate inputs must be present to produce the requested outputs.

The elements of `input` and `output` may be strings or integers representing stokes parameters or correlations. See the Notes for a full list.

### Parameters

`input` [numpy.ndarray] Complex or floating point input data of shape (`dim_1`, ..., `dim_n`, `icorr_1`, ..., `icorr_m`)

`input_schema` [list of str or int] A schema describing the `icorr_1`, ..., `icorr_m` dimension of `input`. Must have the same shape as the last dimensions of `input`.

`output_schema` [list of str or int] A schema describing the `ocorr_1`, ..., `ocorr_n` dimension of the return value.

### Returns

`result` [numpy.ndarray] Result of shape (`dim_1`, ..., `dim_n`, `ocorr_1`, ..., `ocorr_m`) The type may be floating point or promoted to complex depending on the combinations in `output`.

### Notes

Only stokes parameters, linear and circular correlations are currently handled, but the full list of id's and strings as defined in the [CASA documentation](#) is:

```
{} Undefined: 0, I: 1, Q: 2, U: 3, V: 4, RR: 5, RL: 6, LR: 7, LL: 8,
XX: 9, XY: 10, YX: 11, YY: 12, RX: 13, RY: 14, LX: 15, LY: 16,
XR: 17, XL: 18, YR: 19, YL: 20, PP: 21, PQ: 22, QP: 23, QQ:
24, RCircular: 25, LCircular: 26, Linear: 27, Ptotal: 28,
Plinear: 29, PFtotal: 30, PFlinear: 31, Pangle: 32 {}
```

## Cuda

---

`convert(inputs, input_schema, output_schema)`

This function converts forward and backward from stokes I,Q,U,V to both linear XX,XY,YX,YY and circular RR, RL, LR, LL correlations.

---

`africanus.model.coherency.cuda.convert(inputs, input_schema, output_schema)`

This function converts forward and backward from stokes I,Q,U,V to both linear XX,XY,YX,YY and circular RR, RL, LR, LL correlations.

For example, we can convert from stokes parameters to linear correlations:

```
stokes.shape == (10, 4, 4)
corrs = convert(stokes, ["I", "Q", "U", "V"],
                [['XX', 'XY'], ['YX', 'YY']])

assert corrs.shape == (10, 4, 2, 2)
```

Or circular correlations to stokes:

```
vis.shape == (10, 4, 2, 2)

stokes = convert(vis, [['RR', 'RL'], ['LR', 'LL']],
```

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```
[ 'I', 'Q', 'U', 'V'])  
  
assert stokes.shape == (10, 4, 4)
```

`input` can output can be arbitrarily nested or ordered lists, but the appropriate inputs must be present to produce the requested outputs.

The elements of `input` and `output` may be strings or integers representing stokes parameters or correlations. See the Notes for a full list.

### Parameters

**input** [cupy.ndarray] Complex or floating point input data of shape (`dim_1`, ..., `dim_n`, `icorr_1`, ..., `icorr_m`)

**input\_schema** [list of str or int] A schema describing the `icorr_1`, ..., `icorr_m` dimension of `input`. Must have the same shape as the last dimensions of `input`.

**output\_schema** [list of str or int] A schema describing the `ocorr_1`, ..., `ocorr_n` dimension of the return value.

### Returns

**result** [cupy.ndarray] Result of shape (`dim_1`, ..., `dim_n`, `ocorr_1`, ..., `ocorr_m`) The type may be floating point or promoted to complex depending on the combinations in `output`.

### Notes

Only stokes parameters, linear and circular correlations are currently handled, but the full list of id's and strings as defined in the [CASA documentation](#) is:

```
{} Undefined: 0, I: 1, Q: 2, U: 3, V: 4, RR: 5, RL: 6, LR: 7, LL: 8,  
XX: 9, XY: 10, YX: 11, YY: 12, RX: 13, RY: 14, LX: 15, LY: 16,  
XR: 17, XL: 18, YR: 19, YL: 20, PP: 21, PQ: 22, QP: 23, QQ:  
24, RCircular: 25, LCircular: 26, Linear: 27, Ptotal: 28,  
Plinear: 29, PFtotal: 30, PFlinear: 31, Pangle: 32 {}
```

### Dask

<code>convert</code> ( <code>input</code> , <code>input_schema</code> , <code>output_schema</code> )	This function converts forward and backward from stokes I,Q,U,V to both linear XX,XY,YX,YY and circular RR, RL, LR, LL correlations.
--	--

`africanus.model.coherency.dask.convert`(`input`, `input_schema`, `output_schema`)

This function converts forward and backward from stokes I,Q,U,V to both linear XX,XY,YX,YY and circular RR, RL, LR, LL correlations.

For example, we can convert from stokes parameters to linear correlations:

```
stokes.shape == (10, 4, 4)  
corrs = convert(stokes, ["I", "Q", "U", "V"],  
[[ "XX", "XY"], [ "YX", "YY"]])
```

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```
assert corrs.shape == (10, 4, 2, 2)
```

Or circular correlations to stokes:

```
vis.shape == (10, 4, 2, 2)

stokes = convert(vis, [['RR', 'RL'], ['LR', 'LL']],
                 ['I', 'Q', 'U', 'V'])

assert stokes.shape == (10, 4, 4)
```

`input` can `output` can be arbitrarily nested or ordered lists, but the appropriate inputs must be present to produce the requested outputs.

The elements of `input` and `output` may be strings or integers representing stokes parameters or correlations. See the Notes for a full list.

#### Parameters

`input` [dask.array.Array] Complex or floating point input data of shape (`dim_1`, ..., `dim_n`, `icorr_1`, ..., `icorr_m`)  
`input_schema` [list of str or int] A schema describing the `icorr_1`, ..., `icorr_m` dimension of `input`. Must have the same shape as the last dimensions of `input`.  
`output_schema` [list of str or int] A schema describing the `ocorr_1`, ..., `ocorr_n` dimension of the return value.

#### Returns

`result` [dask.array.Array] Result of shape (`dim_1`, ..., `dim_n`, `ocorr_1`, ..., `ocorr_m`) The type may be floating point or promoted to complex depending on the combinations in `output`.

#### Notes

Only stokes parameters, linear and circular correlations are currently handled, but the full list of id's and strings as defined in the [CASA documentation](#) is:

```
{} Undefined: 0, I: 1, Q: 2, U: 3, V: 4, RR: 5, RL: 6, LR: 7, LL: 8,
XX: 9, XY: 10, YX: 11, YY: 12, RX: 13, RY: 14, LX: 15, LY: 16,
XR: 17, XL: 18, YR: 19, YL: 20, PP: 21, PQ: 22, QP: 23, QQ:
24, RCircular: 25, LCircular: 26, Linear: 27, Ptotal: 28,
Plinear: 29, PFtotal: 30, PFlinear: 31, Pangle: 32 {}
```

## 5.6.2 Spectral Model

Functionality for computing a Spectral Model.

### Numpy

---

<code>spectral_model(stokes, spi, ref_freq, frequency)</code>	Compute a spectral model, per polarisation.
---	---

---

`africanus.model.spectral.spectral_model(stokes, spi, ref_freq, frequency, base=0)`

Compute a spectral model, per polarisation.

$$I(\lambda) = I_0 \prod_{i=1} (\lambda/\lambda_0)^{\alpha_i} \quad (5.25)$$

$$\ln(I(\lambda)) = \sum_{i=0} \alpha_i \ln(\lambda/\lambda_0)^i \text{ where } \alpha_0 = \ln I_0 \quad (5.26)$$

$$\log_{10}(I(\lambda)) = \sum_{i=0} \alpha_i \log_{10}(\lambda/\lambda_0)^i \text{ where } \alpha_0 = \log_{10} I_0 \quad (5.27)$$

(5.28)

### Parameters

**stokes** [numpy.ndarray] Stokes parameters of shape (source,) or (source, pol). If a pol dimension is present, then it must also be present on spi.

**spi** [numpy.ndarray] Spectral index of shape (source, spi-comps) or (source, spi-comps, pol).

**ref\_freq** [numpy.ndarray] Reference frequencies of shape (source,)

**frequencies** [numpy.ndarray] Frequencies of shape (chan,)

**base** [{"std", "log", "log10"} or {0, 1, 2} or list.] string or corresponding enumeration specifying the polynomial base. Defaults to 0.

If a list is provided, a polynomial base can be specified for each stokes parameter or polarisation in the pol dimension.

string specification of the base is only supported in python 3. while the corresponding integer enumerations are supported on all python versions.

### Returns

**spectral\_model** [numpy.ndarray] Spectral Model of shape (source, chan) or (source, chan, pol).

### Task

---

<code>spectral_model(stokes, spi, ref_freq, ...[, ...])</code>	Compute a spectral model, per polarisation.
--	---

---

`africanus.model.spectral.dask.spectral_model(stokes, spi, ref_freq, frequencies, base=0)`

Compute a spectral model, per polarisation.

$$I(\lambda) = I_0 \prod_{i=1} (\lambda/\lambda_0)^{\alpha_i} \quad (5.29)$$

$$\ln(I(\lambda)) = \sum_{i=0} \alpha_i \ln(\lambda/\lambda_0)^i \text{ where } \alpha_0 = \ln I_0 \quad (5.30)$$

$$\log_{10}(I(\lambda)) = \sum_{i=0} \alpha_i \log_{10}(\lambda/\lambda_0)^i \text{ where } \alpha_0 = \log_{10} I_0 \quad (5.31)$$

(5.32)

### Parameters

**stokes** [dask.array.Array] Stokes parameters of shape (source,) or (source, pol). If a pol dimension is present, then it must also be present on spi.

**spi** [dask.array.Array] Spectral index of shape (source, spi-comps) or (source, spi-comps, pol).

**ref\_freq** [dask.array.Array] Reference frequencies of shape (source,)

**frequencies** [dask.array.Array] Frequencies of shape (chan,)

**base** [{“std”, “log”, “log10”} or {0, 1, 2} or list.] string or corresponding enumeration specifying the polynomial base. Defaults to 0.

If a list is provided, a polynomial base can be specified for each stokes parameter or polarisation in the pol dimension.

string specification of the base is only supported in python 3. while the corresponding integer enumerations are supported on all python versions.

### Returns

**spectral\_model** [dask.array.Array] Spectral Model of shape (source, chan) or (source, chan, pol).

## 5.6.3 Spectral Index

Functionality related to the spectral index.

For example, we may want to compute the spectral indices of components in a sky model defined by

$$I(\nu) = I(\nu_0) \left( \frac{\nu}{\nu_0} \right)^\alpha$$

where  $\nu$  are frequencies at which we want to construct the intensity of a Stokes I image and the  $\nu_0$  is the corresponding reference frequency. The spectral index  $\alpha$  determines how quickly the intensity grows or decays as a function of frequency. Given a list of model image components (preferably with the residuals added back in) we can recover the corresponding spectral indices and reference intensities using the `fit_spi_components()` function. This will also return a lower bound on the associated uncertainties on these components.

## Numpy

---

<code>fit_spi_components(data, weights, freqs, freq0)</code>	Computes the spectral indices and the intensity at the reference frequency of a spectral index model:
<code>africanus.model.spi.fit_spi_components(data, weights, freqs, freq0, alphai=None, I0i=None, beam=None, tol=0.0001, maxiter=100)</code>	

Computes the spectral indices and the intensity at the reference frequency of a spectral index model:

$$I(\nu) = A(\nu)I(\nu_0) \left( \frac{\nu}{\nu_0} \right)^\alpha$$

where  $I(\nu)$  is the apparent source spectrum,  $A(\nu)$  is the beam model for each component as a function of frequency.

### Parameters

- data** [numpy.ndarray] array of shape (comps, chan) The noisy data as a function of frequency.
- weights** [numpy.ndarray] array of shape (chan,) Inverse of variance on each frequency axis.
- freqs** [numpy.ndarray] frequencies of shape (chan,)
- freq0** [float] Reference frequency
- alphai** [numpy.ndarray, optional] array of shape (comps,) Initial guess for the alphas. Defaults to -0.7.
- I0i** [numpy.ndarray, optional] array of shape (comps,) Initial guess for the intensities at the reference frequency. Defaults to 1.0.
- beam\_comps** [numpy.ndarray, optional] array of shape (comps, chan) Power beam for each component as a function of frequency.
- tol** [float, optional] Solver absolute tolerance (optional). Defaults to 1e-6.
- maxiter** [int, optional] Solver maximum iterations (optional). Defaults to 100.
- dtype** [np.dtype, optional] Datatype of result. Should be either np.float32 or np.float64. Defaults to np.float64.

### Returns

- out** [numpy.ndarray] array of shape (4, comps) The fitted components arranged as [alphas, alphavars, I0s, I0vars]

## Dask

---

<code>fit_spi_components(data, weights, freqs, freq0)</code>	Computes the spectral indices and the intensity at the reference frequency of a spectral index model:
<code>africanus.model.spi.dask.fit_spi_components(data, weights, freqs, freq0, alphai=None, I0i=None, beam=None, tol=1e-05, maxiter=100)</code>	

Computes the spectral indices and the intensity at the reference frequency of a spectral index model:

$$I(\nu) = A(\nu)I(\nu_0) \left( \frac{\nu}{\nu_0} \right)^\alpha$$

where  $I(\nu)$  is the apparent source spectrum,  $A(\nu)$  is the beam model for each component as a function of frequency.

#### Parameters

**data** [dask.array.Array] array of shape (comps, chan) The noisy data as a function of frequency.

**weights** [dask.array.Array] array of shape (chan,) Inverse of variance on each frequency axis.

**freqs** [dask.array.Array] frequencies of shape (chan,)

**freq0** [float] Reference frequency

**alphai** [dask.array.Array, optional] array of shape (comps,) Initial guess for the alphas. Defaults to -0.7.

**I0i** [dask.array.Array, optional] array of shape (comps,) Initial guess for the intensities at the reference frequency. Defaults to 1.0.

**beam\_comps** [dask.array.Array, optional] array of shape (comps, chan) Power beam for each component as a function of frequency.

**tol** [float, optional] Solver absolute tolerance (optional). Defaults to 1e-6.

**maxiter** [int, optional] Solver maximum iterations (optional). Defaults to 100.

**dtype** [np.dtype, optional] Datatype of result. Should be either np.float32 or np.float64. Defaults to np.float64.

#### Returns

**out** [dask.array.Array] array of shape (4, comps) The fitted components arranged as [alphas, alphavars, I0s, I0vars]

## 5.6.4 Source Morphology

Shape functions for different Source Morphologies

### Numpy

---

<code>gaussian(uvw, frequency, shape_params)</code>	Computes the Gaussian Shape Function.
---	---------------------------------------

---

`africanus.model.shape.gaussian(uvw, frequency, shape_params)`  
Computes the Gaussian Shape Function.

$$\begin{aligned}\lambda' &= 2\lambda\pi \\ r &= \frac{e_{min}}{e_{maj}} \\ u_1 &= (u e_{maj} \cos(\alpha) - v e_{maj} \sin(\alpha))r\lambda' \\ v_1 &= (u e_{maj} \sin(\alpha) - v e_{maj} \cos(\alpha))\lambda' \\ \text{shape} &= e^{(-u_1^2 - v_1^2)}\end{aligned}$$

where:

- $u$  and  $v$  are the UV coordinates and  $\lambda$  the frequency.
- $e_{maj}$  and  $e_{min}$  are the major and minor axes and  $\alpha$  the position angle.

**Parameters**

**uvw** [numpy.ndarray] UVW coordinates of shape (row, 3)

**frequency** [numpy.ndarray] frequencies of shape (chan,)

**shape\_param** [numpy.ndarray] Gaussian Shape Parameters of shape (source, 3) where the second dimension contains the (*emajor*, *eminor*, *angle*) parameters describing the shape of the Gaussian

**Returns**

**gauss\_shape** [numpy.ndarray] Shape parameters of shape (source, row, chan)

**Dask**


---

***gaussian*(uvw, frequency, shape\_params)**

Computes the Gaussian Shape Function.

---

**africanus.model.shape.dask.gaussian(uvw, frequency, shape\_params)**

Computes the Gaussian Shape Function.

$$\begin{aligned}\lambda' &= 2\lambda\pi \\ r &= \frac{e_{min}}{e_{maj}} \\ u_1 &= (u e_{maj} \cos(\alpha) - v e_{maj} \sin(\alpha))r\lambda' \\ v_1 &= (u e_{maj} \sin(\alpha) - v e_{maj} \cos(\alpha))\lambda' \\ \text{shape} &= e^{(-u_1^2 - v_1^2)}\end{aligned}$$

where:

- $u$  and  $v$  are the UV coordinates and  $\lambda$  the frequency.
- $e_{maj}$  and  $e_{min}$  are the major and minor axes and  $\alpha$  the position angle.

**Parameters**

**uvw** [dask.array.Array] UVW coordinates of shape (row, 3)

**frequency** [dask.array.Array] frequencies of shape (chan,)

**shape\_param** [dask.array.Array] Gaussian Shape Parameters of shape (source, 3) where the second dimension contains the (*emajor*, *eminor*, *angle*) parameters describing the shape of the Gaussian

**Returns**

**gauss\_shape** [dask.array.Array] Shape parameters of shape (source, row, chan)

## 5.6.5 WSClean Spectral Model

Utilities for creating a spectral model from a wsclean component file.

### Numpy

---

<code>load(filename)</code>	Loads wsclean component model.
<code>spectra(I, coeffs, log_poly, ref_freq, frequency)</code>	Produces a spectral model from a polynomial expansion of a wsclean file model.

---

`africanus.model.wsclean.load(filename)`

Loads wsclean component model.

```
sources = load("components.txt")
sources = dict(sources) # Convert to dictionary

I = sources["I"]
ref_freq = sources["ReferenceFrequency"]
```

See the [WSClean Component List](#) for further details.

#### Parameters

**filename** [str or iterable] Filename of wsclean model file or iterable producing the lines of the file.

#### Returns

**list of (name, list of values) tuples** list of column (name, value) tuples

See also:

`africanus.model.wsclean.spectra`

`africanus.model.wsclean.spectra(I, coeffs, log_poly, ref_freq, frequency)`

Produces a spectral model from a polynomial expansion of a wsclean file model. Depending on how `log_poly` is set ordinary or logarithmic polynomials are used to produce the expansion:

$$\begin{aligned} flux(\lambda) &= I_0 + \sum_{c=0} \text{coeffs}(c)(\lambda/\lambda_{ref} - 1)^{c+1} \\ flux(\lambda) &= \exp \left( \log I_0 + \sum_{c=0} \text{coeffs}(c) \log(\lambda/\lambda_{ref})^{c+1} \right) \end{aligned}$$

See the [WSClean Component List](#) for further details.

#### Parameters

**I** [numpy.ndarray] flux density in Janskys at the reference frequency of shape (source, )

**coeffs** [numpy.ndarray] Polynomial coefficients for each source of shape (source, comp)

**log\_poly** [numpy.ndarray or bool] boolean array of shape (source, ) indicating whether logarithmic (True) or ordinary (False) polynomials should be used.

**ref\_freq** [numpy.ndarray] Source reference frequencies of shape (source, )

**frequency** [numpy.ndarray] frequencies of shape (chan, )

**Returns**

**spectral\_model** [numpy.ndarray] Spectral Model of shape (source, chan)

See also:

[`africanus.model.wsclean.load`](#)

**Task**


---

<code>spectra(stokes, spi, log_si, ref_freq, frequency)</code>	Produces a spectral model from a polynomial expansion of a wsclean file model.
--	--

---

`africanus.model.wsclean.dask.spectra(stokes, spi, log_si, ref_freq, frequency)`

Produces a spectral model from a polynomial expansion of a wsclean file model. Depending on how `log_poly` is set ordinary or logarithmic polynomials are used to produce the expansion:

$$\begin{aligned} flux(\lambda) &= I_0 + \sum_{c=0} \text{coeffs}(c)(\lambda/\lambda_{ref} - 1)^{c+1} \\ flux(\lambda) &= \exp \left( \log I_0 + \sum_{c=0} \text{coeffs}(c) \log(\lambda/\lambda_{ref})^{c+1} \right) \end{aligned}$$

See the [WSClean Component List](#) for further details.

**Parameters**

**I** [dask.array.Array] flux density in Janskys at the reference frequency of shape (source, )

**coeffs** [dask.array.Array] Polynomial coefficients for each source of shape (source, comp)

**log\_poly** [dask.array.Array or bool] boolean array of shape (source, ) indicating whether logarithmic (True) or ordinary (False) polynomials should be used.

**ref\_freq** [dask.array.Array] Source reference frequencies of shape (source, )

**frequency** [dask.array.Array] frequencies of shape (chan, )

**Returns**

**spectral\_model** [dask.array.Array] Spectral Model of shape (source, chan)

See also:

[`africanus.model.wsclean.load`](#)

## 5.7 Averaging

Routines for averaging visibility data.

### 5.7.1 Time and Channel Averaging

The routines in this section average row-based samples by:

1. Averaging samples of consecutive **time** values into bins defined by a period of **time\_bin\_secs** seconds.
2. Averaging channel data into equally sized bins of **chan\_bin\_size**.

In order to achieve this, a **baseline x time** ordering is established over the input data where **baseline** corresponds to the unique (**ANTENNA1**, **ANTENNA2**) pairs and **time** corresponds to the unique, monotonically increasing **TIME** values associated with the rows of a Measurement Set.

Baseline	T0	T1	T2	T3	T4
(0, 0)	0.1	0.2	0.3	0.4	0.5
(0, 1)	0.1	0.2	0.3	0.4	0.5
(0, 2)	0.1	0.2	X	0.4	0.5
(1, 1)	0.1	0.2	0.3	0.4	0.5
(1, 2)	0.1	0.2	0.3	0.4	0.5
(2, 2)	0.1	0.2	0.3	0.4	0.5

It is possible for times or baselines to be missing. In the above example, T2 is missing for baseline (0, 2).

**Warning:** The above requires unique lexicographical combinations of (TIME, ANTENNA1, ANTENNA2). This can usually be achieved by suitably partitioning input data on indexing rows, DATA\_DESC\_ID and SCAN\_NUMBER in particular.

For each baseline, adjacent time's are assigned to a bin if  $h_c - h_e/2 - (l_c - l_e/2) < \text{time\_bin\_secs}$ , where  $h_c$  and  $l_c$  are the upper and lower time and  $h_e$  and  $l_e$  are the upper and lower intervals, taken from the **INTERVAL** column. Note that no distinction is made between flagged and unflagged data when establishing the endpoints in the bin.

The reason for this is that the [Measurement Set v2.0 Specification](#) specifies that **TIME** and **INTERVAL** columns are defined as containing the *nominal* time and period at which the visibility was sampled. This means that their values includie valid, flagged and missing data. Thus, averaging a regular high-resolution **baseline x htime** grid should produce a regular low-resolution **baseline x ltime** grid (**htime > ltime**) in the presence of bad data.

By contrast, other columns such as **TIME\_CENTROID** and **EXPOSURE** contain the *effective* time and period as they exclude missing and bad data. Their increased accuracy, and therefore variability means that they are unsuitable for establishing a grid over the data.

To summarise, the averaged times in each bin establish a map:

- from possibly unordered input rows.
- to a reduced set of output rows ordered by averaged (TIME, ANTENNA1, ANTENNA2).

#### Flagged Data Handling

Both **FLAG\_ROW** and **FLAG** columns may be supplied to the averager, but they should be consistent with each other. The averager will throw an exception if this is not the case, rather than making an assumption as to which is correct.

When provided with flags, the averager will output averages for bins that are completely flagged.

Part of the reason for this is that the specifies that the **TIME** and **INTERVAL** columns represent the *nominal* time and interval values. This means that they should represent valid as well as flagged or missing data in their computation.

By contrast, most other columns such as **TIME\_CENTROID** and **EXPOSURE**, contain the *effective* values and should only include valid, unflagged data.

To support this:

1. **TIME** and **INTERVAL** are averaged using both flagged and unflagged samples.
2. Other columns, such as **TIME\_CENTROID** are handled as follows:
  1. If the bin contains some unflagged data, only this data is used to calculate average.
  2. If the bin is completely flagged, the average of all samples (which are all flagged) will be used.
  3. In both cases, a completely flagged bin will have its flag set.
4. To support the two cases, twice the memory of the output array is required to track both averages, but only one array of merged values is returned.

## Guarantees

1. Averaged output data will be lexicographically ordered by (**TIME**, **ANTENNA1**, **ANTENNA2**)
2. **TIME** and **INTERVAL** columns always contain the *nominal* average and sum and therefore contain both and missing or unflagged data.
3. Other columns will contain the *effective* average and will contain only valid data *except* when all data in the bin is flagged.
4. Completely flagged bins will be set as flagged in both the *nominal* and *effective* case.
5. Certain columns are averaged, while others are summed, or simply assigned to the last value in the bin in the case of antenna indices.
6. **Visibility data** is averaged by multiplying and dividing by **WEIGHT\_SPECTRUM** or **WEIGHT** or natural weighting, in order of priority.

$$\frac{\sum v_i w_i}{\sum w_i}$$

7. **SIGMA\_SPECTRUM** is averaged by multiplying and dividing by **WEIGHT\_SPECTRUM** or **WEIGHT** or natural weighting, in order of priority and availability.

**SIGMA** is only averaged with **WEIGHT** or natural weighting.

$$\sqrt{\frac{\sum w_i^2 \sigma_i^2}{(\sum w_i)^2}}$$

The following table summarizes the handling of each column in the main Measurement Set table:

Column	Unflagged/Flagged sample handling	Aggregation Method	Required
TIME	Nominal	Mean	Yes
INTERVAL	Nominal	Sum	Yes
ANTENNA1	Nominal	Assigned to Last Input	Yes
ANTENNA2	Nominal	Assigned to Last Input	Yes
TIME_CENTROID	Effective	Mean	No
EXPOSURE	Effective	Sum	No
FLAG_ROW	Effective	Set if All Inputs Flagged	No
UVW	Effective	Mean	No
WEIGHT	Effective	Sum	No
SIGMA	Effective	Weighted Mean	No
DATA (vis)	Effective	Weighted Mean	No
FLAG	Effective	Set if All Inputs Flagged	No
WEIGHT_SPECTRUM	Effective	Sum	No
SIGMA_SPECTRUM	Effective	Weighted Mean	No

The following SPECTRAL\_WINDOW sub-table columns are averaged as follows:

Column	Aggregation Method
CHAN_FREQ	Mean
CHAN_WIDTH	Sum
EFFECTIVE_BW	Sum
RESOLUTION	Sum

## Dask Implementation

The dask implementation chunks data up by row and channel and averages each chunk independently of values in other chunks. This should be kept in mind if one wishes to maintain a particular ordering in the output dask arrays.

Typically, Measurement Set data is monotonically ordered in time. To maintain this guarantee in output dask arrays, the chunks will need to be separated by distinct time values. Practically speaking this means that the first and second chunk should not both contain value time 0.1, for example.

## Numpy

<code>time_and_channel1</code> (time, interval, antenna1, ...)	Averages in time and channel.
<code>bda</code> (time, interval, antenna1, antenna2[, ...])	Averages in time and channel, dependent on baseline length.

```
africanus.averaging.time_and_channel1(time, interval, antenna1, antenna2, time_centroid=None,  
exposure=None, flag_row=None, uvw=None, weight=None,  
sigma=None, chan_freq=None, chan_width=None,  
effective_bw=None, resolution=None, visibilities=None, flag=None,  
weight_spectrum=None, sigma_spectrum=None,  
time_bin_secs=1.0, chan_bin_size=1)
```

Averages in time and channel.

### Parameters

`time` [numpy.ndarray] Time values of shape (row, ).

**interval** [numpy.ndarray] Interval values of shape (row, ).

**antenna1** [numpy.ndarray] First antenna indices of shape (row, )

**antenna2** [numpy.ndarray] Second antenna indices of shape (row, )

**time\_centroid** [numpy.ndarray, optional] Time centroid values of shape (row, )

**exposure** [numpy.ndarray, optional] Exposure values of shape (row, )

**flag\_row** [numpy.ndarray, optional] Flagged rows of shape (row, ).

**uvw** [numpy.ndarray, optional] UVW coordinates of shape (row, 3).

**weight** [numpy.ndarray, optional] Weight values of shape (row, corr).

**sigma** [numpy.ndarray, optional] Sigma values of shape (row, corr).

**chan\_freq** [numpy.ndarray, optional] Channel frequencies of shape (chan, ).

**chan\_width** [numpy.ndarray, optional] Channel widths of shape (chan, ).

**effective\_bw** [numpy.ndarray, optional] Effective channel bandwidth of shape (chan, ).

**resolution** [numpy.ndarray, optional] Effective channel resolution of shape (chan, ).

**visibilities** [numpy.ndarray or tuple of numpy.ndarray, optional] Visibility data of shape (row, chan, corr). Tuples of visibilities arrays may be supplied, in which case tuples will be output.

**flag** [numpy.ndarray, optional] Flag data of shape (row, chan, corr).

**weight\_spectrum** [numpy.ndarray, optional] Weight spectrum of shape (row, chan, corr).

**sigma\_spectrum** [numpy.ndarray, optional] Sigma spectrum of shape (row, chan, corr).

**time\_bin\_secs** [float, optional] Maximum summed interval in seconds to include within a bin. Defaults to 1.0.

**chan\_bin\_size** [int, optional] Number of bins to average together. Defaults to 1.

#### Returns

**namedtuple** A namedtuple whose entries correspond to the input arrays. Output arrays will be None if the inputs were None.

#### Notes

The implementation currently requires unique lexicographical combinations of (TIME, ANTENNA1, ANTENNA2). This can usually be achieved by suitably partitioning input data on indexing rows, DATA\_DESC\_ID and SCAN\_NUMBER in particular.

```
africanus.averaging.bda(time, interval, antenna1, antenna2, time_centroid=None, exposure=None,  
    flag_row=None, uvw=None, weight=None, sigma=None, chan_freq=None,  
    chan_width=None, effective_bw=None, resolution=None, visibilities=None,  
    flag=None, weight_spectrum=None, sigma_spectrum=None, max_uvw_dist=None,  
    max_fov=3.0, decorrelation=0.98, time_bin_secs=None, min_nchan=1)
```

Averages in time and channel, dependent on baseline length.

#### Parameters

**time** [numpy.ndarray] Time values of shape (row, ).

**interval** [numpy.ndarray] Interval values of shape (row, ).

**antenna1** [numpy.ndarray] First antenna indices of shape (row, )

**antenna2** [numpy.ndarray] Second antenna indices of shape (row, )

**time\_centroid** [numpy.ndarray, optional] Time centroid values of shape (row, )

**exposure** [numpy.ndarray, optional] Exposure values of shape (row, )

**flag\_row** [numpy.ndarray, optional] Flagged rows of shape (row, ).

**uvw** [numpy.ndarray, optional] UVW coordinates of shape (row, 3).

**weight** [numpy.ndarray, optional] Weight values of shape (row, corr).

**sigma** [numpy.ndarray, optional] Sigma values of shape (row, corr).

**chan\_freq** [numpy.ndarray, optional] Channel frequencies of shape (chan, ).

**chan\_width** [numpy.ndarray, optional] Channel widths of shape (chan, ).

**effective\_bw** [numpy.ndarray, optional] Effective channel bandwidth of shape (chan, ).

**resolution** [numpy.ndarray, optional] Effective channel resolution of shape (chan, ).

**visibilities** [numpy.ndarray or tuple of numpy.ndarray, optional] Visibility data of shape (row, chan, corr). Tuples of visibilities arrays may be supplied, in which case tuples will be output.

**flag** [numpy.ndarray, optional] Flag data of shape (row, chan, corr).

**weight\_spectrum** [numpy.ndarray, optional] Weight spectrum of shape (row, chan, corr).

**sigma\_spectrum** [numpy.ndarray, optional] Sigma spectrum of shape (row, chan, corr).

**max\_uv\_dist** [float, optional] Maximum UVW distance. Will be inferred from the UVW coordinates if not supplied.

**max\_fov** [float] Maximum Field of View Radius. Defaults to 3 degrees.

**decorrelation** [float] Acceptable amount of decorrelation. This is a floating point value between 0.0 and 1.0.

**time\_bin\_secs** [float, optional] Maximum number of seconds worth of data that can be aggregated into a bin. Defaults to None in which case the value is only bounded by the decorrelation factor and the field of view.

**min\_nchan** [int, optional] Minimum number of channels in an averaged sample. Useful in cases where imagers expect at least *min\_nchan* channels. Defaults to 1.

#### Returns

**namedtuple** A namedtuple whose entries correspond to the input arrays. Output arrays will be None if the inputs were None. See the Notes for an explanation of the output formats.

## Notes

In all cases arrays starting with (row, chan) and (row,) dimensions are respectively averaged and expanded into a (rowchan,) dimension, as the number of channels varies per output row.

The output namedtuple contains an *offsets* array of shape (out\_rows + 1,) encoding the starting offsets of each output row, as well as a single entry at the end such that `np.diff(offsets)` produces the number of channels for each output row.

```
avg = bda(...)
time = avg.time[:-1]
out_chans = np.diff(avg.offsets)
```

The implementation currently requires unique lexicographical combinations of (TIME, ANTENNA1, ANTENNA2). This can usually be achieved by suitably partitioning input data on indexing rows, DATA\_DESC\_ID and SCAN\_NUMBER in particular.

## Task

<code>time_and_channel</code> (time, interval, antenna1, ...)	Averages in time and channel.
<code>bda</code> (time, interval, antenna1, antenna2[, ...])	Averages in time and channel, dependent on baseline length.

```
africanus.averaging.dask.time_and_channel(time, interval, antenna1, antenna2, time_centroid=None,
                                             exposure=None, flag_row=None, uvw=None, weight=None,
                                             sigma=None, chan_freq=None, chan_width=None,
                                             effective_bw=None, resolution=None, visibilities=None,
                                             flag=None, weight_spectrum=None, sigma_spectrum=None,
                                             time_bin_secs=1.0, chan_bin_size=1)
```

Averages in time and channel.

### Parameters

- `time` [dask.array.Array] Time values of shape (row,).
- `interval` [dask.array.Array] Interval values of shape (row,).
- `antenna1` [dask.array.Array] First antenna indices of shape (row,)
- `antenna2` [dask.array.Array] Second antenna indices of shape (row,)
- `time_centroid` [dask.array.Array, optional] Time centroid values of shape (row,)
- `exposure` [dask.array.Array, optional] Exposure values of shape (row,)
- `flag_row` [dask.array.Array, optional] Flagged rows of shape (row,).
- `uvw` [dask.array.Array, optional] UVW coordinates of shape (row, 3).
- `weight` [dask.array.Array, optional] Weight values of shape (row, corr).
- `sigma` [dask.array.Array, optional] Sigma values of shape (row, corr).
- `chan_freq` [dask.array.Array, optional] Channel frequencies of shape (chan,).
- `chan_width` [dask.array.Array, optional] Channel widths of shape (chan,).
- `effective_bw` [dask.array.Array, optional] Effective channel bandwidth of shape (chan,).
- `resolution` [dask.array.Array, optional] Effective channel resolution of shape (chan,).

**visibilities** [dask.array.Array or tuple of dask.array.Array, optional] Visibility data of shape (row, chan, corr). Tuples of visibilities arrays may be supplied, in which case tuples will be output.

**flag** [dask.array.Array, optional] Flag data of shape (row, chan, corr).

**weight\_spectrum** [dask.array.Array, optional] Weight spectrum of shape (row, chan, corr).

**sigma\_spectrum** [dask.array.Array, optional] Sigma spectrum of shape (row, chan, corr).

**time\_bin\_secs** [float, optional] Maximum summed interval in seconds to include within a bin. Defaults to 1.0.

**chan\_bin\_size** [int, optional] Number of bins to average together. Defaults to 1.

#### Returns

**namedtuple** A namedtuple whose entries correspond to the input arrays. Output arrays will be None if the inputs were None.

#### Notes

The implementation currently requires unique lexicographical combinations of (TIME, ANTENNA1, ANTENNA2). This can usually be achieved by suitably partitioning input data on indexing rows, DATA\_DESC\_ID and SCAN\_NUMBER in particular.

```
africanus.averaging.dask.bda(time, interval, antenna1, antenna2, time_centroid=None, exposure=None,
                               flag_row=None, uvw=None, weight=None, sigma=None, chan_freq=None,
                               chan_width=None, effective_bw=None, resolution=None, visibilities=None,
                               flag=None, weight_spectrum=None, sigma_spectrum=None,
                               max_uvw_dist=None, max_fov=3.0, decorrelation=0.98, time_bin_secs=None,
                               min_nchan=1, format='flat')
```

Averages in time and channel, dependent on baseline length.

#### Parameters

**time** [dask.array.Array] Time values of shape (row, ).

**interval** [dask.array.Array] Interval values of shape (row, ).

**antenna1** [dask.array.Array] First antenna indices of shape (row, )

**antenna2** [dask.array.Array] Second antenna indices of shape (row, )

**time\_centroid** [dask.array.Array, optional] Time centroid values of shape (row, )

**exposure** [dask.array.Array, optional] Exposure values of shape (row, )

**flag\_row** [dask.array.Array, optional] Flagged rows of shape (row, ).

**uvw** [dask.array.Array, optional] UVW coordinates of shape (row, 3).

**weight** [dask.array.Array, optional] Weight values of shape (row, corr).

**sigma** [dask.array.Array, optional] Sigma values of shape (row, corr).

**chan\_freq** [dask.array.Array, optional] Channel frequencies of shape (chan, ).

**chan\_width** [dask.array.Array, optional] Channel widths of shape (chan, ).

**effective\_bw** [dask.array.Array, optional] Effective channel bandwidth of shape (chan, ).

**resolution** [dask.array.Array, optional] Effective channel resolution of shape (chan, ).

**visibilities** [dask.array.Array or tuple of dask.array.Array, optional] Visibility data of shape (row, chan, corr). Tuples of visibilities arrays may be supplied, in which case tuples will be output.

**flag** [dask.array.Array, optional] Flag data of shape (row, chan, corr).

**weight\_spectrum** [dask.array.Array, optional] Weight spectrum of shape (row, chan, corr).

**sigma\_spectrum** [dask.array.Array, optional] Sigma spectrum of shape (row, chan, corr).

**max\_uvw\_dist** [float, optional] Maximum UVW distance. Will be inferred from the UVW coordinates if not supplied.

**max\_fov** [float] Maximum Field of View Radius. Defaults to 3 degrees.

**decorrelation** [float] Acceptable amount of decorrelation. This is a floating point value between 0.0 and 1.0.

**time\_bin\_secs** [float, optional] Maximum number of seconds worth of data that can be aggregated into a bin. Defaults to None in which case the value is only bounded by the decorrelation factor and the field of view.

**min\_nchan** [int, optional] Minimum number of channels in an averaged sample. Useful in cases where imagers expect at least *min\_nchan* channels. Defaults to 1.

### Returns

**namedtuple** A namedtuple whose entries correspond to the input arrays. Output arrays will be None if the inputs were None. See the Notes for an explanation of the output formats.

### Notes

In all cases arrays starting with (row, chan) and (row,) dimensions are respectively averaged and expanded into a (rowchan,) dimension, as the number of channels varies per output row.

The output namedtuple contains an *offsets* array of shape (out\_rows + 1,) encoding the starting offsets of each output row, as well as a single entry at the end such that `np.diff(offsets)` produces the number of channels for each output row.

```
avg = bda(...)  
time = avg.time[avg.offsets[:-1]]  
out_chans = np.diff(avg.offsets)
```

The implementation currently requires unique lexicographical combinations of (TIME, ANTENNA1, ANTENNA2). This can usually be achieved by suitably partitioning input data on indexing rows, DATA\_DESC\_ID and SCAN\_NUMBER in particular.

## 5.8 Utilities

### 5.8.1 Command Line

---

<code>parse_python_assigns(assign_str)</code>	Parses a string, containing assign statements into a dictionary.
---	--

---

`africanus.util.cmdline.parse_python_assigns(assign_str)`

Parses a string, containing assign statements into a dictionary.

```
data = parse_python_assigns("beta=5.6; l=[2,3], s='hello, world'")  
  
assert data == {  
    'beta': 5.6,  
    'l': [2, 3],  
    's': 'hello, world'  
}
```

#### Parameters

**assign\_str: str** Assignment string. Should only contain assignment statements assigning python literals or builtin function calls, to variable names. Multiple assignment statements should be separated by semi-colons.

#### Returns

**dict** Dictionary { name: value } containing assignment results.

### 5.8.2 Requirements Handling

---

<code>requires_optional(*requirements)</code>	Decorator returning either the original function, or a dummy function raising a <code>MissingPackageException</code> when called, depending on whether the supplied <code>requirements</code> are present.
---	--

---

`africanus.util.requirements.requires_optional(*requirements)`

Decorator returning either the original function, or a dummy function raising a `MissingPackageException` when called, depending on whether the supplied `requirements` are present.

If packages are missing and called within a test, the dummy function will call `pytest.skip()`.

Used in the following way:

```
try:  
    from scipy import interpolate  
except ImportError as e:  
    # https://stackoverflow.com/a/29268974/1611416, pep 3110 and 344  
    scipy_import_error = e  
else:  
    scipy_import_error = None  
  
@requires_optional('scipy', scipy_import_error)
```

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```
def function(*args, **kwargs):
    return interpolate(...)
```

#### Parameters

**requirements** [iterable of string, None or ImportError] Sequence of package names required by the decorated function. ImportError exceptions (or None, indicating their absence) may also be supplied and will be immediately re-raised within the decorator. This is useful for tracking down problems in user import logic.

#### Returns

**callable** Either the original function if all requirements are available or a dummy function that throws a MissingPackageException or skips a pytest.

### 5.8.3 Shapes

<code>africanus.util.shapes.aggregate_chunks(chunks, max_chunks)</code>	Aggregate dask chunks together into chunks no larger than <code>max_chunks</code> .
<code>corr_shape(ncorr, corr_shape)</code>	Returns the shape of the correlations, given <code>ncorr</code> and the type of correlation shape requested

`africanus.util.shapes.aggregate_chunks(chunks, max_chunks)`

Aggregate dask chunks together into chunks no larger than `max_chunks`.

```
chunks, max_c = ((3,4,6,3,6,7), (1,1,1,1,1,1)), (10,3)
expected = ((7,9,6,7), (2,2,1,1))
assert aggregate_chunks(chunks, max_c) == expected
```

#### Parameters

**chunks** [sequence of tuples or tuple]

**max\_chunks** [sequence of ints or int]

#### Returns

**sequence of tuples or tuple**

`africanus.util.shapes.corr_shape(ncorr, corr_shape)`

Returns the shape of the correlations, given `ncorr` and the type of correlation shape requested

#### Parameters

**ncorr** [integer] Number of correlations

**corr\_shape** [{‘flat’, ‘matrix’}] Shape of output correlations

#### Returns

**tuple** Shape tuple describing the correlation dimensions

- If `flat` returns `(ncorr,)`
- If `matrix` returns
  - `(1,)` if `ncorr == 1`

- (2,) if ncorr == 2
- (2,2) if ncorr == 4

## 5.8.4 Beams

---

<code>beam_filenames(filename_schema, corr_types)</code>	Returns a dictionary of beam filename pairs, keyed on correlation, from the cartesian product of correlations and real, imaginary pairs
<code>beam_grids(header[, l_axis, m_axis])</code>	Extracts the FITS indices and grids for the beam dimensions in the supplied FITS header.

---

`africanus.util.beams.beam_filenames(filename_schema, corr_types)`

Returns a dictionary of beam filename pairs, keyed on correlation, from the cartesian product of correlations and real, imaginary pairs

Given `beam_$(corr)_$(reim).fits` returns:

```
{  
    'xx' : ['beam_xx_re.fits', 'beam_xx_im.fits'],  
    'xy' : ['beam_xy_re.fits', 'beam_xy_im.fits'],  
    ...  
    'yy' : ['beam_yy_re.fits', 'beam_yy_im.fits'],  
}
```

Given `beam_$(CORR)_$(REIM).fits` returns:

```
{  
    'xx' : ['beam_XX_RE.fits', 'beam_XX_IM.fits'],  
    'xy' : ['beam_XY_RE.fits', 'beam_XY_IM.fits'],  
    ...  
    'yy' : ['beam_YY_RE.fits', 'beam_YY_IM.fits']),  
}
```

### Parameters

`filename_schema` [str] String containing the filename schema.

`corr_types` [list of integers] list of integers defining the correlation type.

### Returns

`dict` Dictionary of schema {correlation : (refile, imfile)} mapping correlations to real and imaginary filename pairs

`africanus.util.beams.beam_grids(header, l_axis=None, m_axis=None)`

Extracts the FITS indices and grids for the beam dimensions in the supplied FITS header. Specifically the axes specified by

1. L or X CTYPE
2. M or Y CTYPE
3. FREQ CTYPE

If the first two axes have a negative sign, such as -L, the grid will be inverted.

Any grids corresponding to axes with a CUNIT type of DEG will be converted to radians.

#### Parameters

**header** [Header or dict] FITS header object.

**l\_axis** [str] FITS axis interpreted as the L axis.  $L$  and  $X$  are sensible values here.  $-L$  will invert the coordinate system on that axis.

**m\_axis** [str] FITS axis interpreted as the M axis.  $M$  and  $Y$  are sensible values here.  $-M$  will invert the coordinate system on that axis.

#### Returns

**tuple** Returns ((l\_axis, l\_grid), (m\_axis, m\_grid), (freq\_axis, freq\_grid)) where the axis is the FORTRAN indexed FITS axis (1-indexed) and grid contains the values at each pixel along the axis.

## 5.8.5 Code

<code>format_code(code)</code>	Formats some code with line numbers
<code>memoize_on_key(key_fn)</code>	Memoize based on a key function supplied by the user.

`africanus.util.code.format_code(code)`

Formats some code with line numbers

#### Parameters

**code** [str] Code

#### Returns

**str** Code prefixed with line numbers

`class africanus.util.code.memoize_on_key(key_fn)`

Memoize based on a key function supplied by the user. The key function should return a custom key for memoizing the decorated function, based on the arguments passed to it.

In the following example, the arguments required to generate the `_generate_phase_delay_kernel` function are the types of the `lm`, `uvw` and `frequency` arrays, as well as the number of correlations, `ncorr`.

The supplied `key_fn` produces a unique key based on these types and the number of correlations, which is used to cache the generated function.

```
def key_fn(lm, uvw, frequency, ncorrs=4):
    """
    Produce a unique key for the arguments of
    _generate_phase_delay_kernel
    """
    return (lm.dtype, uvw.dtype, frequency.dtype, ncorrs)

_code_template = jinja2.Template('''
#define ncorrs {{ncorrs}}


__global__ void phase_delay(
    const {{lm_type}} * lm,
    const {{uvw_type}} * uvw,
    const {{freq_type}} * frequency,
```

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```

{{out_type}} * out)
{
    ...
}
''')

_type_map = {
    np.float32: 'float',
    np.float64: 'double'
}

@memoize_on_key(key_fn)
def _generate_phase_delay_kernel(lm, uvw, frequency, ncorrs=4):
    """ Generate the phase delay kernel """
    out_dtype = np.result_type(lm.dtype, uvw.dtype, frequency.dtype)
    code = _code_template.render(lm_type=_type_map[lm.dtype],
                                 uvw_type=_type_map[uvw.dtype],
                                 freq_type=_type_map[frequency.dtype],
                                 ncorrs=ncorrs)
    return cp.RawKernel(code, "phase_delay")

```

## Methods

<code>__call__(fn)</code>	Call self as a function.
---------------------------	--------------------------

## 5.8.6 dask

<code>EstimatingProgressBar([minimum, width, dt, out])</code>	Progress Bar that displays elapsed time as well as an estimate of total time taken.
---	---

`class africanus.util.dask_util.EstimatingProgressBar(minimum=0, width=42, dt=1.0, out=sys.stdout)`

Progress Bar that displays elapsed time as well as an estimate of total time taken.

When starting a dask computation, the bar examines the graph and determines the number of chunks contained by a dask collection.

During computation the number of completed chunks and their the total time taken to complete them are tracked. The average derived from these numbers are used to estimate total compute time, relative to the current elapsed time.

The bar is not particularly accurate and will underestimate near the beginning of computation and seems to slightly overestimate during the buk of computation. However, it may be more accurate than the default dask task bar which tracks number of tasks completed by total tasks.

### Parameters

**minimum** [int, optional] Minimum time threshold in seconds before displaying a progress bar.  
Default is 0 (always display)

**width** [int, optional] Width of the bar, default is 42 characters.

**dt** [float, optional] Update resolution in seconds, default is 1.0 seconds.

## 5.8.7 CUDA

---

<code>grids(dims, blocks)</code>	Determine the grid size, given space dimensions sizes and blocks
----------------------------------	--

---

`africanus.util.cuda.grids(dims, blocks)`

Determine the grid size, given space dimensions sizes and blocks

### Parameters

**dims** [tuple of ints] ( $x, y, z$ ) tuple

### Returns

**tuple** ( $x, y, z$ ) grid size tuple

## 5.8.8 Patterns

---

<code>Multiton(*args, **kwargs)</code>	General Multiton metaclass
<code>LazyProxy(fn, *args, **kwargs)</code>	Lazy instantiation of a proxied object.
<code>LazyProxyMultiton(*args, **kwargs)</code>	Combination of a <code>LazyProxy</code> with a <code>Multiton</code>

---

`class africanus.util.patterns.Multiton(*args, **kwargs)`

General Multiton metaclass

Implementation of the `Multiton` pattern, which always returns a unique object for a unique set of arguments provided to a class constructor. For example, in the following, only a single instance of *A* with argument *I* is ever created.

```
class A(metaclass=Multiton):
    def __init__(self, *args, **kw):
        self.args = args
        self.kw = kw

    assert A(1) is A(1)
    assert A(1, "bob") is not A(1)
```

This is useful for ensuring that only a single instance of a heavy-weight resource such as files, sockets, thread/process pools or database connections is created in a single process, for a unique set of arguments.

### Notes

Instantiation of object instances is thread-safe.

`class africanus.util.patterns.LazyProxy(fn, *args, **kwargs)`

Lazy instantiation of a proxied object.

A `LazyProxy` proxies an object which is lazily instantiated on first use. It is primarily useful for embedding references to heavy-weight resources in a dask graph, so they can be pickled and sent to other workers without immediately instantiating those resources.

To this end, the proxy takes as arguments:

1. a class or factory function that instantiates the desired resource.
2. `*args` and `**kwargs` that should be supplied to the instantiator.

Listing 1: The function and arguments for creating a file are wrapped in a LazyProxy. It is only instantiated when `f.write` is called.

```
f = LazyProxy(open, "test.txt", mode="r")
f.write("Hello World!")
f.close()
```

In addition to the class/factory function, it is possible to specify a Finaliser supplied to `weakref.finalize` that is called to cleanup the resource when the LazyProxy is garbage collected. In this case, the first argument should be a tuple of two elements: the factory and the finaliser.

```
# LazyProxy defined with factory function and finaliser function
def finalise_file(file):
    file.close()

f2 = LazyProxy((open, finalise_file), "test.txt", mode="r")

class WrappedFile:
    def __init__(self, *args, **kwargs):
        self.handle = open(*args, **kwargs)

    def close(self):
        self.handle.close()

# LazyProxy defined with class
f1 = LazyProxy(WrappedFile, WrappedFile.close), "test.txt", mode="r")
```

LazyProxy objects are designed to be embedded in `dask.array.blockwise()` calls. For example:

```
# Specify the start and length of each range
file_ranges = np.array([[0, 5], [5, 10], [15, 5] [20, 10]])
# Chunk each range individually
da_file_ranges = dask.array(file_ranges, chunks=(1, 2))
# Reference a binary file
file_proxy = LazyProxy(open, "data.dat", "rb")

def _read(file_proxy, file_range):
    # Seek to range start and read the length of data
    start, length = file_range
    file_proxy.seek(start)
    return np.asarray(file_proxy.read(length), np.uint8)

data = da.blockwise(_read, "x",
                    # Embed the file_proxy in the graph
                    file_proxy, None,
                    # Pass each file range to the _read
                    da_file_ranges, "xy",
                    # output chunks should have the length
                    # of each range
                    adjust_chunks={"x": tuple(file_ranges[:, 1])},
```

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```
concatenate=True)

print(data.compute(processes=True))
```

### Parameters

- fn** [class or callable or tuple] A callable object that used to create the proxied object. In tuple form, this should consist of two callables. The first should create the proxied object and the second should be a finaliser that performs cleanup on the proxied object when the LazyProxy is garbage collected: it is passed directly to `weakref.finalize`.
- \*args** [tuple] Positional arguments passed to the callable object specified in *fn* that will create the proxied object. The contents of *\*args* should be pickleable.
- \*\*kwargs** [dict] Keyword arguments passed to the callable object specified in *fn* that will create the proxied object. The contents of *\*\*kwargs* should be pickleable.

### Notes

- Instantiation of the proxied object is thread-safe.
- LazyProxy's are configured to never instantiate within `dask.array.blockwise()` and `dask.blockwise.blockwise()` calls.

```
class africanus.util.patterns.LazyProxyMultiton(*args, **kwargs)
```

Combination of a `LazyProxy` with a `Multiton`

Ensures that only a single `LazyProxy` is ever created for the given constructor arguments.

```
class A:
    def __init__(self, value):
        self.value = value

assert LazyProxyMultiton("foo") is LazyProxyMultiton("foo")
```

See `LazyProxy` and `Multiton` for further details

## 5.9 Calibration

This module provides basic radio interferometry calibration utilities. Calibration is the process of estimating the  $2 \times 2$  Jones matrices which describe transformations of the signal as it propagates from source to observer. Currently, all utilities assume a discretised form of the radio interferometer measurement equation (RIME) as described in [Radio Interferometer Measurement Equation](#).

Calibration is usually divided into three phases viz.

- First generation calibration (1GC): using an external calibrator to infer the gains during the target observation. Sometimes also referred to as calibrator transfer
- Second generation calibration (2GC): using a partially incomplete sky model to perform direction independent calibration. Also known as direction independent self-calibration.
- Third generation calibration (3GC): using a partially incomplete sky model to perform direction dependent calibration. Also known as direction dependent self-calibration.

On top of these three phases, there are usually three possible calibration scenarios. The first is when both the Jones terms and the visibilities are assumed to be diagonal. In this case the two correlations can be calibrated separately and it is referred to as `diag-diag` calibration. The second case is when the Jones matrices are assumed to be diagonal but the visibility data are full  $2 \times 2$  matrices. This is referred to as `diag` calibration. The final scenario is when both the full  $2 \times 2$  Jones matrices and the full  $2 \times 2$  visibilities are used for calibration. This is simply referred to as calibration. The specific scenario is determined from the shapes of the input gains and the input data.

This module also provides a number of utilities which are useful for calibration.

### 5.9.1 Utils

#### Numpy

<code>corrupt_vis</code> (time_bin_indices, ...)	Corrupts model visibilities with arbitrary Jones terms.
<code>residual_vis</code> (time_bin_indices, ...)	Computes residual visibilities given model visibilities and gains solutions.
<code>correct_vis</code> (time_bin_indices, ...)	Apply inverse of direction independent gains to visibilities to generate corrected visibilities.
<code>compute_and_corrupt_vis</code> (time_bin_indices, ...)	Corrupts time variable component model with arbitrary Jones terms.

`africanus.calibration.utils.corrupt_vis`(*time\_bin\_indices*, *time\_bin\_counts*, *antenna1*, *antenna2*, *jones*, *model*)

Corrupts model visibilities with arbitrary Jones terms.

##### Parameters

- `time_bin_indices`** [numpy.ndarray] The start indices of the time bins of shape (utime)
- `time_bin_counts`** [numpy.ndarray] The counts of unique time in each time bin of shape (utime)
- `antenna1`** [numpy.ndarray] First antenna indices of shape (row, ).
- `antenna2`** [numpy.ndarray] Second antenna indices of shape (row, )
- `jones`** [numpy.ndarray] Gains of shape (time, ant, chan, dir, corr) or (time, ant, chan, dir, corr, corr).
- `model`** [numpy.ndarray] Model data values of shape (row, chan, dir, corr) or (row, chan, dir, corr, corr).

##### Returns

- `vis`** [numpy.ndarray] visibilities of shape (time, ant, chan, dir, corr) or (time, ant, chan, dir, corr, corr).

`africanus.calibration.utils.residual_vis`(*time\_bin\_indices*, *time\_bin\_counts*, *antenna1*, *antenna2*, *jones*, *vis*, *flag*, *model*)

Computes residual visibilities given model visibilities and gains solutions.

##### Parameters

- `time_bin_indices`** [numpy.ndarray] The start indices of the time bins of shape (utime)
- `time_bin_counts`** [numpy.ndarray] The counts of unique time in each time bin of shape (utime)
- `antenna1`** [numpy.ndarray] First antenna indices of shape (row, ).

**antenna2** [numpy.ndarray] Second antenna indices of shape (row,)

**jones** [numpy.ndarray] Gain solutions of shape (time, ant, chan, dir, corr) or (time, ant, chan, dir, corr, corr).

**vis** [numpy.ndarray] Data values of shape (row, chan, corr). or (row, chan, corr, corr).

**flag** [numpy.ndarray] Flag data of shape (row, chan, corr) or (row, chan, corr, corr)

**model** [numpy.ndarray] Model data values of shape (row, chan, dir, corr) or (row, chan, dir, corr, corr).

**Returns**

**residual** [numpy.ndarray] Residual visibilities of shape (time, ant, chan, dir, corr) or (time, ant, chan, dir, corr, corr).

```
africanus.calibration.utils.correct_vis(time_bin_indices, time_bin_counts, antenna1, antenna2, jones,
                                         vis, flag)
```

Apply inverse of direction independent gains to visibilities to generate corrected visibilities. For a measurement model of the form

$$V_{pq} = G_p X_{pq} G_q^H + n_{pq}$$

the corrected visibilities are defined as

$$C_{pq} = G_p^{-1} V_{pq} G_q^{-H}$$

The corrected visibilities therefore have a non-trivial noise contribution. Note it is only possible to form corrected data from direction independent gains solutions so the `dir` axis on the `jones` terms should always be one.

**Parameters**

**time\_bin\_indices** [numpy.ndarray] The start indices of the time bins of shape (utime).

**time\_bin\_counts** [numpy.ndarray] The counts of unique time in each time bin of shape (utime).

**antenna1** [numpy.ndarray] Antenna 1 index used to look up the antenna Jones for a particular baseline with shape (row, ).

**antenna2** [numpy.ndarray] Antenna 2 index used to look up the antenna Jones for a particular baseline with shape (row, ).

**jones** [numpy.ndarray] Gain solutions of shape (time, ant, chan, dir, corr) or (time, ant, chan, dir, corr, corr).

**vis** [numpy.ndarray] Data values of shape (row, chan, corr) or (row, chan, corr, corr).

**flag** [numpy.ndarray] Flag data of shape (row, chan, corr) or (row, chan, corr, corr).

**Returns**


---

**corrected\_vis** [numpy.ndarray] True visibilities of shape (row, chan, corr\_1, corr\_2)

```
africanus.calibration.utils.compute_and_corrupt_vis(time_bin_indices, time_bin_counts, antenna1,
                                                    antenna2, jones, model, uvw, freq, lm)
```

Corrupts time variable component model with arbitrary Jones terms. Currently only time variable point source models are supported.

### Parameters

**time\_bin\_indices** [numpy.ndarray] The start indices of the time bins of shape (utime)

**time\_bin\_counts** [numpy.ndarray] The counts of unique time in each time bin of shape (utime)

**antenna1** [numpy.ndarray] First antenna indices of shape (row, ).

**antenna2** [numpy.ndarray] Second antenna indices of shape (row, )

**jones** [numpy.ndarray] Gains of shape (utime, ant, chan, dir, corr) or (utime, ant, chan, dir, corr, corr).

**model** [numpy.ndarray] Model image as a function of time with shape (utime, chan, dir, corr) or (utime, chan, dir, corr, corr).

**uvw** [numpy.ndarray] uvw coordinates of shape (row, 3)

**lm** [numpy.ndarray] Source lm coordinates as a function of time (utime, dir, 2)

### Returns

**vis** [numpy.ndarray] visibilities of shape (row, chan, corr) or (row, chan, corr, corr).

## Dask

<code>corrupt_vis(time_bin_indices, ...)</code>	Corrupts model visibilities with arbitrary Jones terms.
<code>residual_vis(time_bin_indices, ...)</code>	Computes residual visibilities given model visibilities and gains solutions.
<code>correct_vis(time_bin_indices, ...)</code>	Apply inverse of direction independent gains to visibilities to generate corrected visibilities.
<code>compute_and_corrupt_vis(time_bin_indices, ...)</code>	Corrupts time variable component model with arbitrary Jones terms.

`africanus.calibration.utils.dask.corrupt_vis(time_bin_indices, time_bin_counts, antenna1, antenna2, jones, model)`

Corrupts model visibilities with arbitrary Jones terms.

### Parameters

**time\_bin\_indices** [dask.array.Array] The start indices of the time bins of shape (utime)

**time\_bin\_counts** [dask.array.Array] The counts of unique time in each time bin of shape (utime)

**antenna1** [dask.array.Array] First antenna indices of shape (row, ).

**antenna2** [dask.array.Array] Second antenna indices of shape (row, )

**jones** [dask.array.Array] Gains of shape (time, ant, chan, dir, corr) or (time, ant, chan, dir, corr, corr).

**model** [dask.array.Array] Model data values of shape (row, chan, dir, corr) or (row, chan, dir, corr, corr).

### Returns

**vis** [dask.array.Array] visibilities of shape (time, ant, chan, dir, corr) or (time, ant, chan, dir, corr, corr).

---

```
africanus.calibration.utils.dask.residual_vis(time_bin_indices, time_bin_counts, antenna1, antenna2,
                                              jones, vis, flag, model)
```

Computes residual visibilities given model visibilities and gains solutions.

#### Parameters

- time\_bin\_indices** [dask.array.Array] The start indices of the time bins of shape (utime)
- time\_bin\_counts** [dask.array.Array] The counts of unique time in each time bin of shape (utime)
- antenna1** [dask.array.Array] First antenna indices of shape (row,).
- antenna2** [dask.array.Array] Second antenna indices of shape (row,)
- jones** [dask.array.Array] Gain solutions of shape (time, ant, chan, dir, corr) or (time, ant, chan, dir, corr, corr).
- vis** [dask.array.Array] Data values of shape (row, chan, corr). or (row, chan, corr, corr).
- flag** [dask.array.Array] Flag data of shape (row, chan, corr) or (row, chan, corr, corr)
- model** [dask.array.Array] Model data values of shape (row, chan, dir, corr) or (row, chan, dir, corr, corr).

#### Returns

- residual** [dask.array.Array] Residual visibilities of shape (time, ant, chan, dir, corr) or (time, ant, chan, dir, corr, corr).

```
africanus.calibration.utils.dask.correct_vis(time_bin_indices, time_bin_counts, antenna1, antenna2,
                                              jones, vis, flag)
```

Apply inverse of direction independent gains to visibilities to generate corrected visibilities. For a measurement model of the form

$$V_{pq} = G_p X_{pq} G_q^H + n_{pq}$$

the corrected visibilities are defined as

$$C_{pq} = G_p^{-1} V_{pq} G_q^{-H}$$

The corrected visibilities therefore have a non-trivial noise contribution. Note it is only possible to form corrected data from direction independent gains solutions so the `dir` axis on the `jones` terms should always be one.

#### Parameters

- time\_bin\_indices** [dask.array.Array] The start indices of the time bins of shape (utime).
- time\_bin\_counts** [dask.array.Array] The counts of unique time in each time bin of shape (utime).
- antenna1** [dask.array.Array] Antenna 1 index used to look up the antenna Jones for a particular baseline with shape (row,).
- antenna2** [dask.array.Array] Antenna 2 index used to look up the antenna Jones for a particular baseline with shape (row,).
- jones** [dask.array.Array] Gain solutions of shape (time, ant, chan, dir, corr) or (time, ant, chan, dir, corr, corr).
- vis** [dask.array.Array] Data values of shape (row, chan, corr) or (row, chan, corr, corr).

**flag** [dask.array.Array] Flag data of shape (row, chan, corr) or (row, chan, corr, corr).

#### Returns

---

**corrected\_vis** [dask.array.Array] True visibilities of shape (row, chan, corr\_1, corr\_2)

africanus.calibration.utils.dask.compute\_and\_corrupt\_vis(time\_bin\_indices, time\_bin\_counts,  
antenna1, antenna2, jones, model, uvw,  
freq, lm)

Corrupts time variable component model with arbitrary Jones terms. Currently only time variable point source models are supported.

#### Parameters

**time\_bin\_indices** [dask.array.Array] The start indices of the time bins of shape (utime)

**time\_bin\_counts** [dask.array.Array] The counts of unique time in each time bin of shape (utime)

**antenna1** [dask.array.Array] First antenna indices of shape (row, ).

**antenna2** [dask.array.Array] Second antenna indices of shape (row, )

**jones** [dask.array.Array] Gains of shape (utime, ant, chan, dir, corr) or (utime, ant, chan, dir, corr, corr).

**model** [dask.array.Array] Model image as a function of time with shape (utime, chan, dir, corr) or (utime, chan, dir, corr, corr).

**uvw** [dask.array.Array] uvw coordinates of shape (row, 3)

**lm** [dask.array.Array] Source lm coordinates as a function of time (utime, dir, 2)

#### Returns

**vis** [dask.array.Array] visibilities of shape (row, chan, corr) or (row, chan, corr, corr).

## 5.9.2 Phase only

### Numpy

<code>compute_jhr</code> (time_bin_indices, ...)	Computes the residual projected in to gain space.
<code>compute_jhj</code> (time_bin_indices, ...)	Computes the diagonal of the Hessian required to perform phase-only maximum likelihood calibration.
<code>compute_jhj_and_jhr</code> (time_bin_indices, ...)	Computes the diagonal of the Hessian and the residual locally projected in to gain space.
<code>gauss_newton</code> (time_bin_indices, ...[, tol, ...])	Performs phase-only maximum likelihood calibration using a Gauss-Newton optimisation algorithm.

africanus.calibration.phase\_only.compute\_jhr(time\_bin\_indices, time\_bin\_counts, antenna1, antenna2, jones, residual, model, flag)

Computes the residual projected in to gain space.

#### Parameters

**time\_bin\_indices** [numpy.ndarray] The start indices of the time bins of shape (utime)

**time\_bin\_counts** [numpy.ndarray] The counts of unique time in each time bin of shape (utime)

**antenna1** [numpy.ndarray] First antenna indices of shape (row,).

**antenna2** [numpy.ndarray] Second antenna indices of shape (row,.)

**jones** [numpy.ndarray] Gain solutions of shape (time, ant, chan, dir, corr) or (time, ant, chan, dir, corr, corr).

**residual** [numpy.ndarray] Residual values of shape (row, chan, corr). or (row, chan, corr, corr).

**model** [numpy.ndarray] Model data values of shape (row, chan, dir, corr) or (row, chan, dir, corr, corr).

**flag** [numpy.ndarray] Flag data of shape (row, chan, corr) or (row, chan, corr, corr)

#### Returns

**jhr** [numpy.ndarray] The residual projected into gain space shape (time, ant, chan, dir, corr) or (time, ant, chan, dir, corr, corr).

africanus.calibration.phase\_only.**compute\_jhj**(time\_bin\_indices, time\_bin\_counts, antenna1, antenna2, jones, model, flag)

Computes the diagonal of the Hessian required to perform phase-only maximum likelihood calibration. Currently assumes scalar or diagonal inputs.

#### Parameters

**time\_bin\_indices** [numpy.ndarray] The start indices of the time bins of shape (utime)

**time\_bin\_counts** [numpy.ndarray] The counts of unique time in each time bin of shape (utime)

**antenna1** [numpy.ndarray] First antenna indices of shape (row,).

**antenna2** [numpy.ndarray] Second antenna indices of shape (row,.)

**jones** [numpy.ndarray] Gain solutions of shape (time, ant, chan, dir, corr) or (time, ant, chan, dir, corr, corr).

**model** [numpy.ndarray] Model data values of shape (row, chan, dir, corr) or (row, chan, dir, corr, corr).

**flag** [numpy.ndarray] Flag data of shape (row, chan, corr) or (row, chan, corr, corr)

#### Returns

**jhj** [numpy.ndarray] The diagonal of the Hessian of shape (time, ant, chan, dir, corr) or (time, ant, chan, dir, corr, corr).

africanus.calibration.phase\_only.**compute\_jhj\_and\_jhr**(time\_bin\_indices, time\_bin\_counts, antenna1, antenna2, jones, residual, model, flag)

Computes the diagonal of the Hessian and the residual locally projected in to gain space.

#### Parameters

**time\_bin\_indices** [numpy.ndarray] The start indices of the time bins of shape (utime)

**time\_bin\_counts** [numpy.ndarray] The counts of unique time in each time bin of shape (utime)

**antenna1** [numpy.ndarray] First antenna indices of shape (row,).

**antenna2** [numpy.ndarray] Second antenna indices of shape (row,)

**jones** [numpy.ndarray] Gain solutions of shape (time, ant, chan, dir, corr) or (time, ant, chan, dir, corr, corr).

**residual** [numpy.ndarray] Residual values of shape (row, chan, corr). or (row, chan, corr, corr).

**model** [numpy.ndarray] Model data values of shape (row, chan, dir, corr) or (row, chan, dir, corr, corr).

**flag** [numpy.ndarray] Flag data of shape (row, chan, corr) or (row, chan, corr, corr)

#### Returns

**jhj** [numpy.ndarray] The diagonal of the Hessian of shape (time, ant, chan, dir, corr) or (time, ant, chan, dir, corr, corr).

**jhr** [numpy.ndarray] Residuals projected into signal space of shape (time, ant, chan, dir, corr) or (time, ant, chan, dir, corr, corr).

africanus.calibration.phase\_only.**gauss\_newton**(time\_bin\_indices, time\_bin\_counts, antenna1, antenna2, jones, vis, flag, model, weight, tol=0.0001, maxiter=100)

Performs phase-only maximum likelihood calibration using a Gauss-Newton optimisation algorithm. Currently only DIAG mode is supported.

#### Parameters

**time\_bin\_indices** [numpy.ndarray] The start indices of the time bins of shape (utime)

**time\_bin\_counts** [numpy.ndarray] The counts of unique time in each time bin of shape (utime)

**antenna1** [numpy.ndarray] First antenna indices of shape (row,).

**antenna2** [numpy.ndarray] Second antenna indices of shape (row,).

**jones** [numpy.ndarray] Gain solutions of shape (time, ant, chan, dir, corr) or (time, ant, chan, dir, corr, corr).

**vis** [numpy.ndarray] Data values of shape (row, chan, corr) or (row, chan, corr, corr).

**flag** [numpy.ndarray] Flag data of shape (row, chan, corr) or (row, chan, corr, corr).

**model** [numpy.ndarray] Model data values of shape (row, chan, dir, corr) or (row, chan, dir, corr, corr).

**weight** [numpy.ndarray] Weight spectrum of shape (row, chan, corr). If the channel axis is missing weights are duplicated for each channel.

**tol: float, optional** The tolerance of the solver. Defaults to 1e-4.

**maxiter: int, optional** The maximum number of iterations. Defaults to 100.

#### Returns

**gains** [numpy.ndarray] Gain solutions of shape (time, ant, chan, dir, corr) or shape (time, ant, chan, dir, corr, corr)

**jhj** [numpy.ndarray] The diagonal of the Hessian of shape (time, ant, chan, dir, corr) or shape (time, ant, chan, dir, corr, corr)

**jhr** [numpy.ndarray] Residuals projected into gain space of shape (time, ant, chan, dir, corr) or shape (time, ant, chan, dir, corr, corr).

**k:** int Number of iterations (will equal maxiter if not converged)

## Task

<code>compute_jhr(time_bin_indices, ...)</code>	Computes the residual projected in to gain space.
<code>compute_jhj(time_bin_indices, ...)</code>	Computes the diagonal of the Hessian required to perform phase-only maximum likelihood calibration.

`africanus.calibration.phase_only.dask.compute_jhr(time_bin_indices, time_bin_counts, antennal, antenna2, jones, residual, model, flag)`

Computes the residual projected in to gain space.

### Parameters

**time\_bin\_indices** [dask.array.Array] The start indices of the time bins of shape (utime)

**time\_bin\_counts** [dask.array.Array] The counts of unique time in each time bin of shape (utime)

**antenna1** [dask.array.Array] First antenna indices of shape (row, ).

**antenna2** [dask.array.Array] Second antenna indices of shape (row, )

**jones** [dask.array.Array] Gain solutions of shape (time, ant, chan, dir, corr) or (time, ant, chan, dir, corr, corr).

**residual** [dask.array.Array] Residual values of shape (row, chan, corr). or (row, chan, corr, corr).

**model** [dask.array.Array] Model data values of shape (row, chan, dir, corr) or (row, chan, dir, corr, corr).

**flag** [dask.array.Array] Flag data of shape (row, chan, corr) or (row, chan, corr, corr)

### Returns

**jhr** [dask.array.Array] The residual projected into gain space shape (time, ant, chan, dir, corr) or (time, ant, chan, dir, corr, corr).

`africanus.calibration.phase_only.dask.compute_jhj(time_bin_indices, time_bin_counts, antennal, antenna2, jones, model, flag)`

Computes the diagonal of the Hessian required to perform phase-only maximum likelihood calibration. Currently assumes scalar or diagonal inputs.

### Parameters

**time\_bin\_indices** [dask.array.Array] The start indices of the time bins of shape (utime)

**time\_bin\_counts** [dask.array.Array] The counts of unique time in each time bin of shape (utime)

**antenna1** [dask.array.Array] First antenna indices of shape (row, ).

**antenna2** [dask.array.Array] Second antenna indices of shape (row, )

**jones** [dask.array.Array] Gain solutions of shape (time, ant, chan, dir, corr) or (time, ant, chan, dir, corr, corr).

**model** [dask.array.Array] Model data values of shape (row, chan, dir, corr) or (row, chan, dir, corr, corr).

**flag** [dask.array.Array] Flag data of shape (row, chan, corr) or (row, chan, corr, corr)

#### Returns

**jhj** [dask.array.Array] The diagonal of the Hessian of shape (time, ant, chan, dir, corr) or (time, ant, chan, dir, corr, corr).

## 5.10 Linear Algebra

This module contains specialised linear algebra tools that are not currently available in the python standard scientific libraries.

### 5.10.1 Kronecker tools

A kronecker matrix is matrix that can be written as a kronecker matrix of the individual matrices i.e.

$$\begin{aligned} K &= K_0 \\ &\otimes K_1 \\ &\otimes K_2 \\ &\otimes \dots \\ &\cdots \end{aligned}$$

Matrices which exhibit this structure can exploit properties of the kronecker product to avoid explicitly expanding the matrix  $K$ . This module implements some common linear algebra operations which leverages this property for computational gains and a reduced memory footprint.

#### Numpy

<code>kron_matvec(A, b)</code>	Computes the matrix vector product of a kronecker matrix in linear time.
<code>kron_cholesky(A)</code>	Computes the Cholesky decomposition of a kronecker matrix as a kronecker matrix of Cholesky factors.

#### africanus.linalg.`kron_matvec`(A, b)

Computes the matrix vector product of a kronecker matrix in linear time. Assumes A consists of kronecker product of square matrices.

#### Parameters

**A** [numpy.ndarray] An array of arrays holding matrices [K0, K1, ...] where  $A = K_0 \otimes K_1 \otimes \dots$

**b** [numpy.ndarray] The right hand side vector

#### Returns

**x** [numpy.ndarray] The result of `A.dot(b)`

#### africanus.linalg.`kron_cholesky`(A)

Computes the Cholesky decomposition of a kronecker matrix as a kronecker matrix of Cholesky factors.

**Parameters**

**A** [numpy.ndarray] An array of arrays holding matrices [ $K_0, K_1, \dots$ ] where  $A = K_0 \otimes K_1 \otimes \dots$

**Returns**

**L** [numpy.ndarray] An array of arrays holding matrices [ $L_0, L_1, \dots$ ] where  $L = L_0 \otimes L_1 \otimes \dots$  and each  $L_i = \text{cholesky}(K_i)$

## 5.11 Gaussian processes

This module provides a collection of tools that are useful when performing Gaussian process regression.

### 5.11.1 Numpy

<code>africanus.gps.abs_diff(x, xp)</code>	Gets matrix of differences between $D$ -dimensional vectors $x$ and $xp$ i.e.
<code>africanus.gps.exponential_squared(x, xp, sigmaf, l[, pspec])</code>	Create exponential squared covariance function between $D$ dimensional vectors $x$ and $x_p$ i.e.

`africanus.gps.abs_diff(x, xp)`

Gets matrix of differences between  $D$ -dimensional vectors  $x$  and  $xp$  i.e.

$$X_{ij} = |x_i - x_j|$$

**Parameters**

**x** [numpy.ndarray] Array of inputs of shape (N, D).

**xp** [numpy.ndarray] Array of inputs of shape (Np, D).

**Returns**

**XX** [numpy.ndarray] Array of differences of shape (N, Np).

`africanus.gps.exponential_squared(x, xp, sigmaf, l, pspec=False)`

Create exponential squared covariance function between  $D$  dimensional vectors  $x$  and  $x_p$  i.e.

$$k(x, x_p) = \sigma_f^2 \exp\left(-\frac{(x - x_p)^2}{2l^2}\right)$$

**Parameters**

**x** [numpy.ndarray] Array of shape (N, D).

**xp** [numpy.ndarray] Array of shape (Np, D).

**sigmaf** [float] The signal variance hyper-parameter

**l** [float] The length scale hyper-parameter

**Returns**

**K** [numpy.ndarray] Array of shape (N, Np)

## 5.12 Fused Radio Interferometer Measurement Equation

### 5.12.1 Radio Interferometer Measurement Equation

The Radio Interferometer Measurement Equation (RIME) describes the response of an interferometer to a sky model. As described in [A full-sky Jones formalism](#), a RIME could be written as follows:

$$V_{pq} = G_p \left( \sum_s E_{ps} L_p K_{ps} B_s K_{qs}^H L_q^H E_{qs}^H \right) G_q^H$$

where for antenna  $p$  and  $q$ , and source  $s$ :

- $G_p$  represents direction-independent effects.
- $E_{ps}$  represents direction-dependent effects.
- $L_p$  represents the feed rotation.
- $K_{ps}$  represents the phase delay term.
- $B_s$  represents the brightness matrix.

The RIME is more formally described in the following four papers:

- I. [A full-sky Jones formalism](#)
- II. [Calibration and direction-dependent effects](#)
- III. [Addressing direction-dependent effects in 21cm WSRT observations of 3C147](#)
- IV. [A generalized tensor formalism](#)

### 5.12.2 The Fused RIME

The RIME poses a number of implementation challenges which focus on flexibility, speed and ease of use.

Firstly, the RIME can be composed of many terms representing various physical effects. It is useful for scientist to be able to specify many different terms in the above [Equation](#), for example.

Secondly, the computational complexity of the RIME  $O(S \times V)$  where S is the number of source and V is the number of visibilities. This is computationally expensive relative to degridding strategies.

Thirdly, it should be as easy as possible to define the RIME, but not at the cost of the previous two constraints.

The Fused RIME therefore implements a “RIME Compiler” using [Numba](#) for speed, which compiles a RIME Specification defined by a number of *Terms* into a single, optimal unit of execution.

### 5.12.3 A Simple Example

In the following example, we will define a simple RIME using the Fused RIME API to define terms for computing:

1. the Phase Delay.
2. the Brightness Matrix.

## The RIME Specification

The specification for this RIME is as follows:

```
rime_spec = RimeSpecification("(Kpq, Bpq): [I,Q,U,V] -> [XX,XY,YX,YY]",  
                             terms={"K": Phase},  
                             transformers=[LMTransformer])
```

(Kpq, Bpq) specifies the onion including the Phase Delay and Brightness more formally defined [here](#), while the pq in both terms signifies that they are calculated per-baseline. [I, Q, U, V] -> [XX, XY, YX, YY] defines the stokes to correlation conversion within the RIME and also identifies whether the RIME is handling linear or circular feeds. terms={"K": Phase} indicates that the K term is implemented as a custom Phase term, described in the next section. Finally, LMTransformer is a Transformer that precomputes lm coordinates for use by all terms.

## Custom Phase Term

Within the RIME, each term is *sampled* at an individual source, row and channel.

Therefore each term must provide a sampling function that will provide the necessary data for multiplication within the RIME. Consider the following Phase Term:

```
from africanus.experimental.rime.fused.terms.core import Term  
  
class Phase(Term):  
    def sampler(self):  
        def phase_sample(state, s, r, t, f1, f2, a1, a2, c):  
            p = state.real_phase[s, r] * state.chan_freq[c]  
            return np.cos(p) + np.sin(p)*1j  
  
        return phase_sample
```

This may look simple: we compute the complex phase by multiplying the real phase at each source and row by the channel frequency and return the complex exponential of this value.

However, questions remain: What is the *state* object and how do we know that the *real\_phase* and *chan\_freq* are members? To answer this, we must define (and understand) a second method defined on the *Phase* term, called *init\_fields*.

```
import numba  
from africanus.experimental.rime.fused.terms.core import Term  
  
class Phase(Term):  
    def init_fields(self, typingctx, lm, uvw, chan_freq):  
        # Given the numba types of the lm, uvw and chan_freq  
        # arrays, derive a unified output numba type  
        numba_type = typingctx.unify_types(lm.dtype,  
                                         uvw.dtype,  
                                         chan_freq.dtype)  
  
        # Define the type of new fields on the state object  
        # in this case a 2D Numba array with dtype numba_type  
        fields = [("real_phase", numba_type[:, :, :])]  
  
        def real_phase(lm, uvw, chan_freq):  
            """Compute the real_phase upfront, instead of in
```

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```

the sampling function"""
real_phase = np.empty((lm.shape[0], uvw.shape[0]), numba_type)

for s in range(lm.shape[0]):
    l, m = lm[s]
    n = 1.0 - l**2 - m**2
    n = np.sqrt(0.0 if n <= 0.0 else n) - 1.0

    for r in range(uvw.shape[0]):
        u, v, w = uvw[r]
        real_phase[s, r] = -2.0*np.pi*(l*u + m*v + n*w)/3e8

return real_phase

# Return the new field definition and
# the function for creating it
return fields, real_phase

```

`init_fields` serves multiple purposes:

1. It requests input for the Phase term. The above definition of `init_fields` signifies that the Phase term desires the `lm`, `uvw` and `chan_freq` arrays. Additionally, these arrays will be stored on the `state` object provided to the sampling function.
2. It supports reasoning about Numba types in a manner similar to `numba.generated_jit()`. The `lm`, `uvw` and `chan_freq` arguments contain the Numba types of the variables supplied to the RIME, while the `typingctx` argument contains a Numba Typing Context which can be useful for reasoning about these types. For example `typingctx.unify_types(lm.dtype, uvw.dtype, chan_freq.dtype)` returns a type with sufficient precision, given the input types, similar to `numpy.result_type()`.
3. It allows the user to define new fields, as well as a function for defining those fields on the `state` object. The above definition of `init_fields` returns a list of `(name, type)` tuples defining the new field names and their types, while `real_phase` defines the creation of this new field.

This is useful for optimising the sampling function by pre-computing values. For example, it is wasteful to compute the real phase for each source, row and channel.

Returning to our definition of the Phase Term sampling function, we can see that it uses the new field `real_phase` defined in `init_fields`, as well as the `chan_freq` array requested in `init_fields` to compute a complex exponential.

```

class Phase(Term):
    def sampler(self):
        def phase_sample(state, s, r, t, f1, f2, a1, a2, c):
            p = state.real_phase[s, r] * state.chan_freq[c]
            return np.cos(p) + np.sin(p)*1j

    return phase_sample

```

## Transformers

Using `Term.init_fields()`, we can precompute data for use in sampling functions, within a single `Term`. However, sometimes we wish to precompute data for use by multiple `Terms`. This can be achieved through the use of `Transformers`. A good example of data that it is useful to precompute for multiple `Terms` are `lm` coordinates, which are in turn, derived from `phase_dir` and `radec` which are the phase centre of an observation and the position of a source, respectively. In the following code snippet, `LMTransformer.init_fields`

```
from africanus.experimental.rime.fused.transformers import Transformer

class LMTransformer(Transformer):
    # Must specify list of outputs produced by this transformer on the
    # OUTPUTS class attribute
    OUTPUTS = ["lm"]

    def init_fields(self, typingctx, ra, dec, phase_dir):
        # Type and provide method for initialising the lm output
        dt = typingctx.unify_types(ra.dtype, phase_dir.dtype)
        fields = [("lm", dt[:, :])]

    def lm(ra, dec, phase_dir):
        lm = np.empty_like(ra)
        pc_ra = phase_dir[0]
        pc_dec = phase_dir[1]

        sin_pc_dec = np.sin(pc_dec)
        cos_pc_dec = np.cos(pc_dec)

        for s in range(ra.shape[0]):
            da = ra[s, 0] - pc_ra
            sin_ra_delta = np.sin(da)
            cos_ra_delta = np.cos(da)

            sin_dec = np.sin(dec[s, 1])
            cos_dec = np.cos(dec[s, 1])

            lm[s, 0] = cos_dec * sin_ra_delta
            lm[s, 1] = sin_dec * cos_pc_dec - cos_dec * sin_pc_dec * cos_ra_delta

        return lm

    return fields, lm
```

The `lm` array will be available on the `state` object and as a valid input for `Term.init_fields()`.

## Invoking the RIME

We then invoke the RIME by passing in the `RimeSpecification`, as well as a dataset containing the required arguments:

```
from africanus.experimental.rime.fused.core import rime
import numpy as np

dataset = {
    "radec": np.random.random((10, 2))*1e-5,
    "phase_dir": np.random.random((2,))*1e-5,
    "uvw": np.random.random((100, 3))*1e5,
    "chan_freq": np.linspace(.856e9, 2*.856e9, 16),
    ...,
    "stokes": np.random.random((10, 4)),
    # other required data
}

rime_spec = RimeSpecification("(Kpq, Bpq)",
                               terms={"K": Phase},
                               transformers=LMTransformer)
model_visibilities = rime(rime_spec, dataset)
```

## Dask Support

Dask wrappers are provided for the `africanus.experimental.rime.fused.core.rime()` function. In order to support this, both `Term` and `Transformer` classes need to supply a `dask_schema` function which is used to define the schema for each supplied argument, which in turn is supplied to a `dask.array.blockwise()` call.

The schema should be a tuple of dimension string names. In particular, the `rime` function assigns special meaning to `source`, `row`, `chan` and `corr` – These names are associated with individual sources (fields) and Measurement Set rows, channels and correlations, respectively. Dask Array chunking is supported along these dimensions in the sense that the `rime` will be computed for each chunk along these dimensions.

---

**Note:** Chunks in dimensions other than `source`, `row`, `chan` and `corr` will be contracted into a single array within the `rime` function. It is recommended that other dimensions contain a single chunk, or contain small quantities of data relative to the special dimensions.

---

Therefore, `Phase.dask_schema` could be implemented as follows:

```
class Phase(Term):
    def dask_schema(self, lm, uvw, chan_freq):
        assert lm.ndim == 2
        assert uvw.ndim == 2
        assert chan_freq.ndim == 1

        return {
            "lm": ("source", "lm-component"),
            "uvw": ("row", "uvw-component"),
            "chan_freq": ("chan",),
        }
```

The `dask_schema` for a `Transformer` is slightly different as, in addition a schema for the inputs, it must also provide an `array_like` variable describing the number of dimensions and data type of the output arrays. The `array_like` variables are in turn passed into `Term.dask_schema`. Thus, `LMTransformer.dask_schema` could be implemented as follows;

```
class LMTransformer(Transformer):
    OUTPUTS = ["lm"]

    def dask_schema(self, phase_dir, radec):
        dt = np.result_type(phase_dir.dtype, radec.dtype)
        return ({
            "phase_dir": ("radec-component",),
            "radec": ("source", "radec-component"),
        },
        {
            "lm": np.empty((0, 0), dtype=dt)
        })
```

Then, in a paradigm very similar to the non-dask case, we create a `RimeSpecification` and supply it, along with a dictionary or dataset of dask arrays, to the `rime()` function. This will produce a dask array representing the model visibilities.

```
from africanus.experimental.rime.fused.dask import rime
import dask.array as da
import numpy as np

dataset = {
    "radec": da.random.random((10, 2), chunks=(2, 2))*1e-5,
    "phase_dir": da.random.random((2,), chunks=(2,))*1e-5,
    "uvw": da.random.random((100, 3), chunks=(10, 3))*1e5,
    "chan_freq": da.linspace(.856e9, 2*.856e9, 16, chunks=(4,)),
    ...,
    "stokes": da.random.random((10, 4), chunks=(2, 4)),
    # other required data
}

rime_spec = RimeSpecification("(Kpq, Bpq)",
                               terms={"K": Phase},
                               transformers=LMTransformer)
model_visibilities = rime(rime_spec, dataset)
model_visibilities.compute()
```

## 5.12.4 API

```
class africanus.experimental.rime.fused.specification.RimeSpecification(specification,
                           terms=None,
                           transformers=None)
```

Defines a unique Radio Interferometer Measurement Equation (RIME)

The RIME is composed of a number of Jones Terms, which are multiplied together and combined to produce model visibilities.

The `RimeSpecification` specifies the order of these Jones Terms and supports custom Jones terms specified by the user.

One of the simplest RIME's that can be expressed involve a Phase (Kpq) and a Brightness (Bpq) term. The specification for this RIME is as follows:

```
rime_spec = RimeSpecification("(Kpq, Bpq): [I,Q,U,V] -> [XX,XY,YX,YY]")
```

(Kpq, Bpq) specifies the onion more formally defined [here](#), while [I,Q,U,V] -> [XX,XY,YX,YY] defines the stokes to correlation conversion within the RIME. It also identifies whether the RIME is handling linear or circular feeds.

### Term Configuration

The pq in Kpq and Bpq signifies that their values are calculated per-baseline. It is possible to specify per-antenna terms: Kp and Kq for example which represent left (ANTENNA1) and right (ANTENNA2) terms respectively. Note that the hermitian transpose of the right term is automatically performed and does not need to be implemented in the Term itself. Thus, for example, (Kp, Bpq, Kq) specifies a RIME where the Phase Term is separated into left and right terms, while the Brightness Matrix is calculated per-baseline.

### Stokes to Correlation Mapping

[I,Q,U,V] -> [XX,XY,YX,YY] specifies a mapping from four stokes parameters to four correlations. Both linear [XX,XY,YX,YY] and circular [RR,RL,LR,LL] feed types are supported. A variety of mappings are possible:

```
[I,Q,U,V] -> [XX,XY,YX,YY]
[I,Q] -> [XX,YY]
[I,Q] -> [RR,LL]
```

### Custom Terms

Custom Term classes implemented by a user can be added to the RIME as follows:

```
class CustomJones(Term):
    ...
spec = RimeSpecification("(Apq,Kpq,Bpq)", terms={"A": CustomJones})
```

### Parameters

**specification** [str] A string specifying the terms in the RIME and the stokes to correlation conversion.

**terms** [dict of str or Terms] A map describing custom *Term* implementations. If one has defined a custom Gaussian Term class, for use in RIME (Cpq, Kpq, Bpq), this should be supplied as `terms={"C": Gaussian}`. strings can be supplied for predefined RIME classes.

**transformers** [list of Transformers] A list of *Transformer* classes.

## class africanus.experimental.rime.fused.terms.core.Term

Base class for Terms which describe parts of the Fused RIME. Implementors of a RIME Term should inherit from it.

A Term is an object that defines how a term in the RIME should be sampled to produce the Jones Terms that make up the RIME. It therefore defines a sampling function, which in turn depends on arbitrary inputs for performing the sampling.

A high degree of flexibility and leeway is afforded when implementing a Term. It might be implemented by merely indexing an array of Jones Matrices, or by implementing some computational model describing the Jones Terms.

```
class Phase(Term):
    def __init__(self, configuration):
        super().__init__(configuration)
```

**init\_fields**(*self*, *typing\_ctx*, *arg1*, ..., *argn*, *kwarg1=None*, ..., *kwargn=None*)

Requests inputs to the RIME term, ensuring that they are stored on a `state` object supplied to the sampling function and allows for new fields to be initialised and stored on the `state` object.

Requested inputs *arg1*...*argn* are required to be passed to the Fused RIME by the caller and are supplied to `init_fields` as Numba types. *kwarg1*...*kwargn* are optional – if omitted by the caller, their default types (and values) will be supplied.

`init_fields` should return a (`fields`, `function`) tuple. `fields` should be a list of the form `[(name, numba_type)]`, while `function` should be a function of the form `fn(arg1, ..., argn, kwarg1=None, ..., kwargn=None)` and should return the variables of the type defined in `fields`. Note that it's signature therefore matches that of `init_fields` from after the `typingctx` argument. See the [Simple Example](#).

**Parameters**

- `typingctx` – A Numba typing context.
- `arg1...argn` – Required RIME inputs for this Term.
- `kwarg1...kwargn` – Optional RIME inputs for this Term. Types here should be simple: ints, floats, complex numbers and strings are ideal.

**Return type** `tuple`

**Returns** A (`fields`, `function`) tuple.

**Warning:** The function returned by `init_fields` must be compileable in Numba's `nopython` mode.

**sampler**(*self*)

Return a sampling function of the following form:

```
def sampler(self):
    def sample(state, s, r, t, f1, f2, a1, a2, c):
        ...
    return sample
```

**Parameters**

- `state` – A state object containing the inputs requested by all `Term` objects in the RIME, as well as any fields created by `Term.init_fields`.
- `s` – Source index.
- `r` – Row index.
- `t` – Time index.
- `f1` – Feed 1 index.
- `f2` – Feed 2 index.
- `a1` – Antenna 1 index.
- `a2` – Antenna2 index.

- **c** – Channel index.

**Return type** scalar or a tuple

**Returns** a scalar or a tuple of two scalars or a tuple of four scalars.

**Warning:** The sampling function returned by `sampler` must be compileable in Numba's `nopython` mode.

**dask\_schema**(*self*, *arg1*, ..., *argn*, *kwargs1=None*, ..., *kwargn=None*)

#### Parameters

- **arg1...argn** – Required RIME inputs for this Transformer.
- **kwarg1...kwargn** – Optional RIME inputs for this Transformer. Types here should be simple: ints, floats, complex numbers and strings are ideal.

**Return type** `dict`

**Returns** A dictionary of the form `{name: schema}` defining the `blockwise()` dimension schema of each supplied argument and keyword argument.

**class** `africanus.experimental.rime.fused.transformers.core.Transformer`

Base class for precomputing data for consumption by `Term`'s.

#### OUTPUTS

This class attributes should contain names of the outputs produced by the Transformer class. This should correspond to the fields produced by `Transformer.init_fields()`.

**init\_fields**(*self*, *typing\_ctx*, *arg1*, ..., *argn*, *kwargs1=None*, ..., *kwargn=None*)

Requests inputs to the Transformer, and specifies new fields and the function for creating them on the `state` object. Functionally, this method behaves exactly the same as the `init_fields()` method, the difference being that the outputs are available to all Terms.

**Return type** `tuple`

**Returns** A (`fields`, `function`) tuple.

**Warning:** The function returned by `init_fields` must be compileable in Numba's `nopython` mode.

**dask\_schema**(*self*, *arg1*, ..., *argn*, *kwargs1=None*, ..., *kwargn=None*)

**Return type** `tuple`

**Returns**

A (`inputs`, `outputs`) tuple.

`inputs` should be a dictionary of the form `{name: schema}` where `schema` is a dimension schema suitable for use in `dask.array.blockwise()`. A suitable schema for visibility data would be `(row, chan, corr)`, while a uvw coordinate schema could be `(row, uvw-component)`.

`outputs` should be a dictionary of the form `{name: array_like}`, where `array_like` is an object with `dtype` and `ndim` attributes. A suitable `array_like` for lm data could be `np.empty((0,0), dtype=np.float64)`.

## Predefined Terms

```
class africanus.experimental.rime.fused.terms.phase.Phase(configuration)
    Phase Delay Term
```

### Attributes

**configuration**

```
class africanus.experimental.rime.fused.terms.brightness.Brightness(configuration, stokes,
corrs)
```

Brightness Matrix Term

### Attributes

**configuration**

```
class africanus.experimental.rime.fused.terms.gaussian.Gaussian(configuration)
```

Gaussian Amplitude Term

### Attributes

**configuration**

```
class africanus.experimental.rime.fused.terms.feed_rotation.FeedRotation(configuration,
feed_type)
```

Feed Rotation Term

### Attributes

**configuration**

```
class africanus.experimental.rime.fused.terms.cube_dde.BeamCubeDDE(configuration, corrs)
```

Voxel Beam Cube Term

### Attributes

**configuration**

## Predefined Transformers

```
class africanus.experimental.rime.fused.transformers.lm.LMTransformer
```

```
class africanus.experimental.rime.fused.transformers.parangle.ParallacticTransformer(process_pool)
```

## 5.12.5 Numpy

---

`rime(rime_spec, *args, **kw)`

Evaluates the Radio Interferometer Measurement Equation (RIME), given the Specification of the RIME `rime_spec`, as well as the inputs to the RIME given in `*args` and `**kwargs`.

---

`africanus.experimental.rime.fused.core.rime(rime_spec, *args, **kw)`

Evaluates the Radio Interferometer Measurement Equation (RIME), given the Specification of the RIME `rime_spec`, as well as the inputs to the RIME given in `*args` and `**kwargs`.

## 5.12.6 Dask

---

<code>rime(rime_spec, *args, **kw)</code>	Like <a href="#">rime()</a> , but for a dask paradigm
---	---

---

`africanus.experimental.rime.fused.dask.rime(rime_spec, *args, **kw)`

Like [rime\(\)](#), but for a dask paradigm

## **CONTRIBUTING**

Contributions are welcome, and they are greatly appreciated! Every little bit helps, and credit will always be given.

You can contribute in many ways:

### **6.1 Types of Contributions**

#### **6.1.1 Report Bugs**

Report bugs at <https://github.com/ska-sa/codex-africanus/issues>.

If you are reporting a bug, please include:

- Your operating system name and version.
- Any details about your local setup that might be helpful in troubleshooting.
- Detailed steps to reproduce the bug.

#### **6.1.2 Fix Bugs**

Look through the GitHub issues for bugs. Anything tagged with “bug” and “help wanted” is open to whoever wants to implement it.

#### **6.1.3 Implement Features**

Look through the GitHub issues for features. Anything tagged with “enhancement” and “help wanted” is open to whoever wants to implement it.

#### **6.1.4 Write Documentation**

Codex Africanus could always use more documentation, whether as part of the official Codex Africanus docs, in doc-strings, or even on the web in blog posts, articles, and such.

### 6.1.5 Submit Feedback

The best way to send feedback is to file an issue at <https://github.com/ska-sa/codex-africanus/issues>.

If you are proposing a feature:

- Explain in detail how it would work.
- Keep the scope as narrow as possible, to make it easier to implement.
- Remember that this is a volunteer-driven project, and that contributions are welcome :)

## 6.2 Get Started!

Ready to contribute? Here's how to set up *codex-africanus* for local development.

1. Fork the *codex-africanus* repo on GitHub.
2. Clone your fork locally:

```
$ git clone git@github.com:your_name_here/codex-africanus.git
```

3. Install your local copy into a virtualenv. Assuming you have virtualenvwrapper installed, this is how you set up your fork for local development:

```
$ mkvirtualenv codex-africanus
$ cd codex-africanus/
$ pip install -e .
```

4. Create a branch for local development:

```
$ git checkout -b name-of-your-bugfix-or-feature
```

Now you can make your changes locally.

5. When you're done making changes, check that your changes pass the test cases, fixup your PEP8 compliance, and check for any code style issues:

```
$ py.test -v africanus $ autopep8 -r -i africanus $ flake8 africanus $ pycodestyle africanus
```

To get autopep8 and pycodestyle, just pip install them into your virtualenv.

6. Commit your changes and push your branch to GitHub:

```
$ git add .
$ git commit -m "Your detailed description of your changes."
$ git push origin name-of-your-bugfix-or-feature
```

7. Submit a pull request through the GitHub website.

## 6.3 Pull Request Guidelines

Before you submit a pull request, check that it meets these guidelines:

1. The pull request should include tests.
2. If the pull request adds functionality, the docs should be updated. Put your new functionality into a function with a docstring, and add the feature to the list in HISTORY.rst.
3. The pull request should work for Python 2.7, 3.5 and 3.6. Check [https://travis-ci.org/ska-sa/codex-africanus/pull\\_requests](https://travis-ci.org/ska-sa/codex-africanus/pull_requests) and make sure that the tests pass for all supported Python versions.

## 6.4 Tips

To run the tests:

```
$ py.test -vvv africanus/
```

## 6.5 Deploying

A reminder for the maintainers on how to deploy.

1. Update HISTORY.rst with the intended release number Z.Y.X and commit to git.
2. Bump the version number with bumpversion. This creates a new git commit, as well as an annotated tag Z.Y.X for the release. If your current version is Z.Y.W and the new version is Z.Y.X call:

```
$ python -m pip install bump2version
$ bump2version --current-version Z.Y.W --new-version Z.Y.X patch
```

3. Push the release commit and new tag up:

```
$ git push --follow-tags
```

4. Travis should automatically deploy the tagged release to PyPI if the automated tests pass.



## CREDITS

### 7.1 Development Lead

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---

**CHAPTER  
EIGHT**

---

**HISTORY**

## 8.1 X.Y.Z (YYYY-MM-DD)

- Document Fused RIME ([GH#270](#))
- Add Multiton, LazyProxy and LazyProxyMultiton patterns ([GH#269](#))

## 8.2 0.3.2 (2022-13-01)

- Support numba >= 0.54 ([GH#264](#))
- Fused RIME ([GH#239](#))
- Restrict numba version to <= 0.54.0 ([GH#259](#))
- BDA fix typos in numba wrapper ([GH#261](#))
- BDA Time-smearing fixes ([GH#253](#))

## 8.3 0.3.1 (2021-09-09)

- Handle empty spectral indices in WSClean Model ([GH#258](#))
- Support missing data during BDA ([GH#252](#))

## 8.4 0.3.0 (2021-09-09)

- Deprecate Python 3.6 support, add Python 3.9 support ([GH#248](#))
- Using `contextlib.suppress` instead of deprecated `dask.util.ignoring` in `EstimatingProgressBar` ([GH#256](#))
- Disallow numba 0.54.0 ([GH#254](#))
- Upgrade to CuPy 9.0 and fix template encoding ([GH#251](#))
- Parse and zero spectral models containing ‘nan’ and ‘inf’ in wsClean model files ([GH#250](#))
- Clarify \_wrapper names ([GH#247](#))
- Baseline-Dependent Time-and-Channel Averaging ([GH#173](#), [GH#243](#))

## 8.5 0.2.10 (2021-02-09)

- Don't let dof go to zero during spi fitting ([GH#240](#))
- Add support for Shapelets and Zernike Polynomials ([GH#231](#))
- Add beam model during SPI fitting ([GH#238](#))
- Add double accumulation option and Hessian function to wgridder ([GH#237](#))
- Upgrade ducc0 to version 0.8.0 ([GH#236](#))
- Add mindet to avoid div0 errors in spi fitter and fix alpha and I0 variance estimates ([GH#234](#))

## 8.6 0.2.9 (2020-12-15)

- Upgrade ducc0 to version 0.7.0 ([GH#233](#))
- Fix manually specifying wgridder precision ([GH#230](#))

## 8.7 0.2.8 (2020-10-08)

- Fix NoneType issue in wgridder when weights are None ([GH#228](#))
- Bounding hull geometric and image manipulation routines ([GH#192](#), [GH#154](#))
- Fix row chunk chunking in Perley Polyhedron Degridder Dask Interface

## 8.8 0.2.7 (2020-09-23)

- Deprecate old griddler and filters ([GH#224](#))
- Upgrade to ducc0 0.6.0 ([GH#223](#))
- Add Perley Polyhedron Faceting Griddler/Degridder ([GH#202](#), [GH#215](#), [GH#222](#))

## 8.9 0.2.6 (2020-08-07)

- Add wrapper for ducc0.wgridder ([GH#204](#))
- Correct Irregular Grid nesting in BeamAxes ([GH#217](#))

## 8.10 0.2.5 (2020-07-01)

- Convert WSClean Gaussian arcsecond and degree quantities to radians ([GH#206](#))
- Update classifiers and correct license in setup.py to BSD3 ([GH#201](#))

## 8.11 0.2.4 (2020-05-29)

- Support overriding the l and m axis sign in beam\_grids ([GH#199](#))
- Upgrade to python-casacore 3.3.1 ([GH#197](#))
- Upgrade to jax 0.1.68 and jaxlib 0.1.47 ([GH#197](#))
- Upgrade to scipy 1.4.0 ([GH#197](#))
- Use github workflows ([GH#196](#), [GH#197](#), [GH#198](#), [GH#199](#))
- Make CASA parallactic angles thread-safe ([GH#195](#))
- Fix spectral model documentation ([GH#190](#)), to match changes in ([GH#189](#))

## 8.12 0.2.3 (2020-05-14)

- Fix incorrect SPI calculation and make predict defaults MeqTree equivalent ([GH#189](#))
- Depend on pytest-flake8 >= 1.0.6 ([GH#187](#), [GH#188](#))
- MeqTrees Comparison Script Updates ([GH#160](#))
- Improve requirements handling ([GH#187](#))
- Use python-casacore wheels for travis testing, instead of kernsuite packages ([GH#185](#))

## 8.13 0.2.2 (2020-04-09)

- Add a dask Estimating Progress Bar ([GH#182](#), [GH#183](#))

## 8.14 0.2.1 (2020-04-03)

- Update trove to latest master commit ([GH#178](#))
- Added Cubic Spline support ([GH#174](#))
- Depend on python-casacore >= 3.2.0 ([GH#172](#))
- Drop Python 3.5 support and test Python 3.7 ([GH#168](#))
- Implement optimised WSClean predict ([GH#166](#), [GH#167](#), [GH#177](#), [GH#179](#), [GH#180](#), [GH#181](#))
- Simplify dask predict\_vis code ([GH#164](#), [GH#165](#))
- Document and check weight shapes in simple gridded and degridded ([GH#162](#), [GH#163](#))
- Restructuring calibration module ([GH#127](#))

- Upgrade to numba 0.46.0, using new inlining functionality in the RIME and averaging code.
- Modified predict to be compatible with eidos fits headers ([GH#158](#))

## 8.15 0.2.0 (2019-09-30)

- Added standalone SPI fitter ([GH#153](#))
- Fail earlier and explain duplicate averaging rows ([GH#155](#))
- CUDA Beam Implementation ([GH#152](#))
- Fix documentation package versions ([GH#151](#))
- Deprecate experimental w-stacking gridded in favour of nifty gridded ([GH#148](#))
- Expand travis build matrix ([GH#147](#))
- Drop Python 2 support ([GH#146](#), [GH#149](#), [GH#150](#))
- Support the beam in the predict example ([GH#145](#))
- Fix weight indexing in averaging ([GH#144](#))
- Support EFFECTIVE\_BW and RESOLUTION in averaging ([GH#144](#))
- Optimise predict\_vis jones coherency summation ([GH#143](#))
- Remove use of @wraps ([GH#141](#), [GH#142](#))
- Set row chunks to nan in dask averaging code. ([GH#139](#))
- predict\_vis documentation improvements ([GH#135](#), [GH#140](#))
- Upgrade to dask-ms in the examples ([GH#134](#), [GH#138](#))
- Explain how to obtain predict\_vis time\_index argument ([GH#130](#))
- Update RIME predict example to support Tigger LSM's and Gaussians ([GH#129](#))
- Add dask wrappers for the nifty gridded ([GH#116](#), [GH#136](#), [GH#146](#))
- Testing and requirement updates. ([GH#124](#))
- Upgraded DFT kernels to have a correlation axis and added flags for vis\_to\_im. Added predict\_from\_fits example. ([GH#122](#))
- Fixed segfault when using `_unique_internal` on empty ndarrays ([GH#123](#))
- Removed `apply_gains`. Use `africanus.calibration.utils.correct_vis` instead ([GH#118](#))
- Add streams parameter to dask `predict_vis` ([GH#118](#))
- Implement the beam in numba ([GH#112](#))
- Add residual\_vis, correct\_vis, phase\_only\_GN ([GH#113](#))

## 8.16 0.1.8 (2019-05-28)

- Use environment markers in setup.py (GH#110)
- Add *apply\_gains*, a wrapper around *predict\_vis* (GH#108)
- Fix testing extras\_require (GH#107)
- Fix WEIGHT\_SPECTRUM averaging and add more averaging tests (GH#106)

## 8.17 0.1.7 (2019-05-09)

- Even more support for automated travis deploys.

## 8.18 0.1.6 (2019-05-09)

- Support automated travis deploys.

## 8.19 0.1.5 (2019-05-09)

- Predict script enhancements (GH#103) and dask channel chunking fix (GH#104).
- Directly jit DFT functions (GH#100, GH#101)
- Spectral Models (GH#86)
- Fix radec sign conversion in wsclean sky model (GH#96)
- Full Time and Channel Averaging Implementation (GH#80, GH#97, GH#98)
- Support integer seconds in wsclean ra and dec columns (GH#91, GH#93)
- Fix ratio computation in Gaussian Shape (GH#89, GH#90)

## 8.20 0.1.4 (2019-03-11)

- Support *complete* and *complete-cuda* to support non-GPU installs (GH#87)
- Gaussian Shape Parameter Implementation (GH#82, GH#83)
- WSClean Spectral Model (GH#81)
- Compare predict versus MeqTrees (GH#79)
- Time and channel averaging (GH#75)
- cupy implementation of *predict\_vis* (GH#73)
- Introduce transpose in second antenna term of predict (GH#72)
- cupy implementation of *feed\_rotation* (GH#67)
- cupy implementation of *stokes\_convert* kernel (GH#65)
- Add a basic RIME example (GH#64)

- requires\_optional accepts ImportError's for a better debugging experience ([GH#62](#), [GH#63](#))
- Added *fit\_component\_spi* function ([GH#61](#))
- cupy implementation of the *phase\_delay* kernel ([GH#59](#))
- Correct *phase\_delay* argument ordering ([GH#57](#))
- Support dask for *radec\_to\_lmn* and *lmn\_to\_radec*. Also add support for *radec\_to\_lm* and *lm\_to\_radec* ([GH#56](#))
- Added test for dft to test if image space covariance is symmetric([GH#55](#))
- Correct Parallactic Angle Computation ([GH#49](#))
- Enhance visibility predict ([GH#50](#))
- Fix Kaiser Bessel filter and taper ([GH#48](#))
- Stokes/Correlation conversion ([GH#41](#))
- Fix gridding examples ([GH#43](#))
- Add simple dask gridded example ([GH#42](#))
- Implement Kaiser Bessel filter ([GH#38](#))
- Implement W-stacking gridded/degridded ([GH#38](#))
- Use 2D filters by default ([GH#37](#))
- Fixed bug in im\_to\_vis. Added more tests for im\_to\_vis. Removed division by  $n$  since it is trivial to reinstate after the fact. ([GH#34](#))
- Move numba implementations out of API functions. ([GH#33](#))
- Zernike Polynomial Direction Dependent Effects ([GH#18](#), [GH#30](#))
- Added division by  $n$  to DFT. Fixed dask chunking issue. Updated test\_vis\_to\_im\_dask ([GH#29](#)).
- Implement RIME visibility predict ([GH#24](#), [GH#25](#))
- Direct Fourier Transform ([GH#19](#))
- Parallactic Angle computation ([GH#15](#))
- Implement Feed Rotation term ([GH#14](#))
- Swap gridding correlation dimensions ([GH#13](#))
- Implement Direction Dependent Effect beam cubes ([GH#12](#))
- Implement Brightness Matrix Calculation ([GH#9](#))
- Implement RIME Phase Delay term ([GH#8](#))
- Support user supplied grids ([GH#7](#))
- Add dask wrappers to the gridded and degridded ([GH#4](#))
- Add weights to gridded/degridded and remove PSF function ([GH#2](#))

## **8.21 0.1.2 (2018-03-28)**

- First release on PyPI.



---

**CHAPTER  
NINE**

---

**INDICES AND TABLES**

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