# **LuxPy Documentation**

Release 1.9.6

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# **CONTENTS:**

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• License: GPLv3



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# **CHAPTER**

# **ONE**

**LICENSE: GPLV3** 

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# INSTALLATION

# 2.1 Install luxpy

- 1. Install miniconda
  - download the installer from: https://conda.io/miniconda.html or https://repo.continuum.io/miniconda/)
  - e.g. https://repo.continuum.io/miniconda/Miniconda3-latest-Windows-x86\_64.exe
  - Make sure 'conda.exe' can be found on the windows system path, if necessary do a manual add.
- 2. Create a virtual environment with full anaconda distribution by typing the following at the commandline:
  - >> conda create --name py36 python=3.6 anaconda
- 3. Activate the virtual environment:
  - >> activate py36
- 4. **Install pip to virtual environment (just to ensure any packages to be** installed with pip to this virt. env. will be installed here and not globally):
  - >> conda install -n py36 pip
- 5a. Install luxpy package from pypi:
  - >> pip install luxpy
- 5b. Install luxpy package from anaconda:
  - >> conda install -c ksmet1977 luxpy
- **Note** If any errors show up, try and do a manual install of the dependencies: scipy, numpy, pandas, matplotlib and setuptools, either using e.g. >> conda install scipy or >> pip install scipy, and try and reinstall luxpy using pip.

# 2.2 Use of LuxPy package in Spyder IDE

- 6. Install spyder in py36 environment:
  - >> conda install -n py36 spyder
- 7. Run spyder
  - >> spyder
- 8. To import the luxpy package, on Spyder's commandline for the IPython kernel (or in script) type:

```
import luxpy as lx
```

# 2.3 Use of LuxPy package in Jupyter notebook

- 6. Install jupyter in py36 environment:
  - >> conda install -n py36 jupyter
- 7. Start jupyter notebook:
  - >> jupyter notebook
- 8. **Open an existing or new notebook:** e.g. open "luxpy\_basic\_usage.ipynb" for an overview of how to use the LuxPy package.
- 9. To import LuxPy package type:

```
import luxpy as lx
```

# **THREE**

# **IMPORTED (REQUIRED) PACKAGES**

# **3.1 Core**

- import os
- import warnings
- import pathlib
- import importlib
- from collections import OrderedDict as odict
- from mpl\_toolkits.mplot3d import Axes3D
- import colorsys
- · import itertools
- import copy
- import time
- import tkinter
- import ctypes
- import platform
- · import subprocess
- import cProfile
- import pstats
- import io

# 3.2 3e party dependencies (automatic install)

- import numpy as np
- · import pandas as pd
- import matplotlib.pyplot as plt
- · import scipy as sp
- import imageio

# 3.3 3e party dependencies (automatic install on import)

• import pyswarms (when importing particleswarms from math)

# 3.4 3e party dependencies (requiring manual install)

To control Ocean Optics spectrometers with spectro toolbox:

- import seabreeze (conda install -c poehlmann python-seabreeze)
- pip install pyusb (for use with 'pyseabreeze' backend of python-seabreeze)

# **LUXPY PACKAGE STRUCTURE**

# 4.1 Utils sub-package

```
рy
                 • __init__.py
                 • utilities.py
                 • folder_tree.py
           namespace luxpy.utils
luxpy.utils.np2d(data)
      Make a tuple, list or numpy array at least a 2D numpy array.
      Args:
               data
                   tuple, list, ndarray
      Returns:
               returns
                   ndarray with .ndim >= 2
luxpy.utils.np3d(data)
     Make a tuple, list or numpy array at least a 3d numpy array.
      Args:
               data
                   tuple, list, ndarray
      Returns:
               returns
                   ndarray with .ndim >= 3
luxpy.utils.np2dT(data)
      Make a tuple, list or numpy array at least a 2D numpy array and transpose.
      Args:
               data
```

tuple, list, ndarray

**Returns:** 

#### returns

ndarray with .ndim  $\geq 2$  and with transposed axes.

# luxpy.utils.np3dT(data)

Make a tuple, list or numpy array at least a 3d numpy array and transposed first 2 axes.

### **Args:**

#### data

tuple, list, ndarray

#### **Returns:**

#### returns

ndarray with .ndim >= 3 and with first two axes transposed (axis=3 is kept the same).

# luxpy.utils.put\_args\_in\_db(db, args)

Takes the args with not-None input values of a function and overwrites the values of the corresponding keys in dict db. | (args are collected with the built-in function locals(), | See example usage below)

# Args:

db

dict

#### **Returns:**

### returns

dict with the values of specific keys overwritten by the not-None values of corresponding args of a function fcn.

# Example usage:

print('\_db: { }'.format(\_db))

```
luxpy.utils.vec_to_dict(vec=None, dic={}, vsize=None, keys=None)
      Convert dict to vec and vice versa.
      Args:
                vec
                    list or vector array, optional
                dic
                    dict, optional
                vsize
                    list or vector array with size of values of dict, optional
                keys
                    list or vector array with keys in dict (must be provided).
      Returns:
                returns
                    x, vsize
                           x is an array, if vec is None
                          x is a dict, if vec is not None
luxpy.utils.getdata(data, kind='np', columns=None, header=None, sep=',', datatype='S', copy=True,
                          verbosity=True)
      Get data from csv-file or convert between pandas dataframe and numpy 2d-array.
      Args:
                data
                    - str with path to file containing data
                    - ndarray with data
                    - pandas.dataframe with data
                kind
                    str ['np','df'], optional
                    Determines type(:returns:), np: ndarray, df: pandas.dataframe
                columns
                    None or list[str] of column names for dataframe, optional
                header
                    None, optional
                           - None: no header in file
                          - 'infer': infer headers from file
                sep
                    ',' or ' ' or other char, optional
                    Column separator in data file
                datatype'
                    'S',optional
                    Specifies a type of data.
```

```
Is used when creating column headers (:column: is None).
                          -'S': light source spectrum
                          -'R': reflectance spectrum
                          or other.
                copy
                    True, optional
                    Return a copy of ndarray if kind == 'np', or copy of pd.DataFrame if kind == 'df'
                verbosity
                    True, False, optional
                    Print warning when inferring headers from file.
      Returns:
                returns
                    data as ndarray or pandas.dataframe
luxpy.utils.dictkv(keys=None, values=None, ordered=True)
      Easy input of of keys and values into dict.
      Args:
                keys
                    iterable list[str,...] of keys
                values
                    iterable list[...,...,] of values
                ordered
                    True, False, optional
                    True: creates an ordered dict using 'collections.OrderedDict()'
      Returns:
                returns
                    (ordered) dict
luxpy.utils.meshblock(x, y)
      Create a meshed block from x and y.
      (Similar to meshgrid, but axis = 0 is retained).
      To enable fast blockwise calculation.
      Args:
                X
                    ndarray with ndim == 2
                y
                    ndarray with ndim == 2
      Returns:
                X,Y
```

```
2 ndarrays with ndim == 3
                         X.shape = (x.shape[0], y.shape[0], x.shape[1])
                          Y.shape = (x.shape[0], y.shape[0], y.shape[1])
luxpy.utils.asplit(data)
     Split data on last axis
     Args:
               data
                   ndarray
     Returns:
               returns
                   ndarray, ndarray, ...
                         (number of returns is equal data.shape[-1])
luxpy.utils.ajoin(data)
     Join data on last axis.
     Args:
               data
                   tuple (ndarray, ndarray, ...)
     Returns:
               returns
                   ndarray (shape[-1] is equal to tuple length)
luxpy.utils.broadcast_shape(data, target shape=None, expand 2d to 3d=None, axis0 repeats=None,
                                   axis1_repeats=None)
     Broadcasts shapes of data to a target_shape.
     Useful for block/vector calc. when numpy fails to broadcast correctly.
     Args:
               data
                   ndarray
               target shape
                   None or tuple with requested shape, optional
                         - None: returns unchanged :data:
               expand_2d_to_3d
                   None (do nothing) or ..., optional
                   If ndim == 2, expand from 2 to 3 dimensions
               axis0_repeats
                   None or number of times to repeat axis=0, optional
                         - None: keep axis=0 same size
               axis1_repeats
```

None or number of times to repeat axis=1, optional

- None: keep axis=1 same size

```
Returns:
                returns
                    reshaped ndarray
luxpy.utils.todim(x, tshape, add_axis=1, equal_shape=False)
      Expand x to dims that are broadcast-compatable with shape of another array.
      Args:
                X
                    ndarray
                tshape
                    tuple with target shape
                add_axis
                    1, optional
                    Determines where in x.shape an axis should be added
                equal_shape
                    False or True, optional
                    True: expand :x: to identical dimensions (speficied by :tshape:)
      Returns:
                returns
                    ndarray broadcast-compatable with tshape.
luxpy.utils.write_to_excel(filename, df, sheet_name='Sheet1', startrow=None, truncate_sheet=False,
                                  **to_excel_kwargs)
      Writes a DataFrame to an existing Excel file into a specified sheet. | If [filename] doesn't exist, then this function
      will create it.
      Args:
                filename
                    File path or existing ExcelWriter
                    (Example: '/path/to/file.xlsx')
                df
                    dataframe to save to workbook
                sheet_name
                    Name of sheet which will contain DataFrame.
                    (default: 'Sheet1')
                startrow
                    upper left cell row to dump data frame.
                    Per default (startrow=None) calculate the last row
                    in the existing DF and write to the next row...
                truncate_sheet
```

```
truncate (remove and recreate) [sheet_name]
                                               before writing DataFrame to Excel file
                                      to_excel_kwargs
                                                arguments which will be passed to DataFrame.to_excel()
                                                [can be dictionary]
              Returns: None
              Notes: Copied from https://stackoverflow.com/questions/20219254/how-to-write-to-an-existing-excel-file-without-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-overwriting-ove
luxpy.utils.show_luxpy_tree(omit=['.pyc', '__pycache__', '.txt', '.dat', '.csv', '.npz', '.png', '.jpg', '.md', '.pdf',
                                                                                      '.ini', '.log', '.rar', 'drivers', 'SDK_', 'dll', 'bak'])
              Show luxpy foler tree.
              Args:
                                      omit
                                               List of folders and file-extensions to omit.
              Returns: None
luxpy.utils.is_importable(string, try_pip_install=False)
              Check if string is importable/loadable. If it doesn't then try to 'pip install' it using subprocess. Returns None if
              succesful, otherwise throws and error or outputs False.
              Args:
                                      string
                                               string with package or module name
                                      try_pip_install
                                               False, optional
                                               True: try pip installing it using subprocess
              Returns:
                                      success
                                                True if importable, False if not.
luxpy.utils.get_function_kwargs(f)
              Get dictionary of a function's keyword arguments and their default values.
              Args:
                                     f
                                                function name
              Returns:
                                      dict
                                               Dict with the function's keyword arguments and their default values
                                               Is empty if there are no defaults (i.e. f.__defaults__ or f.__kwdefaults__ are None).
luxpy.utils.profile_fcn(fcn, profile=True, sort_stats='tottime', output_file=None)
              Profile or time a function fcn.
              Args:
                                      fcn
```

```
function to be profiled or timed (using time.time() difference)
                profile
                    True, optional
                    Profile the function, otherwise only time it.
                sort\_stats
                    'tottime', optional
                    Sort profile results according to sort_stats ('tottime', 'cumtime',...)
                output_file
                    None, optional
                    If not None: output result to output_file.
      Return:
                ps
                    Profiler output
luxpy.utils.unique(array, sort=True)
      Get unique elements from array.
      Args:
                array
                    array to get unique elements from.
                sort
                    True, optional
                    If True: get sorted unique elements.
      Returns:
                unique_array
                    ndarray with (sorted) unique elements.
luxpy.utils.save_pkl(filename, obj)
      Save an object in a pickle file.
      Args:
                filename
                    str with filename of pickle file.
                obj
                    python object to save
      Returns:
                None
luxpy.utils.load_pkl(filename)
      Load the object in a pickle file.
      Args:
                filename
                    str with filename of pickle file.
```

#### **Returns:**

obj

loaded python object

Given a directory Path object print a visual tree structure

#### **References:**

1. https://stackoverflow.com/questions/9727673/list-directory-tree-structure-in-python

# 4.2 Math sub-package

рy

- \_\_init\_\_.py
- · basics.py
- minimizebnd.py
- · mupolymodel.py
- Pyswarms\_particleswarm.py
- pymoo\_nsga\_ii.py

namespace luxpy.math

### 4.2.1 Module with useful math functions

```
normalize_3x3_matrix() Normalize 3x3 matrix M to xyz0 -> [1,1,1]

line_intersect()

Line intersections of series of two line segments a and b.

https://stackoverflow.com/questions/3252194/numpy-and-line-intersections

positive_arctan() Calculates the positive angle (0°-360° or 0 - 2*pi rad.) from x and y.

dot23() Dot product of a 2-d ndarray with a (N x K x L) 3-d ndarray using einsum().

check_symmetric() Checks if A is symmetric.

check_posdef() Checks positive definiteness of a matrix via Cholesky.

symmM_to_posdefM()

Converts a symmetric matrix to a positive definite one.

Two methods are supported:

* 'make': A Python/Numpy port of Muhammad Asim Mubeen's matlab function Spd_Mat.m

(https://nl.mathworks.com/matlabcentral/fileexchange/
45873-positive-definite-matrix)
```

\* 'nearest': A Python/Numpy port of John D'Errico's 'nearestSPD' MATLAB code. (https://stackoverflow.com/questions/43238173/ python-convert-matrix-to-positive-semi-definite)

**bvgpdf**() Evaluate bivariate Gaussian probability density function (BVGPDF) at (x,y) with center mu and inverse covariance matric, sigmainv.

**mahalanobis2**() Evaluate the squared mahalanobis distance with center mu and shape and orientation determined by sigmainv.

rms() Calculates root-mean-square along axis.

geomean() Calculates geometric mean along axis.

### polyarea()

Calculates area of polygon.

(First coordinate should also be last)

erf(), erfinv() erf-function and its inverse, direct import from scipy.special

**cart2pol()** Converts Cartesian to polar coordinates.

pol2cart() Converts polar to Cartesian coordinates.

cart2spher() Converts Cartesian to spherical coordinates.

spher2cart() Converts spherical to Cartesian coordinates.

magnitude\_v() Calculates magnitude of vector.

angle\_v1v2() Calculates angle between two vectors.

#### histogram()

Histogram function that can take as bins either the center (cfr. matlab hist) or bin-edges.

v\_to\_cik() Calculate 2x2 '(covariance matrix)^-1' elements cik from v-format ellipse descriptor

cik to v() Calculate v-format ellipse descriptor from 2x2 'covariance matrix'^-1 cik.

**minimizebnd()** scipy.minimize() that allows contrained parameters on unconstrained methods(port of Matlab's fminsearchbnd). Starting, lower and upper bounds values can also be provided as a dict.

**DEMO** Module for Differential Evolutionary Multi-objective Optimization (DEMO).

vec3 Module for spherical vector coordinates.

fmod() Floating point modulus, e.g.: fmod(theta, np.pi \* 2) would keep an angle in [0, 2pi]b

**fit\_ellipse()** Fit an ellipse to supplied data points.

**fit\_cov\_ellipse()** Fit an covariance ellipse to supplied data points.

interp1() Perform a 1-dimensional linear interpolation (wrapper around scipy.interpolate.InterpolatedUnivariateSpline).

**ndinterp1**() Perform n-dimensional interpolation using Delaunay triangulation.

**ndinterp1\_scipy()** Perform n-dimensional interpolation using Delaunay triangulation (wrapper around scipy.interpolate.LinearNDInterpolator)

**box\_m()** Performs a Box M test on covariance matrices.

pitman\_morgan() Pitman-Morgan Test for the difference between correlated variances with paired samples. mupolymod Module for Multivariate Polynomial Model Optimization (2D, 3D)

# **NOT IMPORTED in math-namespace (to minimize dependencies)**

```
pyswarms_particleswarm Module with particleswarm() function for global minimization using particle swarms (wrapper around pyswarms.single.GlobalBestPSO))
```

pymoo\_nsga\_ii Module with nsga\_ii() function for pareto-optimal boundary minimization using Non-Dominated-Sort-Genetic-Algorithm NSGA-II (wrapper around pymoo.NSGAII))

```
luxpy.math.normalize_3x3_matrix(M, xyz0=array([[1.0, 1.0, 1.0]]))
      Normalize 3x3 matrix M to xyz0 -> [1,1,1]
      If M.shape == (1,9): M is reshaped to (3,3)
      Args:
                 M
                     ndarray((3,3) \text{ or } ndarray((1,9))
                 xyz0
                     2darray, optional
      Returns:
                 returns
                     normalized matrix such that M*xyz0 = [1,1,1]
luxpy.math.symmM_to_posdefM(A=None, atol=1e-09, rtol=1e-09, method='make', forcesymm=True)
      Convert a symmetric matrix to a positive definite one.
      Args:
                 A
                     ndarray
                 atol
                     float, optional
                     The absolute tolerance parameter (see Notes of numpy.allclose())
                 rtol
                     float, optional
                     The relative tolerance parameter (see Notes of numpy.allclose())
                 method
                      'make' or 'nearest', optional (see notes for more info)
                 forcesymm
                     True or False, optional
                     If A is not symmetric, force symmetry using:
```

```
Returns:
                  returns
                       ndarray with positive-definite matrix.
      Notes on supported methods: 1. 'make': A Python/Numpy port of Muhammad Asim Mubeen's matlab func-
             tion Spd_Mat.m 2. 'nearest': A Python/Numpy port of John D'Errico's 'nearestSPD MATLAB code.
             <a href="https://stackoverflow.com/questions/43238173/python-convert-matrix-to-positive-semi-definite">https://stackoverflow.com/questions/43238173/python-convert-matrix-to-positive-semi-definite</a>
luxpy.math.check_symmetric(A, atol=1e-09, rtol=1e-09)
      Check if A is symmetric.
      Args:
                  A
                       ndarray
                  atol
                       float, optional
                       The absolute tolerance parameter (see Notes of numpy.allclose())
                  rtol
                       float, optional
                       The relative tolerance parameter (see Notes of numpy.allclose())
      Returns:
                  returns
                       Bool
                       True: the array is symmetric within the given tolerance
luxpy.math.check_posdef(A, atol=1e-09, rtol=1e-09)
      Checks positive definiteness of a matrix via Cholesky.
      Args:
                  A
                       ndarray
                  atol
                       float, optional
                       The absolute tolerance parameter (see Notes of numpy.allclose())
                  rtol
                       float, optional
                       The relative tolerance parameter (see Notes of numpy.allclose())
      Returns:
                  returns
                       Bool
                       True: the array is positive-definite within the given tolerance
luxpy.math.positive_arctan(x, y, htype='deg')
      Calculate positive angle (0^{\circ}-360° or 0 - 2*pi rad.) from x and y.
```

A = numpy.triu(A) + numpy.triu(A).T - numpy.diag(numpy.diag(A))

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# **Args:** $\mathbf{X}$ ndarray of x-coordinates y ndarray of y-coordinates htype 'deg' or 'rad', optional - 'deg': hue angle between $0^{\circ}$ and $360^{\circ}$ - 'rad': hue angle between 0 and 2pi radians **Returns:** returns ndarray of positive angles. luxpy.math.line\_intersect(a1, a2, b1, b2) Line intersections of series of two line segments a and b. **Args:** a1 ndarray (.shape = (N,2)) specifying end-point 1 of line a **a2** ndarray (.shape = (N,2)) specifying end-point 2 of line a b1 ndarray (.shape = (N,2)) specifying end-point 1 of line b **b2** ndarray (.shape = (N,2)) specifying end-point 2 of line b **Note:** N is the number of line segments a and b. **Returns:** returns ndarray with line-intersections (.shape = (N,2)) **References:** 1. https://stackoverflow.com/questions/3252194/numpy-and-line-intersections luxpy.math.histogram(a, bins=10, bin\_center=False, range=None, normed=False, weights=None, density=None) Histogram function that can take as bins either the center (cfr. matlab hist) or bin-edges. Args: bin\_center False, optional False: if :bins: int, str or sequence of scalars: default to numpy.histogram (uses bin edges).

True: if :bins: is a sequence of scalars:

```
bins (containing centers) are transformed to edges
and nump.histogram is run.
Mimicks matlab hist (uses bin centers).
```

Note: For other armuments and output, see ?numpy.histogram

#### **Returns:**

#### returns

ndarray with histogram

luxpy.math.pol2cart(theta, r=None, htype='deg')

Convert Cartesion to polar coordinates.

# Args:

#### theta

float or ndarray with theta-coordinates

r

None or float or ndarray with r-coordinates, optional If None, r-coordinates are assumed to be in :theta:.

# htype

```
'deg' or 'rad, optional
Intput type of :theta:.
```

### **Returns:**

### returns

(float or ndarray of x, float or ndarray of y) coordinates

```
luxpy.math.cart2pol(x, y=None, htype='deg')
```

Convert Cartesion to polar coordinates.

# **Args:**

X

float or ndarray with x-coordinates

y

None or float or ndarray with x-coordinates, optional If None, y-coordinates are assumed to be in :x:.

### htype

```
'deg' or 'rad, optional
Output type of theta.
```

### **Returns:**

### returns

(float or ndarray of theta, float or ndarray of r) values

luxpy.math.spher2cart(theta, phi, r=1.0, deg=True)

Convert spherical to cartesian coordinates.

#### Args:

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```
Float, int or ndarray
                      Angle with positive z-axis.
                  phi
                      Float, int or ndarray
                      Angle around positive z-axis starting from x-axis.
                  r
                      1, optional
                      Float, int or ndarray
                      radius
      Returns:
                  x, y, z
                      tuple of floats, ints or ndarrays
                      Cartesian coordinates
luxpy.math.cart2spher(x, y, z, deg=True)
      Convert cartesian to spherical coordinates.
      Args:
                  x, y, z
                      tuple of floats, ints or ndarrays
                      Cartesian coordinates
      Returns:
                  theta
                      Float, int or ndarray
                      Angle with positive z-axis.
                  phi
                      Float, int or ndarray
                      Angle around positive z-axis starting from x-axis.
                  r
                      1, optional
                      Float, int or ndarray
                      radius
luxpy.math.bvgpdf(x, y=None, mu=None, sigmainv=None)
      Evaluate bivariate Gaussian probability density function (BVGPDF)
      Args:
                  \mathbf{X}
                      scalar or list or ndarray (.ndim = 1 or 2) with
                      x(y)-coordinates at which to evaluate bivariate Gaussian PD.
                  y
                      None or scalar or list or ndarray (.ndim = 1) with
                      y-coordinates at which to evaluate bivariate Gaussian PD, optional.
```

theta

```
If:y: is None,:x: should be a 2d array.
```

#### mu

None or ndarray (.ndim = 2) with center coordinates of

bivariate Gaussian PD, optional.

None defaults to ndarray([0,0]).

### sigmainv

None or ndarray with 'inverse covariance matrix', optional

Determines the shape and orientation of the PD.

None default to numpy.eye(2).

#### **Returns:**

#### returns

ndarray with magnitude of BVGPDF(x,y)

luxpy.math.mahalanobis2(x, y=None, z=None, mu=None, sigmainv=None)

Evaluate the squared mahalanobis distance

# **Args:**

X

scalar or list or ndarray (.ndim = 1 or 2) with x(y)-coordinates at which to evaluate the mahalanobis distance squared.

y

None or scalar or list or ndarray (.ndim = 1) with y-coordinates at which to evaluate the mahalanobis distance squared, optional.

If :y: is None, :x: should be a 2d array.

Z

None or scalar or list or ndarray (.ndim = 1) with z-coordinates at which to evaluate the mahalanobis distance squared, optional.

If :z: is None & :y: is None, then :x: should be a 2d array.

#### mu

None or ndarray (.ndim = 1) with center coordinates of the mahalanobis ellipse, optional. None defaults to zeros(2) or zeros(3).

### sigmainv

None or ndarray with 'inverse covariance matrix', optional

Determines the shape and orientation of the PD.

None default to np.eye(2) or eye(3).

### **Returns:**

# returns

ndarray with magnitude of mahalanobis2(x,y[,z])

luxpy.math.dot23(A, B, keepdims=False)

Dot product of a 2-d ndarray with a (N x K x L) 3-d ndarray using einsum().

### **Args:**

A

```
ndarray (.shape = (M,N))
                 В
                     ndarray (.shape = (N,K,L))
      Returns:
                 returns
                     ndarray (.shape = (M,K,L))
luxpy.math.rms(data, axis=0, keepdims=False)
      Calculate root-mean-square along axis.
      Args:
                 data
                     list of values or ndarray
                 axis
                     0, optional
                     Axis along which to calculate rms.
                 keepdims
                     False or True, optional
                     Keep original dimensions of array.
      Returns:
                 returns
                     ndarray with rms values.
luxpy.math.geomean(data, axis=0, keepdims=False)
      Calculate geometric mean along axis.
      Args:
                 data
                     list of values or ndarray
                 axis
                     0, optional
                     Axis along which to calculate geomean.
                 keepdims
                     False or True, optional
                     Keep original dimensions of array.
      Returns:
```

# returns

ndarray with geomean values.

luxpy.math.polyarea(x, y)

Calculates area of polygon.

First coordinate should also be last.

```
Args:
                 X
                     ndarray of x-coordinates of polygon vertices.
                 y
                     ndarray of x-coordinates of polygon vertices.
      Returns:
                 returns
                     float (area or polygon)
luxpy.math.magnitude_v(v)
      Calculates magnitude of vector.
      Args:
                     ndarray with vector
      Returns:
                 magnitude
                     ndarray
luxpy.math.angle_v1v2(v1, v2, htype='deg')
      Calculates angle between two vectors.
      Args:
                 v1
                     ndarray with vector 1
                 v2
                     ndarray with vector 2
                 htype
                      'deg' or 'rad', optional
                     Requested angle type.
      Returns:
                 ang
                     ndarray
luxpy.math.v_to_cik(v, inverse=False)
      Calculate 2x2 '(covariance matrix)^-1' elements cik
      Args:
                     (Nx5) np.ndarray
                     ellipse parameters [Rmax,Rmin,xc,yc,theta]
                 inverse
                     If True: return inverse of cik.
```

```
Returns:
                 cik
                      'Nx2x2' (covariance matrix)^-1
      Notes:
            cik is not actually a covariance matrix,
            only for a Gaussian or normal distribution!
luxpy.math.cik_to_v(cik, xyc=None, inverse=False)
      Calculate v-format ellipse descriptor from 2x2 'covariance matrix'^-1 cik
      Args:
                 cik
                      'Nx2x2' (covariance matrix)^-1
                 inverse
                      If True: input is inverse of cik.
      Returns:
                      (Nx5) np.ndarray
                      ellipse parameters [Rmax,Rmin,xc,yc,theta]
      Notes:
            cik is not actually the inverse covariance matrix,
            only for a Gaussian or normal distribution!
luxpy.math.fmod(x, y)
      Floating point modulus
      e.g., fmod(theta, np.pi * 2) would keep an angle in [0, 2pi]
      Args:
                 \mathbf{X}
                      angle to restrict
                 y
                      end of interval [0, y] to restrict to
      Returns:
                 r floating point modulus
luxpy.math.remove_outliers(data, alpha=0.01)
      Remove multivariate outliers from data when outside of alpha-level confidence ellipsoid.
      Args:
                 data
                      Nxp ndarray with multivariate data (N samples, p variables)
                 alpha
```

```
0.01, optional
                      Significance level of confidence ellipsoid marking the boundary for outliers.
      Return:
                 data
                      (N-... x p) ndarray with multivariate data; outliers removed.
luxpy.math.fit_ellipse(xy, center_on_mean_xy=False)
      Fit an ellipse to supplied data points.
      Args:
                 хy
                      coordinates of points to fit (Nx2 array)
                 center_on_mean_xy
                      False, optional
                      Center ellipse on mean of xy
                      (otherwise it might be offset due to solving
                      the contrained minization problem: aT*S*a, see ref below.)
      Returns:
                      vector with ellipse parameters [Rmax,Rmin, xc,yc, theta (rad.)]
      Reference: 1. Fitzgibbon, A.W., Pilu, M., and Fischer R.B., Direct least squares fitting of ellipsees, Proc. of the
            13th Internation Conference on Pattern Recognition, pp 253–257, Vienna, 1996.
luxpy.math.fit_cov_ellipse(xy, alpha=0.05, pdf='chi2', SE=False, robust=False, robust alpha=0.01)
      Fit covariance ellipse to xy data.
      Args:
                 хy
                      coordinates of points to fit (Nx2 array)
                 alpha
                      0.05, optional
                      alpha significance level
                      (e.g alpha = 0.05 for 95% confidence ellipse)
                 pdf
                      chi2, optional
                      - 'chi2': Rescale using Chi2-distribution
                      - 't': Rescale using Student t-distribution
                      - 'norm': Rescale using normal-distribution
                      - None: don't rescale using pdf, use alpha as scalefactor (cfr. alpha* 1SD or alpha * 1SE)
                 SE
                      False, optional
                      If false, fit standard error ellipse at alpha significance level
                      If true, fit standard deviation ellipse at alpha significance level
                 robust
```

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```
If True: remove outliers beyond the confidence ellipsoid before calculating
                             the covariances.
                  robust_alpha
                      0.01, optional
                      Significance level of confidence ellipsoid marking the boundary for outliers.
      Returns:
                  V
                      vector with ellipse parameters [Rmax,Rmin, xc,yc, theta (rad.)]
luxpy.math.in_hull(p, hull)
      Test if points in p are in hull
      Args:
                  p
                      NxK coordinates of N points in K dimensions
                  hull
                      Either a scipy.spatial.Delaunay object or the MxK array of the
                      coordinates of M points in K dimensions for which Delaunay
                      triangulation will be computed
      Returns:
                  bool
                      boolean ndarray with True for in-gamut and False for out-of-gamut points
luxpy.math.interp1\_sprague5(x, y, xn, extrap=(nan, nan))
      Perform a 1-dimensional 5th order Sprague interpolation.
      Args:
                  X
                      ndarray with n-dimensional coordinates.
                  y
                      ndarray with values at coordinates in x.
                  xn
                      ndarray of new coordinates.
                  extrap
                      (np.nan, np.nan) or string, optional
                      If tuple: fill with values in tuple (\langle x[0], \rangle x[-1])
                      If string: ('zeros', 'linear', 'nearest', 'nearest-up', 'zero', 'slinear', 'quadratic', 'cubic',
                       'previous', 'next')
                             for more info on the other options see: scipy.interpolate.interp1d?
      Returns:
                  yn
                      ndarray with values at new coordinates in xn.
```

False, optional

```
luxpy.math.interp1(X, Y, Xnew, kind='linear', ext='extrapolate', w=None, bbox=[None, None],
                        check_finite=False)
      Perform a 1-dimensional linear interpolation (wrapper around scipy.interpolate.InterpolatedUnivariateSpline).
      Args:
                 \mathbf{X}
                      ndarray with n-dimensional coordinates (last axis represents dimension)
                 Y
                      ndarray with values at coordinates in X
                 Xnew
                      ndarray of new coordinates (last axis represents dimension)
                 kind
                      str or int, optional
                      if str: kind is 'translated' to an int value for input to
                      scipy.interpolate.InterpolatedUnivariateSpline()
                      supported options for str: 'linear', 'quadratic', 'cubic', 'quartic', 'quintic'
                 other args
                      see scipy.interpolate.InterpolatedUnivariateSpline()
      Returns:
                 Ynew
                      ndarray with new values at coordinates in Xnew
luxpy.math.ndinterp1(X, Y, Xnew)
      Perform nd-dimensional linear interpolation using Delaunay triangulation.
      Args:
                 X
                      ndarray with n-dimensional coordinates (last axis represents dimension).
                 Y
                      ndarray with values at coordinates in X.
                 Xnew
                      ndarray of new coordinates (last axis represents dimension).
                      When outside of the convex hull of X, then a best estimate is
                      given based on the closest vertices.
      Returns:
                 Ynew
                      ndarray with new values at coordinates in Xnew.
luxpy.math.ndinterp1_scipy(X, Y, Xnew, fill_value=nan, rescale=False)
      Perform a n-dimensional linear interpolation (wrapper around scipy.interpolate.LinearNDInterpolator).
      Args:
                 X
                      ndarray with n-dimensional coordinates (last axis represents dimension)
```

### Y

ndarray with values at coordinates in X

# **Xnew**

ndarray of new coordinates (last axis represents dimension)

#### fill\_value

float, optional

Value used to fill in for requested points outside of the convex hull of the input points. If not provided, then the default is nan.

#### rescale

bool, optional

Rescale points to unit cube before performing interpolation.

This is useful if some of the input dimensions have

incommensurable units and differ by many orders of magnitude.

#### **Returns:**

#### Ynew

ndarray with new values at coordinates in Xnew

luxpy.math.box\_m(\*X, ni=None, verbosity=0, robust=False, robust\_alpha=0.01)

Perform Box's M test (p>=2) to check equality of covariance matrices or Bartlett's test (p==1) for equality of variances.

### Args:

 $\mathbf{X}$ 

A number (k groups) or list of 2d-ndarrays (rows: samples, cols: variables) with data. or a number of 2d-ndarrays with covariance matrices (supply ni!)

ni

None, optional

If None: X contains data, else, X contains covariance matrices.

# verbosity

0, optional

If 1: print results.

#### robust

False, optional

If True: remove outliers beyond the confidence ellipsoid before calculating the covariances.

#### robust\_alpha

0.01, optional

Significance level of confidence ellipsoid marking the boundary for outliers.

# **Returns:**

#### statistic

F or chi2 value (see len(dfs))

```
pval
                     p-value
                 df
                     degrees of freedom.
                     if len(dfs) == 2: F-test was used.
                     if len(dfs) == 1: chi2 approx. was used.
     Notes:
              1. If p==1: Reduces to Bartlett's test for equal variances.
              2. If (ni>20).all() & (p<6) & (k<6): then a more appropriate chi2 test is used in a some cases.
luxpy.math.pitman_morgan(X, Y, verbosity=0)
     Pitman-Morgan Test for the difference between correlated variances with paired samples.
     Args:
                 X,Y
                     ndarrays with data.
                 verbosity
                     0, optional
                     If 1: print results.
     Returns:
                 tval
                     statistic
                 pval
                     p-value
                 df
                     degree of freedom.
                 ratio
                     variance ratio var1/var2 (with var1 > var2).
     Note:
              1. Based on Gardner, R.C. (2001). Psychological Statistics Using SPSS for Windows. New Jersey,
                 Prentice Hall.
              2. Python port from matlab code by Janne Kauttonen (https://nl.mathworks.com/matlabcentral/
                 fileexchange/67910-pitmanmorgantest-x-y; accessed Sep 26, 2019)
luxpy.math.stress(DE, DV, axis=0, max_scale=100)
     Calculate STandardize-Residual-Sum-of-Squares (STRESS).
     Args:
                 DE, DV
                     ndarrays of data to be compared.
                 axis
                     0, optional
```

axis with samples

### max\_scale

100, optional

Maximum of scale.

#### **Returns:**

#### stress

nadarray with stress value(s).

**Reference:** 1. Melgosa, M., García, P. A., Gómez-Robledo, L., Shamey, R., Hinks, D., Cui, G., & Luo, M. R. (2011). Notes on the application of the standardized residual sum of squares index for the assessment of intra- and inter-observer variability in color-difference experiments. Journal of the Optical Society of America A, 28(5), 949–953.

luxpy.math.stress\_F\_test(stressA, stressB, N, alpha=0.05)

Perform F-test on significance of difference between STRESS A and STRESS B.

## **Args:**

# stressA, stressB

ndarray with stress(es) values for A and B

N

int or ndarray with number of samples used to determine stress values.

# alpha

0.05, optional significance level

## **Returns:**

## **Fstats**

Dictionary with keys:

- 'p': p-values
- 'F': F-values
- 'Fc': critcal values
- 'H': string reporting on significance of A compared to B.

luxpy.math.mean\_distance\_weighted(x, axis=0, keepdims=False, center\_x=False, rtol=0.001, max\_iter=100, cnt=0, mu=None, mu0=0)

Recursively calculate distance weighted mean.

# Args:

X

ndarray with data

axis

dimension along which to take mean

# keepdims

False, optional

If True: keep dimension of original ndarray

 $center\_x$ 

True, optional Center data first.

#### rtol

1e-3, optional

Relative tolerance on recursive mean values. If two sequential mean values differ less than this amount, the recursion stops.

#### max iter

100, optional

Maximum amount of recursions. If this number is reached the recursion stops, even when rtol is not yet achieved. (to avoid getting stuck in an infinite loop when the recursion doesn't converge)

#### cnt,mu,mu0

Needed for passing values across recursions to be able to stop them. DO NOT CHANGE.

#### **Returns:**

## mu\_dw

distance weighted mean of the array

luxpy.math.minimizebnd(fun, x0, args=(), method='Nelder-Mead', use\_bnd=True, bounds=(None, None), options=None, x0\_vsize=None, x0\_keys=None, \*\*kwargs)

Minimization function that allows for bounds on any type of method in SciPy's minimize function by transforming the parameters values (see Matlab's fminsearchbnd).

Starting values, and lower and upper bounds can also be provided as a dict.

#### Args:

# $\mathbf{x0}$

parameter starting values

If x0\_keys is None then :x0: is vector else, :x0: is dict and

x0\_size should be provided with length/size of values for each of
the keys in :x0: to convert it to a vector.

# use bnd

True, optional

False: omits bounds and defaults to regular minimize function.

# bounds

(lower, upper), optional

Tuple of lists or dicts (x0\_keys is None) of lower and upper bounds for each of the parameters values.

## **kwargs**

allows input for other type of arguments (e.g. in OutputFcn)

**Note:** For other input arguments, see ?scipy.optimize.minimize()

## **Returns:**

res

dict with minimize() output.

Additionally, function value, fval, of solution is also in :res:, as well as a vector or dict (if x0 was dict)

with final solutions (res['x'])

# 4.2.2 vec3/

рy

- \_\_init\_\_.py
- vec3.py

namespace luxpy.math

# 4.2.3 **DEMO**/

рy

- \_\_init\_\_.py
- DEMO.py
- demo\_opt.py

namespace luxpy.math

# 4.3 Spectrum sub-package

рy

- \_\_init\_\_.py
- spdx\_ietm2714.py
- basics/
  - **–** \_\_init\_\_.py
  - cmf.py
  - spectral.py
  - spectral\_databases.py

namespace luxpy

# 4.3.1 spectrum: sub-package supporting basic spectral calculations

# spectrum/cmf.py

# luxpy.\_CMF

```
Dict with keys 'types' and x x are dicts with keys 'bar', 'K', 'M'
```

- \* luxpy.\_CMF[x]['bar'] = numpy array with CMFs for type x between 360 nm and 830 nm (has shape: (4,471))
- \*  $luxpy.\_CMF[x]['K'] = Constant converting Watt to lumen for CMF type x.$
- \* luxpy.\_CMF[x]['M'] = XYZ to LMS conversion matrix for CMF type x.

  Matrix is numpy array with shape: (3,3)
- \* luxpy.\_CMF[x]['N'] = XYZ to RGB conversion matrix for CMF type x. Matrix is numpy array with shape: (3,3)

#### Notes:

- 1. All functions have been expanded (when necessary) using zeros to a full 360-830 range. This way those wavelengths do not contribute in the calculation, AND are not extrapolated using the closest known value, as per CIE recommendation.
- 2. There is no XYZ to LMS conversion matrices defined for the 1931 2° Judd corrected (1951) cmf sets. The Hunt-Pointer-Estevez conversion matrix of the 1931 2° is therefore used as an approximation!
- 3. The XYZ to LMS conversion matrix M for the Judd-Vos XYZ CMFs is the one that converts to the 1979 Smith-Pokorny cone fundamentals.
- 4. The XYZ to LMS conversion matrix for the 1964 10° XYZ CMFs is set to the one of the CIE 2006 10° cone fundamentals, as not matrix has been officially defined for this CMF set.
- 4. The K lm to Watt conversion factors for the Judd and Judd-Vos cmf sets have been set to 683.002 lm/W (same as for standard 1931 2°).
- 5. The 1951 scoptopic V' function has been replicated in the 3 xbar, ybar, zbar columns to obtain a data format similar to the photopic color matching functions. This way V' can be called in exactly the same way as other V functions can be called from the X,Y,Z cmf sets. The K value has been set to 1700.06 lm/W and the conversion matrix has been filled with NaN's.
- 6. The '2015\_x' (with x = 2 or 10) are the same XYZ-CMFs as stored in '2006\_x'.
- 7. \_CMF[x]['M'] for x equal to '2006\_2' (='2015\_2') or '2006\_10' (='2015\_10') is NOT normalized to illuminant E! These are the original matrices as defined by [1] & [2].

8. \_CMF[x]['N'] stores known or calculated conversion matrices from xyz to rgb. If not available, N has been filled with NaNs.

# spectrum/spectral.py

- \_WL3 Default wavelength specification in vector-3 format: numpy.array([start, end, spacing])
- **\_INTERP\_TYPES** Dict with interpolation types associated with various types of spectral data according to CIE recommendation:
- \_S\_INTERP\_TYPE Interpolation type for light source spectral data
- \_R\_INTERP\_TYPE Interpolation type for reflective/transmissive spectral data
- \_C\_INTERP\_TYPE Interpolation type for CMF and cone-fundamental spectral data
- **getwlr()** Get/construct a wavelength range from a (start, stop, spacing) 3-vector.
- getwld() Get wavelength spacing of numpy.ndarray with wavelengths.
- **spd\_normalize**() Spectrum normalization (supports: area, max, lambda, radiometric, photometric and quantal energy units).
- cie\_interp() Interpolate / extrapolate spectral data following standard [CIE15:2018, "Colorimetry," CIE, Vienna, Austria, 2018.]

## spd()

All-in-one function that can:

- 1. Read spectral data from data file or take input directly as pandas.dataframe or ndarray.
- 2. Convert spd-like data from ndarray to pandas.dataframe and back.
- 3. Interpolate spectral data.
- 4. Normalize spectral data.

xyzbar() Get color matching functions.

vlbar() Get Vlambda function.

get cie mesopic adaptation() Get the mesopic adaptation state according to CIE191:2010

spd\_to\_xyz() Calculates xyz tristimulus values from spectral data.

spd\_to\_ler() Calculates Luminous efficacy of radiation (LER) from spectral data.

spd\_to\_power() Calculate power of spectral data in radiometric, photometric or quantal energy units.

**detect\_peakwl()** Detect peak wavelengths and fwhm of peaks in spectrum spd.

## spectrum/spectral databases.py

- \_S\_PATH Path to light source spectra data.
- \_R\_PATH Path to with spectral reflectance data
- \_IESTM3015 Database with spectral reflectances related to and light source spectra contained excel calculator of IES TM30-15 publication.
- **\_IESTM3018** Database with spectral reflectances related to and light source spectra contained excel calculator of IES TM30-18 publication.
- \_IESTM3015\_S Database with only light source spectra contained in the IES TM30-15 excel calculator.
- \_IESTM3018\_S Database with only light source spectra contained in the IES TM30-18 excel calculator.

#### CIE ILLUMINANTS

Database with CIE illuminants:

- \* 'E', 'D65', 'A', 'C',
- \* 'F1', 'F2', 'F3', 'F4', 'F5', 'F6', 'F7', 'F8', 'F9', 'F10', 'F11', 'F12'
- \_CIE\_E, \_CIE\_D65, \_CIE\_A, \_CIE\_C, \_CIE\_F4 Some CIE illuminants for easy use.

# \_CRI\_RFL

Database with spectral reflectance functions for various color rendition calculators:

- \* CIE 13.3-1995 (8, 14 munsell samples)
- \* CIE 224:2015 (99 set)
- \* CRI2012 (HL17 & HL1000 spectrally uniform and 210 real samples)
- \* IES TM30 (99, 4880 sepctrally uniform samples)
- \* MCRI (10 familiar object set)
- \* CQS (v7.5 and v9.0 sets)
- \_MUNSELL Database (dict) with 1269 Munsell spectral reflectance functions and Value (V), Chroma (C), hue (h) and (ab) specifications.

# \_RFL

Database (dict) with RFLs, including:

- \* all those in CRI RFL,
- \* the 1269 Matt Munsell samples (see also \_MUNSELL),
- \* the 24 Macbeth ColorChecker samples,
- $\mbox{*}$  the 215 samples proposed by Opstelten, J.J. , 1983, The establishment of a representative set of test colours

for the specification of the colour rendering properties of light sources, CIE-20th session, Amsterdam.

\* the 114120 RFLs from capbone.com/spectral-reflectance-database/

# spectrum/illuminants.py

- **\_BB** Dict with constants for blackbody radiator calculation constant are (c1, c2, n, na, c, h, k).
- \_S012\_DAYLIGHTPHASE ndarray with CIE S0,S1, S2 curves for daylight phase calculation (linearly interpolated to 1 nm).
- **\_CRI\_REF\_TYPES** Dict with blackbody to daylight transition (mixing) ranges for various types of reference illuminants used in color rendering index calculations.
- **blackbody**() Calculate blackbody radiator spectrum.
- **\_DAYLIGHT\_LOCI\_PARAMETERS** dict with parameters for daylight loci for various CMF sets; used by daylightlocus().
- \_DAYLIGHT\_M12\_COEFFS dict with coefficients in weights M1 & M2 for daylight phases for various CMF sets.
- **get\_daylightloci\_parameters**() Get parameters for the daylight loci functions xD(1000/CCT) and yD(xD); used by daylightlocus().
- **get\_daylightphase\_Mi\_coeffs()** Get coefficients of Mi weights of daylight phase for specific cieobs following Judd et al. (1964).
- **\_get\_daylightphase\_Mi\_values()** Get daylight phase coefficients M1, M2 following Judd et al. (1964).
- **\_get\_daylightphase\_Mi()** Get daylight phase coefficients M1, M2 following Judd et al. (1964)
- daylightlocus() Calculates daylight chromaticity from cct.
- daylightphase() Calculate daylight phase spectrum.
- cri\_ref()
  - Calculates a reference illuminant spectrum based on cct for color rendering index calculations.
  - (CIE15:2018, "Colorimetry," CIE, Vienna, Austria, 2018., cie224:2017, CIE 2017 Colour Fidelity Index for accurate scientific use. (2017), ISBN 978-3-902842-61-9., IES-TM-30-15: Method for Evaluating Light Source Color Rendition. New York, NY: The Illuminating Engineering Society of North America.
- spd\_to\_indoor() Convert spd to indoor variant by multiplying it with the CIE spectral transmission for glass.

# spectrum/spdx\_iestm2714.py

- **\_SPDX\_TEMPLATE** template dictionary for SPDX data.
- **read\_spdx()** Read xml file or convert xml string with spdx data to dictionary.
- write\_spdx() Convert spdx dictionary to xml string (and write to .spdx file)

#### References

- 1. CIE15:2018, "Colorimetry," CIE, Vienna, Austria, 2018.
- 2. CIE, and CIE (2006). Fundamental Chromaticity Diagram with Physiological Axes Part I.(Vienna: CIE).
- 3. cie224:2017, CIE 2017 Colour Fidelity Index for accurate scientific use. (2017), ISBN 978-3-902842-61-9.
- **4.** IES-TM-30-15: Method for Evaluating Light Source Color Rendition. New York, NY: The Illuminating Engineering Society of North America.
  - Judd, D. B., MacAdam, D. L., Wyszecki, G., Budde, H. W., Condit, H. R., Henderson, S. T., & Simonds, J. L. (1964). Spectral Distribution of Typical Daylight as a Function of Correlated Color Temperature. J. Opt. Soc. Am., 54(8), 1031–1040. https://doi.org/10.1364/JOSA.54.001031
  - 6. http://www.ies.org/iestm2714

```
luxpy.spectrum.getwlr(wl3=None)
```

Get/construct a wavelength range from a 3-vector (start, stop, spacing).

#### Args:

#### wl3

list[start, stop, spacing], optional (defaults to luxpy.\_WL3)

#### **Returns:**

#### returns

ndarray (.shape = (n,)) with n wavelengths ranging from start to stop, with wavelength interval equal to spacing.

## luxpy.spectrum.getwld(wl)

Get wavelength spacing.

# **Args:**

wl

ndarray with wavelengths

#### **Returns:**

## returns

- float: for equal wavelength spacings
- ndarray (.shape = (n, )): for unequal wavelength spacings

luxpy.spectrum.spd\_normalize(data, norm\_type=None, norm\_f=1, wl=True, cieobs='1931\_2') Normalize a spectral power distribution (SPD).

# **Args:**

#### data

ndarray

## norm\_type

None, optional

```
- 'lambda': make lambda in norm_f equal to 1
                            - 'area': area-normalization times norm_f
                            - 'max': max-normalization times norm f
                            - 'ru': to :norm_f: radiometric units
                            - 'pu': to :norm_f: photometric units
                            - 'pusa': to :norm_f: photometric units (with Km corrected
                                  to standard air, cfr. CIE TN003-2015)
                            - 'qu': to :norm_f: quantal energy units
                 norm_f
                      1, optional
                      Normalization factor that determines the size of normalization
                      for 'max' and 'area'
                      or which wavelength is normalized to 1 for 'lambda' option.
                 wl
                      True or False, optional
                      If True, the first column of data contains wavelengths.
                 cieobs
                      _CIEOBS or str, optional
                      Type of cmf set to use for normalization using photometric units
                      (norm_type == 'pu')
      Returns:
                 returns
                      ndarray with normalized data.
luxpy.spectrum.cie_interp(data, wl_new, kind=None, sprague5_allowed=False,
                                 negative_values_allowed=False, extrap_values='ext', extrap_kind='linear',
                                 extrap_log=False)
      Interpolate / extrapolate spectral data following standard CIE15-2018.
      The kind of interpolation depends on the spectrum type defined in :kind:.
                 data
                      ndarray with spectral data
                      (.shape = (number of spectra + 1, number of original wavelengths))
                 wl_new
                      ndarray with new wavelengths
                 kind
                      None, optional
                            - If :kind: is None, return original data.
                            - If :kind: is a spectrum type (see _INTERP_TYPES), the correct
```

interpolation type is automatically chosen

**Args:** 

```
(The use of the slow(er) 'sprague5' can be toggled on using
                       :sprague5_allowed:).
           - Or :kind: can be any interpolation type supported by
                 scipy.interpolate.interp1d (or luxpy.math.interp1 if nan's are present!!)
                 or can be 'sprague5' (uses luxpy.math.interp1_sprague5).
sprague5 allowed
    False, optional
    If True: When kind is a spectral data type from _INTERP_TYPES['cubic'],
           then a cubic spline interpolation will be used in case of
           unequal wavelength spacings, otherwise a 5th order Sprague will be used.
    If False: always use 'cubic', don't use 'sprague5'.
           This is the default, as differences are minimal and
           use of the 'sprague5' function is a lot slower!
negative_values_allowed
    False, optional
    If False: negative values are clipped to zero.
extrap_values
     'ext', optional
    If 'ext': extrapolate using 'linear' ('cie167:2005'), 'quadratic' ('cie15:2018')
           'nearest' ('cie15:2004') recommended or other (e.g. 'cubic') methods.
    If None: use CIE15:2004 recommended 'nearest value' approach when extrapolating.
    If float or list or ndarray, use those values to fill extrapolated value(s).
extrap kind
     'linear', optional
    Extrapolation method used when :extrap_values: is set to 'ext'.
    Options: 'linear' ('cie167:2005'), 'quadratic' ('cie15:2018'),
           'nearest' ('cie15:2004'), 'cubic'
    CIE15:2018 states that based on a 2017 paper by Wang that 'quadratic' is 'better'.
    However, no significant difference was found between 'quadratic' and 'linear' methods.
    Also see note 1 below, for why the CIE67:2005 recommended 'linear' extrapolation
    is set as the default.
extrap_log
    False, optional
    If True: extrap the log of the spectral values
           (not CIE recommended but in most cases seems to give a
           more realistic estimate, but can sometimes seriously fail,
           especially for the 'quadratic' extrapolation case (see note 1)!!!)
      ndarray of interpolated spectral data.
      (.shape = (number of spectra + 1, number of wavelength in wl_new))
```

**Notes:** 

returns

1. Type of extrapolation: 'quadratic' vs 'linear'; impact of extrapolating log spectral values:

Using a 'linear' or 'quadratic' extrapolation, as mentioned in

CIE167:2005 and CIE15:2018, resp., can lead to extreme large values

when setting :extrap\_log: (not CIE recommended) to True.

A quick test with the IES TM30 spectra (400 nm - 700 nm, 5 nm spacing)

shows that 'linear' is better than 'quadratic' in terms of

mean, median and max DEu'v' with the original spectra (380 nm - 780 nm, 5 nm spacing).

This conferms the recommendation from CIE167:2005 to use 'linear' extrapolation.

Setting :extrap log: to True reduces the median, but inflates the mean due to some

extremely large DEu'v' values. However, the increase in mean and max DEu'v' is much

larger for the 'quadratic' case, suggesting that 'linear' extrapolation

is likely a more suitable recommendation. When using a 1 nm spacing

'linear' is more similar to 'quadratic' when :extrap\_log: is False, otherwise 'linear'

remains the 'best'. Hence the choice to use the CIE167:2005 recommended linear extrapolation as default!

luxpy.spectrum.spd(data=None, interpolation=None, kind='np', wl=None, columns=None, sep=',', header=None, datatype='S', norm\_type=None, norm\_f=None)

All-in-one function that can:

- 1. Read spectral data from data file or take input directly as pandas.dataframe or ndarray.
- 2. Convert spd-like data from ndarray to pandas.dataframe and back.
- 3. Interpolate spectral data.
- 4. Normalize spectral data.

## Args:

#### data

- str with path to file containing spectral data
- ndarray with spectral data
- pandas.dataframe with spectral data

(.shape = (number of spectra + 1, number of original wavelengths))

# interpolation

None, optional

- None: don't interpolate
- str with interpolation type or spectrum type

# kind

str ['np','df'], optional

Determines type(:returns:), np: ndarray, df: pandas.dataframe

wl

None, optional

New wavelength range for interpolation.

Defaults to wavelengths specified by luxpy.\_WL3.

#### columns

- None or list[str] of column names for dataframe, optional

#### header

None or 'infer', optional

- None: no header in file

- 'infer': infer headers from file

## sep

',' or ' ' or other char, optional

Column separator in case :data: specifies a data file.

# datatype'

'S' (light source) or 'R' (reflectance) or other, optional

Specifies a type of spectral data.

Is used when creating column headers when :column: is None.

## norm\_type

None, optional

- 'lambda': make lambda in norm\_f equal to 1
- 'area': area-normalization times norm f
- 'max': max-normalization times norm\_f
- 'ru': to :norm\_f: radiometric units
- 'pu': to :norm\_f: photometric units
- 'pusa': to :norm\_f: photometric units (with Km corrected to standard air, cfr. CIE TN003-2015)
- 'qu': to :norm\_f: quantal energy units

# norm\_f

1, optional

Normalization factor that determines the size of normalization for 'max' and 'area' or which wavelength is normalized to 1 for 'lambda' option.

## **Returns:**

## returns

ndarray or pandas.dataframe with interpolated and/or normalized spectral data.

luxpy.spectrum.xyzbar(cieobs='1931\_2', scr='dict', wl\_new=None, kind='np') Get color matching functions.

**Args:** 

# cieobs

luxpy.\_CIEOBS, optional

Sets the type of color matching functions to load.

scr

'dict' or 'file', optional

Determines whether to load cmfs from file (./data/cmfs/)

or from dict defined in .cmf.py

wl

None, optional

New wavelength range for interpolation.

```
Defaults to wavelengths specified by luxpy._WL3.
                 kind
                       str ['np','df'], optional
                       Determines type(:returns:), np: ndarray, df: pandas.dataframe
     Returns:
                  returns
                       ndarray or pandas.dataframe with CMFs
     References:
               1. CIE15:2018, "Colorimetry," CIE, Vienna, Austria, 2018.
luxpy.spectrum.vlbar(cieobs='1931_2', scr='dict', wl_new=None, kind='np', out=1)
     Get Vlambda functions.
     Args:
                  cieobs
                       str, optional
                       Sets the type of Vlambda function to obtain.
                  scr
                       'dict' or array, optional
                       - 'dict': get from ybar from _CMF
                       - 'array': ndarray in :cieobs:
                       Determines whether to load cmfs from file (./data/cmfs/)
                       or from dict defined in .cmf.py
                       Vlambda is obtained by collecting Ybar.
                  wl
                       None, optional
                       New wavelength range for interpolation.
                       Defaults to wavelengths specified by luxpy._WL3.
                  kind
                       str ['np','df'], optional
                       Determines type(:returns:), np: ndarray, df: pandas.dataframe
                  out
                       1 or 2, optional
                             1: returns Vlambda
                             2: returns (Vlambda, Km)
     Returns:
                  returns
                       dataframe or ndarray with Vlambda of type :cieobs:
     References:
               1. CIE15:2018, "Colorimetry," CIE, Vienna, Austria, 2018.
luxpy.spectrum.vlbar_cie_mesopic(m=[1], wl_new=None, kind='np', out=1, Lp=None, Ls=None, SP=None)
     Get CIE mesopic luminous efficiency function Vmesm according to CIE191:2010
     Args:
                  m
```

```
float or list or ndarray with mesopic adaptation coefficients
                  wl
                       None, optional
                       New wavelength range for interpolation.
                       Defaults to wavelengths specified by luxpy._WL3.
                  out
                       1 or 2, optional
                             1: returns Vmesm
                             2: returns (Vmes, Kmesm)
                 Lp
                       None, optional
                       float or ndarray with photopic adaptation luminance
                       If not None: use this (and SP or Ls) to calculate the
                       mesopic adaptation coefficient
                 Ls
                       None, optional
                       float or ndarray with scotopic adaptation luminance
                       If None: SP must be supplied.
                 SP
                       None, optional
                       S/P ratio
                       If None: Ls must be supplied.
                  Vmes
                       ndarray with mesopic luminous efficiency function
                       for adaptation coefficient(s) m
                  Kmes
                       ndarray with luminous efficacies of 555 nm monochromatic light
                       for for adaptation coefficient(s) m
     Reference: 1. CIE 191:2010 Recommended System for Mesopic Photometry Based on Visual Performance.
           (ISBN 978-3-901906-88-6),
luxpy.spectrum.get_cie_mesopic_adaptation(Lp, Ls=None, SP=None)
     Get the mesopic adaptation state according to CIE191:2010
                 Lp
                       float or ndarray with photopic adaptation luminance
                 Ls
                       None, optional
                       float or ndarray with scotopic adaptation luminance
                       If None: SP must be supplied.
```

Args:

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```
None, optional
                       S/P ratio
                       If None: Ls must be supplied.
      Returns:
                  Lmes
                       mesopic adaptation luminance
                  m
                       mesopic adaptation coefficient
      Reference: 1. CIE 191:2010 Recommended System for Mesopic Photometry Based on Visual Performance.
            (ISBN 978-3-901906-88-6),
luxpy.spectrum.spd_to_xyz(data, relative=True, rfl=None, cieobs='1931_2', K=None, out=None,
                                cie_std_dev_obs=None)
      Calculates xyz tristimulus values from spectral data.
      Args:
                  data
                       ndarray or pandas.dataframe with spectral data
                       (.shape = (number of spectra + 1, number of wavelengths))
                       Note that :data: is never interpolated, only CMFs and RFLs.
                       This way interpolation errors due to peaky spectra are avoided.
                       Conform CIE15-2018.
                  relative
                       True or False, optional
                       Calculate relative XYZ (Yw = 100) or absolute XYZ (Y = Luminance)
                  rfl
                       ndarray with spectral reflectance functions.
                       Will be interpolated if wavelengths do not match those of :data:
                  cieobs
                       luxpy._CIEOBS or str, optional
                       Determines the color matching functions to be used in the
                       calculation of XYZ.
                  K
                       None, optional
                             e.g. K = 683 lm/W for '1931_2' (relative == False)
                             or K = 100/\text{sum}(\text{spd*dl}) (relative == True)
                  out
                       None or 1 or 2, optional
                       Determines number and shape of output. (see :returns:)
                  cie_std_dev_obs
                       None or str, optional
                       - None: don't use CIE Standard Deviate Observer function.
```

- 'f1': use F1 function.

returns

```
If rfl is None:
                              If out is None: ndarray of xyz values
                                    (.shape = (data.shape[0],3))
                             If out == 1: ndarray of xyz values
                                    (.shape = (data.shape[0],3))
                             If out == 2: (ndarray of xyz, ndarray of xyzw) values
                                    Note that xyz == xyzw, with (.shape = (data.shape[0],3))
                        If rfl is not None:
                             If out is None: ndarray of xyz values
                                    (.shape = (rfl.shape[0], data.shape[0], 3))
                             If out == 1: ndarray of xyz values
                                          (.shape = (rfl.shape[0]+1,data.shape[0],3))
                                                The xyzw values of the light source spd are the first set
                                                of values of the first dimension. The following values
                                          along this dimension are the sample (rfl) xyz values.
                                    If out == 2: (ndarray of xyz, ndarray of xyzw) values
                                          with xyz.shape = (rfl.shape[0],data.shape[0],3)
                                          and with xyzw.shape = (data.shape[0],3)
      References:
               1. CIE15:2018, "Colorimetry," CIE, Vienna, Austria, 2018.
luxpy.spectrum.spd_to_ler(data, cieobs='1931_2', K=None)
      Calculates Luminous efficacy of radiation (LER) from spectral data.
      Args:
                  data
                        ndarray or pandas.dataframe with spectral data
                        (.shape = (number of spectra + 1, number of wavelengths))
                        Note that :data: is never interpolated, only CMFs and RFLs.
                        This way interpolation errors due to peaky spectra are avoided.
                        Conform CIE15-2018.
                  cieobs
                        luxpy._CIEOBS, optional
                        Determines the color matching function set used in the
                        calculation of LER. For cieobs = '1931_2' the ybar CMF curve equals
                        the CIE 1924 Vlambda curve.
                  K
                        None, optional
                              e.g. K = 683 \text{ lm/W for '}1931_2'
      Returns:
                  ler
                        ndarray of LER values.
      References:
               1. CIE15:2018, "Colorimetry," CIE, Vienna, Austria, 2018.
luxpy.spectrum.spd_to_power(data, ptype='ru', cieobs='1931 2')
      Calculate power of spectral data in radiometric, photometric or quantal energy units.
      Args:
```

```
data
                        ndarray with spectral data
                  ptype
                        'ru' or str, optional
                        str: - 'ru': in radiometric units
                              - 'pu': in photometric units
                              - 'pusa': in photometric units with Km corrected
                                    to standard air (cfr. CIE TN003-2015)
                              - 'qu': in quantal energy units
                  cieobs
                        _CIEOBS or str, optional
                        Type of cmf set to use for photometric units.
      Returns:
            returns:
                  ndarray with normalized spectral data (SI units)
luxpy.spectrum.detect_peakwl(spd, n=1, verbosity=1, **kwargs)
      Detect primary peak wavelengths and fwhm in spectrum spd.
      Args:
                  spd
                        ndarray with spectral data (2xN).
                        First row should be wavelengths.
                  n
                        1, optional
                        The number of peaks to try to detect in spd.
                  verbosity
                        Make a plot of the detected peaks, their fwhm, etc.
                  kwargs
                        Additional input arguments for scipy.signal.find_peaks.
      Returns:
                  prop
                        list of dictionaries with keys:
                        - 'peaks_idx' : index of detected peaks
                        - 'peaks': peak wavelength values (nm)
                        - 'heights': height of peaks
                        - 'fwhms' : full-width-half-maxima of peaks
                        - 'fwhms_mid' : wavelength at the middle of the fwhm-range of the peaks (if this is
                        different from the values in 'peaks', then their is some non-symmetry in the peaks)
                        - 'fwhms mid heights': height at the middle of the peak
luxpy.spectrum.cri_ref(ccts, wl3=None, ref_type='ciera', mix_range=None, cieobs=None, norm_type=None,
                             norm_f=None, force_daylight_below4000K=False, n=None, daylight_locus=None)
      Calculates a reference illuminant spectrum based on cct for color rendering index calculations .
      Args:
                  ccts
```

list of int/floats or ndarray with ccts.

#### wl3

None, optional

New wavelength range for interpolation.

Defaults to wavelengths specified by luxpy.\_WL3.

## ref\_type

str or list[str], optional

Specifies the type of reference spectrum to be calculated.

Defaults to luxpy. CRI REF TYPE.

If :ref\_type: is list of strings, then for each cct in :ccts:

a different reference illuminant can be specified.

If :ref\_type: == 'spd', then :ccts: is assumed to be an ndarray

of reference illuminant spectra.

#### mix\_range

None or ndarray, optional

Determines the cct range between which the reference illuminant is

a weigthed mean of a Planckian and Daylight Phase spectrum.

Weighthing is done as described in IES TM30:

SPDreference = (Te-T)/(Te-Tb)\*Planckian+(T-Tb)/(Te-Tb)\*daylight with Tb and Te are resp. the starting and end CCTs of the mixing range and whereby the Planckian and Daylight SPDs have been normalized for equal luminous flux.

If None: use the default specified for :ref type:.

Can be a ndarray with shape[0] > 1, in which different mixing ranges will be used for cct in :ccts:.

## cieobs

None, optional

Required for the normalization of the Planckian and Daylight SPDs when calculating a 'mixed' reference illuminant.

Required when calculating daylightphase (adjust locus parameters to cieobs)

If None: CIEOBS will be used.

# norm\_type

None, optional

- 'lambda': make lambda in norm\_f equal to 1
- 'area': area-normalization times norm\_f
- 'max': max-normalization times norm f
- 'ru': to :norm\_f: radiometric units
- 'pu': to :norm\_f: photometric units
- 'pusa': to :norm\_f: photometric units (with Km corrected to standard air, cfr. CIE TN003-2015)
- 'qu': to :norm\_f: quantal energy units

## norm f

1, optional

```
Normalization factor that determines the size of normalization
                         for 'max' and 'area'
                         or which wavelength is normalized to 1 for 'lambda' option.
                  force daylight below4000K
                         False or True, optional
                         Daylight locus approximation is not defined below 4000 K,
                         but by setting this to True, the calculation can be forced to
                         calculate it anyway.
                  n
                         None, optional
                         Refractive index (for use in calculation of blackbody radiators).
                         If None: use the one stored in _BB['n']
                  daylight_locus
                         None, optional
                         dict with xD(T) and yD(xD) parameters to calculate daylight locus
                         for specified cieobs.
                         If None: use pre-calculated values.
                         If 'calc': calculate them on the fly.
                  returns
                         ndarray with reference illuminant spectra.
                         (:returns:[0] contains wavelengths)
      Note: Future versions will have the ability to take a dict as input for ref_type. This way other reference illumi-
            nants can be specified than the ones in _CRI_REF_TYPES.
luxpy.spectrum.blackbody(cct, wl3=None, n=None, relative=True)
      Calculate blackbody radiator spectrum for correlated color temperature (cct).
                  cct
                         int or float
                         (for list of cct values, use cri_ref() with ref_type = 'BB')
                  wl3
                         None, optional
                         New wavelength range for interpolation.
                         Defaults to wavelengths specified by luxpy._WL3.
                  n
                         None, optional
                         Refractive index.
                         If None: use the one stored in _BB['n']
                  relative
                         False, optional
                         True: return relative spectrum normalized to 560 nm
                         False: return absolute spectral radiance (Planck's law; W/(sr.m<sup>2</sup>.nm))
                  returns
```

**Returns:** 

Args:

ndarray with blackbody radiator spectrum

```
(:returns:[0] contains wavelengths)
      References:
               1. CIE15:2018, "Colorimetry," CIE, Vienna, Austria, 2018.
luxpy.spectrum.spd_to_indoor(spd)
      Convert spd to indoor variant by multiplying it with the CIE spectral transmission for glass.
luxpy.spectrum.daylightlocus(cct, force_daylight_below4000K=False, cieobs=None, daylight_locus=None)
      Calculates daylight chromaticity (xD,yD) from correlated color temperature (cct).
      Args:
                  cct
                        int or float or list of int/floats or ndarray
                  force_daylight_below4000K
                        False or True, optional
                        Daylight locus approximation is not defined below 4000 K,
                        but by setting this to True, the calculation can be forced to
                        calculate it anyway.
                  cieobs
                        CMF set corresponding to xD, yD output.
                        If None: use default CIE15-20xx locus for '1931_2'
                        Else: use the locus specified in :daylight_locus:
                  daylight_locus
                        None, optional
                        dict with xD(T) and yD(xD) parameters to calculate daylight locus
                        for specified cieobs.
                        If None: use pre-calculated values.
                        If 'calc': calculate them on the fly.
      Returns:
                  (xD, yD)
                        (ndarray of x-coordinates, ndarray of y-coordinates)
      References:
               1. CIE15:2018, "Colorimetry," CIE, Vienna, Austria, 2018.
luxpy.spectrum.daylightphase(cct, wl3=None, nominal_cct=False, force_daylight_below4000K=False,
                                    verbosity=None, n=None, cieobs=None, daylight_locus=None,
                                    daylight Mi coeffs=None)
      Calculate daylight phase spectrum for correlated color temperature (cct).
      Args:
                  cct
                        int or float
                        (for list of cct values, use cri_ref() with ref_type = 'DL')
                  wl3
                        None, optional
                        New wavelength range for interpolation.
                        Defaults to wavelengths specified by luxpy._WL3.
                  nominal_cct
```

False, optional

If cct is nominal (e.g. when calculating D65): multiply cct first

by 1.4388/1.4380 to account for change in 'c2' in definition of Planckian.

#### cieobs

None or str or ndarray, optional

CMF set to use when calculating coefficients for daylight locus and for M1, M2 weights.

If None: use standard coefficients for CIE 1931 2° CMFs (for Si at 10 nm).

Else: calculate coefficients following Appendix C of CIE15-2004 and Judd (1964).

## force\_daylight\_below4000K

False or True, optional

Daylight locus approximation is not defined below 4000 K,

but by setting this to True, the calculation can be forced to

calculate it anyway.

# verbosity

None, optional

If None: do not print warning when CCT < 4000 K.

n

None, optional

Refractive index (for use in calculation of blackbody radiators).

If None: use the one stored in \_BB['n']

## daylight locus

None, optional

dict with xD(T) and yD(xD) parameters to calculate daylight locus

for specified cieobs.

If None: use pre-calculated values.

If 'calc': calculate them on the fly.

# daylight\_Mi\_coeffs

None, optional

dict with coefficients for M1 & M2 weights for specified cieobs.

If None: use pre-calculated values.

If 'calc': calculate them on the fly.

## **Returns:**

# returns

ndarray with daylight phase spectrum (:returns:[0] contains wavelengths)

## **References:**

- 1. CIE15:2018, "Colorimetry," CIE, Vienna, Austria, 2018.
- 2. Judd, MacAdam, Wyszecki, Budde, Condit, Henderson, & Simonds (1964). Spectral Distribution of Typical Daylight as a Function of Correlated Color Temperature. J. Opt. Soc. Am., 54(8), 1031–1040.

luxpy.spectrum.get\_daylightloci\_parameters(ccts=None, cieobs=None, wl3=[300, 830, 10], verbosity=0)
Get parameters for the daylight loci functions xD(1000/CCT) and yD(xD).
Args:

ccts

**Args:** 

**Returns:** 

Args:

```
None, optional
                        ndarray with CCTs, if None: ccts = np.arange(4000,25000,250)
                  cieobs
                        None or list of str or list of ndarrays, optional
                        CMF sets to determine parameters for.
                        If None: get for all CMFs sets in CMF (except scoptopic and deviate observer)
                  wl3
                        [300,830,10], optional
                        Wavelength range and spacing of daylight phases to be determined
                        from '1931_2'. The default setting results in parameters very close
                        to that in CIE15-2004/2018.
                  verbosity
                        print parameters and make plots.
                  dayloci
                        dict with parameters for each cieobs
                        If cieobs contains ndarrays, then keys in dict will be
                        labeled 'cmf_0', 'cmf_1', ...
luxpy.spectrum.get_daylightphase_Mi_coeffs(cieobs=None, wl3=None, S012_daylightphase=None)
      Get coefficients of Mi weights of daylight phase for specific cieobs
                  cieobs
                        None or str or ndarray or list of str or list of ndarrays, optional
                        CMF set to get coefficients for.
                        If None: get coeffs for all CMFs in _CMF
                  wl3
                        None, optional
                        Wavelength range to interpolate S012_daylightphase to.
                  S012_daylightphase
                        None, optional
                        Daylight phase component functions.
                        If None: use _S012_DAYLIGHTPHASE
                  Mcoeffs
                        Dictionary with i,j,k,i1,j1,k1,i2,j2,k2 for each cieobs in :cieobs:
                        If cieobs contains ndarrays, then keys in dict will be
                        labeled 'cmf 0', 'cmf 1', ...
luxpy.spectrum.read_spdx(spdx)
      Read xml file or convert xml string with spdx data to dictionary.
                  spdx
                        xml string or file with spdx data.
```

```
Returns:
                 spdx_dict
                       spdx data in a dictionary.
luxpy.spectrum.write_spdx(spdx_dict, filename=None)
     Convert spdx dictionary to xml string (and write to .spdx file).
     Args:
                 spdx_dict
                       dictionary with spdx keys (see _SPDX for keys).
                  filename
                       None, optional
                       string with filename to write xml data to.
     Returns:
                 spdx_xml
                       string with xml data in spdx dictionary.
4.3.2 SPD class
           рy
                     · SPD.py
           namespace luxpy
class luxpy.spectrum.SPD.SPD(spd=None, wl=None, ax0iswl=True, dtype='S', wl_new=None,
                                    interp_method='auto', negative_values_allowed=False, extrap_values='ext',
                                    norm type=None, norm f=1, header=None, sep=',')
     read_csv_(file, header=None, sep=',')
           Reads spectral data from file.
            Args:
                       file
                             filename
                       header
                             None or 'infer', optional
                             If 'infer': headers will be inferred from file itself.
                             If None: no headers are expected from file.
                       sep
                             ',', optional
                             Column separator.
            Returns:
                       returns
                             ndarray with spectral data (first row are wavelengths)
           Note: Spectral data in file should be organized in columns with the first column containing the wave-
                 lengths.
     plot(ylabel='Spectrum', wavelength_bar=True, *args, **kwargs)
            Make a plot of the spectral data in SPD instance.
```

# returns handle to current axes. mean() Take mean of all spectra in SPD instance. sum() Sum all spectra in SPD instance. dot(S)Take dot product with instance of SPD. add(S)Add instance of SPD. sub(S)Subtract instance of SPD. $\mathbf{mul}(S)$ Multiply by instance of SPD. div(S)Divide by instance of SPD. pow(n)Raise SPD instance to power n. get\_() Get spd as ndarray in instance of SPD. setwlv(spd) Store spd ndarray in fields wl and values of instance of SPD. getwld\_() Get wavelength spacing of SPD instance. **Returns:** returns float: for equal wavelength spacings ndarray (.shape = (n,)): for unequal wavelength spacings **normalize**(norm\_type=None, norm\_f=1, cieobs='1931\_2') Normalize spectral power distributions in SPD instance. Args: norm\_type None, optional - 'lambda': make lambda in norm\_f equal to 1 - 'area': area-normalization times norm\_f - 'max': max-normalization times norm\_f - 'ru': to :norm\_f: radiometric units - 'pu': to :norm f: photometric units - 'pusa': to :norm\_f: photometric units (with Km corrected to standard air, cfr. CIE TN003-2015) - 'qu': to :norm\_f: quantal energy units norm\_f

1, optional

Determines size of normalization for 'max' and 'area' or which wavelength is normalized to 1 for 'lambda' option.

#### cieobs

CIEOBS or str, optional

Type of cmf set to use for normalization using photometric units (norm\_type == 'pu')

The interpolation type depends on the spectrum type defined in :kind:.

## **Args:**

#### wl new

ndarray with new wavelengths

#### kind

'auto', optional

If :kind: is None, return original data.

If :kind: is a spectrum type (see INTERP TYPES), the correct

interpolation type if automatically chosen.

(The use of the slow(er) 'sprague5' can be toggled on using :sprague5\_allowed:).

If kind = 'auto': use self.dtype

Or :kind: can be any interpolation type supported by

 $scipy.interpolate.interp1d\ (luxpy.math.interp1\ if\ nan's\ are\ present!!)$ 

or can be 'sprague5' (uses luxpy.math.interp1\_sprague5).

# sprague5\_allowed

False, optional

If True: When kind is a spectral data type from \_INTERP\_TYPES['cubic'],

then a cubic spline interpolation will be used in case of

unequal wavelength spacings, otherwise a 5th order Sprague will be used.

If False: always use 'cubic', don't use 'sprague5'.

This is the default, as differences are minimal and use of the 'sprague5' function is a lot slower!

# negative\_values\_allowed

False, optional

If False: negative values are clipped to zero

# extrap\_values

'ext', optional

If 'ext': extrapolate using 'linear' ('cie167:2005' r), 'quadratic' ('cie15:2018')

'nearest' ('cie15:2004') recommended or other (e.g. 'cubic') methods.

If None: use CIE15:2004 recommended 'nearest value' approach when extrapolating.

If float or list or ndarray, use those values to fill extrapolated value(s).

## extrap\_kind

'linear', optional

Extrapolation method used when :extrap\_values: is set to 'ext'.

Options: 'linear' ('cie167:2005'), 'quadratic' ('cie15:2018'),

'nearest' ('cie15:2004'), 'cubic'

CIE15:2018 states that based on a 2017 paper by Wang that 'quadratic' is 'better'.

However, no significant difference was found between 'quadratic' and 'linear' methods.

Also see note 1 below, for why the CIE67:2005 recommended 'linear' extrapolation

is set as the default.

# extrap\_log

False, optional

If True: extrap the log of the spectral values

(not CIE recommended but in most cases seems to give a more realistic estimate, but can sometimes seriously fail, especially for the 'quadratic' extrapolation case (see note 1)!!!)

# **Returns:**

#### returns

ndarray of interpolated spectral data.

(.shape = (number of spectra+1, number of wavelength in wl\_new))

## **Notes:**

1. Type of extrapolation: 'quadratic' vs 'linear'; impact of extrapolating log spectral values:

Using a 'linear' or 'quadratic' extrapolation, as mentioned in

CIE167:2005 and CIE15:2018, resp., can lead to extreme large values

when setting :extrap\_log: (not CIE recommended) to True.

A quick test with the IES TM30 spectra (400 nm - 700 nm, 5 nm spacing)

shows that 'linear' is better than 'quadratic' in terms of

mean, median and max DEu'v' with the original spectra (380 nm - 780 nm, 5 nm spacing).

This conferms the recommendation from CIE167:2005 to use 'linear' extrapolation.

Setting: extrap\_log: to True reduces the median, but inflates the mean due to some

extremely large DEu'v' values. However, the increase in mean and max DEu'v' is much

larger for the 'quadratic' case, suggesting that 'linear' extrapolation

is likely a more suitable recommendation. When using a 1 nm spacing

'linear' is more similar to 'quadratic' when :extrap\_log: is False, otherwise 'linear' remains the 'best'. Hence the choice to use the CIE167:2005 recommended linear extrapolation as default!

to\_xyz(relative=True, rfl=None, cieobs='1931\_2', out=None)

Calculates xyz tristimulus values from spectral data and return as instance of XYZ. **Args:** 

# relative

True or False, optional

Calculate relative XYZ (Yw = 100) or absolute XYZ (Y = Luminance)

```
rfl
```

ndarray with spectral reflectance functions.

Will be interpolated if wavelengths don't match those of :data:

#### cieobs

luxpy.\_CIEOBS, optional

Determines the color matching functions to be used in the calculation of XYZ.

#### out

None or 1 or 2, optional

Determines number and shape of output. (see :returns:)

#### **Returns:**

#### returns

luxpy.XYZ instance with ndarray .value field:

If rfl is None:

If out is None: ndarray of xyz values

(.shape = (data.shape[0],3))

If out == 1: ndarray of xyz values

(.shape = (data.shape[0],3))

If out == 2: (ndarray of xyz, ndarray of xyzw) values

Note that xyz == xyzw, with (.shape=(data.shape[0],3))

If rfl is not None:

If out is None: ndarray of xyz values

(.shape = (rfl.shape[0], data.shape[0], 3))

If out == 1: ndarray of xyz values

(.shape = (rfl.shape[0]+1,data.shape[0],3))

The xyzw values of the light source spd are the first

set of values of the first dimension.

The following values along this dimension are the sample (rfl) xyz values.

If out == 2: (ndarray of xyz, ndarray of xyzw) values with xyz.shape = (rfl.shape[0],data.shape[0],3)

and with xyzw.shape = (data.shape[0],3)

### **References:**

1. CIE15:2018, "Colorimetry," CIE, Vienna, Austria, 2018.

# 4.4 Color sub-package

рy

• \_\_init\_\_.py

namespace luxpy

# 4.4.1 utils/

рy

- \_\_init\_\_.py
- · plotters.py

namespace luxpy

# Module with functions related to plotting of color data

```
get_cmap() Get an idarray of rgb values representing a linearly sampled matplotlib colormap
get_subplot_layout() Calculate layout of multiple subplots.
plot_color_data() Plot color data (local helper function)
plotDL() Plot daylight locus.
plotBB() Plot blackbody locus.
plotSL()
      Plot spectrum locus.
      (plotBB() and plotDL() are also called, but can be turned off).
plotcerulean()
      Plot cerulean (yellow (577 nm) - blue (472 nm)) line
      (Kuehni, CRA, 2014: Table II: spectral lights)
      Kuehni, R. G. (2014). Unique hues and their stimuli—state of the art. Color Research
      & Application, 39(3), 279–287.
plotUH()
      Plot unique hue lines from color space center point xyz0.
      (Kuehni, CRA, 2014: uY,uB,uG: Table II: spectral lights;
      uR: Table IV: Xiao data)
      Kuehni, R. G. (2014). Unique hues and their stimuli—state of the art. Color Research
      & Application, 39(3), 279–287.
plotcircle() Plot one or more concentric circles.
plotellipse() Plot one or more ellipses.
plot_chromaticity_diagram_colors() Plot the chromaticity diagram colors.
plot_spectrum_colors() Plot spd with spectrum colors.
plot_rfl_color_patches() Create (and plot) an image with colored patches representing a set
      of reflectance spectra illuminated by a specified illuminant.
plot_rgb_color_patches() Create (and plot) an image with patches with specified rgb values.
```

```
luxpy.color.utils.get_cmap(N, cmap_name='jet')
```

Get an ndarray of rgba values representing a linearly sampled matplotlib colormap.

Args:

N

Number of rgba values in returned cmap.

#### cmap\_name

'jet', optional

Matplotlib color map name to sample from.

#### **Returns:**

cmap

ndarray with rgba values.

luxpy.color.utils.get\_subplot\_layout(N, min\_1xncols=3)

Calculate layout of multiple subplots.

**Args:** 

N

Number of plots.

# min\_1xncols

Minimum number of columns before splitting over multiple rows.

#### **Returns:**

nrows, ncols

Plot spectrum locus for cieobs in cspace.

# **Args:**

DL

True or False, optional

True plots Daylight Locus as well.

# BBL

True or False, optional

True plots BlackBody Locus as well.

# **D65**

False or True, optional

True plots D65 chromaticity as well.

# **EEW**

False or True, optional

True plots Equi-Energy-White chromaticity as well.

# cctlabels

False or True, optional

Add cct text labels at various points along the blackbody locus.

## axh

None or axes handle, optional

Determines axes to plot data in.

None: make new figure.

#### show

True or False, optional

Invoke matplotlib.pyplot.show() right after plotting

#### cieobs

luxpy.\_CIEOBS or str, optional

Determines CMF set to calculate spectrum locus or other.

## cspace

luxpy.\_CSPACE or str, optional

Determines color space / chromaticity diagram to plot data in.

Note that data is expected to be in specified :cspace:

#### formatstr

'k-' or str, optional

Format str for plotting (see ?matplotlib.pyplot.plot)

# cspace\_pars

{} or dict, optional

Dict with parameters required by color space specified in :cspace:

(for use with luxpy.colortf())

# diagram\_colors

False, optional

True: plot colored chromaticity diagram.

## diagram\_samples

256, optional

Sampling resolution of color space.

# diagram\_opacity

1.0, optional

Sets opacity of chromaticity diagram

# diagram\_lightness

0.25, optional

Sets lightness of chromaticity diagram

## kwargs

additional keyword arguments for use with matplotlib.pyplot.

#### **Returns:**

#### returns

handle to current axes (:show: == False)

luxpy.color.utils.plotDL(ccts=None, cieobs='1931\_2', cspace='Yuv', axh=None, show=True, force\_daylight\_below4000K=False, cspace\_pars={}, formatstr='k-', \*\*kwargs)

Plot daylight locus.

## **Args:**

ccts

None or list[float], optional

None defaults to [4000 K to 1e11 K] in 100 steps on a log10 scale.

### force\_daylight\_below4000K

False or True, optional

CIE daylight phases are not defined below 4000 K.

```
If True plot anyway.
                  axh
                        None or axes handle, optional
                        Determines axes to plot data in.
                        None: make new figure.
                  show
                        True or False, optional
                        Invoke matplotlib.pyplot.show() right after plotting
                  cieobs
                        luxpy._CIEOBS or str, optional
                        Determines CMF set to calculate spectrum locus or other.
                  cspace
                        luxpy. CSPACE or str, optional
                        Determines color space / chromaticity diagram to plot data in.
                        Note that data is expected to be in specified :cspace:
                  formatstr
                        'k-' or str, optional
                        Format str for plotting (see ?matplotlib.pyplot.plot)
                  cspace_pars
                        {} or dict, optional
                        Dict with parameters required by color space specified in :cspace:
                        (for use with luxpy.colortf())
                  kwargs
                        additional keyword arguments for use with matplotlib.pyplot.
      Returns:
                  returns
                        handle to current axes (:show: == False)
luxpy.color.utils.plotBB(ccts=None, cieobs='1931_2', cspace='Yuv', axh=None, cctlabels=True, show=True,
                                cspace_pars={}, formatstr='k-', **kwargs)
      Plot blackbody locus.
      Args:
                  ccts
                        None or list[float], optional
                        None defaults to [1000 to 1e19 K].
                        Range:
                                     [1000,1500,2000,2500,3000,3500,4000,5000,6000,8000,10000]
                              + [15000 K to 1e11 K] in 100 steps on a log10 scale
                  cctlabels
                        True or False, optional
                        Add cct text labels at various points along the blackbody locus.
                  axh
```

```
None or axes handle, optional
                        Determines axes to plot data in.
                        None: make new figure.
                  show
                        True or False, optional
                        Invoke matplotlib.pyplot.show() right after plotting
                  cieobs
                        luxpy._CIEOBS or str, optional
                        Determines CMF set to calculate spectrum locus or other.
                  cspace
                        luxpy._CSPACE or str, optional
                        Determines color space / chromaticity diagram to plot data in.
                        Note that data is expected to be in specified :cspace:
                  formatstr
                        'k-' or str, optional
                        Format str for plotting (see ?matplotlib.pyplot.plot)
                  cspace_pars
                        {} or dict, optional
                        Dict with parameters required by color space specified in :cspace:
                        (for use with luxpy.colortf())
                  kwargs
                        additional keyword arguments for use with matplotlib.pyplot.
      Returns:
                  returns
                        handle to current axes (:show: == False)
luxpy.color.utils.plot_color_data(x, y, z=None, axh=None, show=True, cieobs='1931_2', cspace='Yuv',
                                           formatstr='k-', legend_loc=None, **kwargs)
      Plot color data from x,y [,z].
      Args:
                  X
                        float or ndarray with x-coordinate data
                  y
                        float or ndarray with y-coordinate data
                  Z
                        None or float or ndarray with Z-coordinate data, optional
                        If None: make 2d plot.
                  axh
                        None or axes handle, optional
                        Determines axes to plot data in.
                        None: make new figure.
                  show
```

```
True or False, optional
```

Invoke matplotlib.pyplot.show() right after plotting

## cieobs

luxpy.\_CIEOBS or str, optional

Determines CMF set to calculate spectrum locus or other.

## cspace

luxpy.\_CSPACE or str or None, optional

Determines color space / chromaticity diagram to plot data in.

Note that data is expected to be in specified :cspace:

If None: don't do any formatting of x,y [z] axes.

## formatstr

'k-' or str, optional

Format str for plotting (see ?matplotlib.pyplot.plot)

#### kwargs

additional keyword arguments for use with matplotlib.pyplot.

#### **Returns:**

#### returns

handle to current axes (:show: == False)

Plot cerulean (yellow (577 nm) - blue (472 nm)) line

Kuehni, CRA, 2014:

Table II: spectral lights.

## **Args:**

#### axh

None or axes handle, optional

Determines axes to plot data in.

None: make new figure.

# cieobs

luxpy.\_CIEOBS or str, optional

Determines CMF set to calculate spectrum locus or other.

# cspace

luxpy.\_CSPACE or str, optional

Determines color space / chromaticity diagram to plot data in.

Note that data is expected to be in specified :cspace:

## formatstr

'k-' or str, optional

Format str for plotting (see ?matplotlib.pyplot.plot)

## cspace\_pars

```
{} or dict, optional
                        Dict with parameters required by color space specified in :cspace:
                        (for use with luxpy.colortf())
                  kwargs
                        additional keyword arguments for use with matplotlib.pyplot.
      Returns:
                  returns
                        handle to cerulean line
      References: 1. Kuehni, R. G. (2014). Unique hues and their stimuli—state of the art. Color Research &
            Application, 39(3), 279–287. (see Table II, IV)
luxpy.color.utils.plotUH(xyz0=None, uhues=[0, 1, 2, 3], cieobs='1931_2', cspace='Yuv', axh=None,
                                formatstr=['yo-.', 'bo-.', 'ro-.', 'go-.'], excludefromlegend=", cspace_pars={})
      Plot unique hue lines from color space center point xyz0.
      Kuehni, CRA, 2014:
            uY,uB,uG: Table II: spectral lights;
            uR: Table IV: Xiao data.
      Args:
                  xyz0
                        None, optional
                        Center of color space (unique hue lines are expected to cross here)
                        None defaults to equi-energy-white.
                  uhues
                        [0,1,2,3], optional
                        Unique hue lines to plot [0:'yellow',1:'blue',2:'red',3:'green']
                  axh
                        None or axes handle, optional
                        Determines axes to plot data in.
                        None: make new figure.
                  cieobs
                        luxpy._CIEOBS or str, optional
                        Determines CMF set to calculate spectrum locus or other.
                  cspace
                        luxpy._CSPACE or str, optional
                        Determines color space / chromaticity diagram to plot data in.
                        Note that data is expected to be in specified :cspace:
                  formatstr
                        ['yo-.','bo-.','ro-.','go-.'] or list[str], optional
                        Format str for plotting the different unique lines
                        (see also ?matplotlib.pyplot.plot)
                  excludefromlegend
```

```
" or str, optional
                        To exclude certain hues from axes legend.
                  cspace_pars
                        {} or dict, optional
                        Dict with parameters required by color space specified in :cspace:
                        (for use with luxpy.colortf())
      Returns:
                  returns
                        list[handles] to unique hue lines
      References: 1. Kuehni, R. G. (2014). Unique hues and their stimuli—state of the art. Color Research &
            Application, 39(3), 279–287. (see Table II, IV)
luxpy.color.utils.plotcircle(center=array([[0.0, 0.0]]), radii=array([0, 10, 20, 30, 40, 50]),
                                     angles=array([0, 10, 20, 30, 40, 50, 60, 70, 80, 90, 100, 110, 120, 130, 140,
                                     150, 160, 170, 180, 190, 200, 210, 220, 230, 240, 250, 260, 270, 280, 290, 300,
                                     310, 320, 330, 340]), color='k', linestyle='--', out=None, axh=None,
      Plot one or more concentric circles.
      Args:
                  center
                        np.array([[0.,0.]]) or ndarray with center coordinates, optional
                  radii
                        np.arange(0,60,10) or ndarray with radii of circle(s), optional
                  angles
                        np.arange(0,350,10) or ndarray with angles (°), optional
                  color
                        'k', optional
                        Color for plotting.
                  linestyle
                        '-', optional
                        Linestyle of circles.
                  out
                        None, optional
                        If None: plot circles, return (x,y) otherwise.
luxpy.color.utils.plotellipse(v, cspace_in='Yxy', cspace_out=None, nsamples=100, show=True,
                                      axh=None, line color='darkgray', line style=':', line width=1,
                                      line_marker=", line_markersize=4, plot_center=False, center_marker='o',
                                      center_color='darkgray', center_markersize=4, show_grid=False, llabel=",
                                      label_fontname='Times New Roman', label_fontsize=12, out=None)
      Plot ellipse(s) given in v-format [Rmax,Rmin,xc,yc,theta].
      Args:
                        (Nx5) ndarray
                        ellipse parameters [Rmax,Rmin,xc,yc,theta]
                  cspace_in
```

```
'Yxy', optional
      Color space of v.
      If None: no color space assumed. Axis labels assumed ('x','y').
cspace_out
      None, optional
      Color space to plot ellipse(s) in.
      If None: plot in cspace_in.
nsamples
      100 or int, optional
      Number of points (samples) in ellipse boundary
show
      True or boolean, optional
      Plot ellipse(s) (True) or not (False)
axh
      None, optional
      Ax-handle to plot ellipse(s) in.
      If None: create new figure with axes.
line_color
      'darkgray', optional
      Color to plot ellipse(s) in.
line_style
      ":', optional
      Linestyle of ellipse(s).
line_width'
      1, optional
      Width of ellipse boundary line.
line_marker
      'none', optional
      Marker for ellipse boundary.
line_markersize
      4, optional
      Size of markers in ellipse boundary.
plot_center
      False, optional
      Plot center of ellipse: yes (True) or no (False)
center_color
      'darkgray', optional
      Color to plot ellipse center in.
center_marker
      'o', optional
```

Marker for ellipse center.

#### center\_markersize

4, optional

Size of marker of ellipse center.

## show\_grid

False, optional

Show grid (True) or not (False)

#### llabel

None, optional

Legend label for ellipse boundary.

#### label fontname

'Times New Roman', optional Sets font type of axis labels.

## label fontsize

12, optional

Sets font size of axis labels.

#### out

None, optional

Output of function

If None: returns None. Can be used to output axh of newly created figure axes or to return Yxys an ndarray with coordinates of ellipse boundaries in cspace\_out (shape = (nsamples,3,N))

# **Returns:**

returns None, or whatever set by :out:.

```
luxpy.color.utils.plot_chromaticity_diagram_colors(diagram_samples=256, diagram_opacity=1.0, diagram_lightness=0.25, cieobs='1931_2',
```

cspace='Yxy', cspace\_pars={}, show=True, axh=None, show\_grid=False, label\_fontname='Times New Roman', label\_fontsize=12, \*\*kwargs)

Plot the chromaticity diagram colors.

# **Args:**

# diagram\_samples

256, optional

Sampling resolution of color space.

# diagram\_opacity

1.0, optional

Sets opacity of chromaticity diagram

# diagram\_lightness

0.25, optional

Sets lightness of chromaticity diagram

axh

None or axes handle, optional Determines axes to plot data in.

None: make new figure.

```
show
                        True or False, optional
                        Invoke matplotlib.pyplot.show() right after plotting
                  cieobs
                        luxpy._CIEOBS or str, optional
                        Determines CMF set to calculate spectrum locus or other.
                  cspace
                        luxpy._CSPACE or str, optional
                        Determines color space / chromaticity diagram to plot data in.
                        Note that data is expected to be in specified :cspace:
                  cspace_pars
                        {} or dict, optional
                        Dict with parameters required by color space specified in :cspace:
                        (for use with luxpy.colortf())
                  show_grid
                        False, optional
                        Show grid (True) or not (False)
                  label_fontname
                        'Times New Roman', optional
                        Sets font type of axis labels.
                  label_fontsize
                        12, optional
                        Sets font size of axis labels.
                  kwargs
                        additional keyword arguments for use with matplotlib.pyplot.
      Returns:
luxpy.color.utils.plot_spectrum_colors(spd=None, spdmax=None, wavelength_height=-0.05,
                                                 wavelength_opacity=1.0, wavelength_lightness=1.0,
                                                 cieobs='1931_2', show=True, axh=None, show_grid=False,
                                                 ylabel='Spectral intensity (a.u.)', xlim=None, **kwargs)
      Plot the spectrum colors.
      Args:
                  spd
                        None, optional
                        Spectrum
                  spdmax
                        None, optional
                        max ylim is set at 1.05 or (1+abs(wavelength_height)*spdmax)
```

# wavelength\_opacity 1.0, optional Sets opacity of

Sets opacity of wavelength rectangle.

## wavelength\_lightness

1.0, optional

Sets lightness of wavelength rectangle.

# wavelength\_height

-0.05 or 'spd', optional

Determine wavelength bar height

if not 'spd': x% of spd.max()

#### axh

None or axes handle, optional

Determines axes to plot data in.

None: make new figure.

#### show

True or False, optional

Invoke matplotlib.pyplot.show() right after plotting

#### cieobs

luxpy.\_CIEOBS or str, optional

Determines CMF set to calculate spectrum locus or other.

## show\_grid

False, optional

Show grid (True) or not (False)

## ylabel

'Spectral intensity (a.u.)' or str, optional

Set y-axis label.

#### xlim

None, optional

list or ndarray with xlimits.

## kwargs

additional keyword arguments for use with matplotlib.pyplot.

#### Returns:

luxpy.color.utils.plot\_rfl\_color\_patches(rfl, spd=None, cieobs='1931\_2', patch\_shape=(100, 100), patch\_layout=None, ax=None, show=True)

Create (and plot) an image with colored patches representing a set of reflectance spectra illuminated by a specified illuminant.

## **Args:**

rfl

ndarray with reflectance spectra

#### spd

None, optional

```
If None: _CIE_D65 is used.
                  cieobs
                        '1931_2', optional
                        CIE standard observer to use when converting rfl to xyz.
                  patch_shape
                        (100,100), optional
                        shape of each of the patches in the image
                  patch_layout
                        None, optional
                        If None: layout is calculated automatically to give a 'good' aspect ratio
                  ax
                        None, optional
                        Axes to plot the image in. If None: a new axes is created.
                  show
                        True, optional
                        If True: plot image in axes and return axes handle; else: return ndarray with image.
      Return:
                  ax or :imagae: | Axes is returned if show == True, else: ndarray with rgb image is returned.
luxpy.color.utils.plot_rgb_color_patches(rgb, patch_shape=(100, 100), patch_layout=None, ax=None,
                                                    show=True)
      Create (and plot) an image with patches with specified rgb values.
      Args:
                  rgb
                        ndarray with rgb values for each of the patches
                  patch_shape
                        (100,100), optional
                        shape of each of the patches in the image
                  patch_layout
                        None, optional
                        If None: layout is calculated automatically to give a 'good' aspect ratio
                  ax
                        None, optional
                        Axes to plot the image in. If None: a new axes is created.
                  show
                        True, optional
                        If True: plot image in axes and return axes handle; else: return ndarray with image.
      Return:
                  ax or :imagae: | Axes is returned if show == True, else: ndarray with rgb image is returned.
luxpy.color.utils.plot_cmfs(cmfs, cmf_symbols=['x', 'y', 'z'], cmf_label=", ylabel='Sensitivity',
                                    wavelength_bar=True, colors=['r', 'g', 'b'], axh=None, legend=True, **kwargs)
      Plot CMFs.
```

ndarray with illuminant spectral power distribution

## Args:

## cmfs

ndarray with a set of CMFs.

# cmf\_symbols

['x,'y','z], optional

Symbols of the CMFs

If not a list but a string, the same label will be used for all CMF and the same color will be used ('k' if colors is a list)

## cmf\_label

", optional

Additional label that will be added in front of the cmf symbols.

# ylabel

'Sensitivity', optional label for y-axis.

## wavelength\_bar

True, optional

Add a colored wavelength bar with spectral colors.

## colors

['r','g','b'], optional

Color for plotting each of the individual CMF.

## axh

None, optional

Axes to plot the image in. If None: a new axes is created.

## **kwargs**

additional kwargs for plt.plot().

# **Returns:**

axh

figure axes handle.

# 4.4.2 ctf/

рy

- \_\_init\_\_.py
- colortransformations.py
- · colortf.py

namespace luxpy

## Module with functions related to basic colorimetry

#### **Note**

Note that colorimetric data is always located in the last axis of the data arrays. (See also xyz specification in \_\_doc\_\_ string of luxpy.spd\_to\_xyz())

## colortransforms.py

```
_CSPACE_AXES dict with list[str,str,str] containing axis labels of defined cspaces
_IPT_M Conversion matrix for IPT color space
```

:\_COLORTF\_DEFAULT\_WHITE\_POINT : default white point for colortf (set at Illuminant E)

## **Supported chromaticity / colorspace functions:**

```
* xyz_to_Yxy(), Yxy_to_xyz(): (X,Y,Z) <-> (Y,x,y);

* xyz_to_Yuv(), Yuv_to_Yxy(): (X,Y,Z) <-> CIE 1976 (Y,u',v');

* xyz_to_Yuv76(), Yuv76_to_Yxy(): (X,Y,Z) <-> CIE 1976 (Y,u',v');

* xyz_to_Yuv60(), Yuv60_to_Yxy(): (X,Y,Z) <-> CIE 1960 (Y,u,v);

* xyz_to_xyz(), lms_to_xyz(): (X,Y,Z) <-> (X,Y,Z); for use with colortf()

* xyz_to_lms(), lms_to_xyz(): (X,Y,Z) <-> (L,M,S) cone fundamental responses

* xyz_to_lab(), lab_to_xyz(): (X,Y,Z) <-> CIE 1976 (L*a*b*)

* xyz_to_luv(), luv_to_xyz(): (X,Y,Z) <-> CIE 1976 (L*u*v*)

* xyz_to_Vrb_mb(),Vrb_mb_to_xyz(): (X,Y,Z) <-> (V,r,b); [Macleod & Boyton, 1979]

* xyz_to_ipt(), ipt_to_xyz(): (X,Y,Z) <-> (I,P,T); (Ebner et al, 1998)

* xyz_to_Ydlep(), Ydlep_to_xyz(): (X,Y,Z) <-> (Y,dl, ep);

Y, dominant wavelength (dl) and excitation purity (ep)

* xyz_to_srgb(), srgb_to_xyz(): (X,Y,Z) <-> sRGB; (IEC:61966 sRGB)
```

### References

1. CIE15:2018, "Colorimetry," CIE, Vienna, Austria, 2018. 2. Ebner F, and Fairchild MD (1998). Development and testing of a color space (IPT) with improved hue uniformity. In IS&T 6th Color Imaging Conference, (Scottsdale, Arizona, USA), pp. 8–13. 3. MacLeod DI, and Boynton RM (1979). Chromaticity diagram showing cone excitation by stimuli of equal luminance. J. Opt. Soc. Am. 69, 1183–1186.

ndarray with Yxy chromaticity values
(Y value refers to luminance or luminance factor)

```
(Y value refers to luminance or luminance factor)
     Returns:
                 XYZ
                       ndarray with tristimulus values
luxpy.color.ctf.colortransforms.xyz_to_Yuv(xyz, **kwargs)
     Convert XYZ tristimulus values CIE 1976 Y,u',v' chromaticity values.
     Args:
                 XYZ
                       ndarray with tristimulus values
     Returns:
                 Yuv
                       ndarray with CIE 1976 Y,u',v' chromaticity values
                             (Y value refers to luminance or luminance factor)
luxpy.color.ctf.colortransforms.Yuv_to_xyz(Yuv, **kwargs)
     Convert CIE 1976 Y,u',v' chromaticity values to XYZ tristimulus values.
     Args:
                 Yuv
                       ndarray with CIE 1976 Y,u',v' chromaticity values
                             (Y value refers to luminance or luminance factor)
     Returns:
                 XYZ
                       ndarray with tristimulus values
luxpy.color.ctf.colortransforms.xyz_to_Yuv76(xyz, **kwargs)
     Convert XYZ tristimulus values CIE 1976 Y,u',v' chromaticity values.
     Args:
                 XYZ
                       ndarray with tristimulus values
     Returns:
                 Yuv
                       ndarray with CIE 1976 Y,u',v' chromaticity values
                             (Y value refers to luminance or luminance factor)
luxpy.color.ctf.colortransforms.Yuv76_to_xyz(Yuv, **kwargs)
     Convert CIE 1976 Y,u',v' chromaticity values to XYZ tristimulus values.
     Args:
                 Yuv
                       ndarray with CIE 1976 Y,u',v' chromaticity values
                             (Y value refers to luminance or luminance factor)
     Returns:
                 xyz
                       ndarray with tristimulus values
luxpy.color.ctf.colortransforms.xyz_to_Yuv60(xyz, **kwargs)
     Convert XYZ tristimulus values CIE 1960 Y,u,v chromaticity values.
     Args:
                 XYZ
                       ndarray with tristimulus values
```

ndarray with Yxy chromaticity values

```
Returns:
                 Yuv
                       ndarray with CIE 1960 Y,u,v chromaticity values
                            (Y value refers to luminance or luminance factor)
luxpy.color.ctf.colortransforms.Yuv60_to_xyz(Yuv60, **kwargs)
     Convert CIE 1976 Y,u,v chromaticity values to XYZ tristimulus values.
     Args:
                 Yuv
                       ndarray with CIE 1976 Yu'v' chromaticity values
                            (Y value refers to luminance or luminance factor)
     Returns:
                 XYZ
                       ndarray with tristimulus values
luxpy.color.ctf.colortransforms.xyz_to_wuv(xyz, xyzw=array([[100.0, 100.0, 100.0, 100.0]]), **kwargs)
     Convert XYZ tristimulus values CIE 1964 U*V*W* color space.
     Args:
                 XYZ
                       ndarray with tristimulus values
                 xyzw
                       ndarray with tristimulus values of white point, optional
                            (Defaults to luxpy._COLORTF_DEFAULT_WHITE_POINT)
     Returns:
                 wuv
                       ndarray with W*U*V* values
luxpy.color.ctf.colortransforms.wuv_to_xyz(wuv, xyzw=array([[100.0, 100.0, 100.0]]), **kwargs)
     Convert CIE 1964 U*V*W* color space coordinates to XYZ tristimulus values.
     Args:
                 wuv
                       ndarray with W*U*V* values
                 XYZW
                       ndarray with tristimulus values of white point, optional
                            (Defaults to luxpy._COLORTF_DEFAULT_WHITE_POINT)
     Returns:
                 XYZ
                       ndarray with tristimulus values
luxpy.color.ctf.colortransforms.xyz_to_xyz(xyz, **kwargs)
     Convert XYZ tristimulus values to XYZ tristimulus values.
     Args:
                 XYZ
                       ndarray with tristimulus values
     Returns:
                 XYZ
                       ndarray with tristimulus values
```

```
luxpy.color.ctf.colortransforms.xyz_to_lms(xyz, cieobs='1931_2', M=None, **kwargs)
     Convert XYZ tristimulus values to LMS cone fundamental responses.
     Args:
                 xyz
                       ndarray with tristimulus values
                 cieobs
                       CIEOBS or str, optional
                 M
                       None, optional
                       Conversion matrix for xyz to lms.
                             If None: use the one defined by :cieobs:
     Returns:
                 lms
                       ndarray with LMS cone fundamental responses
luxpy.color.ctf.colortransforms.lms_to_xyz(lms, cieobs='1931_2', M=None, **kwargs)
     Convert LMS cone fundamental responses to XYZ tristimulus values.
     Args:
                 lms
                       ndarray with LMS cone fundamental responses
                 cieobs
                       _CIEOBS or str, optional
                 M
                       None, optional
                       Conversion matrix for xyz to lms.
                             If None: use the one defined by :cieobs:
     Returns:
                 XYZ
                       ndarray with tristimulus values
luxpy.color.ctf.colortransforms.xyz_to_lab(xyz, xyzw=None, cieobs='1931_2', **kwargs)
     Convert XYZ tristimulus values to CIE 1976 L*a*b* (CIELAB) coordinates.
     Args:
                 XYZ
                       ndarray with tristimulus values
                 xyzw
                       None or ndarray with tristimulus values of white point, optional
                       None defaults to xyz of CIE D65 using the :cieobs: observer.
                 cieobs
                       luxpy._CIEOBS, optional
                       CMF set to use when calculating xyzw.
     Returns:
                 lab
                       ndarray with CIE 1976 L*a*b* (CIELAB) color coordinates
```

```
luxpy.color.ctf.colortransforms.lab_to_xyz(lab, xyzw=None, cieobs='1931_2', **kwargs)
     Convert CIE 1976 L*a*b* (CIELAB) color coordinates to XYZ tristimulus values.
     Args:
                 lab
                       ndarray with CIE 1976 L*a*b* (CIELAB) color coordinates
                 xyzw
                       None or ndarray with tristimulus values of white point, optional
                       None defaults to xyz of CIE D65 using the :cieobs: observer.
                 cieobs
                       luxpy._CIEOBS, optional
                       CMF set to use when calculating xyzw.
     Returns:
                 XYZ
                       ndarray with tristimulus values
luxpy.color.ctf.colortransforms.xyz_to_luv(xyz, xyzw=None, cieobs='1931_2', **kwargs)
     Convert XYZ tristimulus values to CIE 1976 L*u*v* (CIELUV) coordinates.
     Args:
                 xyz
                       ndarray with tristimulus values
                 xyzw
                       None or ndarray with tristimulus values of white point, optional
                       None defaults to xyz of CIE D65 using the :cieobs: observer.
                 cieobs
                       luxpy._CIEOBS, optional
                       CMF set to use when calculating xyzw.
     Returns:
                 luv
                       ndarray with CIE 1976 L*u*v* (CIELUV) color coordinates
luxpy.color.ctf.colortransforms.luv_to_xyz(luv, xyzw=None, cieobs='1931_2', **kwargs)
     Convert CIE 1976 L*u*v* (CIELUVB) coordinates to XYZ tristimulus values.
     Args:
                 luv
                       ndarray with CIE 1976 L*u*v* (CIELUV) color coordinates
                 xyzw
                       None or ndarray with tristimulus values of white point, optional
                       None defaults to xyz of CIE D65 using the :cieobs: observer.
                 cieobs
                       luxpy._CIEOBS, optional
                       CMF set to use when calculating xyzw.
     Returns:
                 xyz
                       ndarray with tristimulus values
```

```
luxpy.color.ctf.colortransforms.xyz_to_Vrb_mb(xyz, cieobs='1931_2', scaling=[1, 1], M=None,
                                                         **kwargs)
     Convert XYZ tristimulus values to V,r,b (Macleod-Boynton) color coordinates.
     Macleod Boynton: V = R+G, r = R/V, b = B/V
     Note that R,G,B ~ L,M,S
     Args:
                 xyz
                       ndarray with tristimulus values
                 cieobs
                       luxpy._CIEOBS, optional
                       CMF set to use when getting the default M, which is the xyz to lms conversion matrix.
                 scaling
                       list of scaling factors for r and b dimensions.
                 M
                       None, optional
                       Conversion matrix for going from XYZ to RGB (LMS)
                             If None, :cieobs: determines the M (function does inversion)
     Returns:
                 Vrb
                       ndarray with V,r,b (Macleod-Boynton) color coordinates
     Reference:
              1. MacLeod DI, and Boynton RM (1979). Chromaticity diagram showing cone excitation by stimuli
                 of equal luminance. J. Opt. Soc. Am. 69, 1183-1186.
luxpy.color.ctf.colortransforms.Vrb_mb_to_xyz(Vrb, cieobs='1931_2', scaling=[1, 1], M=None,
                                                         Minverted=False, **kwargs)
     Convert V,r,b (Macleod-Boynton) color coordinates to XYZ tristimulus values.
     Macleod Boynton: V = R+G, r = R/V, b = B/V
     Note that R,G,B ~ L,M,S
     Args:
                 Vrb
                       ndarray with V,r,b (Macleod-Boynton) color coordinates
                 cieobs
                       luxpy._CIEOBS, optional
                       CMF set to use when getting the default M, which is
                       the xyz to lms conversion matrix.
                 scaling
                       list of scaling factors for r and b dimensions.
                 \mathbf{M}
```

None, optional

Conversion matrix for going from XYZ to RGB (LMS)

If None, :cieobs: determines the M (function does inversion)

#### Minverted

False, optional

Bool that determines whether M should be inverted.

#### **Returns:**

**XYZ** 

ndarray with tristimulus values

#### Reference:

1. MacLeod DI, and Boynton RM (1979). Chromaticity diagram showing cone excitation by stimuli of equal luminance. J. Opt. Soc. Am. 69, 1183–1186.

luxpy.color.ctf.colortransforms.xyz\_to\_ipt(xyz, cieobs='1931\_2', xyzw=None, M=None, \*\*kwargs)
Convert XYZ tristimulus values to IPT color coordinates.

I: Lightness axis, P, red-green axis, T: yellow-blue axis.

## Args:

xyz

ndarray with tristimulus values

**XYZW** 

None or ndarray with tristimulus values of white point, optional None defaults to xyz of CIE D65 using the :cieobs: observer.

## cieobs

luxpy.\_CIEOBS, optional

CMF set to use when calculating xyzw for rescaling M (only when not None).

M

None, optional

None defaults to xyz to lms conversion matrix determined by :cieobs:

# **Returns:**

ipt

ndarray with IPT color coordinates

#### Note:

xyz is assumed to be under D65 viewing conditions! If necessary perform chromatic adaptation!

# Reference:

1. Ebner F, and Fairchild MD (1998). Development and testing of a color space (IPT) with improved hue uniformity. In IS&T 6th Color Imaging Conference, (Scottsdale, Arizona, USA), pp. 8–13.

luxpy.color.ctf.colortransforms.ipt\_to\_xyz(ipt, cieobs='1931\_2', xyzw=None, M=None, \*\*kwargs)
Convert XYZ tristimulus values to IPT color coordinates.

I: Lightness axis, P, red-green axis, T: yellow-blue axis.

#### **Args:**

ipt

ndarray with IPT color coordinates

## xyzw

None or ndarray with tristimulus values of white point, optional None defaults to xyz of CIE D65 using the :cieobs: observer.

#### cieobs

luxpy.\_CIEOBS, optional

CMF set to use when calculating xyzw for rescaling Mxyz2lms (only when not None).

M

None, optional

None defaults to xyz to lms conversion matrix determined by:cieobs:

#### **Returns:**

XYZ

ndarray with tristimulus values

#### Note:

xyz is assumed to be under D65 viewing conditions! If necessary perform chromatic adaptation!

#### **Reference:**

1. Ebner F, and Fairchild MD (1998). Development and testing of a color space (IPT) with improved hue uniformity. In IS&T 6th Color Imaging Conference, (Scottsdale, Arizona, USA), pp. 8–13.

Convert XYZ tristimulus values to Y, dominant (complementary) wavelength and excitation purity. **Args:** 

xyz

ndarray with tristimulus values

#### **XYZW**

None or ndarray with tristimulus values of a single (!) native white point, optional None defaults to xyz of CIE D65 using the :cieobs: observer.

#### cieobs

luxpy.\_CIEOBS, optional

CMF set to use when calculating spectrum locus coordinates.

## flip\_axes

False, optional

If True: flip axis 0 and axis 1 in Ydelep to increase speed of loop in function. (single xyzw with is not flipped!)

# $SL_max_lambda$

None or float, optional

Maximum wavelength of spectrum locus before it turns back on itelf in the high wavelength range ( $\sim$ 700 nm)

# **Returns:**

## Ydlep

ndarray with Y, dominant (complementary) wavelength and excitation purity

Convert Y, dominant (complementary) wavelength and excitation purity to XYZ tristimulus values.  $\bf Args:$ 

#### Ydlep

ndarray with Y, dominant (complementary) wavelength and excitation purity

## xyzw

None or narray with tristimulus values of a single (!) native white point, optional None defaults to xyz of CIE D65 using the :cieobs: observer.

### cieobs

luxpy.\_CIEOBS, optional

CMF set to use when calculating spectrum locus coordinates.

## flip\_axes

False, optional

If True: flip axis 0 and axis 1 in Ydelep to increase speed of loop in function. (single xyzw with is not flipped!)

## SL\_max\_lambda

None or float, optional

Maximum wavelength of spectrum locus before it turns back on itelf in the high wavelength range ( $\sim$ 700 nm)

#### **Returns:**

xyz

ndarray with tristimulus values

luxpy.color.ctf.colortransforms.xyz\_to\_srgb(xyz, gamma=2.4, offset=-0.055,  $use\_linear\_part=True$ , M=None, \*\*kwargs)

Calculates IEC:61966 sRGB values from xyz.

## Args:

XYZ

ndarray with relative tristimulus values.

# gamma

2.4, optional

Gamma compression in gamma-function gf(x): see notes

#### offset

-0.055, optional

Offset in gamma-function gf(x): see notes

## use\_linear\_part

True, optional

If False: omit linear part at low RGB values and use gamma function throughout

## M

None, optional

xyz to linear srgb conversion matrix.

If None: use predefined matrix

#### **Returns:**

rgb

ndarray with R,G,B values (uint8).

#### Notes:

- 1. Gamma-function: gf(x) = ((1-offset)\*x\*\*gamma + offset)\*255
- 2. dark values use linear function: lf(x) = x[dark] \* 12.92 \* 255
- 3. To use a pure gamma function, set offset to zero and use\_linear\_part to False.

Calculates xyz from IEC:61966 sRGB values.

## **Args:**

rgb

ndarray with srgb values (uint8).

#### gamma

2.4, optional

Gamma compression in gamma-function gf(x): see notes

## offset

-0.055, optional

Offset in gamma-function gf(x): see notes

## use\_linear\_part

True, optional

If False: omit linear part at low RGB values and use gamma function throughout

#### M

None, optional

xyz to linear srgb conversion matrix

(!!! Don't give inverse matrix as input, function will take inverse of input to M!!!).

If None: use predefined inverse matrix

#### **Returns:**

xyz

ndarray with xyz tristimulus values.

#### **Notes:**

- 1. Gamma-function: gf(x) = ((1-offset)\*x\*\*gamma + offset)\*255
- 2. dark values use linear function: lf(x) = x[dark] \* 12.92 \* 255
- 3. To use a pure gamma function, set offset to zero and use\_linear\_part to False.

## Extension of basic colorimetry module

Global internal variables:

\_COLORTF\_DEFAULT\_WHITE\_POINT ndarray with XYZ values of default white point (equi-energy white) for color transformation if none is supplied.

Functions:

**colortf()** Calculates conversion between any two color spaces ('cspace') for which functions xyz\_to\_cspace() and cspace\_to\_xyz() are defined.

```
luxpy.color.ctf.colortf.colortf(data, tf='Yuv', fwtf={}, bwtf={}, **kwargs)
```

Wrapper function to perform various color transformations.

Args:

data

ndarray

tf

\_CSPACE or str specifying transform type, optional

If tf is for example 'Yuv', it is assumed to be a transformation of type: 'xyz>Yuv'

fwtf

dict with parameters (keys) and values required

by some color transformations for the forward transform:

bwtf

dict with parameters (keys) and values required

by some color transformations for the backward transform:

**Returns:** 

returns

ndarray with data transformed to new color space

**Note:** For the forward transform ('xyz>...'), one can input the keyword arguments specifying the transform parameters directly without having to use the dict:fwtf: (should be empty!) [i.e. kwargs overwrites empty fwtf dict]

# 4.4.3 cct/

рy

- \_\_init\_\_.py
- · cct.py
- cct\_legacy.py
- cctduv\_ohno\_CORM2011.py

namespace luxpy

## cct: Module with functions related to correlated color temperature calculations

These methods supersede earlier methods in cct\_legacy.y (prior to Nov 2021)

- **\_CCT\_MAX** (= 1e11 K), max. value that does not cause overflow problems.
- **\_CCT\_MIN** (= 550 K), min. value that does not cause underflow problems.
- **\_CCT\_FALLBACK\_N** Number of intervals to divide an ndarray with CCTs.
- \_CCT\_FALLBACK\_UNIT Type of scale (units) an ndarray will be subdivided.
- **\_CCT\_LUT\_PATH** Folder with Look-Up-Tables (LUT) for correlated color temperature calculations.
- \_CCT\_LUT Dict with pre-calculated LUTs with structure LUT[mode][cspace][cieobs][lut i].
- \_CCT\_LUT\_CALC Boolean determining whether to force LUT calculation, even if the LUT.pkl files can be found in ./data/cctluts/.
- \_CCT\_LUT\_RESOLUTION\_REDUCTION\_FACTOR number of subdivisions when performing a cascading lut calculation to zoom-in progressively on the CCT (until a certain tolerance is met)
- **\_CCT\_CSPACE** default chromaticity space to calculate CCT and Duv in.
- \_CCT\_CSPACE\_KWARGS nested dict with cspace parameters for forward and backward modes.
- **get\_tcs4**() Get an indarray of Tc's obtained from a list or tuple of tc4 4-vectors.
- calculate\_lut() Function that calculates the LUT for the input ccts.
- **generate\_luts()** Generate a number of luts and store them in a nested dictionary. (Structure: lut[cspace][cieobs][lut type])
- xyz\_to\_cct() Calculates CCT, Duv from XYZ (wraps a variety of methods)
- xyz\_to\_duv() Calculates Duv, (CCT) from XYZ (wrapper around xyz\_to\_cct, but with Duv output.)
- cct\_to\_xyz() Calculates xyz from CCT, Duv by estimating the line perpendicular to the planckian locus (=iso-T line).
- cct\_to\_xyz() Calculates xyz from CCT, Duv [\_CCT\_MIN < CCT < \_CCT\_MAX]</pre>

## xyz\_to\_cct\_mcamy1992()

Calculates CCT from XYZ using Mcamy model:

McCamy, Calvin S. (April 1992). Correlated color temperature as an explicit function of chromaticity coordinates. Color Research & Application. 17 (2): 142–144.

# $xyz\_to\_cct\_hernandez1999()$

Calculate CCT from XYZ using Hernández-Andrés et al. model.

Hernández-Andrés, Javier; Lee, RL; Romero, J (September 20, 1999). Calculating Correlated Color Temperatures Across the Entire Gamut of Daylight and Skylight Chromaticities. Applied Optics. 38 (27): 5703–5709. PMID 18324081.

#### xyz\_to\_cct\_ohno2014()

Calculates CCT, Duv from XYZ using a Ohno's 2014 LUT method.

Ohno Y. (2014) Practical use and calculation of CCT and Duv. Leukos. 2014 Jan 2;10(1):47-55.

```
xyz_to_cct_zhang2019()
```

Calculates CCT, Duv from XYZ using Zhang's 2019 golden-ratio search algorithm Zhang, F. (2019). High-accuracy method for calculating correlated color temperature with a lookup table based on golden section search. Optik, 193, 163018.

## xyz\_to\_cct\_robertson1968()

Calculates CCT, Duv from XYZ using a Robertson's 1968 search method.

Robertson, A. R. (1968). Computation of Correlated Color Temperature and Distribution Temperature. Journal of the Optical Society of America, 58(11), 1528–1535.

## xyz\_to\_cct\_li2016()

Calculates CCT, Duv from XYZ using a Li's 2019 Newton-Raphson method.

Li, C., Cui, G., Melgosa, M., Ruan, X., Zhang, Y., Ma, L., Xiao, K., & Luo, M. R. (2016). Accurate method for computing correlated color temperature. Optics Express, 24(13), 14066–14078.

#### xvz to cct fibonacci()

Calculates CCT, Duv from XYZ using a Fibonacci search method.

cct\_to\_mired() Converts from CCT to Mired scale (or back).

xyz\_to\_cct\_ohno2011() Calculate cct and Duv from CIE 1931 2° xyz following Ohno (CORM 2011).

cct\_legacy module with old (pre Nov 2021 cct conversion functions)

```
luxpy.color.cct.cct_to_mired(data)
```

Convert cct to Mired scale (or back).

**Args:** 

data

ndarray with cct or Mired values.

**Returns:** 

returns

ndarray ((10\*\*6) / data)

Convert XYZ tristimulus values to correlated color temperature (CCT) using the mccamy approximation (!!! only valid for CIE 1931  $2^{\circ}$  input !!!).

Only valid for approx. 3000 < T < 9000, if < 6500, error < 2 K

Args:

xyzw

ndarray of tristimulus values

cieobs

'1931\_2', optional

CMF set used to calculated xyzw.

```
using the 1931 2° CMFs, this is only valid for that CMF set.
                              It can be changed, but will only impact the calculation of Duv and
                              thereby causing a potential mismatch/error. Change at own discretion.
                  out
                        'cct' (or 1), optional
                        Determines what to return.
                        Other options: 'duv' (or -1), 'cct,duv'(or 2), "[cct,duv]" (or -2)
                  wl
                        None, optional
                        Wavelengths used when calculating Planckian radiators when determining Duv.
                              (!!CCT is determined using a fixed set of equations optimized for the 1931 2°
                              CMFS!!)
                  cspace
                        _CCT_SPACE, optional
                        Color space to do calculations in.
                        Options:
                                    - cspace string:
                                          e.g. 'Yuv60' for use with luxpy.colortf()
                                    - tuple with forward (i.e. xyz_to..) [and backward (i.e. ..to_xyz)]
                                    functions
                                          (and an optional string describing the cspace):
                                                e.g. (forward, backward) or (forward, backward, cspace
                                                string) or (forward, cspace string)
                                    - dict with keys: 'fwtf' (foward), 'bwtf' (backward) [, optional: 'str'
                                    (cspace string)]
                              Note: if the backward tf is not supplied, optimization in cct to xyz() is done in
                              the CIE 1976 u'v' diagram
                  cspace_kwargs
                        _CCT_CSPACE_KWARGS, optional
                        Parameter nested dictionary for the forward and backward transforms.
      Returns:
                  cct
                        ndarray of correlated color temperatures estimates
      References: 1. McCamy, Calvin S. (April 1992). "Correlated color temperature as an explicit function of
            chromaticity coordinates". Color Research & Application. 17 (2): 142–144.
luxpy.color.cct.xyz_to_cct_hernandez1999(xyzw, cieobs='1931_2', wl=None, out='cct', cspace='Yuv60',
                                                    cspace_kwargs={'bwtf': {}, 'fwtf': {}})
      Convert XYZ tristimulus values to correlated color temperature (CCT) using the mccamy approximation (!!!
      only valid for CIE 1931 2° input !!!).
      According to paper small error from 3000 - 800 000 K
      Args:
                  xyzw
```

Note: since the parameter values in Mcamy's equation were optimized,

```
ndarray of tristimulus values
cieobs
      '1931_2', optional
      CMF set used to calculated xyzw.
      Note: since the parameter values in the HA equations were optimized,
            using the 1931 2° CMFs, this is only valid for that CMF set.
            It can be changed, but will only impact the calculation of Duv and
            thereby causing a potential mismatch/error. Change at own discretion.
out
      'cct' (or 1), optional
      Determines what to return.
      Other options: 'duv' (or -1), 'cct,duv'(or 2), "[cct,duv]" (or -2)
wl
      None, optional
      Wavelengths used when calculating Planckian radiators when determining Duv.
            (!!CCT is determined using a fixed set of equations optimized for the 1931 2°
            CMFS!!)
cspace
      _CCT_SPACE, optional
      Color space to do calculations in.
      Options:
                  - cspace string:
                        e.g. 'Yuv60' for use with luxpy.colortf()
                  - tuple with forward (i.e. xyz_to..) [and backward (i.e. ..to_xyz)]
                  functions
                        (and an optional string describing the cspace):
                               e.g. (forward, backward) or (forward, backward, cspace
                               string) or (forward, cspace string)
                  - dict with keys: 'fwtf' (foward), 'bwtf' (backward) [, optional: 'str'
                  (cspace string)]
            Note: if the backward tf is not supplied, optimization in cct_to_xyz() is done in
            the CIE 1976 u'v' diagram
cspace_kwargs
      _CCT_CSPACE_KWARGS, optional
      Parameter nested dictionary for the forward and backward transforms.
```

cct

**Returns:** 

ndarray of correlated color temperatures estimates

References: 1. Hernández-Andrés, Javier; Lee, RL; Romero, J (September 20, 1999). Calculating Correlated Color Temperatures Across the Entire Gamut of Daylight and Skylight Chromaticities. Applied Optics. 38 (27), 5703–5709. P

```
luxpy.color.cct.xyz_to_cct_robertson1968(xyzw, cieobs='1931_2', out='cct', is_uv_input=False,
                                                     wl=None, atol=0.1, rtol=1e-05, force_tolerance=True,
                                                     tol method='newton-raphson',
                                                     lut_resolution_reduction_factor=4,
                                                     split_calculation_at_N=25, max_iter=10, cspace='Yuv60',
                                                     cspace_kwargs={'bwtf': {}, 'fwtf': {}}, lut=None,
                                                     luts_dict=None, ignore_wl_diff=False, use_fast_duv=True,
                                                     **kwargs)
      Convert XYZ tristimulus values to correlated color temperature (CCT) and Duv(distance above (> 0) or below (
      < 0) the Planckian locus) using Robertson's 1968 search method.
      Args:
                  XYZW
                        ndarray of tristimulus values
                  cieobs
                        luxpy._CIEOBS, optional
                        CMF set used to calculated xyzw.
                  out
                        'cct' (or 1), optional
                        Determines what to return.
                        Other options: 'duv' (or -1), 'cct,duv' (or 2), "[cct,duv]" (or -2)
                  is_uv_input
                        False, optional
                        If True: xyzw contain uv input data, not xyz data!
                  wl
                        None, optional
                        Wavelengths used when calculating Planckian radiators.
                        If None: use same wavelengths as CMFs in :cieobs:.
                  rtol
                        1e-5, float, optional
                        Stop search when cct a relative tolerance is reached.
                        The relative tolerance is calculated as dCCT/CCT_est,
                        with CCT est the current intermediate estimate in the
                        search and with dCCT the difference between
                        the present and former estimates.
                  atol
                        0.1, optional
                        Stop search when cct a absolute tolerance (K) is reached.
                  force_tolerance
                        True, optional
                        If False: search only using the list of CCTs in the used lut.
                              Only one loop of the full algorithm is performed.
                              Accuracy depends on CCT of test source and the location
                              and spacing of the CCTs in the list.
                        If True: search will use adjacent CCTs to test source to create a new LUT,
```

(repeat the algoritm at higher resolution, progessively zooming in toward the ground-truth) for tol\_method == 'cl'; when tol\_method == 'nr' a newton-raphson method is used.

Because the CCT for multiple source is calculated in one go, the atol and rtol values have to be met for all!

#### tol method

'newton-raphson', optional (Additional) method to try and achieve set tolerances. Options:

- 'cl', 'cascading-lut': use increasingly higher CCT-resolution to 'zoom-in' on the ground-truth.
- 'nr', 'newton-raphson': use the method as described in Li, 2016.

## lut\_resolution\_reduction\_factor

\_CCT\_LUT\_RESOLUTION\_REDUCTION\_FACTOR, optional Number of times the interval spanned by the adjacent Tc in a search or lut method is downsampled (the search process will then start again)

#### max\_iter

\_CCT\_MAX\_ITER, optional

Maximum number of iterations used by the cascading-lut or newton-raphson methods.

## split\_calculation\_at\_N

\_CCT\_SPLIT\_CALC\_AT\_N, optional

Split calculation when xyzw.shape[0] > split\_calculation\_at\_N.

Splitting speeds up the calculation. If None: no splitting is done.

#### lut

None, optional

Look-Up-Table with Ti, u,v,u',v',u",v'',slope values of Planckians.

## Options:

- None: defaults to the lut specified in
- \_CCT\_LUT['robertson1968']['lut\_type\_def'].
- list (lut,lut\_kwargs): use this pre-calculated lut (add additional kwargs for the lut\_generator\_fcn(), defaults to None if omitted)
- tuple: must be key (label) in :luts\_dict: (pre-calculated dict of luts),
   if not: then a new lut will be generated from scratch using the info in the tuple.
- str: must be key (label) in :luts\_dict: (pre-calculated dict of luts)
- ndarray [Nx1]: list of luts for which to generate a lut
- ndarray [Nxn] with n>3: pre-calculated lut (last col must contain slope of the isotemperature lines).

# luts\_dict

None, optional

Dictionary of pre-calculated luts for various cspaces and cmf sets.

Must have structure luts\_dict[cspace][cieobs][lut\_label] with the

lut part of a two-element list [lut, lut\_kwargs]. It must contain

at the top-level a key 'wl' containing the wavelengths of the Planckians used to generate the luts in this dictionary.

If None: luts\_dict defaults to \_CCT\_LUT['robertson1968']['luts']

#### cspace

\_CCT\_SPACE, optional

Color space to do calculations in.

Options:

- cspace string:

e.g. 'Yuv60' for use with luxpy.colortf()

- tuple with forward (i.e. xyz\_to..) [and backward (i.e. ..to\_xyz)] functions

(and an optional string describing the cspace):

e.g. (forward, backward) or (forward, backward, cspace string) or (forward, cspace string)

- dict with keys: 'fwtf' (foward), 'bwtf' (backward) [, optional: 'str' (cspace string)]

Note: if the backward tf is not supplied, optimization in cct\_to\_xyz() is done in the CIE 1976 u'v' diagram

## cspace\_kwargs

\_CCT\_CSPACE\_KWARGS, optional

Parameter nested dictionary for the forward and backward transforms.

## ignore\_wl\_diff

False, optional

When getting a lut from the dictionary, if differences are detected in the wavelengts of the lut and the ones used to calculate any plankcians then a new lut should be generated. Seting this to True ignores these differences and proceeds anyway.

## use\_fast\_duv

\_CCT\_FAST\_DUV, optional

If True: use a fast estimator of the Duv

(one that avoids calculation of Planckians and uses the former best estimate's u,v coordinates. This method is accurate enough when the atol is small enough -> as long as abs(T-T\_former)<=1K the Duv estimate should be ok.)

#### **Returns:**

#### returns

ndarray with:

```
cct: out == 'cct' (or 1)
duv: out == 'duv' (or -1)
cct, duv: out == 'cct,duv' (or 2)
[cct,duv]: out == "[cct,duv]" (or -2)
```

**Note:** 1. Out-of-lut CCTs are encoded as negative CCTs (with as absolute value the value of the closest CCT from the lut.)

**References:** 1. Robertson, A. R. (1968). Computation of Correlated Color Temperature and Distribution Temperature. Journal of the Optical Society of America, 58(11), 1528–1535.

```
2. Li, C., Cui, G., Melgosa, M., Ruan, X., Zhang, Y., Ma, L., Xiao, K., & Luo, M. R. (2016). Accurate method for computing correlated color temperature. Optics Express, 24(13), 14066–14078.
```

```
luxpy.color.cct.xyz_to_cct_ohno2014(xyzw, cieobs='1931_2', out='cct', is_uv_input=False, wl=None, atol=0.1, rtol=1e-05, force_tolerance=True, tol_method='newton-raphson', lut_resolution_reduction_factor=4, duv_parabolic_threshold=0.002, split_calculation_at_N=25, max_iter=10, cspace='Yuv60', cspace_kwargs={'bwtf': {}, 'fwtf': {}}, lut=None, luts_dict=None, ignore_wl_diff=False, use_fast_duv=True, **kwargs)
```

Convert XYZ tristimulus values to correlated color temperature (CCT) and Duv (distance above (>0) or below (<0) the Planckian locus) using Ohno's 2014 method. **Args:** 

#### **xyzw**

ndarray of tristimulus values

#### cieobs

luxpy.\_CIEOBS, optional CMF set used to calculated xyzw.

#### out

'cct' (or 1), optional

Determines what to return.

Other options: 'duv' (or -1), 'cct,duv' (or 2), "[cct,duv]" (or -2)

#### is\_uv\_input

False, optional

If True: xyzw contain uv input data, not xyz data!

## wl

None, optional

Wavelengths used when calculating Planckian radiators.

If None: use same wavelengths as CMFs in :cieobs:.

#### rtol

1e-5, float, optional

Stop search when cct a relative tolerance is reached. The relative tolerance is calculated as dCCT/CCT\_est, with CCT\_est the current intermediate estimate in the search and with dCCT the difference between the present and former estimates.

#### atol

0.1, optional

Stop search when cct a absolute tolerance (K) is reached.

## force\_tolerance

True, optional

If False: search only using the list of CCTs in the used lut.

Only one loop of the full algorithm is performed.

Accuracy depends on CCT of test source and the location

and spacing of the CCTs in the list.

If True: search will use adjacent CCTs to test source to create a new LUT,

(repeat the algoritm at higher resolution, progessively zooming in

toward the ground-truth) for tol\_method == 'cl'; when

tol\_method == 'nr' a newton-raphson method is used.

Because the CCT for multiple source is calculated in one go,

the atol and rtol values have to be met for all!

#### tol method

'newton-raphson', optional

(Additional) method to try and achieve set tolerances.

#### **Options**:

- 'cl', 'cascading-lut': use increasingly higher CCT-resolution to 'zoom-in' on the ground-truth.
- 'nr', 'newton-raphson': use the method as described in Li, 2016.

## lut\_resolution\_reduction\_factor

\_CCT\_LUT\_RESOLUTION\_REDUCTION\_FACTOR, optional

Number of times the interval spanned by the adjacent Tc in a search or lut method is downsampled (the search process will then start again)

## duv\_parabolic\_threshold

0.002, optional

Threshold for use of the parabolic solution

(if larger then use parabolic, else use triangular solution)

# max\_iter

\_CCT\_MAX\_ITER, optional

Maximum number of iterations used by the cascading-lut or newton-raphson methods.

## split\_calculation\_at\_N

\_CCT\_SPLIT\_CALC\_AT\_N, optional

Split calculation when xyzw.shape[0] > split\_calculation\_at\_N.

Splitting speeds up the calculation. If None: no splitting is done.

#### lut

None, optional

Look-Up-Table with Ti, u,v,u',v',u",v",slope values of Planckians.

### Options:

- None: defaults to the lut specified in
- \_CCT\_LUT['ohno2014']['lut\_type\_def'].
- list (lut,lut\_kwargs): use this pre-calculated lut

(add additional kwargs for the lut\_generator\_fcn(), defaults to None if omitted)

- tuple: must be key (label) in :luts\_dict: (pre-calculated dict of luts),
  - if not: then a new lut will be generated from scratch using the info in the tuple.
- str: must be key (label) in :luts\_dict: (pre-calculated dict of luts)
- ndarray [Nx1]: list of luts for which to generate a lut

- ndarray [Nxn] with n>3: pre-calculated lut (last col must contain slope of the isotemperature lines).

## luts\_dict

None, optional

Dictionary of pre-calculated luts for various espaces and cmf sets.

Must have structure luts\_dict[cspace][cieobs][lut\_label] with the lut part of a two-element list [lut, lut\_kwargs]. It must contain at the top-level a key 'wl' containing the wavelengths of the

Planckians used to generate the luts in this dictionary.

If None: luts\_dict defaults to \_CCT\_LUT['ohno2014']['luts']

## cspace

CCT SPACE, optional

Color space to do calculations in.

Options:

- cspace string:

e.g. 'Yuv60' for use with luxpy.colortf()

- tuple with forward (i.e. xyz\_to..) [and backward (i.e. ..to\_xyz)] functions

(and an optional string describing the cspace):

e.g. (forward, backward) or (forward, backward, cspace string) or (forward, cspace string)

- dict with keys: 'fwtf' (foward), 'bwtf' (backward) [, optional: 'str' (cspace string)]

Note: if the backward tf is not supplied, optimization in  $cct\_to\_xyz()$  is done in the CIE 1976 u'v' diagram

# cspace\_kwargs

\_CCT\_CSPACE\_KWARGS, optional

Parameter nested dictionary for the forward and backward transforms.

# ignore\_wl\_diff

False, optional

When getting a lut from the dictionary, if differences are detected in the wavelengts of the lut and the ones used to calculate any plankcians then a new lut should be generated. Seting this to True ignores these differences and proceeds anyway.

#### use\_fast\_duv

\_CCT\_FAST\_DUV, optional

If True: use a fast estimator of the Duv

(one that avoids calculation of Planckians and uses the former best estimate's u,v coordinates. This method is accurate enough when the atol is small enough -> as long as abs(T-T\_former)<=1K the Duv estimate should be ok.)

## **Returns:**

## returns

ndarray with:

```
cct: out == 'cct' (or 1)
                              duv: out == 'duv' (or -1)
                              cct, duv: out == 'cct,duv' (or 2)
                              [cct,duv]: out == "[cct,duv]" (or -2)
      Note: 1. Out-of-lut CCTs are encoded as negative CCTs (with as absolute value the value of the closest CCT
            from the lut.)
      References: 1. Ohno Y. Practical use and calculation of CCT and Duv. Leukos. 2014 Jan 2:10(1):47-55.
            2. Li, C., Cui, G., Melgosa, M., Ruan, X., Zhang, Y., Ma, L., Xiao, K., & Luo, M. R. (2016). Accurate
            method for computing correlated color temperature. Optics Express, 24(13), 14066–14078.
luxpy.color.cct.xyz_to_cct_li2016(xyzw, cieobs='1931_2', out='cct', is_uv_input=False, wl=None,
                                           atol=0.1, rtol=1e-0.5, max iter=10, split calculation at N=2.5,
                                           lut=None, luts_dict=None, ignore_wl_diff=False,
                                           lut_resolution_reduction_factor=4, cspace='Yuv60',
                                           cspace_kwargs={'bwtf': {}, 'fwtf': {}},
                                           first_guess_mode='robertson1968', fgm_kwargs={}, use_fast_duv=True,
                                            **kwargs)
      Convert XYZ tristimulus values to correlated color temperature (CCT) and Duv(distance above (> 0) or below (
      < 0) the Planckian locus) using the Newton-Raphson method described in Li et al. (2016).
                  xyzw
                        ndarray of tristimulus values
                  cieobs
                        luxpy. CIEOBS, optional
                        CMF set used to calculated xyzw.
                  out
                        'cct' (or 1), optional
                        Determines what to return.
                        Other options: 'duv' (or -1), 'cct,duv' (or 2), "[cct,duv]" (or -2)
                  is_uv_input
                        False, optional
                        If True: xyzw contain uv input data, not xyz data!
                  wl
                        None, optional
                        Wavelengths used when calculating Planckian radiators.
                        If None: use same wavelengths as CMFs in :cieobs:.
                  rtol
                        1e-5, float, optional
                        Stop method when cct a relative tolerance is reached.
                        The relative tolerance is calculated as dCCT/CCT est,
                        with CCT_est the current intermediate estimate in the
                        search and with dCCT the difference between
                        the present and former estimates.
                  atol
                        0.1, optional
```

**Args:** 

Stop method when cct a absolute tolerance (K) is reached.

#### max iter

```
_CCT_MAX_ITER, optional
```

Maximum number of iterations used newton-raphson methods.

#### lut\_resolution\_reduction\_factor

\_CCT\_LUT\_RESOLUTION\_REDUCTION\_FACTOR, optional

Number of times the interval spanned by the adjacent Tc in a search or lut method is downsampled (the search process will then start again)

## split calculation at N

```
_CCT_SPLIT_CALC_AT_N, optional
```

Split calculation when xyzw.shape[0] > split\_calculation\_at\_N.

Splitting speeds up the calculation. If None: no splitting is done.

#### lut

None, optional

Look-Up-Table with Ti, u,v,u',v',u",v",slope values of Planckians.

## Options:

- None: defaults to the lut specified in
- \_CCT\_LUT[first\_guess\_mode]['lut\_type\_def'].
- list (lut,lut\_kwargs): use this pre-calculated lut

(add additional kwargs for the lut\_generator\_fcn(), defaults to None if omitted)

- tuple: must be key (label) in :luts\_dict: (pre-calculated dict of luts),

if not: then a new lut will be generated from scratch using the info in the tuple.

- str: must be key (label) in :luts\_dict: (pre-calculated dict of luts)
- ndarray [Nx1]: list of luts for which to generate a lut
- ndarray [Nxn] with n>3: pre-calculated lut (last col must contain slope of the isotemperature lines).

## luts dict

None, optional

Dictionary of pre-calculated luts for various espaces and cmf sets.

Must have structure luts\_dict[cspace][cieobs][lut\_label] with the lut part of a two-element list [lut, lut\_kwargs]. It must contain at the top-level a key 'wl' containing the wavelengths of the Planckians used to generate the luts in this dictionary.

If None: luts\_dict defaults to \_CCT\_LUT[first\_guess\_mode]['luts']

## cspace

\_CCT\_SPACE, optional

Color space to do calculations in.

# Options:

- cspace string:
  - e.g. 'Yuv60' for use with luxpy.colortf()
- tuple with forward (i.e. xyz\_to..) [and backward (i.e. ..to\_xyz)] functions

```
(and an optional string describing the cspace):
```

e.g. (forward, backward) or (forward, backward, cspace string) or (forward, cspace string)

- dict with keys: 'fwtf' (foward), 'bwtf' (backward) [, optional: 'str' (cspace string)]

Note: if the backward tf is not supplied, optimization in cct\_to\_xyz() is done in the CIE 1976 u'v' diagram

## cspace\_kwargs

```
_CCT_CSPACE_KWARGS, optional
```

Parameter nested dictionary for the forward and backward transforms.

## ignore\_wl\_diff

False, optional

When getting a lut from the dictionary, if differences are detected in the wavelengts of the lut and the ones used to calculate any plankcians then a new lut should be generated. Seting this to True ignores these differences and proceeds anyway.

## first\_guess\_mode

'robertson1968', optional

Method used to get an approximate (first guess) estimate of the cct,

after which the newton-raphson method is started.

Options: 'robertson1968', 'ohno2014', 'zhang2019'

## fgm\_kwargs

Dict with keyword arguments for the selected first guess mode.

#### use fast duv

```
_CCT_FAST_DUV, optional
```

If True: use a fast estimator of the Duv

(one that avoids calculation of Planckians and uses the former best estimate's u,v coordinates. This method is accurate enough when the atol is small enough -> as long as abs(T-T\_former)<=1K

the Duv estimate should be ok.)

## **Returns:**

## returns

```
ndarray with:
```

```
cct: out == 'cct' (or 1)
duv: out == 'duv' (or -1)
cct, duv: out == 'cct,duv' (or 2)
[cct,duv]: out == "[cct,duv]" (or -2)
```

**Note:** 1. Out-of-lut (of first\_guess\_mode) CCTs are encoded as negative CCTs (with as absolute value the value of the closest CCT from the lut.)

**References:** 1. Li, C., Cui, G., Melgosa, M., Ruan, X., Zhang, Y., Ma, L., Xiao, K., & Luo, M. R. (2016). Accurate method for computing correlated color temperature. Optics Express, 24(13), 14066–14078.

2. Robertson, A. R. (1968). Computation of Correlated Color Temperature and Distribution Temperature. Journal of the Optical Society of America, 58(11), 1528–1535.

```
luxpy.color.cct.xyz_to_cct_zhang2019(xyzw, cieobs='1931_2', out='cct', is_uv_input=False, wl=None,
                                               atol=0.1, rtol=1e-05, force_tolerance=True,
                                               tol method='newton-raphson', lut resolution reduction factor=4,
                                               split_calculation_at_N=25, max_iter=10, cspace='Yuv60',
                                               cspace_kwargs={'bwtf': {}, 'fwtf': {}}, lut=None, luts_dict=None,
                                               ignore_wl_diff=False, use_fast_duv=True, **kwargs)
      Convert XYZ tristimulus values to correlated color temperature (CCT) and Duv(distance above (> 0) or below (
      < 0) the Planckian locus) using the golden-ratio search method described in Zhang et al. (2019).
      Args:
                  XYZW
                        ndarray of tristimulus values
                  cieobs
                        luxpy._CIEOBS, optional
                        CMF set used to calculated xyzw.
                  out
                        'cct' (or 1), optional
                        Determines what to return.
                        Other options: 'duv' (or -1), 'cct,duv' (or 2), "[cct,duv]" (or -2)
                  is_uv_input
                        False, optional
                        If True: xyzw contain uv input data, not xyz data!
                  wl
                        None, optional
                        Wavelengths used when calculating Planckian radiators.
                        If None: use same wavelengths as CMFs in :cieobs:.
                  rtol
                        1e-5, float, optional
                        Stop search when cct a relative tolerance is reached.
                        The relative tolerance is calculated as dCCT/CCT_est,
                        with CCT_est the current intermediate estimate in the
                        search and with dCCT the difference between
                        the present and former estimates.
                  atol
                        0.1, optional
                        Stop search when cct a absolute tolerance (K) is reached.
                  force_tolerance
                        True, optional
                        If False: search only using the list of CCTs in the used lut.
                              Only one loop of the full algorithm is performed.
                              Accuracy depends on CCT of test source and the location
                              and spacing of the CCTs in the list.
                        If True: search will use adjacent CCTs to test source to create a new LUT,
                              (repeat the algoritm at higher resolution, progessively zooming in
                                    toward the ground-truth) for tol_method == 'cl'; when
```

tol\_method == 'nr' a newton-raphson method is used.

Because the CCT for multiple source is calculated in one go, the atol and rtol values have to be met for all!

## tol\_method

'newton-raphson', optional

(Additional) method to try and achieve set tolerances.

#### Options:

- 'cl', 'cascading-lut': use increasingly higher CCT-resolution to 'zoom-in' on the ground-truth.
- 'nr', 'newton-raphson': use the method as described in Li, 2016.

#### lut\_resolution\_reduction\_factor

## \_CCT\_LUT\_RESOLUTION\_REDUCTION\_FACTOR, optional

Number of times the interval spanned by the adjacent Tc in a search or lut method is downsampled (the search process will then start again)

#### max iter

\_CCT\_MAX\_ITER, optional

Maximum number of iterations used by the cascading-lut or newton-raphson methods.

# split\_calculation\_at\_N

\_CCT\_SPLIT\_CALC\_AT\_N, optional

Split calculation when xyzw.shape[0] > split\_calculation\_at\_N.

Splitting speeds up the calculation. If None: no splitting is done.

## lut

None, optional

Look-Up-Table with Ti, u,v,u',v',u'',v'',slope values of Planckians.

## Options:

- None: defaults to the lut specified in
- \_CCT\_LUT['zhang2019']['lut\_type\_def'].
- list (lut,lut\_kwargs): use this pre-calculated lut

(add additional kwargs for the lut\_generator\_fcn(), defaults to None if omitted)

- tuple: must be key (label) in :luts\_dict: (pre-calculated dict of luts),
  - if not: then a new lut will be generated from scratch using the info in the tuple.
- str: must be key (label) in :luts\_dict: (pre-calculated dict of luts)
- ndarray [Nx1]: list of luts for which to generate a lut
- ndarray [Nxn] with n>3: pre-calculated lut (last col must contain slope of the isotemperature lines).

## luts\_dict

None, optional

Dictionary of pre-calculated luts for various espaces and cmf sets.

Must have structure luts\_dict[cspace][cieobs][lut\_label] with the lut part of a two-element list [lut, lut\_kwargs]. It must contain at the top-level a key 'wl' containing the wavelengths of the Planckians used to generate the luts in this dictionary.

```
If None: luts_dict defaults to _CCT_LUT['zhang2019']['luts']
cspace
      _CCT_SPACE, optional
      Color space to do calculations in.
      Options:
                  - cspace string:
                         e.g. 'Yuv60' for use with luxpy.colortf()
                  - tuple with forward (i.e. xyz_to..) [and backward (i.e. ..to_xyz)]
                  functions
                         (and an optional string describing the cspace):
                               e.g. (forward, backward) or (forward, backward, cspace
                               string) or (forward, cspace string)
                  - dict with keys: 'fwtf' (foward), 'bwtf' (backward) [, optional: 'str'
                  (cspace string)]
            Note: if the backward tf is not supplied, optimization in cct_to_xyz() is done in
            the CIE 1976 u'v' diagram
cspace_kwargs
      _CCT_CSPACE_KWARGS, optional
      Parameter nested dictionary for the forward and backward transforms.
ignore wl diff
      False, optional
      When getting a lut from the dictionary, if differences are
      detected in the wavelengts of the lut and the ones used to calculate any
      plankcians then a new lut should be generated. Seting this to True ignores
      these differences and proceeds anyway.
use_fast_duv
      _CCT_FAST_DUV, optional
      If True: use a fast estimator of the Duv
            (one that avoids calculation of Planckians and uses the former
                  best estimate's u,v coordinates. This method is accurate enough
                  when the atol is small enough -> as long as abs(T-T_former) \le 1K
                  the Duv estimate should be ok.)
returns
      ndarray with:
            cct: out == 'cct' (or 1)
            duv: out == 'duv' (or -1)
            cct, duv: out == 'cct,duv' (or 2)
            [cct,duv]: out == "[cct,duv]" (or -2)
```

Note: 1. Out-of-lut CCTs are encoded as negative CCTs (with as absolute value the value of the closest CCT from the lut.)

**References:** 1. Zhang, F. (2019). High-accuracy method for calculating correlated color temperature with a lookup table based on golden section search. Optik, 193, 163018.

2. Li, C., Cui, G., Melgosa, M., Ruan, X., Zhang, Y., Ma, L., Xiao, K., & Luo, M. R. (2016). Accurate method for computing correlated color temperature. Optics Express, 24(13), 14066–14078.

**Returns:** 

```
luxpy.color.cct.xyz_to_cct_fibonacci(xyzw, cieobs='1931_2', out='cct', is_uv_input=False, wl=None,
                                               atol=0.1, rtol=1e-05, force_tolerance=True,
                                               tol method='newton-raphson', lut resolution reduction factor=4,
                                               split_calculation_at_N=25, max_iter=10, cspace='Yuv60',
                                               cspace_kwargs={'bwtf': {}, 'fwtf': {}}, lut=None, luts_dict=None,
                                               ignore wl diff=False, use fast duv=True, **kwargs)
      Convert XYZ tristimulus values to correlated color temperature (CCT) and Duv(distance above (> 0) or below (
      < 0) the Planckian locus) using a Fibonacci search.
      Args:
                  XYZW
                        ndarray of tristimulus values
                  cieobs
                        luxpy._CIEOBS, optional
                        CMF set used to calculated xyzw.
                  out
                        'cct' (or 1), optional
                        Determines what to return.
                        Other options: 'duv' (or -1), 'cct,duv' (or 2), "[cct,duv]" (or -2)
                  is_uv_input
                        False, optional
                        If True: xyzw contain uv input data, not xyz data!
                  wl
                        None, optional
                        Wavelengths used when calculating Planckian radiators.
                        If None: use same wavelengths as CMFs in :cieobs:.
                  rtol
                        1e-5, float, optional
                        Stop search when cct a relative tolerance is reached.
                        The relative tolerance is calculated as dCCT/CCT est,
                        with CCT_est the current intermediate estimate in the
                        search and with dCCT the difference between
                        the present and former estimates.
                  atol
                        0.1, optional
                        Stop search when cct a absolute tolerance (K) is reached.
                  force_tolerance
                        True, optional
                        If False: search only using the list of CCTs in the used lut.
                              Only one loop of the full algorithm is performed.
                              Accuracy depends on CCT of test source and the location
                              and spacing of the CCTs in the list.
                        If True: search will use adjacent CCTs to test source to create a new LUT,
                              (repeat the algoritm at higher resolution, progessively zooming in
```

toward the ground-truth) for tol\_method == 'cl'; when

tol\_method == 'nr' a newton-raphson method is used.

Because the CCT for multiple source is calculated in one go, the atol and rtol values have to be met for all!

## tol\_method

'newton-raphson', optional

(Additional) method to try and achieve set tolerances.

#### Options:

- 'cl', 'cascading-lut': use increasingly higher CCT-resolution to 'zoom-in' on the ground-truth.
- 'nr', 'newton-raphson': use the method as described in Li, 2016.

#### lut\_resolution\_reduction\_factor

\_CCT\_LUT\_RESOLUTION\_REDUCTION\_FACTOR, optional

Number of times the interval spanned by the adjacent Tc in a search or lut method is downsampled (the search process will then start again)

#### max iter

\_CCT\_MAX\_ITER, optional

Maximum number of iterations used by the cascading-lut or newton-raphson methods.

# split\_calculation\_at\_N

\_CCT\_SPLIT\_CALC\_AT\_N, optional

Split calculation when xyzw.shape[0] > split\_calculation\_at\_N.

Splitting speeds up the calculation. If None: no splitting is done.

## lut

None, optional

Look-Up-Table with Ti, u,v,u',v',u'',v'',slope values of Planckians.

## Options:

- None: defaults to the lut specified in \_CCT\_LUT['fibonacci']['lut\_type\_def'].
- list (lut,lut\_kwargs): use this pre-calculated lut (add additional kwargs for the lut\_generator\_fcn(), defaults to None if omitted)
- tuple: must be key (label) in :luts\_dict: (pre-calculated dict of luts),
   if not: then a new lut will be generated from scratch using the info in the tuple.
- str: must be key (label) in :luts dict: (pre-calculated dict of luts)
- ndarray [Nx1]: list of luts for which to generate a lut
- ndarray [Nxn] with n>3: pre-calculated lut (last col must contain slope of the isotemperature lines).

## luts\_dict

None, optional

Dictionary of pre-calculated luts for various espaces and cmf sets.

Must have structure luts\_dict[cspace][cieobs][lut\_label] with the lut part of a two-element list [lut, lut\_kwargs]. It must contain at the top-level a key 'wl' containing the wavelengths of the Planckians used to generate the luts in this dictionary.

If None: luts\_dict defaults to \_CCT\_LUT['fibonacci']['luts']

#### cspace

```
__CCT_SPACE, optional
Color space to do calculations in.
Options:

- cspace string:
- e.g. 'Yuv60' for use with luxpy.colortf()
- tuple with forward (i.e. xyz_to..) [and backward (i.e. ..to_xyz)]
functions

(and an optional string describing the cspace):
- e.g. (forward, backward) or (forward, backward, cspace
string) or (forward, cspace string)
- dict with keys: 'fwtf' (foward), 'bwtf' (backward) [, optional: 'str'
(cspace string)]

Note: if the backward tf is not supplied, optimization in cct_to_xyz() is done in
the CIE 1976 u'v' diagram
```

## cspace\_kwargs

```
_CCT_CSPACE_KWARGS, optional
```

Parameter nested dictionary for the forward and backward transforms.

## ignore\_wl\_diff

False, optional

When getting a lut from the dictionary, if differences are detected in the wavelengts of the lut and the ones used to calculate any plankcians then a new lut should be generated. Seting this to True ignores these differences and proceeds anyway.

## use\_fast\_duv

```
_CCT_FAST_DUV, optional

If True: use a fast estimator of the Duv

(one that avoids calculation of Planckians and uses the former

best estimate's u,v coordinates. This method is accurate enough

when the atol is small enough -> as long as abs(T-T_former)<=1K

the Duv estimate should be ok.)
```

#### **Returns:**

## returns

```
ndarray with:
     cct: out == 'cct' (or 1)
     duv: out == 'duv' (or -1)
     cct, duv: out == 'cct,duv' (or 2)
     [cct,duv]: out == "[cct,duv]" (or -2)
```

**Note:** 1. Out-of-lut CCTs (or close to) are encoded as negative CCTs (with as absolute value the value of the closest CCT from the lut.)

```
luxpy.color.cct.xyz_to_cct(xyzw, mode='robertson1968', cieobs='1931_2', out='cct', is_uv_input=False,
                                  wl=None, atol=0.1, rtol=1e-10, force_tolerance=True,
                                  tol method='newton-raphson', lut resolution reduction factor=4,
                                  split_calculation_at_N=25, max_iter=10, cspace='Yuv60',
                                  cspace_kwargs={'bwtf': {}}, 'fwtf': {}}, lut=None, luts_dict=None,
                                  ignore_wl_diff=False, duv_parabolic_threshold=0.002,
                                  first guess mode='robertson1968', fgm kwargs={}, use fast duv=True,
                                  **kwargs)
      Convert XYZ tristimulus values to correlated color temperature (CCT) and Duv (distance above (>0) or below
      (<0) the Planckian locus) using a number of modes (methods).
      Args:
                  XYZW
                        ndarray of tristimulus values
                  mode
                        'robertson1968', optional
                        String with name of method to use.
                        Options: 'robertson1968', 'ohno2014', 'li2016', 'zhang2019', 'fibonacci',
                              (also, but see note below: 'mcamy1992', 'hernandez1999')
                        Note: first_guess_mode for li2016 can also be specified using a ':' separator,
                              e.g. 'li2016:robertson1968'
                  cieobs
                        luxpy._CIEOBS, optional
                        CMF set used to calculated xyzw.
                  out
                        'cct' (or 1), optional
                        Determines what to return.
                        Other options: 'duv' (or -1), 'cct,duv' (or 2), "[cct,duv]" (or -2)
                  is_uv_input
                        False, optional
                        If True: xyzw contain uv input data, not xyz data!
                  wl
                        None, optional
                        Wavelengths used when calculating Planckian radiators.
                        If None: use same wavelengths as CMFs in :cieobs:.
                  rtol
                        1e-5, float, optional
                        Stop search when cct a relative tolerance is reached.
                        The relative tolerance is calculated as dCCT/CCT_est,
                        with CCT est the current intermediate estimate in the
                        search and with dCCT the difference between
                        the present and former estimates.
                  atol
                        0.1, optional
                        Stop search when cct a absolute tolerance (K) is reached.
```

#### force tolerance

True, optional

If False: search only using the list of CCTs in the used lut.

Only one loop of the full algorithm is performed.

Accuracy depends on CCT of test source and the location and spacing of the CCTs in the list.

If True: search will use adjacent CCTs to test source to create a new LUT,

(repeat the algoritm at higher resolution, progessively zooming in

toward the ground-truth) for tol\_method == 'cl'; when

tol\_method == 'nr' a newton-raphson method is used.

Because the CCT for multiple source is calculated in one go,

the atol and rtol values have to be met for all!

## tol method

'newton-raphson', optional

(Additional) method to try and achieve set tolerances.

## Options:

- 'cl', 'cascading-lut': use increasingly higher CCT-resolution to 'zoom-in' on the ground-truth. (not for mode == 'li2016')
- 'nr', 'newton-raphson': use the method as described in Li, 2016.

## lut\_resolution\_reduction\_factor

CCT LUT RESOLUTION REDUCTION FACTOR, optional

Number of times the interval spanned by the adjacent Tc in a search or lut method is downsampled (the search process will then start again)

# max iter

\_CCT\_MAX\_ITER, optional

Maximum number of iterations used by the cascading-lut or newton-raphson methods.

## split\_calculation\_at\_N

\_CCT\_SPLIT\_CALC\_AT\_N, optional

Split calculation when xyzw.shape[0] > split calculation at N.

Splitting speeds up the calculation. If None: no splitting is done.

#### lut

None, optional

Look-Up-Table with Ti, u,v,u',v',u'',v'',slope values of Planckians.

## Options:

- None: defaults to the lut specified in \_CCT\_LUT[mode]['lut\_type\_def'].
- list (lut,lut\_kwargs): use this pre-calculated lut (add additional kwargs for the lut\_generator\_fcn(), defaults to None if omitted)
- tuple: must be key (label) in :luts\_dict: (pre-calculated dict of luts),
   if not: then a new lut will be generated from scratch using the info in the tuple.
- str: must be key (label) in :luts\_dict: (pre-calculated dict of luts)
- ndarray [Nx1]: list of luts for which to generate a lut

- ndarray [Nxn] with n>3: pre-calculated lut (last col must contain slope of the isotemperature lines).

## luts\_dict

```
None, optional
```

Dictionary of pre-calculated luts for various espaces and cmf sets.

Must have structure luts\_dict[cspace][cieobs][lut\_label] with the

lut part of a two-element list [lut, lut\_kwargs]. It must contain at the top-level a key 'wl' containing the wavelengths of the

Planckians used to generate the luts in this dictionary.

If None: the default dict for the mode is used

```
(e.g. _CCT_LUT['ohno2014']['lut_type_def'], for mode=='ohno2014').
```

# cspace

```
_CCT_SPACE, optional
```

Color space to do calculations in.

Options:

- cspace string:

e.g. 'Yuv60' for use with luxpy.colortf()

- tuple with forward (i.e. xyz\_to..) [and backward (i.e. ..to\_xyz)] functions

(and an optional string describing the cspace):

e.g. (forward, backward) or (forward, backward, cspace string) or (forward, cspace string)

- dict with keys: 'fwtf' (foward), 'bwtf' (backward) [, optional: 'str' (cspace string)]

Note: if the backward tf is not supplied, optimization in cct\_to\_xyz() is done in the CIE 1976 u'v' diagram

#### cspace\_kwargs

```
_CCT_CSPACE_KWARGS, optional
```

Parameter nested dictionary for the forward and backward transforms.

## ignore\_wl\_diff

False, optional

When getting a lut from the dictionary, if differences are detected in the wavelengts of the lut and the ones used to calculate any plankcians then a new lut should be generated. Seting this to True ignores these differences and proceeds anyway.

## duv\_parabolic\_threshold

```
0.002, optional (cfr. mode == 'ohno2014')
```

Threshold for use of the parabolic solution

(if larger then use parabolic, else use triangular solution)

## first\_guess\_mode

```
'robertson1968', optional (cfr. mode == 'li2016')
```

Method used to get an approximate (first guess) estimate of the cct,

after which the newton-raphson method is started.

Options: 'robertson1968', 'ohno2014', 'zhang2019'

#### use fast duv

CCT FAST DUV, optional

If True: use a fast estimator of the Duv

(one that avoids calculation of Planckians and uses the former

best estimate's u,v coordinates. This method is accurate enough when the atol is small enough -> as long as abs(T-T\_former)<=1K  $\,$ 

the Duv estimate should be ok.)

#### **Returns:**

#### returns

ndarray with:

```
cct: out == 'cct' (or 1)
duv: out == 'duv' (or -1)
cct, duv: out == 'cct,duv' (or 2)
[cct,duv]: out == "[cct,duv]" (or -2)
```

**Note:** 1. Using the 'mcamy1992' and 'hernandez1999' options will result in additional errors when cieobs is different from '1931\_2' as for these options the CCT is determined using a fixed set of equations optimized for the 1931 2° CMFs!! The only impact will be on the calculation of the Duv from the CCT. That does depend on the settings of cieobs and cspace! Change at own discretion. 2. Out-of-lut CCTs are encoded as negative CCTs (with as absolute value the value of the closest CCT from the lut.)

**References:** 1. Robertson, A. R. (1968). Computation of Correlated Color Temperature and Distribution Temperature. Journal of the Optical Society of America, 58(11), 1528–1535.

- 2. Ohno Y. Practical use and calculation of CCT and Duv. Leukos. 2014 Jan 2;10(1):47-55.
- 3. Zhang, F. (2019). High-accuracy method for calculating correlated color temperature with a lookup table based on golden section search. Optik, 193, 163018.
- 3. Li, C., Cui, G., Melgosa, M., Ruan, X., Zhang, Y., Ma, L., Xiao, K., & Luo, M. R. (2016). Accurate method for computing correlated color temperature. Optics Express, 24(13), 14066–14078.
- 4. McCamy, Calvin S. (April 1992). "Correlated color temperature as an explicit function of chromaticity coordinates". Color Research & Application. 17 (2): 142–144.
- 5. Hernández-Andrés, Javier; Lee, RL; Romero, J (September 20, 1999). Calculating Correlated Color Temperatures Across the Entire Gamut of Daylight and Skylight Chromaticities. Applied Optics. 38 (27), 5703–5709. P

Convert correlated color temperature (550 K  $\leq$  CCT  $\leq$  1e11 K) and Duv (distance above (>0) or below (<0) the Planckian locus) to XYZ tristimulus values.

Finds xyzw\_estimated by determining the iso-temperature line

(= line perpendicular to the Planckian locus):

Option 1 (fastest):

First, the angle between the coordinates corresponding to ccts and ccts-cct\_offset are calculated, then 90° is added, and finally

the new coordinates are determined, while taking sign of duv into account.

Option 2 (slowest, about 55% slower):

Calculate the slope of the iso-T-line directly using the Planckian spectrum and its derivative.

Args:

```
ccts
                  ndarray [N,1] of cct values
            duv
                  None or ndarray [N,1] of duv values, optional
                  Note that duv can be supplied together with cct values in :ccts:
                  as ndarray with shape [N,2].
            cct offset
                  None, optional
                  If None: use option 2 (direct iso-T slope calculation, more accurate,
                        but slower: about 1.55 slower)
                  else: use option 1 (estimate slope from 90° + angle of small cct_offset)
            cieobs
                  luxpy._CIEOBS, optional
                  CMF set used to calculated xyzw.
            wl
                  None, optional
                  Wavelengths used when calculating Planckian radiators.
                  If None: use same wavelengths as CMFs in :cieobs:.
            cspace
                  CCT SPACE, optional
                  Color space to do calculations in.
                  Options:
                               - cspace string:
                                     e.g. 'Yuv60' for use with luxpy.colortf()
                               - tuple with forward (i.e. xyz_to..) [and backward (i.e. ..to_xyz)]
                               functions
                                     (and an optional string describing the cspace):
                                           e.g. (forward, backward) or (forward, backward, cspace
                                           string) or (forward, cspace string)
                               - dict with keys: 'fwtf' (foward), 'bwtf' (backward) [, optional: 'str'
                               (cspace string)]
                        Note: if the backward tf is not supplied, optimization in cct_to_xyz() is done in
                        the CIE 1976 u'v' diagram
            cspace_kwargs
                  _CCT_CSPACE_KWARGS, optional
                  Parameter nested dictionary for the forward and backward transforms.
Returns:
```

returns

ndarray with estimated XYZ tristimulus values

**Note:** 1. If duv is not supplied (:ccts:.shape is (N,1) and :duv: is None), source is assumed to be on the Planckian locus. 2. Minimum CCT is 550 K (lower than 550 K, some negative Duv values will result in coordinates outside of the Spectrum Locus !!!)

```
luxpy.color.cct.calculate_lut(ccts, cieobs, wl=None, lut_vars=['T', 'uv', 'uvpp', 'uvpp', 'iso-T-slope'],
                                       cspace='Yuv60', cspace_kwargs={'bwtf': {}, 'fwtf': {}})
      Function that calculates a LUT for the specified calculation method for the input ccts. Calculation is performed
      for CMF set specified in cieobs and in the chromaticity diagram in cspace.
      Args:
                  ccts
                         ndarray [Nx1] or str
                         list of ccts for which to (re-)calculate the LUTs.
                         If str, ccts contains path/filename.dat to list.
                  cieobs
                         None or str, optional
                         str specifying cmf set.
                  wl
                         None, optional
                         Generate luts based on Planckians with wavelengths (range).
                         If None: use same wavelengths as CMFs in :cieobs:.
                  lut_vars
                         ['T','uv','uvp','uvpp','iso-T-slope'], optional
                         Data the lut should contain. Must follow this order
                         and minimum should be ['T']
                  cspace
                         _CCT_SPACE, optional
                         Color space to do calculations in.
                         Options:
                                     - cspace string:
                                           e.g. 'Yuv60' for use with luxpy.colortf()
                                     - tuple with forward (i.e. xyz_to..) [and backward (i.e. ..to_xyz)]
                                     functions
                                            (and an optional string describing the cspace):
                                                  e.g. (forward, backward) or (forward, backward, cspace
                                                  string) or (forward, cspace string)
                                     - dict with keys: 'fwtf' (foward), 'bwtf' (backward) [, optional: 'str'
                                     (cspace string)]
                               Note: if the backward tf is not supplied, optimization in cct_to_xyz() is done in
                               the CIE 1976 u'v' diagram
                  cspace_kwargs
                         _CCT_CSPACE_KWARGS, optional
                         Parameter nested dictionary for the forward and backward transforms.
      Returns:
                  returns
                              lut
                                   ndarray with T, u, v, u', v', u", v", slope (note ':1st deriv., ":2nd deriv.).
```

```
luxpy.color.cct.generate_luts(types=[None], seamless_stitch=True, fallback_unit='K-1', fallback_n=50,
                                       cct min=450, cct max=10000000000000, lut file=None, load=False,
                                       lut path='C:\\Users\\u0032318\\OneDrive - KU
                                       Leuven\\Documents\\Github\\luxpy\\luxpy\\data\\cctluts\\', save_luts=True,
                                       wl=None, cieobs=['1931_2'], lut_vars=['T', 'uv', 'uvp', 'uvp', 'iso-T-slope'],
                                       cspace=['Yuv60'], cspace_kwargs=[{'fwtf': {}}, 'bwtf': {}}], verbosity=0,
                                       lut generator fcn=<function generate lut>, lut generator kwargs={})
      Generate a number of luts and store them in a nested dictionary. Structure: lut[cspace][cieobs][lut type].
      Args:
                  lut_file
                         None, optional
                         string specifying the filename to save the lut (as .pkl) to.
                         If None: don't save anything when generated (i.e. load==False).
                  load
                         True, optional
                         If True: load previously generated dictionary.
                         If False: generate from scratch.
                  lut_path
                         _CCT_LUT_PATH, optional
                         Path to file.
                  wl
                         None, optional
                         Wavelength for Planckian spectrum generation.
                         If None: use same wavelengths as CMFs in :cieobs:.
                  cieobs
                         [_CIEOBS] or list, optional
                         Generate a LUT for each one in the list.
                         If None: generate for all cmfs in _CMF.
                  types
                         [None], optional
                         List of lut specifiers of format [(Tmin,Tmax,Tinterval,unit),...]
                         If units are in MK-1 then the range is also!
                               Unit options are:
                               - '%': equal relative Tc spacing (in %, cfr. (Ti+1 - Ti-1)/Ti-1).
                               - 'K' equal absolute Tc spacing (in K, cfr. (Ti+1 - Ti-1).
                               - '%-1': equal relative reciprocal Tc (MK-1 = mired).
                               - 'K-1': equal absolute reciprocal Tc (MK-1 = mired).
                         If the last element of the list is a bool, then the way the different
                         lists of Tcs generated by each list element can be set. If True:
                         the Tcs will be 'seamlessly' stitched together (this does have an
                         an impact on the min-max range of each Tc set) so that there are no
                         discontinuities in terms of the intervals.
                  seamless_stitch
```

True, optional

When stitching (creating) LUTs composed of several CCT ranges with different intervals, these do not always 'match' well, in the sense that discontinuities might be generated. This can be avoided (at the expense of possibly slightly changed ranges)

by setting the :seamless\_stitch: argument to True. Is overriden when the last element in the lut list is a boolean.

## cct\_max

\_CCT\_MAX, optional

Limit Tc's to a maximum value of cct max

## cct\_min

\_CCT\_MIN, optional

Limit Tc's to a minimum value of cct max

#### fallback unit

\_CCT\_FALLBACK\_UNIT, optional

Unit to fall back on when the input unit in tc4 (of first list) is 'au'. As there is no common distancing of the unit types ['K','%','%-1','K-1'] the Tc's are generated by dividing the min-max range into a number of divisions, specified by the negative 3 element (or when positive or NaN, the number of divisions is set by :fallback\_divisions:)

## fallback n

\_CCT\_FALLBACK\_N, optional

Number of divisions the min-max range is divided into, in the fallback case in which unit=='au' and the 3e 4-vector element is NaN or positive.

# lut\_vars

['T','uv','uvp','uvpp','iso-T-slope'], optional Data the lut should contain. Must follow this order and minimum should be ['T']

#### cspace\_cspace\_kwargs

Lists with the cspace and cspace\_kwargs for which luts will be generated. Default is single chromaticity diagram in \_CCT\_CSPACE.

#### verbosity

0, optional

If > 0: give some intermediate feedback while generating luts.

#### lut generator fcn

generate lut, optional

Lets a user specify his own lut generation function (must output a list of 1 lut). Default is the general function. There is a specific one for Ohno's 2014 method as that one requires a different correction factor for each lut for the parabolic solutions. This optimized value is specified in the

for each lut for the parabolic solutions. This optimized value is specified in the second list index. (see \_generate\_lut\_ohno2014()).

#### lut\_generator\_kwargs

{}, optional

Dict with keyword arguments specific to the (user) lut\_generator\_fcn.

(e.g. {'f\_corr':0.9991} for \_generate\_lut\_ohno2014())

#### **Returns:**

#### dict

Dictionary with luts for the specified mode, cieobs(s) and cspace(s).

Structure: lut[cspace][cieobs][lut type]

At the upper dict level there is also a key 'wl' which contains a dict with keys the cieobs and with values the wavelengths used to calculate the Planckians for each lut for the specified cieobs; as well as a key with the lut\_vars

The luts contains as data the variables as specified in lut\_vars:

- T: (in K)
- uv: chromaticity coordinates of planckians
- uvp: chromaticity coordinates of 1st derivative of the planckians.
- uvpp: chromaticity coordinates of 2nd derivative of the planckians.
- iso-T-slope: slope of isotemperature lines (calculated as in Robertson, 1968).

luxpy.color.cct.get\_tcs4(tc4, uin=None, seamless stitch=True, fallback unit='K-1', fallback n=50) Get an idarray of Tc's obtained from a list or tuple of tc4 4-vectors. **Args:** 

tc4

list or tuple of 4-vectors.

e.g. (tc4\_1, tc4\_2, tc4\_3,...) or (tc4\_1, tc4\_2, tc4\_3,..., bool::seamless\_stitch)

When the last element of the list/tuple is a bool, then this specifies

how the Tc arrays generated for each of the 4-vector elements need to be stitched together. This overrides the seamless\_stitch input argument.

Vector elements are:

[Tmin, Tmax inclusive, Tinterval(or number of intervals), unit]

Unit specifies unit of the Tc interval, i.e. it determines the

type of scale in which the spacing of the Tc are done.

Unit options are:

- '%': equal relative Tc spacing (in %, cfr. (Ti+1 Ti-1)/Ti-1).
- 'K' equal absolute Tc spacing (in K, cfr. (Ti+1 Ti-1).
- '%-1': equal relative reciprocal Tc (MK-1 = mired).
- 'K-1': equal absolute reciprocal Tc (MK-1 = mired).

If the 'interval' element is negative, it actually represents the number of intervals between Tmin, Tmax (included).

# uin

None, optional

Unit of input Tmin, Tmax (by default it is assumed to be the same as the scale 'unit').

#### seamless stitch

True, optional

Determines how the Tc arrays generated for each of the 4-vector elements are stitched together. Is overriden by the presence of a bool as last list/tuple element in :tc4:.

For a seamless stitch, all units for all 4-vectors should be the same!!

## fallback\_unit

```
_CCT_FALLBACK_UNIT, optional
```

Unit to fall back on when the input unit in tc4 (of first list) is 'au'. As there is no common distancing of the unit types ['K','%','%-1','K-1'] the Tc's are generated by dividing the min-max range into a number of divisions, specified by the negative 3 element (or when positive or NaN, the number of divisions is set by :fallback\_divisions:)

## fallback n

# \_CCT\_FALLBACK\_N, optional

Number of divisions the min-max range is divided into, in the fallback case in which unit=='au' and the 3e 4-vector element is NaN or positive.

#### **Returns:**

tcs

ndarray with Tcs

Get an ndarray LUT from various sources.

# **Args:**

#### lut

Look-Up-Table with Ti, u,v,u',v',u",v'',slope values of Planckians, or whatever quantities are specified in lut\_vars ('T','uv' is always part of the lut). Options:

- list: must have two elements: [lut,lut\_kwargs]
- None: lut from luts\_dict with lut\_type\_def as key
- str: lut from luts\_dict at key :lut:
- ndarray [Nxn, with n>1]: precalculated lut (only processing will be to keep it with cct\_min-cct\_max range)
- ndarray [Nx1]: list of Tc's from which a new lut will be calculated.
- tuple of 4-vectors: used as key in luts\_dict or to generate new lut from scratch 4-vector info:
  - + format: e.g. (tc4\_1, tc4\_2, tc4\_3,...) or (tc4\_1, tc4\_2, tc4\_3,..., bool::seamless stitch)
  - + When the last element of the list/tuple is a bool, then this specifies how the Tc arrays generated for each of the 4-vector elements need to be

stitched together. This overrides the seamless $\_$ stitch input argument.

+ Vector elements are:

[Tmin, Tmax inclusive, Tinterval(or number of intervals), unit]

Unit specifies unit of the Tc interval, i.e. it determines the

type of scale in which the spacing of the Tc are done. Unit options are:

- '%': equal relative Tc spacing (in %, cfr. (Ti+1 Ti-1)/Ti-1).
- 'K' equal absolute Tc spacing (in K, cfr. (Ti+1 Ti-1).
- '%-1': equal relative reciprocal Tc (MK-1 = mired).
- 'K-1': equal absolute reciprocal Tc (MK-1 = mired).

If the 'interval' element is negative, it actually represents the number of intervals between Tmin, Tmax (included).

#### uin

None, optional

Unit of input Tmin, Tmax (by default it is assumed to be the same as the scale 'unit') in Tc generation from tuple.

## seamless\_stitch

True, optional

Determines how the Tc arrays generated for each of the 4-vector elements are stitched together. Is overriden by the presence of a bool as last list/tuple element in :tc4:.

For a seamless stitch, all units for all 4-vectors should be the same!!

#### cct max

\_CCT\_MAX, optional

Limit Tc's to a maximum value of cct\_max

#### cct min

\_CCT\_MIN, optional

Limit Tc's to a minimum value of cct\_max

#### fallback unit

```
_CCT_FALLBACK_UNIT, optional
```

Unit to fall back on when the input unit in tc4 (of first list) is 'au'. As there is no common distancing of the unit types ['K','%','%-1','K-1'] the Tc's are generated by dividing the min-max range into a number of divisions, specified by the negative 3 element (or when positive or NaN, the number of divisions is set by :fallback\_divisions:)

#### fallback n

\_CCT\_FALLBACK\_N, optional

Number of divisions the min-max range is divided into, in the fallback case in which unit=='au' and the 3e 4-vector element is NaN or positive.

## resample\_tc4\_array

False, optional

If False: do not resample Tc's of an ndarray input for tc4 else: divide min-max range in fallback\_n intervals. Uses fallback\_unit to determine the scale for the resampling.

#### wl

None, optional

Wavelength for Planckian spectrum generation.

If None: use same wavelengths as CMFs in :cieobs:.

#### cieobs

\_CIEOBS, optional

CMF set used to convert Planckian spectra to chromaticity coordinates

## lut\_type\_def

None, placeholder

Default lut (tuple key) to read from luts\_dict.

# luts\_dict

None, optional

Dictionary of pre-calculated luts for various espaces and cmf sets.

Must have structure luts\_dict[cspace][cieobs][lut\_label] with the lut part of a two-element list [lut, lut\_kwargs]. It must contain at the top-level a key 'wl' containing the wavelengths of the Planckians used to generate the luts in this dictionary.

If None: the default dict for the mode is used

(e.g. \_CCT\_LUT['ohno2014']['lut\_type\_def'], for mode=='ohno2014').

# lut\_vars

```
['T','uv','uvp','uvpp','iso-T-slope'], optional Data the lut should contain. Must follow this order and minimum should be ['T']
```

#### cspace\_cspace\_kwargs

Lists with the cspace and cspace\_kwargs for which luts will be generated. Default is single chromaticity diagram in \_CCT\_CSPACE.

## ignore\_unequal\_wl

False, optional

If True: ignore any differences in the wavelengths used to calculate the lut (cfr. Planckians) from the luts\_dict and the requested wavelengths in :wl:

#### lut\_generator\_fcn

```
_generate_lut, optional
```

Lets a user specify his own lut generation function (must output a list of 1 lut).

Default is the general function. There is a specific one for

Ohno's 2014 method as that one requires a different correction factor

for each lut for the parabolic solutions. This optimized value is specified in the second list index. (see \_generate\_lut\_ohno2014()).

## lut\_generator\_kwargs

```
{}, optional
```

Dict with keyword arguments specific to the (user) lut\_generator\_fcn.

```
(e.g. {'f_corr':0.9991} for _generate_lut_ohno2014())
```

#### **Returns:**

lut

List with an ndarray with in the columns whatever is specified in lut\_vars (Tc and uv are always present!).

Default lut\_vars = ['T','uv','uvp','uvpp','iso-T-slope']

- Tc: (in K)
- u,v: chromaticity coordinates of planckians
- u'v': chromaticity coordinates of 1st derivative of the planckians.
- u",v": chromaticity coordinates of 2nd derivative of the planckians.
- slope of isotemperature lines (calculated as in Robertson, 1968).

#### lut\_kwargs

{}

Dictionary with additional parameters related to the generation of the

luxpy.color.cct.\_generate\_tcs(tc4, uin=None, seamless\_stitch=True, cct\_max=1000000000000,

cct\_min=450, fallback\_unit='K-1', fallback\_n=50, resample\_ndarray=False)

Get an ndarray of Tc's obtained from a list or tuple of tc4 4-vectors (or ndarray).

## **Args:**

tc4

list or tuple of 4-vectors or ndarray.

If ndarray: return tc4 limited to a cct\_min-cct\_max range (do nothing else).

If list/tuple: e.g.  $(tc4_1, tc4_2, tc4_3,...)$  or  $(tc4_1, tc4_2, tc4_3,...,$ 

bool::seamless\_stitch)

When the last element of the list/tuple is a bool, then this specifies

how the Tc arrays generated for each of the 4-vector elements need to be stitched together. This overrides the seamless\_stitch input argument.

Vector elements are:

[Tmin, Tmax inclusive, Tinterval(or number of intervals), unit]

Unit specifies unit of the Tc interval, i.e. it determines the

type of scale in which the spacing of the Tc are done.

Unit options are:

- '%': equal relative Tc spacing (in %, cfr. (Ti+1 Ti-1)/Ti-1).
- 'K' equal absolute Tc spacing (in K, cfr. (Ti+1 Ti-1).
- '%-1': equal relative reciprocal Tc (MK-1 = mired).
- 'K-1': equal absolute reciprocal Tc (MK-1 = mired).

If the 'interval' element is negative, it actually represents

the number of intervals between Tmin, Tmax (included).

#### uin

None, optional

Unit of input Tmin, Tmax (by default it is assumed to be the same as the scale 'unit').

## seamless\_stitch

True, optional

Determines how the Tc arrays generated for each of the 4-vector elements are stitched together. Is overriden by the presence of a

bool as last list/tuple element in :tc4:.

For a seamless stitch, all units for all 4-vectors should be the same!!

#### cct\_max

\_CCT\_MAX, optional

Limit Tc's to a maximum value of cct max

## cct\_min

\_CCT\_MIN, optional

Limit Tc's to a minimum value of cct\_max

#### fallback unit

\_CCT\_FALLBACK\_UNIT, optional

Unit to fall back on when the input unit in tc4 (of first list) is 'au'. As there is no common distancing of the unit types ['K','%','%-1','K-1'] the Tc's are generated by dividing the min-max range into a number of divisions, specified by the negative 3 element (or when

positive or NaN, the number of divisions is set by :fallback\_divisions:)

## fallback\_n

CCT FALLBACK N, optional

Number of divisions the min-max range is divided into, in the fallback case in which unit=='au' and the 3e 4-vector element is NaN or positive.

## resample\_ndarray

False, optional

If False: do not resample Tc's of an ndarray input for tc4 else: divide min-max range in fallback\_n intervals. Uses fallback\_unit to determine the scale for the resampling.

#### **Returns:**

tcs

ndarray with Tcs

Get an ndarray LUT for Tc's obtained from a list or tuple of tc4 4-vectors (or ndarray).

# **Args:**

tc4

list or tuple of 4-vectors or ndarray.

If ndarray: return tc4 limited to a cct\_min-cct\_max range (do nothing else).

If list/tuple: e.g. (tc4\_1, tc4\_2, tc4\_3,...) or (tc4\_1, tc4\_2, tc4\_3,...,

bool::seamless\_stitch)

When the last element of the list/tuple is a bool, then this specifies

how the Tc arrays generated for each of the 4-vector elements need to be stitched together. This overrides the seamless\_stitch input argument.

Vector elements are:

[Tmin, Tmax inclusive, Tinterval(or number of intervals), unit]

Unit specifies unit of the Tc interval, i.e. it determines the type of scale in which the spacing of the Tc are done.

Unit options are:

- '%': equal relative Tc spacing (in %, cfr. (Ti+1 Ti-1)/Ti-1).
- 'K' equal absolute Tc spacing (in K, cfr. (Ti+1 Ti-1).
- '%-1': equal relative reciprocal Tc (MK-1 = mired).
- 'K-1': equal absolute reciprocal Tc (MK-1 = mired).

If the 'interval' element is negative, it actually represents the number of intervals between Tmin, Tmax (included).

#### uin

None, optional

Unit of input Tmin, Tmax (by default it is assumed to be the same as the scale 'unit').

## seamless\_stitch

True, optional

Determines how the Tc arrays generated for each of the 4-vector elements are stitched together. Is overriden by the presence of a bool as last list/tuple element in :tc4:.

For a seamless stitch, all units for all 4-vectors should be the same!!

#### cct\_max

\_CCT\_MAX, optional

Limit Tc's to a maximum value of cct\_max

## cct min

\_CCT\_MIN, optional

Limit Tc's to a minimum value of cct\_max

# fallback\_unit

```
_CCT_FALLBACK_UNIT, optional
```

Unit to fall back on when the input unit in tc4 (of first list) is 'au'. As there is no common distancing of the unit types ['K','%','%-1','K-1'] the Tc's are generated by dividing the min-max range into a number of divisions, specified by the negative 3 element (or when positive or NaN, the number of divisions is set by :fallback\_divisions:)

# fallback\_n

```
_CCT_FALLBACK_N, optional
```

Number of divisions the min-max range is divided into, in the fallback case in which unit=='au' and the 3e 4-vector element is NaN or positive.

# resample\_tc4\_array

False, optional

If False: do not resample Tc's of an ndarray input for tc4 else: divide min-max range in fallback\_n intervals. Uses fallback\_unit to determine the scale for the resampling.

#### wl

None, optional

Wavelength for Planckian spectrum generation.

If None: use same wavelengths as CMFs in :cieobs:.

#### cieobs

[\_CIEOBS] or list, optional

Generate a LUT for each one in the list.

If None: generate for all cmfs in \_CMF.

#### lut vars

['T','uv','uvp','uvpp','iso-T-slope'], optional Data the lut should contain. Must follow this order

and minimum should be ['T']

## cspace,cspace\_kwargs

Lists with the cspace and cspace\_kwargs for which luts will be generated. Default is single chromaticity diagram in CCT CSPACE.

## **Returns:**

#### lut

List with an ndarray with in the columns whatever is specified in

lut vars (Tc and uv are always present!).

Default lut\_vars = ['T','uv','uvp','uvpp','iso-T-slope']

- Tc: (in K)
- u,v: chromaticity coordinates of planckians
- u'v': chromaticity coordinates of 1st derivative of the planckians.
- u",v": chromaticity coordinates of 2nd derivative of the planckians.
- slope of isotemperature lines (calculated as in Robertson, 1968).

# lut\_kwargs

{},

Dictionary with additional parameters related to the generation of the

```
luxpy.color.cct._generate_lut_ohno2014(lut, uin=None, seamless_stitch=True, fallback_unit='K-1',
                                              fallback_n=50, resample_ndarray=False,
                                              cct_max=1000000000000.0, cct_min=450, luts_dict=None,
                                              lut_type_def=None, lut_vars=['T', 'uv'], cieobs='1931_2',
                                              cspace_str=None, wl=None, ignore_unequal_wl=False,
```

lut\_generator\_fcn=<function \_generate\_lut>,

lut\_generator\_kwargs={}, cspace='Yuv60', cspace\_kwargs={'bwtf': {}, 'fwtf': {}}, f\_corr=None,

ignore\_f\_corr\_is\_None=False, ignore\_wl\_diff=False, \*\*kwargs)

# Lut generator function for ohno2014.

## Args:

see docstring for \_generate\_lut

## f\_corr

Tc,x correction factor for the parabolic solution in Ohno2014.

If None, it will be recalculated (note that it depends on the lut) for increased accuracy.

# ignore\_f\_corr\_is\_None

If True, ignore f\_corr is None, i.e. don't re-calculate f\_corr.

## **Returns:**

lut

an ndarray with the lut

dict

a dictionary with the (re-optmized) value for f\_corr and for ignore\_f\_cor\_is\_None.)

# luxpy.color.cct.xyz\_to\_cct\_ohno2011(xyz)

Calculate cct and Duv from CIE 1931 2° xyz following Ohno (2011).

Args:

XYZ

ndarray with CIE 1931 2° X,Y,Z tristimulus values

#### **Returns:**

cct, duv

ndarrays with correlated color temperatures and distance to blackbody locus in CIE 1960 uv

**References:** 1. Ohno, Y. (2011). Calculation of CCT and Duv and Practical Conversion Formulae. CORM 2011 Conference, Gaithersburg, MD, May 3-5, 2011

## 4.4.4 cat/

рy

- \_\_init\_\_.py
- · chromaticadaptation.py

namespace luxpy.cat

# cat: Module supporting chromatic adaptation transforms (corresponding colors)

```
_WHITE_POINT default adopted white point
```

\_LA default luminance of the adaptation field

MCATS default chromatic adaptation sensor spaces

- 'hpe': Hunt-Pointer-Estevez: R. W. G. Hunt, The Reproduction of Colour: Sixth Edition, 6th ed. Chichester, UK: John Wiley & Sons Ltd, 2004.
- 'cat02': from ciecam02: CIE159-2004, "A Colour Apperance Model for Color Management System: CIECAM02," CIE, Vienna, 2004.
- 'cat02-bs': cat02 adjusted to solve yellow-blue problem (last line = [0 0 1]): Brill MH, Süsstrunk S. Repairing gamut problems in CIECAM02: A progress report. Color Res Appl 2008;33(5), 424–426.
- 'cat02-jiang': cat02 modified to solve yb-probem + purple problem: Jun Jiang, Zhifeng Wang,M. Ronnier Luo,Manuel Melgosa,Michael H. Brill,Changjun Li, Optimum solution of the CIECAM02 yellow–blue and purple problems, Color Res Appl 2015: 40(5), 491-503.

- · 'kries'
- 'judd-1945': from CIE16-2004, Eq.4, a23 modified from 0.1 to 0.1020 for increased accuracy
- 'bfd': bradford transform: G. D. Finlayson and S. Susstrunk, "Spectral sharpening and the Bradford transform," 2000, vol. Proceeding, pp. 236–242.
- 'sharp': sharp transform: S. Süsstrunk, J. Holm, and G. D. Finlayson, "Chromatic adaptation performance of different RGB sensors," IS&T/SPIE Electronic Imaging 2001: Color Imaging, vol. 4300. San Jose, CA, January, pp. 172–183, 2001.
- 'cmc': C. Li, M. R. Luo, B. Rigg, and R. W. G. Hunt, "CMC 2000 chromatic adaptation transform: CMCCAT2000," Color Res. Appl., vol. 27, no. 1, pp. 49–58, 2002.
- 'ipt': F. Ebner and M. D. Fairchild, "Development and testing of a color space (IPT) with improved hue uniformity," in IS&T 6th Color Imaging Conference, 1998, pp. 8–13.
- 'lms':
- 'bianco': S. Bianco and R. Schettini, "Two new von Kries based chromatic adaptation transforms found by numerical optimization," Color Res. Appl., vol. 35, no. 3, pp. 184–192, 2010.
- 'bianco-pc': S. Bianco and R. Schettini, "Two new von Kries based chromatic adaptation transforms found by numerical optimization," Color Res. Appl., vol. 35, no. 3, pp. 184–192, 2010.
- 'cat16': C. Li, Z. Li, Z. Wang, Y. Xu, M. R. Luo, G. Cui, M. Melgosa, M. H. Brill, and M. Pointer, "Comprehensive color solutions: CAM16, CAT16, and CAM16-UCS," Color Res. Appl., p. n/a–n/a.

check\_dimensions() Check if dimensions of data and xyzw match.

## get\_transfer\_function()

Calculate the chromatic adaptation diagonal matrix transfer function Dt.

Default = 'vonkries' (others: 'rlab', see Fairchild 1990)

## smet2017\_D()

Calculate the degree of adaptation based on chromaticity.

Smet, K.A.G.\*, Zhai, Q., Luo, M.R., Hanselaer, P., (2017), Study of chromatic adaptation using memory color matches, Part II: colored illuminants. Opt. Express, 25(7), pp. 8350-8365

## get\_degree\_of\_adaptation()

Calculates the degree of adaptation.

D passes either right through or D is calculated following some D-function (Dtype) published in literature (cat02, cat16, cmccat, smet2017) or set manually.

- parse\_x1x2\_parameters() local helper function that parses input parameters and makes them the target\_shape for easy calculation
- **apply()** Calculate corresponding colors by applying a von Kries chromatic adaptation transform (CAT), i.e. independent rescaling of 'sensor sensitivity' to data to adapt from current adaptation conditions (1) to the new conditions (2).

```
luxpy.color.cat.check_dimensions(data, xyzw, caller='cat.apply()')
     Check if dimensions of data and xyzw match.
     Does nothing when they do, but raises error if dimensions don't match.
     Args:
                  data
                       ndarray with color data.
                  xyzw
                       ndarray with white point tristimulus values.
                  caller
                       str with caller function for error handling, optional
     Returns:
                  returns
                       ndarray with input color data,
                       Raises error if dimensions don't match.
luxpy.color.cat.get_transfer_function(cattype='vonkries', catmode='1>0>2', lmsw1=None,
                                               lmsw2=None, lmsw0=array([[100, 100, 100]]), D10=1.0,
                                               D20=1.0, La1=100.0, La2=100.0, La0=100.0)
     Calculate the chromatic adaptation diagonal matrix transfer function Dt.
     Args:
                 cattype
                        'vonkries' (others: 'rlab', see Farchild 1990), optional
                  catmode
                       '1>0>2, optional
                             -'1>0>2': Two-step CAT
                                   from illuminant 1 to baseline illuminant 0 to illuminant 2.
                             -'1>0': One-step CAT
                                   from illuminant 1 to baseline illuminant 0.
                             -'0>2': One-step CAT
                                   from baseline illuminant 0 to illuminant 2.
                 lmsw1
                       None, depending on :catmode: optional
                 lmsw2
                       None, depending on :catmode: optional
                 lmsw0
                       _WHITE_POINT, optional
                 D10
                       1.0, optional
                       Degree of adaptation for ill. 1 to ill. 0
                 D20
```

```
Degree of adaptation for ill. 2 to ill. 0
                  La1
                        luxpy._LA, optional
                        Adapting luminance under ill. 1
                  La2
                        luxpy._LA, optional
                        Adapting luminance under ill. 2
                  La<sub>0</sub>
                        luxpy._LA, optional
                        Adapting luminance under baseline ill. 0
      Returns:
                  Dt
                        ndarray (diagonal matrix)
luxpy.color.cat.get_degree_of_adaptation(Dtype=None, **kwargs)
      Calculates the degree of adaptation according to some function published in literature.
      Args:
                  Dtype
                        None, optional
                              If None: kwargs should contain 'D' with value.
                              If 'manual: kwargs should contain 'D' with value.
                        If 'cat02' or 'cat16': kwargs should contain keys 'F' and 'La'.
                              Calculate D according to CAT02 or CAT16 model:
                                    D = F*(1-(1/3.6)*numpy.exp((-La-42)/92))
                        If 'cmc': kwargs should contain 'La', 'La0'(or 'La2') and 'order'
                              for 'order' = '1>0': 'La' is set La1 and 'La0' to La0.
                              for 'order' = '0>2': 'La' is set La0 and 'La0' to La1.
                              for 'order' = '1>2': 'La' is set La1 and 'La2' to La0.
                              D is calculated as follows:
                                    D = 0.08*numpy.log10(La1+La0)+0.76-0.45*(La1-La0)/(La1+La0)
                        If 'smet2017': kwargs should contain 'xyzw' and 'Dmax'
                              (see Smet2017_D for more details).
                        If "? user defined", then D is calculated by:
                              D = ndarray(eval(:Dtype:))
      Returns:
                  D
                        ndarray with degree of adaptation values.
      Notes:
               1. D passes either right through or D is calculated following some D-function (Dtype) published in
                  literature.
               2. D is limited to values between zero and one
               3. If kwargs do not contain the required parameters, an exception is raised.
luxpy.color.cat.smet2017_D(xyzw, Dmax=None)
      Calculate the degree of adaptation based on chromaticity following Smet et al. (2017)
      Args:
                  XYZW
```

1.0, optional

```
ndarray with white point data (CIE 1964 10° XYZs!!)
```

#### **Dmax**

None or float, optional

Defaults to 0.6539 (max D obtained under experimental conditions, but probably too low due to dark surround leading to incomplete chromatic adaptation even for neutral illuminants

resulting in background luminance (fov~ $50\hat{A}^{\circ}$ ) of 760 cd/m $\hat{A}^{2}$ ))

#### **Returns:**

D

ndarray with degrees of adaptation

**References:** 1. Smet, K.A.G.\*, Zhai, Q., Luo, M.R., Hanselaer, P., (2017), Study of chromatic adaptation using memory color matches, Part II: colored illuminants, Opt. Express, 25(7), pp. 8350-8365.

luxpy.color.cat.parse\_x1x2\_parameters(x, target\_shape, catmode, expand\_2d\_to\_3d=None, default=[1.0, 1.0])

Parse input parameters x and make them the target\_shape for easy calculation.

Input in main function can now be a single value valid for all xyzw or an array with a different value for each xyzw.

## **Args:**

X

list[float, float] or ndarray

#### target\_shape

tuple with shape information

## catmode

```
'1>0>2, optional
```

-'1>0>2': Two-step CAT

from illuminant 1 to baseline illuminant 0 to illuminant 2.

-'1>0': One-step CAT

from illuminant 1 to baseline illuminant 0.

-'0>2': One-step CAT

from baseline illuminant 0 to illuminant 2.

## expand\_2d\_to\_3d

None, optional

[will be removed in future, serves no purpose]

Expand:x: from 2 to 3 dimensions.

## default

[1.0,1.0], optional

Default values for :x:

#### **Returns:**

#### returns

(ndarray, ndarray) for x10 and x20

```
luxpy.color.cat.apply(data, n_step=2, catmode=None, cattype='vonkries', xyzw1=None, xyzw2=None,
                            xyzw0=None, D=None, mcat=['cat02'], normxyz0=None, outtype='xyz', La=None,
                            F=None, Dtype=None)
      Calculate corresponding colors by applying a von Kries chromatic adaptation transform (CAT), i.e. independent
      rescaling of 'sensor sensitivity' to data to adapt from current adaptation conditions (1) to the new conditions (2).
      Args:
                  data
                        ndarray of tristimulus values (can be NxMx3)
                  n_step
                        2, optional
                        Number of step in CAT (1: 1-step, 2: 2-step)
                  catmode
                        None, optional
                              - None: use :n_step: to set mode: 1 = 1 \ge 2, 2 \le 1 \ge 0 \ge 2
                              -'1>0>2': Two-step CAT
                                    from illuminant 1 to baseline illuminant 0 to illuminant 2.
                              -'1>2': One-step CAT
                                    from illuminant 1 to illuminant 2.
                              -'1>0': One-step CAT
                                    from illuminant 1 to baseline illuminant 0.
                              -'0>2': One-step CAT
                                    from baseline illuminant 0 to illuminant 2.
                  cattype
                        'vonkries' (others: 'rlab', see Farchild 1990), optional
                  xyzw1
                        None, depending on :catmode: optional (can be Mx3)
                  xyzw2
                        None, depending on :catmode: optional (can be Mx3)
                  xyzw0
                        None, depending on :catmode: optional (can be Mx3)
                  D
                        None, optional
                        Degrees of adaptation. Defaults to [1.0, 1.0].
                  La
                        None, optional
                        Adapting luminances.
                        If None: xyz values are absolute or relative.
                        If not None: xyz are relative.
                  \mathbf{F}
                        None, optional
                        Surround parameter(s) for CAT02/CAT16 calculations
                              (:Dtype: == 'cat02' or 'cat16')
```

Defaults to [1.0, 1.0].

```
Dtype
                        None, optional
                        Type of degree of adaptation function from literature
                        See luxpy.cat.get_degree_of_adaptation()
                  mcat
                        [_MCAT_DEFAULT], optional
                        List[str] or List[ndarray] of sensor space matrices for each
                              condition pair. If len(:mcat:) == 1, the same matrix is used.
                  normxyz0
                        None, optional
                        Set of xyz tristimulus values to normalize the sensor space matrix to.
                  outtype
                        'xyz' or 'lms', optional
                              - 'xyz': return corresponding tristimulus values
                              - 'lms': return corresponding sensor space excitation values
                                    (e.g. for further calculations)
      Returns:
                  returns
                        ndarray with corresponding colors
      Reference: 1. Smet, K. A. G., & Ma, S. (2020). Some concerns regarding the CAT16 chromatic adaptation
            transform. Color Research & Application, 45(1), 172–177.
luxpy.color.cat.apply_vonkries1(xyz, xyzw1, xyzw2, D=1, mcat=None, invmcat=None, in_type='xyz',
                                        out_type='xyz', use_Yw=False)
      Apply a 1-step von kries chromatic adaptation transform.
      Args:
                  XYZ
                        ndarray with sample tristimulus or cat-sensor values
                  xyzw1
                        ndarray with white point tristimulus or cat-sensor values of illuminant 1
                  xyzw2
                        ndarray with white point tristimulus or cat-sensor values of illuminant 2
                  D
                        1, optional
                        Degree of chromatic adaptation
                  mcat
                        None, optional
                        Specifies CAT sensor space.
                        - options:

    None defaults to luxpy.cat._MCAT_DEFAULT

                              - str: see see luxpy.cat._MCATS.keys() for options
                                    (details on type, ?luxpy.cat)
```

```
- ndarray: matrix with sensor primaries
```

```
invmcat
```

None, optional

Pre-calculated inverse mcat.

If None: calculate inverse of mcat.

## in\_type

'xyz', optional

Input type ('xyz', 'rgb') of data in xyz, xyzw1, xyzw2

## out\_type

'xyz', optional

Output type ('xyz', 'rgb') of corresponding colors

## use\_Yw

False, optional

Use CAT version with Yw factors included (but this results in potential wrong predictions, see Smet & Ma (2020)).

#### **Returns:**

xyzc

ndarray with corresponding colors.

**Reference:** 1. Smet, K. A. G., & Ma, S. (2020). Some concerns regarding the CAT16 chromatic adaptation transform. Color Research & Application, 45(1), 172–177.

Apply a 2-step von kries chromatic adaptation transform.

## **Args:**

xyz

ndarray with sample tristimulus or cat-sensor values

# xyzw1

ndarray with white point tristimulus or cat-sensor values of illuminant 1

#### xyzw2

ndarray with white point tristimulus or cat-sensor values of illuminant 2

# xyzw0

None, optional

ndarray with white point tristimulus or cat-sensor values of baseline illuminant 0

None: defaults to EEW.

D

[1,1], optional

Degree of chromatic adaptations (Ill.1->Ill.0, Ill.2.->Ill.0)

#### mcat

None, optional

Specifies CAT sensor space.

- options:

- None defaults to luxpy.cat.\_MCAT\_DEFAULT

```
- str: see see luxpy.cat._MCATS.keys() for options
                                    (details on type, ?luxpy.cat)
                              - ndarray: matrix with sensor primaries
                  invmcat
                        None, optional
                        Pre-calculated inverse mcat.
                        If None: calculate inverse of mcat.
                  in_type
                        'xyz', optional
                        Input type ('xyz', 'rgb') of data in xyz, xyzw1, xyzw2
                  out_type
                        'xyz', optional
                        Output type ('xyz', 'rgb') of corresponding colors
                  use_Yw
                        False, optional
                        Use CAT version with Yw factors included (but this results in
                        potential wrong predictions, see Smet & Ma (2020)).
      Returns:
                  xyzc
                        ndarray with corresponding colors.
      Reference: 1. Smet, K. A. G., & Ma, S. (2020). Some concerns regarding the CAT16 chromatic adaptation
            transform. Color Research & Application, 45(1), 172–177.
luxpy.color.cat.apply_vonkries(xyz, xyzw1, xyzw2, xyzw0=None, D=1, n_step=2, catmode='1>0>2',
                                       mcat=None, invmcat=None, in_type='xyz', out_type='xyz', use_Yw=False)
      Apply a 1-step or 2-step von kries chromatic adaptation transform.
      Args:
                  XYZ
                        ndarray with sample tristimulus or cat-sensor values
                  xyzw1
                        ndarray with white point tristimulus or cat-sensor values of illuminant 1
                  xyzw2
                        ndarray with white point tristimulus or cat-sensor values of illuminant 2
                  xyzw0
                        None, optional
                        ndarray with white point tristimulus or cat-sensor values of baseline illuminant 0
                        None: defaults to EEW.
                  D
                        [1,1], optional
                        Degree of chromatic adaptations (III.1->III.0, III.2.->III.0)
                  n_step
                        2, optional
                        Number of step in CAT (1: 1-step, 2: 2-step)
```

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#### catmode

```
None, optional
```

- None: use :n\_step: to set mode:  $1 = 1 \ge 2, 2 \le 1 \ge 0 \ge 2$
- -'1>0>2': Two-step CAT

from illuminant 1 to baseline illuminant 0 to illuminant 2.

-'1>2': One-step CAT

from illuminant 1 to illuminant 2.

-'1>0': One-step CAT

from illuminant 1 to baseline illuminant 0.

-'0>2': One-step CAT

from baseline illuminant 0 to illuminant 2.

#### mcat

None, optional

Specifies CAT sensor space.

- options:
  - None defaults to luxpy.cat.\_MCAT\_DEFAULT
  - str: see see luxpy.cat.\_MCATS.keys() for options (details on type, ?luxpy.cat)
  - ndarray: matrix with sensor primaries

#### invmcat

None, optional

Pre-calculated inverse mcat.

If None: calculate inverse of mcat.

## in\_type

'xyz', optional

Input type ('xyz', 'rgb') of data in xyz, xyzw1, xyzw2

## out\_type

'xyz', optional

Output type ('xyz', 'rgb') of corresponding colors

## use\_Yw

False, optional

Use CAT version with Yw factors included (but this results in potential wrong predictions, see Smet & Ma (2020)).

#### **Returns:**

xyzc

ndarray with corresponding colors.

**Reference:** 1. Smet, K. A. G., & Ma, S. (2020). Some concerns regarding the CAT16 chromatic adaptation transform. Color Research & Application, 45(1), 172–177.

luxpy.color.cat.apply\_ciecat94(xyz, xyzw, xyzwr=None, E=1000, Er=1000, Yb=20, D=1, cat94\_old=True) Calculate corresponding color tristimulus values using the CIECAT94 chromatic adaptation transform. Args:

xyz

ndarray with sample 1931 2° XYZ tristimulus values under the test illuminant

```
xyzw
                   ndarray with white point tristimulus values of the test illuminant
            xyzwr
                   None, optional
                   ndarray with white point tristimulus values of the reference illuminant
                   None defaults to D65.
            \mathbf{E}
                   100, optional
                   Illuminance (lx) of test illumination
            Er
                   63.66, optional
                   Illuminance (lx) of the reference illumination
            Yb
                   20, optional
                   Relative luminance of the adaptation field (background)
            D
                   1, optional
                   Degree of chromatic adaptation.
                   For object colours D = 1,
                   and for luminous colours (typically displays) D=0
Returns:
            xyzc
                   ndarray with corresponding tristimlus values.
Reference:
```

1. CIE160-2004. (2004). A review of chromatic adaptation transforms (Vols. CIE160-200). CIE.

# 4.4.5 cam/

рy

- \_\_init\_\_.py
- · colorappearancemodels.py
- helpers.py
- · utils.py
- ciecam02.py
- · cam02ucs.py
- ciecam16.py
- · cam16ucs.py
- cam15u
- sww2016.py
- · cam18sl.py

- · camjabz.py
- zcam.py
- cmf\_translator\_sww2021

namespace luxpy.cam

# cam: sub-package with color appearance models

# \_UNIQUE\_HUE\_DATA

database of unique hues with corresponding Hue quadratures and eccentricity factors for ciecam02, ciecam16, ciecam97s, cam15u, cam18sl)

## \_SURROUND\_PARAMETERS

database of surround param. c, Nc, F and FLL for ciecam02, ciecam16, ciecam97s and cam15u.

## \_NAKA\_RUSHTON\_PARAMETERS

database with parameters (n, sig, scaling and noise)

for the Naka-Rushton function:

NK(x) = sign(x) \* scaling \* ((abs(x)\*\*n) / ((abs(x)\*\*n) + (sig\*\*n))) + noise

#### CAM UCS PARAMETERS

database with parameters specifying the conversion

from ciecamX to:

camXucs (uniform color space),

camXlcd (large color diff.),

camXscd (small color diff).

- \_CAM15U\_PARAMETERS database with CAM15u model parameters.
- **CAM SWW16 PARAMETERS** cam sww16 model parameters.
- \_CAM18SL\_PARAMETERS database with CAM18sl model parameters
- **\_CAM\_DEFAULT\_WHITE\_POINT** Default internal reference white point (xyz)
- \_CAM\_DEFAULT\_CONDITIONS Default CAM model parameters for model.
- **CAM AXES** dict with list[str,str,str] containing axis labels of defined cspaces.

**deltaH()** Compute a hue difference, dH = 2\*C1\*C2\*sin(dh/2).

naka\_rushton() applies a Naka-Rushton function to the input

hue\_angle() calculates a positive hue angle

**hue\_quadrature()** calculates the Hue quadrature from the hue.

## ciecam02()

calculates ciecam02 output

N. Moroney, M. D. Fairchild, R. W. G. Hunt, C. Li, M. R. Luo, and T. Newman, "The CIECAM02 color appearance model," IS&T/SID Tenth Color Imaging Conference. p. 23, 2002.

cam16()

## calculates cam16 output

C. Li, Z. Li, Z. Wang, Y. Xu, M. R. Luo, G. Cui, M. Melgosa, M. H. Brill, and M. Pointer, "Comprehensive color solutions: CAM16, CAT16, and CAM16-UCS," Color Res. Appl., p. n/a–n/a.

#### cam02ucs()

calculates ucs (or lcd, scd) output based on ciecam02

(forward + inverse available)

M. R. Luo, G. Cui, and C. Li, "Uniform colour spaces based on CIECAM02 colour appearance model," Color Res. Appl., vol. 31, no. 4, pp. 320–330, 2006.

#### cam16ucs()

calculates ucs (or lcd, scd) output based on cam16

(forward + inverse available)

C. Li, Z. Li, Z. Wang, Y. Xu, M. R. Luo, G. Cui, M. Melgosa, M. H. Brill, and M. Pointer, "Comprehensive color solutions: CAM16, CAT16, and CAM16-UCS," Color Res. Appl.

## cam15u()

calculates the output for the CAM15u model for self-luminous unrelated stimuli.

M. Withouck, K. A. G. Smet, W. R. Ryckaert, and P. Hanselaer, "Experimental driven modelling of the color appearance of unrelated self-luminous stimuli: CAM15u," Opt. Express, vol. 23, no. 9, pp. 12045–12064, 2015.

M. Withouck, K. A. G. Smet, and P. Hanselaer, (2015), "Brightness prediction of different sized unrelated self-luminous stimuli," Opt. Express, vol. 23, no. 10, pp. 13455–13466.

#### cam\_sww16()

A simple principled color appearance model based on a mapping of the Munsell color system.

Smet, K. A. G., Webster, M. A., & Whitehead, L. A. (2016). "A simple principled approach for modeling and understanding uniform color metrics." Journal of the Optical Society of America A, 33(3), A319–A331.

# cam18sl()

calculates the output for the CAM18sl model for self-luminous related stimuli.

Hermans, S., Smet, K. A. G., & Hanselaer, P. (2018). "Color appearance model for self-luminous stimuli." Journal of the Optical Society of America A, 35(12), 2000–2009.

camXucs() Wraps ciecam02(), ciecam16(), cam02ucs(), cam16ucs().

#### specific wrappers in the 'xyz\_to\_cspace()' and 'cpsace\_to\_xyz()' format

```
'xyz_to_jabM_ciecam02', 'jabM_ciecam02_to_xyz', 'xyz_to_jabC_ciecam02', 'jabC_ciecam02_to_xyz', 'xyz_to_jabM_ciecam16', 'jabM_ciecam16_to_xyz', 'xyz_to_jabC_ciecam16', 'jabC_ciecam16_to_xyz', 'xyz_to_jabz', 'jabz_to_xyz', 'xyz_to_jabM_camjabz', 'jabM_camjabz_to_xyz', 'xyz_to_jabC_camjabz', 'jabC_camjabz_to_xyz',
```

'xyz\_to\_jab\_cam02ucs', 'jab\_cam02ucs\_to\_xyz',

'xyz\_to\_jab\_cam02lcd', 'jab\_cam02lcd\_to\_xyz',

```
'xyz_to_jab_cam16ucs', 'jab_cam16ucs_to_xyz',
                  'xyz_to_jab_cam16lcd', 'jab_cam16lcd_to_xyz',
                  'xyz_to_jab_cam16scd', 'jab_cam16scd_to_xyz',
                  'xyz_to_qabW_cam15u', 'qabW_cam15u_to_xyz',
                  'xyz_to_lab_cam_sww16','lab_cam_sww16_to_xyz',
                  'xyz_to_qabM_cam18sl', 'qabM_cam18sl_to_xyz',
                  'xyz_to_qabS_cam18sl', 'qabS_cam18sl_to_xyz',
            _update_parameter_dict() Get parameter dict and update with values in args dict
            setup default adaptation field() Setup a default illuminant adaptation field with Lw =
                  100 cd/m<sup>2</sup> for selected CIE observer.
            _massage_input_and_init_output() Redimension input data to ensure most they have the
                  appropriate sizes for easy and efficient looping.
            massage output data to original shape() Massage output data to restore original shape
                  of original CAM input.
            get absolute xyz xyzw() Calculate absolute xyz tristimulus values of stimulus and white
                  point from spectral input or convert relative xyz values to absolute ones.
            _simple_cam() An example CAM illustration the usage of the functions in luxpy.cam.helpers
Module for CAM "front-end" cmf adaptation
           translate cmfI to cmfS()
                  Using smooth RGB primaries, translate input data (spectral or tristimlus)
                  for an indivual observer to the expected tristimulus values for a standard observer.
           get conversion matrix()
                  Using smooth RGB primaries, get the 'translator' matrix to convert
                  tristimulus values calculated using an individual observer's
                  color matching functions (cmfs) to those calculated using the cmfs of
                  a standard observer.
           get rgb smooth prims() Get smooth R, G, B primaries with specified wavelength range
            _R,_G,_B precalculated smooth primaries with [360,830,1] wavelength range.
luxpy.color.cam.hue_angle(a, b, htype='deg')
     Calculate positive hue angle (0^{\circ}-360° or 0 - 2*pi rad.) from opponent signals a and b.
     Args:
                  a
                        ndarray of a-coordinates
                 b
                        ndarray of b-coordinates
                 htype
                        'deg' or 'rad', optional
                              - 'deg': hue angle between 0° and 360°
                              - 'rad': hue angle between 0 and 2pi radians
     Returns:
```

'xyz\_to\_jab\_cam02scd', 'jab\_cam02scd\_to\_xyz',

```
returns
```

```
ndarray of positive hue angles.
```

```
luxpy.color.cam.naka_rushton(data, sig=2.0, n=0.73, scaling=1.0, noise=0.0, forward=True) Apply a Naka-Rushton response compression (n) and an adaptive shift (sig).
```

```
NK(x) = sign(x) * scaling * ((abs(x)**n) / ((abs(x)**n) + (sig**n))) + noise
```

# Args:

## data

float or ndarray

sig

2.0, optional

Semi-saturation constant. Value for which NK(:data:) is 1/2

n

0.73, optional

Compression power.

## scaling

1.0, optional

Maximum value of NK-function.

#### noise

0.0, optional

Cone excitation noise.

## forward

True, optional

True: do NK(x)

False: do NK(x)\*\*(-1).

## **Returns:**

#### returns

float or ndarray with NK-(de)compressed input :x:

luxpy.color.cam.deltaH(h1, C1, h2=None, C2=None, htype='deg') Compute a hue difference, dH = 2\*C1\*C2\*sin(dh/2)

**Args:** 

h1

hue for sample 1 (or hue difference if h2 is None)

**C1** 

chroma of sample 1 (or prod C1\*C2 if C2 is None)

h2

hue angle of sample 2 (if None, then h1 contains a hue difference)

**C2** 

chroma of sample 2

```
htype
                        'deg' or 'rad', optional
                              - 'deg': hue angle between 0^{\circ} and 360^{\circ}
                              - 'rad': hue angle between 0 and 2pi radians
      Returns:
                  returns
                        ndarray of deltaH values.
luxpy.color.cam.hue_quadrature(h, unique_hue_data=None, forward=True)
      Get hue quadrature H from hue h.
      Args:
                  h
                        float or ndarray [(N,) or (N,1)] with:
                              - hue angle data in degrees (!) if forward == True.
                              - Hue quadrature data if forward = False
                  unique_hue data
                        None or dict, optional
                              - None: defaults to:
                                           {'hues': 'red yellow green blue red'.split(),
                                     'i': np.arange(5.0),
                                     'hi':[20.14, 90.0, 164.25,237.53,380.14],
                                     'ei':[0.8,0.7,1.0,1.2,0.8],
                                     'Hi':[0.0,100.0,200.0,300.0,400.0]}
                              - dict: user specified unique hue data
                                     (same structure as above)
                  forward
                        True, optional
                        If true: input h is hue angle, else it is Hue quadrature
      Returns:
                  Н
                        ndarray of Hue quadrature value(s) (forward == True) or of hue angle values(s)
                        (foward == False).
luxpy.color.cam._update_parameter_dict(args, parameters={}, cieobs='2006_10',
                                                  match_conversionmatrix_to_cieobs=False,
                                                  Mxyz2lms_whitepoint=None)
      Get parameter dict and update with values in args dict.
            Also replace the xyz-to-lms conversion matrix with the one corresponding
            to cieobs and normalize it to illuminant E.
      Args:
                  args
                        dictionary with updated values.
                        (get by placing 'args = locals().copy()' immediately after the start
                        of the function from which the update is called,
                        see simple cam() code for an example.)
                  parameters
```

dictionary with all (adjustable) parameter values used by the model

#### cieobs

String with the CIE observer CMFs (one of \_CMF['types'] of the input data Is used to get the Mxyz2lms matrix when match\_conversionmatrix\_to\_cieobs == True)

## match\_conversionmatrix\_to\_cieobs

False, optional

If False: keep the Mxyz2lms in the parameters dict

# Mxyz2lms\_whitepoint

None, optional

If not None: update the Mxyz2lms key in the parameters dict so that the conversion matrix is the one in \_CMF[cieobs]['M'], in other such that it matches the cieobs of the input data.

#### **Returns:**

# parameters

updated dictionary with model parameters for further use in the CAM.

**Notes:** For an example on the use, see code \_simple\_cam() (type: \_simple\_cam??)

luxpy.color.cam.\_setup\_default\_adaptation\_field(dataw=None, Lw=100, cie\_illuminant='D65', inputtype='xyz', relative=True, cieobs='2006 10')

Setup a default illuminant adaptation field with  $Lw = 100 \text{ cd/m}^2$  for selected CIE observer.

# **Args:**

## dataw

None or ndarray, optional

Input tristimulus values or spectral data of white point.

None defaults to the use of the illuminant specified in :cie\_illuminant:.

# cie\_illuminant

'D65', optional

String corresponding to one of the illuminants (keys)

in luxpy.\_CIE\_ILLUMINANT

If ndarray, then use this one.

This is ONLY USED WHEN dataw is NONE!!!

## Lw

100.0, optional

Luminance (cd/m<sup>2</sup>) of white point.

## inputtype

'xyz' or 'spd', optional

Specifies the type of input:

tristimulus values or spectral data for the forward mode.

#### relative

True or False, optional

True: xyz tristimulus values are relative (Yw = 100)

## cieobs

```
_CAM_DEFAULT_CIEOBS, optional
```

CMF set to use to perform calculations where spectral data

is involved (inputtype == 'spd'; dataw = None)

Other options: see luxpy.\_CMF['types']

#### **Returns:**

# dataw

Ndarray with default adaptation field data (spectral or xyz)

**Notes:** For an example on the use, see code \_simple\_cam() (type: \_simple\_cam??)

## Args:

#### data

ndarray with input tristimulus values

or spectral data

or input color appearance correlates

Can be of shape: (N [, xM], x 3), whereby:

N refers to samples and M refers to light sources.

Note that for spectral input shape is  $(N \times (M+1) \times W)$ 

#### dataw

None or ndarray, optional

Input tristimulus values or spectral data of white point.

None defaults to the use of CIE illuminant C.

## inputtype

```
'xyz' or 'spd', optional
```

Specifies the type of input:

tristimulus values or spectral data for the forward mode.

## direction

```
'forward' or 'inverse', optional
- 'forward': xyz -> cam
- 'inverse': cam -> xyz
```

#### n out

3, optional

output size of last dimension of camout

(e.g.  $n_{out}=3$  for j,a,b output or  $n_{out}=5$  for J,M,h,a,b output)

## **Returns:**

## data

ndarray with reshaped data

#### dataw

ndarray with reshaped dataw

## camout

```
NaN filled ndarray for output of CAMv (camout.shape[-1] == Nout)
                  originalshape
                        original shape of data
      Notes: For an example on the use, see code _simple_cam() (type: _simple_cam??)
luxpy.color.cam._massage_output_data_to_original_shape(data, originalshape)
      Massage output data to restore original shape of original CAM input.
      Notes: For an example on the use, see code simple cam() (type: simple cam??)
luxpy.color.cam._get_absolute_xyz_xyzw(data, dataw, i=0, Lw=100, direction='forward',
                                                  cieobs='2006_10', inputtype='xyz', relative=True)
      Calculate absolute xyz tristimulus values of stimulus and white point from spectral input or convert relative xyz
      values to absolute ones.
      Args:
                  data
                        ndarray with input tristimulus values
                        or spectral data
                        or input color appearance correlates
                        Can be of shape: (N [, xM], x 3), whereby:
                        N refers to samples and M refers to light sources.
                        Note that for spectral input shape is (N \times (M+1) \times W)
                  dataw
                        None or ndarray, optional
                        Input tristimulus values or spectral data of white point.
                        None defaults to the use of CIE illuminant C.
                  i
                        0, optional
                        row number in data and dataw ndarrays
                        (for loops across illuminant dimension after dimension reshape
                        with _massage_output_data_to_original_shape).
                  Lw
                        100.0, optional
                        Luminance (cd/m<sup>2</sup>) of white point.
                  inputtype
                        'xyz' or 'spd', optional
                        Specifies the type of input:
                              tristimulus values or spectral data for the forward mode.
                  direction
                        'forward' or 'inverse', optional
                              -'forward': xyz -> cam
                              -'inverse': cam -> xyz
                  relative
                        True or False, optional
```

True: xyz tristimulus values are relative (Yw = 100)

```
cieobs
                        _CAM_DEFAULT_CIEOBS, optional
                        CMF set to use to perform calculations where spectral data is involved (inputtype ==
                        'spd'; dataw = None)
                        Other options: see luxpy._CMF['types']
      Returns:
                  xyzti
                        in forward mode: ndarray with relative or absolute sample xyz for data[i]
                        in inverse mode: None
                  xyzwi
                        ndarray with relative or absolute white point for dataw[i]
                  xyzw_abs
                        ndarray with absolute xyz for white point for dataw[i]
      Notes: For an example on the use, see code _simple_cam() (type: _simple_cam??)
luxpy.color.cam._simple_cam(data, dataw=None, Lw=100.0, relative=True, inputtype='xyz',
                                    direction='forward', cie_illuminant='D65', parameters={'Mxyz2lms':
                                    array([[0.38971, 0.68898, - 0.07868], [- 0.22981, 1.1834, 0.04641], [0.0, 0.0,
                                    1.0]]), 'cA': 1, 'ca': array([1, -1, 0]), 'cb': array([0.16667, 0.16667, -
                                    0.33333]), 'n': 0.333333333333333}, cieobs='2006_10',
                                    match to conversionmatrix to cieobs=True)
      An example CAM illustration the usage of the functions in luxpy.cam.helpers
      Note that this example uses NO chromatic adaptation
      and SIMPLE compression, opponent and correlate processing.
      THIS IS ONLY FOR ILLUSTRATION PURPOSES!!!
      Args:
                  data
                        ndarray with input:
                              - tristimulus values
                        or
                              - spectral data
                        or
                              - input color appearance correlates
                        Can be of shape: (N [, xM], x 3), whereby:
                        N refers to samples and M refers to light sources.
                        Note that for spectral input shape is (N \times (M+1) \times W)
                  dataw
                        None or ndarray, optional
                        Input tristimulus values or spectral data of white point.
                        None defaults to the use of :cie_illuminant:
                  cie_illuminant
                        'D65', optional
                        String corresponding to one of the illuminants (keys)
```

```
in luxpy._CIE_ILLUMINANT
      If ndarray, then use this one.
      This is ONLY USED WHEN dataw is NONE!!!
Lw
      100.0, optional
      Luminance (cd/m<sup>2</sup>) of white point.
relative
      True or False, optional
      True: data and dataw input is relative (i.e. Yw = 100)
parameters
      {'cA': 1, 'ca':np.array([1,-1,0]), 'cb':(1/3)*np.array([0.5,0.5,-1]),
            'n': 1/3, 'Mxyz2lms': _CMF['1931_2']['M'].copy()}
      Dict with model parameters
      (For illustration purposes of match_conversionmatrix_to_cieobs,
            the conversion matrix luxpy._CMF['1931_2']['M'] does NOT match
            the default observer specification of the input data in :cieobs: !!!)
inputtype
      'xyz' or 'spd', optional
      Specifies the type of input:
            tristimulus values or spectral data for the forward mode.
direction
      'forward' or 'inverse', optional
            -'forward': xyz -> cam
            -'inverse': cam -> xyz
cieobs
      '2006_10', optional
      CMF set to use to perform calculations where spectral data
      is involved (inputtype == 'spd'; dataw = None)
      Other options: see luxpy._CMF['types']
match_conversionmatrix_to_cieobs
      True, optional
      When changing to a different CIE observer, change the xyz_to_lms
      matrix to the one corresponding to that observer.
      Set to False to keep the one in the parameter dict!
returns
      ndarray with:
      - color appearance correlates (:direction: == 'forward')
      - XYZ tristimulus values (:direction: == 'inverse')
```

**Returns:** 

```
luxpy.color.cam.ciecam02(data, xyzw=array([[100.0, 100.0, 100.0]]), Yw=None, outin='J,aM,bM',
                               conditions=None, naka_rushton_parameters=None, unique_hue_data=None,
                               forward=True, yellowbluepurplecorrect=False, mcat='cat02')
     Run CIECAM02 color appearance model in forward or backward modes.
     Args:
                  data
                        ndarray with relative sample xyz values (forward mode) or J'a'b' coordinates (inverse
                        mode)
                 xyzw
                        ndarray with relative white point tristimulus values
                  Yw
                        None, optional
                        Luminance factor of white point.
                        If None: xyz (in data) and xyzw are entered as relative tristimulus values
                             (normalized to Yw = 100).
                        If not None: input tristimulus are absolute and Yw is used to
                             rescale the absolute values to relative ones
                             (relative to a reference perfect white diffuser
                                   with Ywr = 100).
                        Yw can be < 100 for e.g. paper as white point. If Yw is None, it
                        is assumed that the relative Y-tristimulus value in xyzw
                        represents the luminance factor Yw.
                  conditions
                        None, optional
                        Dictionary with viewing condition parameters for:
                                   La, Yb, D and surround.
                             surround can contain:
                                   - str (options: 'avg','dim','dark') or
                                   - dict with keys c, Nc, F.
                        None results in:
                              { 'La':100, 'Yb':20, 'D':1, 'surround':'avg'}
                  naka_rushton_parameters
                        None, optional
                        If None: use _NAKA_RUSHTON_PARAMETERS
                  unique_hue_data
                        None, optional
                        If None: use _UNIQUE_HUE_DATA
                 forward
                        True, optional
                        If True: run in CAM in forward mode, else: inverse mode.
                  outin
                        'J,aM,bM', optional
                        String with requested output (e.g. "J,aM,bM,M,h") [Forward mode]
```

- attributes: 'J': lightness,'Q': brightness,

```
'M': colorfulness,'C': chroma, 's': saturation,
                              'h': hue angle, 'H': hue quadrature/composition,
                        String with inputs in data [inverse mode].
                        Input must have data.shape[-1]==3 and last dim of data must have
                        the following structure for inverse mode:
                              * data[...,0] = J \text{ or } O,
                              * data[...,1:] = (aM,bM) or (aC,bC) or (aS,bS) or (M,h) or (C,h), ...
                  yellowbluepurplecorrect
                        False, optional
                        If False: don't correct for yellow-blue and purple problems in ciecam02.
                        If 'brill-suss':
                              for yellow-blue problem, see:
                                    - Brill [Color Res Appl, 2006; 31, 142-145] and
                                    - Brill and Süsstrunk [Color Res Appl, 2008; 33, 424-426]
                        If 'jiang-luo':
                              for yellow-blue problem + purple line problem, see:
                                    - Jiang, Jun et al. [Color Res Appl 2015: 40(5), 491-503]
                  mcat
                        'cat02', optional
                        Specifies CAT sensor space.
                        - options:
                              - None defaults to 'cat02'
                                    (others e.g. 'cat02-bs', 'cat02-jiang',
                                    all trying to correct gamut problems of original cat02 matrix)
                              - str: see see luxpy.cat._MCATS.keys() for options
                                    (details on type, ?luxpy.cat)
                              - ndarray: matrix with sensor primaries
      Returns:
                  camout
                        ndarray with color appearance correlates (forward mode)
                        XYZ tristimulus values (inverse mode)
      References: 1. N. Moroney, M. D. Fairchild, R. W. G. Hunt, C. Li, M. R. Luo, and T. Newman, (2002), "The
            CIECAM02 color appearance model," IS&T/SID Tenth Color Imaging Conference. p. 23, 2002.
luxpy.color.cam.xyz_to_jabM_ciecam02(data, xyzw=array([[100.0, 100.0, 100.0]]), Yw=None,
                                               conditions=None, naka_rushton_parameters=None,
                                               unique_hue_data=None, yellowbluepurplecorrect=False,
                                               mcat='cat02', **kwargs)
      Wrapper function for ciecam02 forward mode with J,aM,bM output.
      For help on parameter details: ?luxpy.cam.ciecam02
```

```
luxpy.color.cam.jabM_ciecam02_to_xyz(data, xyzw=array([[100.0, 100.0, 100.0]]), Yw=None,
                                            conditions=None, naka rushton parameters=None,
                                            unique hue data=None, yellowbluepurplecorrect=False,
                                            mcat='cat02', **kwargs)
     Wrapper function for ciecam02 inverse mode with J,aM,bM input.
     For help on parameter details: ?luxpy.cam.ciecam02
luxpy.color.cam.xyz_to_jabC_ciecam02(data, xyzw=array([[100.0, 100.0, 100.0]]), Yw=None,
                                            conditions=None, naka_rushton_parameters=None,
                                            unique_hue_data=None, yellowbluepurplecorrect=False,
                                            mcat='cat02', **kwargs)
     Wrapper function for ciecam02 forward mode with J,aC,bC output.
     For help on parameter details: ?luxpy.cam.ciecam02
luxpy.color.cam.jabC_ciecam02_to_xyz(data, xyzw=array([[100.0, 100.0, 100.0]]), Yw=None,
                                            conditions=None, naka rushton parameters=None,
                                            unique_hue_data=None, yellowbluepurplecorrect=False,
                                            mcat='cat02', **kwargs)
     Wrapper function for ciecam02 inverse mode with J,aC,bC input.
     For help on parameter details: ?luxpy.cam.ciecam02
luxpy.color.cam.cam02ucs(data, xyzw=array([[100.0, 100.0, 100.0]]), Yw=None, conditions=None,
                             naka_rushton_parameters=None, unique_hue_data=None, ucstype='ucs',
                             forward=True, yellowbluepurplecorrect=False, mcat='cat02')
     Run the CAM02-UCS[,-LCD,-SDC] color appearance difference model in forward or backward modes.
     Args:
                 data
                      ndarray with sample xyz values (forward mode) or J'a'b' coordinates (inverse mode)
                 xyzw
                      ndarray with white point tristimulus values
                 conditions
                      None, optional
                      Dictionary with viewing conditions.
                      None results in:
                            {'La':100, 'Yb':20, 'D':1, 'surround':'avg'}
                      For more info see luxpy.cam.ciecam02()?
                 naka_rushton_parameters
                      None, optional
                      If None: use NAKA RUSHTON PARAMETERS
```

unique\_hue\_data

None, optional

```
If None: use _UNIQUE_HUE_DATA
                  ucstype
                        'ucs', optional
                        String with type of color difference appearance space
                        options: 'ucs', 'scd', 'lcd'
                  forward
                        True, optional
                        If True: run in CAM in forward mode, else: inverse mode.
                  yellowbluepurplecorrect
                        False, optional
                        If False: don't correct for yellow-blue and purple problems in ciecam02.
                        If 'brill-suss':
                              for yellow-blue problem, see:
                                    - Brill [Color Res Appl, 2006; 31, 142-145] and
                                    - Brill and Süsstrunk [Color Res Appl, 2008; 33, 424-426]
                        If 'jiang-luo':
                              for yellow-blue problem + purple line problem, see:
                                    - Jiang, Jun et al. [Color Res Appl 2015: 40(5), 491-503]
                  mcat
                        'cat02', optional
                        Specifies CAT sensor space.
                        - options:
                              - None defaults to 'cat02'
                                    (others e.g. 'cat02-bs', 'cat02-jiang',
                                    all trying to correct gamut problems of original cat02 matrix)
                              - str: see see luxpy.cat._MCATS.keys() for options
                                    (details on type, ?luxpy.cat)
                              - ndarray: matrix with sensor primaries
      Returns:
                  camout
                        ndarray with J'a'b' coordinates (forward mode)
                              or
                        XYZ tristimulus values (inverse mode)
      References: 1. M.R. Luo, G. Cui, and C. Li, 'Uniform colour spaces based on CIECAM02 colour appearance
            model,' Color Res. Appl., vol. 31, no. 4, pp. 320–330, 2006.
luxpy.color.cam.xyz_to_jab_cam02ucs(data, xyzw=array([[100.0, 100.0, 100.0]]), Yw=None,
                                             conditions=None, naka_rushton_parameters=None,
                                             unique_hue_data=None, yellowbluepurplecorrect=None,
                                             mcat='cat02', **kwargs)
      Wrapper function for cam02ucs forward mode with J,aM,bM output.
      For help on parameter details: ?luxpy.cam.cam02ucs
```

```
luxpy.color.cam.jab_cam02ucs_to_xyz(data, xyzw=array([100.0, 100.0, 100.0])), Yw=None,
                                          conditions=None, naka rushton parameters=None,
                                          unique hue data=None, yellowbluepurplecorrect=None,
                                          mcat='cat02', **kwargs)
     Wrapper function for cam02ucs inverse mode with J,aM,bM input.
     For help on parameter details: ?luxpy.cam.cam02ucs
luxpy.color.cam.xyz_to_jab_cam02lcd(data, xyzw=array([[100.0, 100.0, 100.0]]), Yw=None,
                                          conditions=None, naka_rushton_parameters=None,
                                          unique_hue_data=None, yellowbluepurplecorrect=None,
                                          mcat='cat02', **kwargs)
     Wrapper function for cam02ucs forward mode with J,aMp,bMp output and ucstype = lcd.
     For help on parameter details: ?luxpy.cam.cam02ucs
luxpy.color.cam.jab_cam02lcd_to_xyz(data, xyzw=array([[100.0, 100.0, 100.0]]), Yw=None,
                                          conditions=None, naka rushton parameters=None,
                                          unique_hue_data=None, yellowbluepurplecorrect=None,
                                          mcat='cat02', **kwargs)
     Wrapper function for cam02ucs inverse mode with J,aMp,bMp input and ucstype = lcd.
     For help on parameter details: ?luxpy.cam.cam02ucs
luxpy.color.cam.xyz_to_jab_cam02scd(data, xyzw=array([[100.0, 100.0, 100.0]]), Yw=None,
                                          conditions=None, naka_rushton_parameters=None,
                                          unique_hue_data=None, yellowbluepurplecorrect=None,
                                          mcat='cat02', **kwargs)
     Wrapper function for cam02ucs forward mode with J,aMp,bMp output and ucstype = scd.
     For help on parameter details: ?luxpy.cam.cam02ucs
luxpy.color.cam.jab_cam02scd_to_xyz(data, xyzw=array([[100.0, 100.0, 100.0]]), Yw=None,
                                          conditions=None, naka rushton parameters=None,
                                          unique hue data=None, yellowbluepurplecorrect=None,
                                          mcat='cat02', **kwargs)
     Wrapper function for cam02ucs inverse mode with J,aMp,bMp input and ucstype = scd.
     For help on parameter details: ?luxpy.cam.cam02ucs
luxpy.color.cam.ciecam16(data, xyzw=array([[100.0, 100.0, 100.0]]), Yw=None, outin='J,aM,bM',
                             conditions=None, naka_rushton_parameters=None, unique_hue_data=None,
                             forward=True, mcat='cat16')
     Run CIECAM16 color appearance model in forward or backward modes.
```

```
Args:
            data
                  ndarray with relative sample xyz values (forward mode) or J'a'b' coordinates (inverse
                  mode)
            XYZW
                  ndarray with relative white point tristimulus values
            Yw
                  None, optional
                  Luminance factor of white point.
                  If None: xyz (in data) and xyzw are entered as relative tristimulus values
                        (normalized to Yw = 100).
                  If not None: input tristimulus are absolute and Yw is used to
                        rescale the absolute values to relative ones
                        (relative to a reference perfect white diffuser
                              with Ywr = 100).
                  Yw can be < 100 for e.g. paper as white point. If Yw is None, it
                  is assumed that the relative Y-tristimulus value in xyzw
                  represents the luminance factor Yw.
            conditions
                  None, optional
                  Dictionary with viewing condition parameters for:
                              La, Yb, D and surround.
                        surround can contain:
                              - str (options: 'avg','dim','dark') or
                              - dict with keys c, Nc, F.
                  None results in:
                        {'La':100, 'Yb':20, 'D':1, 'surround':'avg'}
            naka rushton parameters
                  None, optional
                  If None: use _NAKA_RUSHTON_PARAMETERS
            unique_hue_data
                  None, optional
                  If None: use _UNIQUE_HUE_DATA
            forward
                  True, optional
```

If True: run in CAM in forward mode, else: inverse mode.

#### outin

```
'J,aM,bM', optional
String with requested output (e.g. "J,aM,bM,M,h") [Forward mode]
- attributes: 'J': lightness,'Q': brightness,
      'M': colorfulness,'C': chroma, 's': saturation,
      'h': hue angle, 'H': hue quadrature/composition,
String with inputs in data [inverse mode].
```

```
Input must have data.shape[-1]==3 and last dim of data must have
                       the following structure for inverse mode:
                             * data[...,0] = J \text{ or } Q,
                             * data[...,1:] = (aM,bM) or (aC,bC) or (aS,bS) or (M,h) or (C,h), ...
                 mcat
                       'cat16', optional
                       Specifies CAT sensor space.
                       - options:
                             - None defaults to 'cat16'
                             - str: see see luxpy.cat._MCATS.keys() for options
                                   (details on type, ?luxpy.cat)
                             - ndarray: matrix with sensor primaries
     Returns:
                 camout
                       ndarray with color appearance correlates (forward mode)
                       XYZ tristimulus values (inverse mode)
     References: 1. C. Li, Z. Li, Z. Wang, Y. Xu, M. R. Luo, G. Cui, M. Melgosa, M. H. Brill, and M. Pointer, (2017),
           "Comprehensive color solutions: CAM16, CAT16, and CAM16-UCS," Color Res. Appl., p. n/a-n/a.
luxpy.color.cam.xyz_to_jabM_ciecam16(data, xyzw=array([[100.0, 100.0, 100.0]]), Yw=None,
                                             conditions=None, naka_rushton_parameters=None,
                                             unique_hue_data=None, mcat='cat16', **kwargs)
     Wrapper function for ciecam16 forward mode with J,aM,bM output.
     For help on parameter details: ?luxpy.cam.ciecam16
luxpy.color.cam.jabM_ciecam16_to_xyz(data, xyzw=array([[100.0, 100.0, 100.0]]), Yw=None,
                                             conditions=None, naka_rushton_parameters=None,
                                             unique_hue_data=None, mcat='cat16', **kwargs)
     Wrapper function for ciecam16 inverse mode with J,aM,bM input.
     For help on parameter details: ?luxpy.cam.ciecam16
luxpy.color.cam.xyz_to_jabC_ciecam16(data, xyzw=array([[100.0, 100.0, 100.0]]), Yw=None,
                                             conditions=None, naka_rushton_parameters=None,
                                             unique_hue_data=None, mcat='cat16', **kwargs)
     Wrapper function for ciecam16 forward mode with J,aC,bC output.
     For help on parameter details: ?luxpy.cam.ciecam16
luxpy.color.cam.jabC_ciecam16_to_xyz(data, xyzw=array([[100.0, 100.0, 100.0]]), Yw=None,
                                             conditions=None, naka_rushton_parameters=None,
                                             unique_hue_data=None, mcat='cat16', **kwargs)
     Wrapper function for ciecam16 inverse mode with J,aC,bC input.
```

```
For help on parameter details: ?luxpy.cam.ciecam16
luxpy.color.cam.cam16ucs(data, xyzw=array([[100.0, 100.0, 100.0]]), Yw=None, conditions=None,
                               naka_rushton_parameters=None, unique_hue_data=None, ucstype='ucs',
                              forward=True, mcat='cat16')
     Run the CAM16-UCS[,-LCD,-SDC] color appearance difference model in forward or backward modes.
     Args:
                 data
                       ndarray with sample xyz values (forward mode) or J'a'b' coordinates (inverse mode)
                 xyzw
                       ndarray with white point tristimulus values
                 conditions
                       None, optional
                       Dictionary with viewing conditions.
                       None results in:
                             {'La':100, 'Yb':20, 'D':1, 'surround':'avg'}
                       For more info see luxpy.cam.ciecam16()?
                 naka_rushton_parameters
                       None, optional
                       If None: use _NAKA_RUSHTON_PARAMETERS
                 unique_hue_data
                       None, optional
                       If None: use _UNIQUE_HUE_DATA
                 ucstype
                       'ucs', optional
                       String with type of color difference appearance space
                       options: 'ucs', 'scd', 'lcd'
                 forward
                       True, optional
                       If True: run in CAM in forward mode, else: inverse mode.
                 mcat
                       'cat16', optional
                       Specifies CAT sensor space.
                       - options:
                             - None defaults to 'cat16'
                             - str: see see luxpy.cat._MCATS.keys() for options
                                   (details on type, ?luxpy.cat)
                             - ndarray: matrix with sensor primaries
     Returns:
                 camout
                       ndarray with J'a'b' coordinates (forward mode)
```

or

```
XYZ tristimulus values (inverse mode)
```

**References:** 1. M.R. Luo, G. Cui, and C. Li, 'Uniform colour spaces based on CIECAM02 colour appearance model,' Color Res. Appl., vol. 31, no. 4, pp. 320–330, 2006.

luxpy.color.cam.xyz\_to\_jab\_cam16ucs(data, xyzw=array([[100.0, 100.0, 100.0]]), Yw=None, conditions=None, naka\_rushton\_parameters=None, unique\_hue\_data=None, mcat='cat16', \*\*kwargs)

Wrapper function for cam16ucs forward mode with J,aM,bM output.

For help on parameter details: ?luxpy.cam.cam16ucs

luxpy.color.cam.jab\_cam16ucs\_to\_xyz(data, xyzw=array([[100.0, 100.0, 100.0]]), Yw=None, conditions=None, naka\_rushton\_parameters=None, unique\_hue\_data=None, mcat='cat16', \*\*kwargs')

Wrapper function for cam16ucs inverse mode with J,aM,bM input.

For help on parameter details: ?luxpy.cam.cam16ucs

luxpy.color.cam.xyz\_to\_jab\_cam16lcd(data, xyzw=array([[100.0, 100.0, 100.0]]), Yw=None, conditions=None, naka\_rushton\_parameters=None, unique\_hue\_data=None, mcat='cat16', \*\*kwargs')

Wrapper function for cam16ucs forward mode with J,aMp,bMp output and ucstype = lcd.

For help on parameter details: ?luxpy.cam.cam16ucs

luxpy.color.cam.jab\_cam16lcd\_to\_xyz(data, xyzw=array([[100.0, 100.0, 100.0]]), Yw=None, conditions=None, naka\_rushton\_parameters=None, unique\_hue\_data=None, mcat='cat16', \*\*kwargs)

Wrapper function for cam16ucs inverse mode with J,aMp,bMp input and ucstype = lcd.

For help on parameter details: ?luxpy.cam.cam16ucs

luxpy.color.cam.xyz\_to\_jab\_cam16scd(data, xyzw=array([[100.0, 100.0, 100.0]]), Yw=None, conditions=None, naka\_rushton\_parameters=None, unique\_hue\_data=None, mcat='cat16', \*\*kwargs')

Wrapper function for cam16ucs forward mode with J,aMp,bMp output and ucstype = scd.

For help on parameter details: ?luxpy.cam.cam16ucs

luxpy.color.cam.jab\_cam16scd\_to\_xyz(data, xyzw=array([[100.0, 100.0, 100.0]]), Yw=None, conditions=None, naka\_rushton\_parameters=None, unique\_hue\_data=None, mcat='cat16', \*\*kwargs')

Wrapper function for cam16ucs inverse mode with J,aMp,bMp input and ucstype = scd.

For help on parameter details: ?luxpy.cam.cam16ucs luxpy.color.cam.zcam(data, xyzw=None, outin='J,aM,bM', cieobs='1931\_2', conditions=None, forward=True, mcat='cat02', apply\_cat\_to\_whitepoint=False, \*\*kwargs) Run the Jz,az,bz based color appearance model in forward or backward modes. Args: data ndarray with relative sample xyz values (forward mode) or J'a'b' coordinates (inverse mode) **XYZW** ndarray with relative white point tristimulus values None defaults to D65 cieobs \_CIEOBS, optional CMF set to use when calculating :xyzw: if this is None. conditions None, optional Dictionary with viewing condition parameters for: La, Yb, D and surround. surround can contain: - str (options: 'avg','dim','dark') or - dict with keys c, Nc, F. None results in: { 'La':100, 'Yb':20, 'D':1, 'surround':'avg'} forward True, optional If True: run in CAM in forward mode, else: inverse mode. outin 'J,aM,bM', optional String with requested output (e.g. "J,aM,bM,M,h") [Forward mode] - attributes: 'J': lightness,'Q': brightness, 'M': colorfulness,'C': chroma, 's': saturation, 'h': hue angle, 'H': hue quadrature/composition, 'Wz': whiteness, 'Kz':blackness, 'Sz': saturation, 'V': vividness String with inputs in data [inverse mode]. Input must have data.shape[-1]==3 and last dim of data must have

mcat

'cat02', optional

Specifies CAT sensor space.

the following structure for inverse mode: \* data[...,0] = J or Q,

\* data[...,1:] = (aM,bM) or (aC,bC) or (aS,bS) or (M,h) or (C,h),...

- options:

- None defaults to 'cat02'
- str: see see luxpy.cat.\_MCATS.keys() for options (details on type, ?luxpy.cat)
- ndarray: matrix with sensor primaries

# apply\_cat\_to\_whitepoint

False, optional

Apply a CAT to the white point.

However, ZCAM as published doesn't do this for some reason.

#### **Returns:**

#### camout

ndarray with color appearance correlates (forward mode)

or

XYZ tristimulus values (inverse mode)

- **References:** 1. Safdar, M., Cui, G., Kim, Y. J., and Luo, M. R.(2017). Perceptually uniform color space for image signals including high dynamic range and wide gamut. Opt. Express, vol. 25, no. 13, pp. 15131–15151, Jun. 2017.
  - 2. Safdar, M., Hardeberg, J., Cui, G., Kim, Y. J., and Luo, M. R.(2018). A Colour Appearance Model based on Jzazbz Colour Space, 26th Color and Imaging Conference (2018), Vancouver, Canada, November 12-16, 2018, pp96-101.
  - 3. Safdar, M., Hardeberg, J.Y., Luo, M.R. (2021) ZCAM, a psychophysical model for colour appearance prediction, Optics Express. 29(4), 6036-6052, <a href="https://doi.org/10.1364/OE.413659">https://doi.org/10.1364/OE.413659</a>

luxpy.color.cam.xyz\_to\_jabz(xyz, ztype='jabz', use\_zcam\_parameters=False, \*\*kwargs)

Convert XYZ tristimulus values to Jz,az,bz color coordinates.

### Args:

**XYZ** 

ndarray with absolute tristimulus values (Y in cd/m<sup>2</sup>!)

# ztype

'jabz', optional

String with requested return:

Options: 'jabz', 'iabz'

### use\_zcam\_parameters

False, optional

ZCAM uses a slightly different values (see notes)

# **Returns:**

### jabz

ndarray with Jz (or Iz), az, bz color coordinates

### **Notes:**

- 1. :xyz: is assumed to be under D65 viewing conditions! If necessary perform chromatic adaptation!
- 2a. Jz represents the 'lightness' relative to a D65 white with luminance =  $10000 \text{ cd/m}^2$  (note that Jz that not exactly equal 1 for this high value, but rather for  $102900 \text{ cd/m}^2$ )
- 2b. az, bz represent respectively a red-green and a yellow-blue opponent axis (but note that a D65 shows a small offset from (0,0))
- 3. ZCAM: calculates Iz as M' epsilon (instead L'/2 + M'/2 as in Iz,az,bz color space!).

```
Reference: 1. Safdar, M., Cui, G., Kim, Y. J., and Luo, M. R. (2017). Perceptually uniform color space for image signals including high dynamic range and wide gamut. Opt. Express, vol. 25, no. 13, pp. 15131–15151, June 2017.
```

2. Safdar, M., Hardeberg, J.Y., Luo, M.R. (2021) ZCAM, a psychophysical model for colour appearance prediction, Optics Express. 29(4), 6036-6052, <a href="https://doi.org/10.1364/OE.413659">https://doi.org/10.1364/OE.413659</a>

```
luxpy.color.cam.jabz_to_xyz(jabz, ztype='jabz', use_zcam_parameters=False, **kwargs) Convert Jz,az,bz color coordinates to XYZ tristimulus values.
```

# Args:

# jabz

ndarray with Jz,az,bz color coordinates

### ztype

'jabz', optional

String with requested return:

Options: 'jabz', 'iabz'

# use\_zcam\_parameters

False, optional

ZCAM uses a slightly different values (see notes)

#### **Returns:**

xyz

ndarray with tristimulus values

#### Note:

- 1. :xyz: is assumed to be under D65 viewing conditions! If necessary perform chromatic adaptation!
- 2a. Jz represents the 'lightness' relative to a D65 white with luminance = 10000 cd/m<sup>2</sup> (note that Jz that not exactly equal 1 for this high value, but rather for 102900 cd/m2)
- 2b. az, bz represent respectively a red-green and a yellow-blue opponent axis (but note that a D65 shows a small offset from (0,0))
- 3. ZCAM: calculates Iz as M' epsilon (instead L'/2 + M'/2 as in Iz,az,bz color space!).
- **Reference:** 1. Safdar, M., Cui, G., Kim, Y. J., and Luo, M. R. (2017). Perceptually uniform color space for image signals including high dynamic range and wide gamut. Opt. Express, vol. 25, no. 13, pp. 15131–15151, June, 2017.
  - 2. Safdar, M., Hardeberg, J.Y., Luo, M.R. (2021) ZCAM, a psychophysical model for colour appearance prediction, Optics Express. 29(4), 6036-6052, <a href="https://doi.org/10.1364/OE.413659">https://doi.org/10.1364/OE.413659</a>

```
luxpy.color.cam.xyz_to_jabM_zcam(data, xyzw='_CIE_D65', cieobs='1931_2', conditions=None, mcat='cat02', apply_cat_to_whitepoint=False, **kwargs)

Wrapper function for zcam forward mode with J,aM,bM output.
```

For help on parameter details: ?luxpy.cam.zcam

```
luxpy.color.cam.jabM_zcam_to_xyz(data, xyzw='_CIE_D65', cieobs='1931_2', conditions=None, mcat='cat02', apply_cat_to_whitepoint=False, **kwargs)
Wrapper function for zcam inverse mode with J,aM,bM input.
```

For help on parameter details: ?luxpy.cam.zcam

```
luxpy.color.cam.xyz_to_jabC_zcam(data, xyzw='_CIE_D65', cieobs='1931_2', conditions=None,
                                         mcat='cat02', apply_cat_to_whitepoint=False, **kwargs)
     Wrapper function for zeam forward mode with J,aC,bC output.
     For help on parameter details: ?luxpy.cam.zcam
luxpy.color.cam.jabC_zcam_to_xyz(data, xyzw='_CIE_D65', cieobs='1931_2', conditions=None,
                                         mcat='cat02', apply_cat_to_whitepoint=False, **kwargs)
     Wrapper function for zcam inverse mode with J,aC,bC input.
     For help on parameter details: ?luxpy.cam.zcam
luxpy.color.cam.cam15u(data, fov=10.0, inputtype='xyz', direction='forward', outin='Q,aW,bW',
                            parameters=None)
     Convert between CIE 2006 10° XYZ tristimulus values (or spectral data) and CAM15u color appearance corre-
     Args:
                 data
                       ndarray of CIE 2006 10° XYZ tristimulus values or spectral data
                             or color appearance attributes
                 fov
                       10.0, optional
                       Field-of-view of stimulus (for size effect on brightness)
                 inputtpe
                        'xyz' or 'spd', optional
                       Specifies the type of input:
                             tristimulus values or spectral data for the forward mode.
                 direction
                        'forward' or 'inverse', optional
                             -'forward': xyz -> cam15u
                             -'inverse': cam15u -> xyz
                 outin
                        'Q,aW,bW' or str, optional
                        'Q,aW,bW' (brightness and opponent signals for amount-of-neutral)
                             other options: 'Q,aM,bM' (colorfulness) and 'Q,aS,bS' (saturation)
                       Str specifying the type of
                             input (:direction: == 'inverse') and
                             output (:direction: == 'forward')
                 parameters
                       None or dict, optional
                       Set of model parameters.
                             - None: defaults to luxpy.cam._CAM15U_PARAMETERS
```

(see references below)

**Returns:** 

returns

ndarray with color appearance correlates (:direction: == 'forward')

or

XYZ tristimulus values (:direction: == 'inverse')

**References:** 1. M. Withouck, K. A. G. Smet, W. R. Ryckaert, and P. Hanselaer, "Experimental driven modelling of the color appearance of unrelated self-luminous stimuli: CAM15u," Opt. Express, vol. 23, no. 9, pp. 12045–12064, 2015. 2. M. Withouck, K. A. G. Smet, and P. Hanselaer, (2015), "Brightness prediction of different sized unrelated self-luminous stimuli," Opt. Express, vol. 23, no. 10, pp. 13455–13466.

luxpy.color.cam.xyz\_to\_qabW\_cam15u(xyz, fov=10.0, parameters=None, \*\*kwargs) Wrapper function for cam15u forward mode with 'Q,aW,bW' output.

For help on parameter details: ?luxpy.cam.cam15u

luxpy.color.cam.qabW\_cam15u\_to\_xyz(qab, fov=10.0, parameters=None, \*\*kwargs) Wrapper function for cam15u inverse mode with 'Q,aW,bW' input.

For help on parameter details: ?luxpy.cam.cam15u

A simple principled color appearance model based on a mapping of the Munsell color system.

This function implements the JOSA A (parameters = 'JOSA') published model.

### **Args:**

# data

ndarray with input tristimulus values or spectral data or input color appearance correlates

Can be of shape: (N [, xM], x 3), whereby:

N refers to samples and M refers to light sources.

Note that for spectral input shape is (N x (M+1) x wl)

# dataw

None or ndarray, optional

Input tristimulus values or spectral data of white point.

None defaults to the use of CIE illuminant C.

Yb

20.0, optional

Luminance factor of background (perfect white diffuser, Yw = 100)

Lw

```
Luminance (cd/m<sup>2</sup>) of white point.
            Ccwb
                  None, optional
                  Degree of cognitive adaptation (white point balancing)
                  If None: use [..,..] from parameters dict.
            relative
                  True or False, optional
                  True: xyz tristimulus values are relative (Yw = 100)
            parameters
                  'JOSA' or str or dict, optional
                  Dict with model parameters.
                        - str: 'JOSA', 'best-fit-JOSA' or 'best-fit-all-Munsell'
                        - dict: user defined model parameters
                              (dict should have same structure)
            inputtype
                  'xyz' or 'spd', optional
                  Specifies the type of input:
                        tristimulus values or spectral data for the forward mode.
            direction
                  'forward' or 'inverse', optional
                        -'forward': xyz -> cam_sww_2016
                        -'inverse': cam_sww_2016 -> xyz
            cieobs
                  '2006 10', optional
                  CMF set to use to perform calculations where spectral data
                  is involved (inputtype == 'spd'; dataw = None)
                  Other options: see luxpy._CMF['types']
            match conversionmatrix to cieobs
                  When changing to a different CIE observer, change the xyz_to_lms
                  matrix to the one corresponding to that observer. If False: use
                  the one set in parameters or _CAM_SWW16_PARAMETERS
Returns:
            returns
                  ndarray with color appearance correlates (:direction: == 'forward')
                        or
                  XYZ tristimulus values (:direction: == 'inverse')
Notes:
     This function implements the JOSA A (parameters = 'JOSA')
      published model.
      With:
            1. A correction for the parameter
```

400.0, optional

```
in Eq.4 of Fig. 11: 0.952 -> -0.952
```

2. The delta\_ac and delta\_bc white-balance shifts in Eq. 5e & 5f should be: -0.028 & 0.821

(cfr. Ccwb = 0.66 in:

ab\_test\_out = ab\_test\_int - Ccwb\*ab\_gray\_adaptation\_field\_int))

**References:** 1. Smet, K. A. G., Webster, M. A., & Whitehead, L. A. (2016). A simple principled approach for modeling and understanding uniform color metrics. Journal of the Optical Society of America A, 33(3), A319–A331.

 $\label{locam_swv16} \verb|luxpy.color.cam.xyz_to_lab_cam_sww16| (xyz, xyzw=None, Yb=20.0, Lw=400.0, Ccwb=None, relative=True, parameters='JOSA', inputtype='xyz', cieobs='2006_10', **kwargs') \\$ 

Wrapper function for cam\_sww16 forward mode with 'xyz' input.

For help on parameter details: ?luxpy.cam.cam\_sww16

luxpy.color.cam.lab\_cam\_sww16\_to\_xyz(lab, xyzw=None, Yb=20.0, Lw=400.0, Ccwb=None, relative=True, parameters='JOSA', inputtype='xyz', cieobs='2006\_10', \*\*kwargs')
Wrapper function for cam\_sww16 inverse mode with 'xyz' input.

For help on parameter details: ?luxpy.cam.cam\_sww16

luxpy.color.cam.cam18s1(data, datab=None, Lb=[100], fov=10.0, inputtype='xyz', direction='forward', outin='Q,aS,bS', parameters=None)

Convert between CIE 2006 10° XYZ tristimulus values (or spectral data) and CAM18sl color appearance correlates.

Args:

data

ndarray of CIE 2006  $10^{\circ}$  absolute XYZ tristimulus values or spectral data or color appearance attributes of stimulus

datab

ndarray of CIE 2006  $10^\circ$  absolute XYZ tristimulus values or spectral data of stimulus background

Lb

[100], optional

Luminance (cd/m<sup>2</sup>) value(s) of background(s) calculated using the CIE 2006 10°

(only used in case datab == None and the background is assumed to be an Equal-Energy-White)

fov

10.0, optional

Field-of-view of stimulus (for size effect on brightness)

inputtpe

```
Specifies the type of input:
                              tristimulus values or spectral data for the forward mode.
                  direction
                        'forward' or 'inverse', optional
                              -'forward': xyz -> cam18sl
                              -'inverse': cam18sl -> xyz
                  outin
                        'Q,aS,bS' or str, optional
                        'Q,aS,bS' (brightness and opponent signals for saturation)
                              other options: 'Q,aM,bM' (colorfulness)
                                    (Note that 'Q,aW,bW' would lead to a Cartesian
                                          a,b-coordinate system centered at (1,0)
                        Str specifying the type of
                              input (:direction: == 'inverse') and
                              output (:direction: == 'forward')
                  parameters
                        None or dict, optional
                        Set of model parameters.
                              - None: defaults to luxpy.cam._CAM18SL_PARAMETERS
                                    (see references below)
      Returns:
                  returns
                        ndarray with color appearance correlates (:direction: == 'forward')
                        XYZ tristimulus values (:direction: == 'inverse')
      Notes:
            * Instead of using the CIE 1964 10° CMFs in some places of the model,
                  the CIE 2006 10° CMFs are used througout, making it more self_consistent.
                  This has an effect on the k scaling factors (now different those in CAM15u)
                  and the illuminant E normalization for use in the chromatic adaptation transform.
                  (see future erratum to Hermans et al., 2018)
            * The paper also used an equation for the amount of white W, which is
                  based on a Q value not expressed in 'bright' ('cA' = 0.937 instead of 123).
                  This has been corrected for in the luxpy version of the model, i.e.
                  _CAM18SL_PARAMETERS['cW'][0] has been changed from 2.29 to 1/11672.
                  (see future erratum to Hermans et al., 2018)
            * Default output was 'Q,aW,bW' prior to March 2020, but since this
                  is an a,b Cartesian system centered on (1,0), the default output
                  has been changed to 'Q,aS,bS'.
      References: 1. Hermans, S., Smet, K. A. G., & Hanselaer, P. (2018). "Color appearance model for self-
            luminous stimuli." Journal of the Optical Society of America A, 35(12), 2000–2009.
luxpy.color.cam.xyz_to_qabM_cam18s1(xyz, xyzb=None, Lb=[100], fov=10.0, parameters=None, **kwargs)
      Wrapper function for cam18sl forward mode with 'Q,aM,bM' output.
```

'xyz' or 'spd', optional

For help on parameter details: ?luxpy.cam.cam18sl

luxpy.color.cam.qabM\_cam18sl\_to\_xyz(qab, xyzb=None, Lb=[100], fov=10.0, parameters=None, \*\*kwargs) Wrapper function for cam18sl inverse mode with 'Q,aM,bM' input.

For help on parameter details: ?luxpy.cam.cam18sl

luxpy.color.cam.**xyz\_to\_qabS\_cam18sl**(*xyz*, *xyzb=None*, *Lb=[100]*, *fov=10.0*, *parameters=None*, \*\*kwargs) Wrapper function for cam18sl forward mode with 'Q,aS,bS' output.

For help on parameter details: ?luxpy.cam.cam18sl

luxpy.color.cam.qabS\_cam18sl\_to\_xyz(qab, xyzb=None, Lb=[100], fov=10.0, parameters=None, \*\*kwargs) Wrapper function for cam18sl inverse mode with 'Q,aS,bS' input.

For help on parameter details: ?luxpy.cam.cam18sl

Wraps ciecam02(), ciecam16(), cam02ucs(), cam16ucs().

**Args:** 

camtype

\_DEFAULT\_TYPE, optional String specifying the cam-model.

**Notes:** 

- 1. To call ciecam02() or ciecam16(): set ucstype to None!!!
- 2. For more info on other input arguments, see doc-strings of those functions.

# 4.4.6 deltaE/

рy

- \_\_init\_\_.py
- colordifferences.py
- · discriminationellipses.py
- · frieleellipses.py
- macadamellipses.py

namespace luxpy.deltaE

### Module for color difference calculations

```
process DEi() Process color difference input DEi for output (helper fnc).
            DE_camucs() Calculate color appearance difference DE using camucs type model.
           DE_2000() Calculate DE2000 color difference.
           DE_cspace() Calculate color difference DE in specific color space.
            get_macadam_ellipse() Estimate n-step MacAdam ellipse at CIE x,y coordinates
            get_brown1957_ellipse() Estimate n-step Brown (1957) ellipse at CIE x,y coordinates.
            get gij fmc() Get gij matrices describing the discrimination ellipses for Yxy using FMC-1
                  or FMC-2.
            get fmc discrimination ellipse() Get n-step discrimination ellipse(s) in v-format (R,r, xc,
                  yc, theta) for Yxy using FMC-1 or FMC-2.
luxpy.color.deltaE.deltaH(h1, C1, h2=None, C2=None, htype='deg')
      Compute a hue difference, dH = 2*C1*C2*sin(dh/2)
      Args:
                  h1
                        hue for sample 1 (or hue difference if h2 is None)
                  C1
                        chroma of sample 1 (or prod C1*C2 if C2 is None)
                  h2
                        hue angle of sample 2 (if None, then h1 contains a hue difference)
                  C2
                        chroma of sample 2
                  htype
                        'deg' or 'rad', optional
                              - 'deg': hue angle between 0^{\circ} and 360^{\circ}
                              - 'rad': hue angle between 0 and 2pi radians
      Returns:
                  returns
                        ndarray of deltaH values.
luxpy.color.deltaE.DE_camucs(xyzt, xyzr, DEtype='jab', avg=None, avg_axis=0, out='DEi',
                                    xyzwt=array([[100.0, 100.0, 100.0]]), xyzwr=array([[100.0, 100.0, 100.0]]),
                                    Ywt=None, conditionst={'D': 1.0, 'Dtype': None, 'La': 100.0, 'Yb': 20.0,
                                     'surround': 'avg'}, Ywr=None, conditionsr={'D': 1.0, 'Dtype': None, 'La':
                                     100.0, 'Yb': 20.0, 'surround': 'avg'}, camtype='ciecam02', ucstype='ucs',
                                    mcat=None, outin='J,aM,bM', yellowbluepurplecorrect=False, **kwargs)
      Calculate color appearance difference DE using camucs type model.
      Args:
                  xyzt
                        ndarray with tristimulus values of test data.
                  xyzr
                        ndarray with tristimulus values of reference data.
```

```
DEtype
                        'jab' or str, optional
                        Options:
                               - 'jab' : calculates full color difference over all 3 dimensions.
                               - 'ab': calculates chromaticity difference.
                               - 'j' : calculates lightness or brightness difference
                                     (depending on :outin:).
                               - 'j,ab': calculates both 'j' and 'ab' options
                                     and returns them as a tuple.
                  avg
                        None, optional
                        None: don't calculate average DE,
                               otherwise use function handle in :avg:.
                  avg_axis
                        axis to calculate average over, optional
                  out
                        'DEi' or str, optional
                        Requested output.
                  camtype
                        luxpy.cam._CAM_DEFAULT_TYPE, optional
                        Str specifier for CAM type to use, options: 'ciecam02' or 'ciecam16'.
                  ucstype
                         'ucs' or 'lcd' or 'scd', optional
                        Str specifier for which type of color attribute compression
                               parameters to use:
                                     -'ucs': uniform color space,
                                     -'lcd': large color differences,
                                     -'scd': small color differences
      Note: For the other input arguments, see ?luxpy.cam.camucs_structure.
      Returns:
                  returns
                        ndarray with DEi [, DEa] or other as specified by :out:
luxpy.color.deltaE.DE2000(xyzt, xyzr, dtype='xyz', DEtype='jab', avg=None, avg axis=0, out='DEi',
                                 xyzwt=None, xyzwr=None, KLCH=None)
      Calculate DE2000 color difference.
      Args:
                  xyzt
                        ndarray with tristimulus values of test data.
                  xyzr
                        ndarray with tristimulus values of reference data.
                  dtype
                         'xyz' or 'lab', optional
```

```
xyzwt
                        None or ndarray, optional
                               White point tristimulus values of test data
                               None defaults to the one set in lx.xyz_to_lab()
                  xyzwr
                        None or ndarray, optional
                               Whitepoint tristimulus values of reference data
                               None defaults to the one set in lx.xyz to lab()
                  DEtype
                         'jab' or str, optional
                        Options:
                               - 'jab' : calculates full color difference over all 3 dimensions.
                               - 'ab': calculates chromaticity difference.
                               - 'j' : calculates lightness or brightness difference
                                     (depending on :outin:).
                               - 'j,ab': calculates both 'j' and 'ab' options
                                     and returns them as a tuple.
                  KLCH
                        None, optional
                        Weigths for L, C, H
                        None: default to [1,1,1]
                  avg
                        None, optional
                        None: don't calculate average DE,
                               otherwise use function handle in :avg:.
                  avg_axis
                        axis to calculate average over, optional
                  out
                         'DEi' or str, optional
                        Requested output.
      Note: For the other input arguments, see specific color space used.
      Returns:
                  returns
                        ndarray with DEi [, DEa] or other as specified by :out:
      References: 1. Sharma, G., Wu, W., & Dalal, E. N. (2005). The CIEDE2000 color-difference formula: Imple-
            mentation notes, supplementary test data, and mathematical observations. Color Research & Application,
            30(1), 21–30.
luxpy.color.deltaE.DE_cspace(xyzt, xyzr, dtype='xyz', tf='Yuv', DEtype='jab', avg=None, avg_axis=0,
                                     out='DEi', xyzwt=None, xyzwr=None, fwtft={}, fwtfr={}, KLCH=None,
                                     camtype='ciecam02', ucstype='ucs')
      Calculate color difference DE in specific color space.
      Args:
```

Specifies data type in :xyzt: and :xyzr:.

```
xyzt
      ndarray with tristimulus values of test data.
xyzr
      ndarray with tristimulus values of reference data.
dtype
      'xyz' or 'jab', optional
      Specifies data type in :xyzt: and :xyzr:.
xyzwt
      None or ndarray, optional
            White point tristimulus values of test data
            None defaults to the one set in :fwtft:
            or else to the default of cspace.
xyzwr
      None or ndarray, optional
            Whitepoint tristimulus values of reference data
                   None defaults to the one set in non-empty: fwtfr:
                   or else to default of cspace.
tf
      _CSPACE, optional
      Color space to use for color difference calculation.
fwtft
      {}, optional
      Dict with parameters for forward transform from xyz to cspace for test data.
fwtfr
      {}, optional
      Dict with parameters for forward transform
      from xyz to cspace for reference data.
KLCH
      None, optional
      Weigths for L, C, H
      None: default to [1,1,1]
      KLCH is not used when tf == 'camucs'.
DEtype
      'jab' or str, optional
      Options:
            - 'jab' : calculates full color difference over all 3 dimensions.
            - 'ab': calculates chromaticity difference.
            - 'j' : calculates lightness or brightness difference
                   (depending on :outin:).
            - 'j,ab': calculates both 'j' and 'ab' options
                   and returns them as a tuple.
```

```
avg
                        None, optional
                        None: don't calculate average DE,
                              otherwise use function handle in :avg:.
                  avg axis
                        axis to calculate average over, optional
                  out
                        'DEi' or str, optional
                        Requested output.
                  camtype
                        luxpy.cam._CAM_DEFAULT_TYPE, optional
                        Str specifier for CAM type to use, options: 'ciecam02' or 'ciecam16'.
                        Only when DEtype == 'camucs'.
                  ucstype
                        'ucs' or 'lcd' or 'scd', optional
                        Str specifier for which type of color attribute compression
                        parameters to use:
                              -'ucs': uniform color space,
                              -'lcd', large color differences,
                              -'scd': small color differences
                        Only when DEtype == 'camucs'.
      Note: For the other input arguments, see specific color space used.
                  returns
                        ndarray with DEi [, DEa] or other as specified by :out:
luxpy.color.deltaE.get_discrimination_ellipse(Yxy=array([[100.0, 0.33333, 0.33333]]), etype='fmc2',
                                                          nsteps=10, k_neighbours=3, average_cik=True,
                                                          Y=None, brown1957_weighted=True)
      Get discrimination ellipse(s) in v-format (R,r, xc, yc, theta) for Yxy using an interpolation of the MacAdam
      ellipses or using FMC-1 or FMC-2.
                  Yxy
                        2D ndarray with [Y,]x,y coordinate centers.
                        If Yxy.shape[-1]==2: Y is added using the value from the Y-input argument.
                  etype
                        'fmc2', optional
                        Type color discrimination ellipse estimation to use.
                        options: 'macadam', 'fmc1', 'fmc2'
                              - 'macadam': interpolate covariance matrices of closest MacAdam ellipses (see:
                              get_macadam_ellipse?).
                              - 'fmc1': use FMC-1 from ref 2 (see get_fmc_discrimination_ellipse?).
                              - 'fmc2': use FMC-1 from ref 3 (see get fmc discrimination ellipse?).
                              - 'brown1957': interpolate covariance matrices of closest Brown1957 ellipses
                              (see: get_brown1957_ellipse?).
```

Returns:

Args:

### nsteps

10, optional

Set multiplication factor for ellipses

(nsteps=1 corresponds to approximately 1 MacAdam step,

for FMC-2, Y also has to be 10.69, see note below).

# brown1957\_weighted

True, optional

If True: use weighted averages from Table III in Brown 1957 paper, else use the straight averages.

# k\_neighbours

3, optional

Only for option 'macadam'.

Number of nearest ellipses to use to calculate ellipse at xy

# average\_cik

True, optional

Only for option 'macadam'.

If True: take distance weighted average of inverse

'covariance ellipse' elements cik.

If False: average major & minor axis lengths and ellipse orientation angles directly.

Y

None, optional

Only for option 'fmc2' (see note below).

If not None: Y = 10.69 and overrides values in Yxy.

# Note:

1. FMC-2 is almost identical to FMC-1 is Y = 10.69!; see [3]

# **References:**

- 1. MacAdam DL. Visual Sensitivities to Color Differences in Daylight\*. J Opt Soc Am. 1942;32(5):247-274.
- 2. Chickering, K.D. (1967), Optimization of the MacAdam-Modified 1965 Friele Color-Difference Formula, 57(4):537-541
- 3. Chickering, K.D. (1971), FMC Color-Difference Formulas: Clarification Concerning Usage, 61(1):118-122
- 4. Brown, WRJ. (1957). Color Discrimination of Twelve Observers\*. Journal of the Optical Society of America, 47(2), 137–143.

luxpy.color.deltaE.get\_macadam\_ellipse(xy=None, k\_neighbours=3, nsteps=10, average\_cik=True)

Estimate n-step MacAdam ellipse at CIE x,y coordinates xy by calculating average inverse covariance ellipse of the k\_neighbours closest ellipses.

# Args:

хy

None or ndarray, optional

If None: output Macadam ellipses, if not None: xy are the CIE xy coordinates for which ellipses will be estimated.

# k\_neighbours

3, optional

Number of nearest ellipses to use to calculate ellipse at xy

#### nsteps

10, optional

Set number of MacAdam steps of ellipse.

# average\_cik

True, optional

If True: take distance weighted average of inverse

'covariance ellipse' elements cik.

If False: average major & minor axis lengths and

ellipse orientation angles directly.

#### **Returns:**

#### v\_mac\_est

estimated MacAdam ellipse(s) in v-format [Rmax,Rmin,xc,yc,theta]

#### **References:**

1. MacAdam DL. Visual Sensitivities to Color Differences in Daylight\*. J Opt Soc Am. 1942;32(5):247-274.

Estimate n-step Brown1957 ellipse at CIE x,y coordinates xy by calculating average inverse covariance ellipse of the k\_neighbours closest ellipses.

# **Args:**

хy

None or ndarray, optional

If None: output Brown1957 ellipses, if not None: xy are the

CIE xy coordinates for which ellipses will be estimated.

# weighted

True, optional

If True: use weighted averages from Table III in Brown 1957 paper, else use the straight averages.

# k\_neighbours

3, optional

Number of nearest ellipses to use to calculate ellipse at xy

# nsteps

10, optional

Set number of steps of ellipse.

# average\_cik

True, optional

If True: take distance weighted average of inverse

'covariance ellipse' elements cik.

If False: average major & minor axis lengths and

ellipse orientation angles directly.

### **Returns:**

#### v\_brown\_est

estimated Brown1957 ellipse(s) in v-format [Rmax,Rmin,xc,yc,theta]

# **References:** 1. Brown, W.R.J. (1957). Color Discrimination of Twelve Observers\*. Journal of the Optical Society of America, 47(2), 137–143. https://doi.org/10.1364/JOSA.47.000137 luxpy.color.deltaE.get\_gij\_fmc(Yxy, etype='fmc2', ellipsoid=True, Y=None, cspace='Yxy') Get gij matrices describing the discrimination ellipses/ellipsoids for Yxy or xyz using FMC-1 or FMC-2. Args: Yxy 2D ndarray with [Y,]x,y coordinate centers. If Yxy.shape[-1]==2: Y is added using the value from the Y-input argument. etype 'fmc2', optional Type of FMC color discrimination equations to use (see references below). options: 'fmc1', fmc2' Y None, optional Only affects FMC-2 (see note below). If not None: Y = 10.69 and overrides values in Yxy. ellipsoid True, optional If True: return ellipsoids, else return ellipses (only if cspace == 'Yxy')! cspace 'Yxy', optional Return coefficients for Yxy-ellipses/ellipsoids ('Yxy') or XYZ ellipsoids ('xyz') Note: 1. FMC-2 is almost identical to FMC-1 is Y = 10.69!; see [2] **References:** 1. Chickering, K.D. (1967), Optimization of the MacAdam-Modified 1965 Friele Color-Difference Formula, 57(4), p.537-541 2. Chickering, K.D. (1971), FMC Color-Difference Formulas: Clarification Concerning Usage, 61(1), p.118-122 luxpy.color.deltaE.get\_fmc\_discrimination\_ellipse(Yxy=array([[100.0, 0.33333, 0.33333]]), etype='fmc2', Y=None, nsteps=10) Get discrimination ellipse(s) in v-format (R,r, xc, yc, theta) for Yxy using FMC-1 or FMC-2. Args: Yxy 2D ndarray with [Y,]x,y coordinate centers. If Yxy.shape[-1]==2: Y is added using the value from the Y-input argument. etype 'fmc2', optional Type of FMC color discrimination equations to use (see references below).

options: 'fmc1', fmc2'

Only affects FMC-2 (see note below).

None, optional

Y

If not None: Y = 10.69 and overrides values in Yxy.

#### nsteps

10, optional

Set multiplication factor for ellipses

(nsteps=1 corresponds to approximately 1 MacAdam step,

for FMC-2, Y also has to be 10.69, see note below).

#### Note:

1. FMC-2 is almost identical to FMC-1 is Y = 10.69!; see [2]

### **References:**

- 1. Chickering, K.D. (1967), Optimization of the MacAdam-Modified 1965 Friele Color-Difference Formula, 57(4), p.537-541
- 2. Chickering, K.D. (1971), FMC Color-Difference Formulas: Clarification Concerning Usage, 61(1), p.118-122

luxpy.color.deltaE.discrimination\_hotelling\_t2(Yxy1, Yxy2, etype='fmc2', ellipsoid=True, Y1=None, Y2=None, cspace='Yxy')

Check 'significance' of difference using Hotelling's T2 test on the centers Yxy1 and Yxy2 and their associate FMC-1/2 discrimination ellipses.

### Args:

# Yxy1, Yxy2

2D ndarrays with [Y,]x,y coordinate centers.

If Yxy.shape[-1]==2: Y is added using the value from the Y-input argument.

# etype

'fmc2', optional

Type of FMC color discrimination equations to use (see references below).

options: 'fmc1', fmc2'

## Y1, Y2

None, optional

Only affects FMC-2 (see note below).

If not None: Yi = 10.69 and overrides values in Yxyi.

# ellipsoid

True, optional

If True: return ellipsoids, else return ellipses (only if cspace == 'Yxy')!

# cspace

'Yxy', optional

Return coefficients for Yxy-ellipses/ellipsoids ('Yxy') or XYZ ellipsoids ('xyz')

#### **Returns:**

p

Chi-square based p-value

Т2

T2 test statistic (= mahalanobis distance on summed standard error cov. matrices)

Steps: 1. For each center coordinate, the standard error covariance matrix gij^-1 = Si/ni is determined using the FMC-1 or FMC-2 equations (see refs. 1 & 2). 2. Calculate sum of covariance matrices: SIG = S1/n1 + S2/n2 = gij1^-1 + gij2^-1 3. These are then used in Hotelling's T2 test: T2 = (xy1 - xy2).T\*(SIG^-1)\*(xy1\_xy2) 4. The T2 statistic is then tested against a Chi-square distribution with 2 or 3 degrees of freedom.

#### **References:**

- 1. Chickering, K.D. (1967), Optimization of the MacAdam-Modified 1965 Friele Color-Difference Formula, 57(4):537-541
- 2. Chickering, K.D. (1971), FMC Color-Difference Formulas: Clarification Concerning Usage, 61(1):118-122

# 4.4.7 whiteness/

рy

- \_\_init\_\_.py
- smet\_white\_loci.py

namespace luxpy

# Module with Smet et al. (2018) neutral white loci

- \_UW\_NEUTRALITY\_PARAMETERS\_SMET2014 dict with parameters of the unique white models in Smet et al. (2014)
- xyz\_to\_neutrality\_smet2018() Calculate degree of neutrality using the unique white model in Smet et al. (2014) or the normalized (max = 1) degree of chromatic adaptation model from Smet et al. (2017).
- cct\_to\_neutral\_loci\_smet2018() Calculate the most neutral appearing Duv10 in and the degree of neutrality for a specified CCT using the models in Smet et al. (2018).

### References

- 1. Smet, K. A. G. (2018). Two Neutral White Illumination Loci Based on Unique White Rating and Degree of Chromatic Adaptation. LEUKOS, 14(2), 55–67.
- 2. Smet, K., Deconinck, G., & Hanselaer, P., (2014), Chromaticity of unique white in object mode. Optics Express, 22(21), 25830–25841.
- 3. Smet, K.A.G.\*, Zhai, Q., Luo, M.R., Hanselaer, P., (2017), Study of chromatic adaptation using memory color matches, Part II: colored illuminants, Opt. Express, 25(7), pp. 8350-8365.

Added August 02, 2019.

luxpy.color.whiteness.xyz\_to\_neutrality\_smet2018(xyz10, nlocitype='uw', uw\_model='Linvar')

Calculate degree of neutrality using the unique white model in Smet et al. (2014) or the normalized (max = 1) degree of chromatic adaptation model from Smet et al. (2017).

Args:

### xyz10

ndarray with CIE 1964 10° xyz tristimulus values.

# nlocitype

'uw', optional

'uw': use unique white models published in Smet et al. (2014).

'ca': use degree of chromatic adaptation model from Smet et al. (2017).

### uw\_model

'Linvar', optional

Use Luminance invariant unique white model from Smet et al. (2014). Other options: 'L200' (200 cd/m<sup>2</sup>), 'L1000' (1000 cd/m<sup>2</sup>) and 'L2000' (2000 cd/m<sup>2</sup>).

#### **Returns:**

N

ndarray with calculated neutrality

**References:** 1. Smet, K., Deconinck, G., & Hanselaer, P., (2014), Chromaticity of unique white in object mode. Optics Express, 22(21), 25830–25841.

2. Smet, K.A.G., Zhai, Q., Luo, M.R., Hanselaer, P., (2017), Study of chromatic adaptation using memory color matches, Part II: colored illuminants, Opt. Express, 25(7), pp. 8350-8365.

# luxpy.color.whiteness.cct\_to\_neutral\_loci\_smet2018(cct, nlocitype='uw', out='duv,D')

Calculate the most neutral appearing Duv10 in and the degree of neutrality for a specified CCT using the models in Smet et al. (2018).

# Args:

cct10

ndarray CCT

# nlocitype

'uw', optional

'uw': use unique white models published in Smet et al. (2014).

'ca': use degree of chromatic adaptation model from Smet et al. (2017).

out

'duv,D', optional

Specifies requested output (other options: 'duv', 'D').

### **Returns:**

duv

ndarray with most neutral Duv10 value corresponding to the cct input.

D

ndarray with the degree of neutrality at (cct, duv).

**References:** 1. Smet, K.A.G., (2018), Two Neutral White Illumination Loci Based on Unique White Rating and Degree of Chromatic Adaptation. LEUKOS, 14(2), 55–67.

#### **Notes:**

- 1. Duv is specified in the CIE 1960 u10v10 chromatity diagram as the models were developed using CIE 1964  $10^{\circ}$  tristimulus, chromaticity and CCT values.
- 2. The parameter +0.0172 in Eq. 4b should be -0.0172.

# 4.4.8 cri/

рy

- \_\_init\_\_.py
- · colorrendition.py
- /utils/
- \_\_init\_\_.py
- init\_cri\_defaults\_database.py
- DE\_scalers.py

- helpers.py
- graphics.py

### /indices/

- \_\_init\_\_.py
- indices.py
- cie\_wrappers.py
- iestm30\_wrappers.py
- cri2012.py
- mcri.py
- cqs.py
- fci.py
- thorntoncpi.py

#### /iestm30/

- \_\_init\_\_.py
- metrics.py
- graphics.py
- metrics\_fast.py

#### /VFPX/

- \_\_inint\_\_.py
- vectorshiftmodel.py
- pixelshiftmodel.py
- VF\_PX\_models.py

namespace luxpy.cri

cri: sub-package suppporting color rendition calculations (colorrendition.py)

utils/init cri defaults database.py

```
_CRI_TYPE_DEFAULT Default cri_type.
_CRI_DEFAULTS
```

**default parameters for color fidelity and gamut area metrics** (major dict has 9 keys (04-Jul-2017): sampleset [str/dict], ref\_type [str], cieobs [str], avg [fcn handle], scale [dict], cspace [dict], catf [dict], rg pars [dict], cri specific pars [dict])

## • Supported cri-types:

- 'ciera', 'ciera-8', 'ciera-14', 'cierf',
- 'iesrf', 'iesrf-tm30-15', 'iesrf-tm30-18', 'iesrf-tm30-20',
- 'cri2012','cri2012-hl17','cri2012-hl1000','cri2012-real210',
- 'mcri',

- 'cqs-v7.5','cqs-v9.0'
- 'fci'
- 'thornton\_cpi'

process\_cri\_type\_input() load a cri\_type dict but overwrites any keys that have a non-None
input in calling function.

# utils/DE scalers.py

# linear\_scale()

Linear color rendering index scale from CIE13.3-1974/1995:

Rfi,a = 
$$100 - c1*DEi$$
,a. (c1 = 4.6)

#### log scale()

Log-based color rendering index scale from Davis & Ohno (2009):

Rfi,a = 
$$10 * ln(exp((100 - c1*DEi,a)/10) + 1)$$

### psy scale()

Psychometric based color rendering index scale from Smet et al. (2013):

Rfi,a = 
$$100 * (2 / (\exp(c1*abs(DEi,a)**(c2) + 1))) ** c3$$

# utils/helpers.py

- **\_get\_hue\_bin\_data**() Slice gamut spanned by the sample jabt, jabr and calculate hue-bin data.
- \_hue\_bin\_data\_to\_rxhj() Calculate hue bin measures: Rcshj, Rhshj, Rfhj, DEhj
- \_hue\_bin\_data\_to\_rfi() Get sample color differences DEi and calculate color fidelity values Rfi.
- \_hue\_bin\_data\_to\_rg() Calculates gamut area index, Rg.
- spd\_to\_jab\_t\_r() Calculates jab color values for a sample set illuminated with test source
  and its reference illuminant.
- spd\_to\_rg() Calculates the color gamut index of spectral data for a sample set illuminated with test source (data) with respect to some reference illuminant.
- spd\_to\_DEi() Calculates color difference (~fidelity) of spectral data between sample set illuminated with test source (data) and some reference illuminant.
- **optimize\_scale\_factor()** Optimize scale\_factor of cri-model in cri\_type such that average Rf for a set of light sources is the same as that of a target-cri (default: 'ciera')
- spd\_to\_cri() Calculates the color rendering fidelity index (CIE Ra, CIE Rf, IES Rf, CRI2012 Rf) of spectral data. Can also output Rg, Rfhi, Rcshi, Rhshi, cct, duy, ...

# utils/graphics.py

```
plot_hue_bins() Makes basis plot for Color Vector Graphic (CVG).
plot_ColorVectorGraphic() Plots Color Vector Graphic (see IES TM30).
```

# indices/indices.py

# wrapper\_functions\_for\_fidelity\_type\_metrics

```
spd_to_ciera(): CIE 13.3 1995 version
spd_to_ciera_133_1995(): CIE 13.3 1995 version
spd_to_cierf(): latest version
spd_to_cierf_224_2017(): CIE224-2017 version

spd_to_iesrf(): latest version
spd_to_iesrf_tm30(): latest version
spd_to_iesrf_tm30_15(): TM30-15 version
spd_to_iesrf_tm30_18(): TM30-18 version
spd_to_iesrf_tm30_20(): TM30-20 version (= TM30-18)

spd_to_cri2012()
spd_to_cri2012_hl17()
spd_to_cri2012_hl1000()
spd_to_cri2012_real210()
```

# wrapper\_functions\_for\_gamut\_area\_metrics

```
spd_to_iesrg(): latest version
spd_to_iesrg_tm30(): latest version
spd_to_iesrg_tm30_15(): TM30-15 version
spd_to_iesrg_tm30_18(): TM30-18 version
spd_to_iesrg_tm30_20(): TM30-20 version (= TM30-18)
```

### indices/mcri.py

# spd\_to\_mcri()

```
Calculates the memory color rendition index, Rm:
K. A. G. Smet, W. R. Ryckaert, M. R. Pointer, G. Deconinck, and P. Hanselaer, (2012)
"A memory colour quality metric for white light sources,"
Energy Build., vol. 49, no. C, pp. 216–225.
```

# indices/cqs.py

```
spd_to_cqs()
versions 7.5 and 9.0 are supported.
W. Davis and Y. Ohno,
"Color quality scale," (2010),
Opt. Eng., vol. 49, no. 3, pp. 33602–33616.
```

# iestm30/graphics.py

```
spd_to_ies_tm30_metrics() Calculates IES TM30 metrics from spectral data
```

plot\_cri\_graphics() Plots graphical information on color rendition properties based on spectral data input or dict with pre-calculated measures.

\_tm30\_process\_spd() Calculate all required parameters for plotting from spd using cri.spd\_to\_cri()

plot\_tm30\_cvg() Plot TM30 Color Vector Graphic (CVG).

plot\_tm30\_Rfi() Plot Sample Color Fidelity values (Rfi).

plot\_tm30\_Rxhj() Plot Local Chroma Shifts (Rcshj), Local Hue Shifts (Rhshj) and Local Color Fidelity values (Rfhj).

plot\_tm30\_Rcshj() Plot Local Chroma Shifts (Rcshj).

plot\_tm30\_Rhshj() Plot Local Hue Shifts (Rhshj).

plot\_tm30\_Rfhj() Plot Local Color Fidelity values (Rfhj).

plot\_tm30\_spd() Plot test SPD and reference illuminant, both normalized to the same luminous power.

plot\_tm30\_report() Plot a figure with an ANSI/IES-TM30 color rendition report.

plot\_cri\_graphics() Plots graphical information on color rendition properties based on spectral data input or dict with pre-calculated measures (cusom design). Includes Metameric uncertainty index Rt and vector-fields of color rendition shifts.

### iestm30/metrics.py

# iestm30/metrics\_fast.py

- \_cri\_ref() Calculate multiple reference illuminant spectra based on ccts for color rendering index calculations.
- **\_xyz\_to\_jab\_cam02ucs()** Calculate CAM02-UCS J'a'b' coordinates from xyz tristimulus values of sample and white point.

spd\_to\_tm30() Calculate tm30 measures from spd.

• Created for faster spectral optimization based on ANSI/IES-TM30 measures

### **VFPX**

:Module\_for\_VectorField\_and\_Pixelation\_CRI models.

• see ?luxpy.cri.VFPX

luxpy.color.cri.linear\_scale(data, scale\_factor=[4.6], scale\_max=100.0)

Linear color rendering index scale from CIE13.3-1974/1995:

Rfi,a = 
$$100 - c1*DEi$$
,a. ( $c1 = 4.6$ )

### **Args:**

#### data

float or list[floats] or ndarray

#### scale\_factor

[4.6] or list[float] or ndarray, optional

Rescales color differences before subtracting them from :scale\_max:

### scale\_max

100.0, optional

Maximum value of linear scale

#### **Returns:**

#### returns

float or list[floats] or ndarray

**References:** 1. CIE13.3-1995, "Method of Measuring and Specifying Colour Rendering Properties of Light Sources," CIE, Vienna, Austria, 1995.,ISBN 978-3-900734-57-2

luxpy.color.cri.log\_scale(data, scale\_factor=[6.73], scale\_max=100.0)

Log-based color rendering index scale from Davis & Ohno (2009):

Rfi,a = 
$$10 * ln(exp((100 - c1*DEi,a)/10) + 1)$$
.

# Args:

### data

float or list[floats] or ndarray

# scale factor

[6.73] or list[float] or ndarray, optional

Rescales color differences before subtracting them from :scale\_max:

Note that the default value is the one from cie-224-2017.

### scale\_max

100.0, optional

Maximum value of linear scale

### **Returns:**

#### returns

float or list[floats] or ndarray

**References:** 1. W. Davis and Y. Ohno, "Color quality scale," (2010), Opt. Eng., vol. 49, no. 3, pp. 33602–33616. 2. CIE224:2017. CIE 2017 Colour Fidelity Index for accurate scientific use. Vienna, Austria: CIE. (2017).

luxpy.color.cri.psy\_scale(data, scale\_factor=[0.018181818181818, 1.5, 2.0], scale\_max=100.0)
Psychometric based color rendering index scale from CRI2012:

Rfi,a =  $100 * (2 / (\exp(c1*abs(DEi,a)**(c2) + 1))) ** c3.$ 

# Args:

### data

float or list[floats] or ndarray

### scale\_factor

[1/55, 3/2, 2.0] or list[float] or ndarray, optional

Rescales color differences before subtracting them from :scale\_max:

Note that the default value is the one from (Smet et al. 2013, LRT).

### scale\_max

100.0, optional

Maximum value of linear scale

### **Returns:**

#### returns

float or list[floats] or ndarray

**References:** 1. Smet, K., Schanda, J., Whitehead, L., & Luo, R. (2013). CRI2012: A proposal for updating the CIE colour rendering index. Lighting Research and Technology, 45, 689–709.

luxpy.color.cri.\_get\_hue\_bin\_data(jabt, jabr, start\_hue=0, nhbins=16, normalized\_chroma\_ref=100)
 Slice gamut spanned by the sample jabt, jabr and calculate hue-bin data.
 Args:

# jabt

ndarray with jab sample data under test illuminant

### jabr

ndarray with jab sample data under reference illuminant

### start\_hue

0.0 or float, optional

Hue angle to start bin slicing

### nhbins

None or int, optional

- None: defaults to using the sample hues themselves as 'bins'.

In other words, the number of bins will be equal to the number of samples.

- float: number of bins to slice the sample gamut in.

# normalized\_chroma\_ref

100.0 or float, optional

Controls the size (chroma/radius) of the normalization circle/gamut.

#### **Returns:**

#### dict

Dictionary with keys:

- 'jabt', 'jabr': ndarrays with jab sample data under test & ref. illuminants
- 'DEi': ndarray with sample jab color difference between test and ref.
- 'Ct', 'Cr': chroma for each sample under test and ref.
- 'ht', 'hr': hue angles (rad.) for each sample under test and ref.
- 'ht idx', 'hr idx': hue bin indices for each sample under test and ref.
- 'jabt\_hj', 'jabr\_hj': ndarrays with hue-bin averaged jab's under test & ref. illuminants
- 'DE\_hj' : ndarray with average sample DE in each hue bin
- 'jabt\_hj\_closed', 'jabr\_hj\_closed': ndarrays with hue-bin averaged jab's under test & ref. illuminants (closed gamut: 1st == last)
- 'jabtn\_hj', 'jabrn\_hj': ndarrays with hue-bin averaged and normalized jab's under test & ref. illuminants
- 'jabtn\_hj\_closed', 'jabrn\_hj\_closed': ndarrays with hue-bin and normalized averaged jab's under test & ref. illuminants (closed gamut: 1st == last)
- 'ht\_hj', 'hr\_hj': hues (rad.) for each hue bin for test and ref.
- 'Ct\_hj', 'Cr\_hj': chroma for each hue bin for test and ref.
- 'Ctn\_hj' : normalized chroma for each hue bin for test (ref = normalized\_chroma\_ref)
- 'nhbins': number of hue bins
- 'start hue' : start hue for bin slicing
- 'normalized\_chroma\_ref': normalized chroma value for ref.
- 'dh': hue-angle arcs (°)
- 'hue\_bin\_edges': hue bin edge (rad)
- 'hbinnrs': hue bin indices for each sample under ref. (= hr\_idx)

Calculates jab color values for a sample set illuminated with test source SPD and its reference illuminant. **Args:** 

St

ndarray with spectral data (can be multiple SPDs, first axis are the wavelengths)

out

'jabt,jabr' or str, optional Specifies requested output (e.g.'jabt,jabr' or 'jabt,jabr,cct,duv')

wl

None, optional

Wavelengths (or [start, end, spacing]) to interpolate the spds in St to.

None: default to no interpolation

### cri\_type

\_CRI\_TYPE\_DEFAULT or str or dict, optional

-'str: specifies dict with default cri model parameters

```
(for supported types, see luxpy.cri._CRI_DEFAULTS['cri_types'])
```

- dict: user defined model parameters

(see e.g. luxpy.cri.\_CRI\_DEFAULTS['cierf']

for required structure)

Note that any non-None input arguments to the function will override default values in cri\_type dict.

# sampleset

None or ndarray or str, optional

Specifies set of spectral reflectance samples for cri calculations.

- None defaults to standard set for metric in cri\_type.
- ndarray: user defined set of spectral reflectance functions (.shape = (N+1, number of wavelengths);

first axis are wavelengths)

# ref\_type

None or str or ndarray, optional

Specifies type of reference illuminant type.

- None: defaults to metric\_specific reference illuminant in accordance with cri\_type.
- str: 'BB': Blackbody radiatiors,

'DL': daylightphase,

'ciera': used in CIE CRI-13.3-1995,

'cierf': used in CIE 224-2017,

'iesrf': used in TM30-15, ...

- ndarray: user defined reference SPD

# cieobs

None or dict, optional

Specifies which CMF sets to use for the calculation of the sample

XYZs and the CCT (for reference illuminant calculation).

None defaults to the one specified in :cri\_type: dict.

- key: 'xyz': str specifying CMF set for calculating xyz of samples and white
- key: 'cct': str specifying CMF set for calculating cct

#### cspace

None or dict, optional

Specifies which color space to use.

None defaults to the one specified in :cri\_type: dict.

- key: 'type': str specifying color space used to calculate color differences in.
- key: 'xyzw': None or ndarray with white point of color space
   If None: use xyzw of test / reference (after chromatic adaptation, if specified)
- other keys specify other possible parameters needed for color space calculation,

see lx.cri.\_CRI\_DEFAULTS['iesrf']['cspace'] for details.

catf

```
None or dict, optional
                        Perform explicit CAT before converting to color space coordinates.
                               - None: don't apply a cat (other than perhaps the one built
                                     into the colorspace)
                               - dict: with CAT parameters:
                                     - key: 'D': ndarray with degree of adaptation
                                     - key: 'mcat': ndarray with sensor matrix specification
                                     - key: 'xyzw': None or ndarray with white point
                                           None: use xyzw of reference otherwise transform both
                                                 test and ref to xyzw
                  cri_specific_pars
                        None or dict, optional
                        Specifies other parameters specific to type of cri
                        (e.g. maxC for CQS calculations)
                               - None: default to the one specified in :cri_type: dict.
                               - dict: user specified parameters.
                                     For its use, see for example:
                                           luxpy.cri._CRI_DEFAULTS['mcri']['cri_specific_pars']
      Returns:
                  returns
                        (ndarray, ndarray)
                        with jabt and jabr data for :out: 'jabt,jabr'
                        Other output is also possible by changing the :out: str value.
luxpy.color.cri.spd_to_rg(St, cri_type='ies-tm30', out='Rg', wl=None, sampleset=None, ref_type=None,
                                  cieobs=None, avg=None, cspace=None, catf=None, cri_specific_pars=None,
                                  rg_pars=None, fit_gamut_ellipse=False)
      Calculates the color gamut index, Rg, of spectral data.
      Args:
                  St
                        ndarray with spectral data
                        (can be multiple SPDs, first axis are the wavelengths)
                  out
                        'Rg' or str, optional
                        Specifies requested output (e.g. 'Rg,cct,duv')
                  wl
                        None, optional
                        Wavelengths (or [start, end, spacing]) to interpolate the SPDs to.
                        None: default to no interpolation
                  cri type
                         _CRI_TYPE_DEFAULT or str or dict, optional
                               - 'str: specifies dict with default cri model parameters
                                     (for supported types, see luxpy.cri._CRI_DEFAULTS['cri_types'])
```

- dict: user defined model parameters

(see e.g. luxpy.cri.\_CRI\_DEFAULTS['cierf']

for required structure)

Note that any non-None input arguments to the function will override default values in cri\_type dict.

# sampleset

None or ndarray or str, optional

Specifies set of spectral reflectance samples for cri calculations.

- None defaults to standard set for metric in cri\_type.
- ndarray: user defined set of spectral reflectance functions
   (.shape = (N+1, number of wavelengths);

first axis are wavelengths)

# ref\_type

None or str or ndarray, optional

Specifies type of reference illuminant type.

- None: defaults to metric\_specific reference illuminant in accordance with cri\_type.
- str: 'BB' : Blackbody radiatiors,

'DL': daylightphase,

'ciera': used in CIE CRI-13.3-1995,

'cierf': used in CIE 224-2017,

'iesrf': used in TM30-15, ...

- ndarray: user defined reference SPD

#### cieobs

None or dict, optional

Specifies which CMF sets to use for the calculation of the sample

XYZs and the CCT (for reference illuminant calculation).

None defaults to the one specified in :cri\_type: dict.

- key: 'xyz': str specifying CMF set for calculating xyz of samples and white
- key: 'cct': str specifying CMF set for calculating cct

## cspace

None or dict, optional

Specifies which color space to use.

None defaults to the one specified in :cri\_type: dict.

- key: 'type': str specifying color space used to calculate color differences in.
- other keys specify other possible parameters needed for color space calculation,

see lx.cri.\_CRI\_DEFAULTS['iesrf']['cspace'] for details.

## catf

None or dict, optional

Perform explicit CAT before converting to color space coordinates.

- None: don't apply a cat (other than perhaps the one built into the colorspace)
- dict: with CAT parameters:
  - key: 'D': ndarray with degree of adaptation
  - key: 'mcat': ndarray with sensor matrix specification
  - key: 'xyzw': None or ndarray with white point

None: use xyzw of reference otherwise transform both test and ref to xyzw

# cri\_specific\_pars

None or dict, optional

Specifies other parameters specific to type of cri

(e.g. maxC for CQS calculations)

- None: default to the one specified in :cri\_type: dict.
- dict: user specified parameters.

For its use, see for example:

luxpy.cri.\_CRI\_DEFAULTS['mcri']['cri\_specific\_pars']

# rg\_pars

None or dict, optional

Dict containing specifying parameters for slicing the gamut.

Dict structure:

{'nhbins' : None, 'start\_hue' : 0,

'normalize\_gamut' : False, 'normalized\_chroma\_ref': 100.0}

- key: 'nhbins': int, number of hue bins to slice gamut (None use the one specified in :cri\_type: dict).
- key: 'start\_hue': float (°), hue at which to start slicing
- key: 'normalize\_gamut': True or False:
   normalize gamut or not before calculating a gamut
   area index Rg.
- key: 'normalized\_chroma\_ref': 100.0 or float, optional Controls the size (chroma/radius)
   of the normalization circle/gamut.
- key 'use\_bin\_avg\_DEi': True or False

Note that following IES-TM30 DEhj from gamut\_slicer() is obtained by averaging the DEi per hue bin (True), and NOT by averaging the jabt and jabr per hue bin and then calculating the DEhj (False).

#### avg

None or fcn handle, optional

Averaging function (handle) for color differences, DEi

(e.g. numpy.mean, .math.rms, .math.geomean)

None use the one specified in :cri\_type: dict.

#### scale

None or dict, optional

Specifies scaling of color differences to obtain CRI.

- None use the one specified in :cri type: dict.
- dict: user specified dict with scaling parameters.
  - key: 'fcn': function handle to type of cri scale,
    - e.g
    - \* linear() scale -> (100 scale factor\*DEi),
    - \* log\_scale -> (cfr. Ohno's CQS),
    - \* psy\_scale (Smet et al.'s cri2012,See: LRT 2013)
  - key: 'cfactor': factors used in scaling function,

If None:

Scaling factor value(s) will be optimized to minimize the rms between the Rf's of the requested metric and the target metric specified in:

- key: 'opt\_cri\_type': str

- \* str: one of the preset \_CRI\_DEFAULTS
- \* dict: user specifed

(dict must contain all keys as normal)

Note that if key not in :scale: dict, then 'opt\_cri\_type' is added with default setting = 'ciera'.

 key: 'opt\_spd\_set': ndarray with set of light source spds used to optimize cfactor.
 Note that if key not in :scale: dict, then default = 'F1-F12'.

## fit\_gamut\_ellipse

fit ellipse to normalized color gamut

(extract from function using out; also stored in hue bin data['gamut ellipse fit'])

### **Returns:**

## returns

float or ndarray with Rg for :out: 'Rg'

Other output is also possible by changing the :out: str value.

E.g. out == 'Rg,data' would output an ndarray with Rg values

and a dictionary :data: with keys:

'St', 'Sr', 'cct', 'duv', 'hue\_bin\_data'

'xyzti', xyzti, 'xyztw', 'xyzri', 'xyzrw'

**References:** 1. IES TM30, Method for Evaluating Light Source Color Rendition. New York, NY: The Illuminating Engineering Society of North America.

2. A. David, P. T. Fini, K. W. Houser, Y. Ohno, M. P. Royer, K. A. G. Smet, M. Wei, and L. Whitehead, "Development of the IES method for evaluating the color rendition of light sources," Opt. Express, vol. 23, no. 12, pp. 15888–15906, 2015.

luxpy.color.cri.spd\_to\_DEi(St, cri\_type='ies-tm30', out='DEi', wl=None, sampleset=None, ref\_type=None, cieobs=None, avg=None, cspace=None, catf=None, cri\_specific\_pars=None)

Calculates color differences (~fidelity), DEi, of spectral data.

Args:

```
St
      ndarray with spectral data
      (can be multiple SPDs, first axis are the wavelengths)
out
      'DEi' or str, optional
      Specifies requested output (e.g. 'DEi,DEa,cct,duv')
wl
      None, optional
      Wavelengths (or [start, end, spacing]) to interpolate the spds in St to.
      None: default to no interpolation
cri_type
      _CRI_TYPE_DEFAULT or str or dict, optional
            - 'str: specifies dict with default cri model parameters
                   (for supported types, see luxpy.cri._CRI_DEFAULTS['cri_types'])
            - dict: user defined model parameters
                   (see e.g. luxpy.cri._CRI_DEFAULTS['cierf']
                   for required structure)
      Note that any non-None input arguments to the function will override default values in
      cri_type dict.
sampleset
      None or ndarray or str, optional
      Specifies set of spectral reflectance samples for cri calculations.
            - None defaults to standard set for metric in cri_type.
            - ndarray: user defined set of spectral reflectance functions
                   (.shape = (N+1, number of wavelengths);
                         first axis are wavelengths)
ref_type
      None or str or ndarray, optional
      Specifies type of reference illuminant type.
            - None: defaults to metric_specific reference illuminant in
                   accordance with cri_type.
            - str: 'BB': Blackbody radiatiors,
                   'DL': daylightphase,
                   'ciera': used in CIE CRI-13.3-1995,
                   'cierf': used in CIE 224-2017,
                   'iesrf': used in TM30-15, ...
            - ndarray: user defined reference SPD
cieobs
      None or dict, optional
      Specifies which CMF sets to use for the calculation of the sample
      XYZs and the CCT (for reference illuminant calculation).
      None defaults to the one specified in :cri_type: dict.
            - key: 'xyz': str specifying CMF set for calculating xyz
                   of samples and white
            - key: 'cct': str specifying CMF set for calculating cct
cspace
```

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None or dict, optional

Specifies which color space to use.

None defaults to the one specified in :cri\_type: dict.

- key: 'type': str specifying color space used to calculate color differences in.
- key: 'xyzw': None or ndarray with white point of color space If None: use xyzw of test / reference (after chromatic adaptation, if specified)
- other keys specify other possible parameters needed for color space calculation,

see lx.cri.\_CRI\_DEFAULTS['iesrf']['cspace'] for details.

#### catf

None or dict, optional

Perform explicit CAT before converting to color space coordinates.

- None: don't apply a cat (other than perhaps the one built into the colorspace)
- dict: with CAT parameters:
  - key: 'D': ndarray with degree of adaptation
  - key: 'mcat': ndarray with sensor matrix specification
  - key: 'xyzw': None or ndarray with white point

None: use xyzw of reference otherwise transform both test and ref to xyzw

# cri\_specific\_pars

None or dict, optional

Specifies other parameters specific to type of cri

(e.g. maxC for CQS calculations)

- None: default to the one specified in :cri\_type: dict.
- dict: user specified parameters.

For its use, see for example:

luxpy.cri.\_CRI\_DEFAULTS['mcri']['cri\_specific\_pars']

### **Returns:**

## returns

float or ndarray with DEi for :out: 'DEi'

Other output is also possible by changing the :out: str value.

```
luxpy.color.cri.optimize_scale_factor(cri_type, opt_scale_factor, scale_fcn, avg,
                                            rf_from_avg_rounded_rfi)
```

Optimize scale\_factor of cri-model in cri\_type such that average Rf for a set of light sources is the same as that of a target-cri (default: 'ciera').

# Args:

# cri\_type

str or dict

- 'str: specifies dict with default cri model parameters
  - (for supported types, see luxpy.cri.\_CRI\_DEFAULTS['cri\_types'])
- dict: user defined model parameters

(see e.g. luxpy.cri.\_CRI\_DEFAULTS['cierf']

for required structure)

opt scale

```
True or False
                        True: optimize scaling-factor, else do nothing and use value of
                        scaling-factor in :scale: dict.
                  scale_fcn
                        function handle to type of cri scale,
                        e.g.
                               * linear()_scale -> (100 - scale_factor*DEi),
                              * log_scale -> (cfr. Ohno's CQS),
                              * psy_scale (Smet et al.'s cri2012,See: LRT 2013)
                  avg
                        None or fcn handle
                        Averaging function (handle) for color differences, DEi
                        (e.g. numpy.mean, .math.rms, .math.geomean)
                        None use the one specified in :cri_type: dict.
      Returns:
                  scaling_factor
                        ndarray
luxpy.color.cri.spd_to_cri(St, cri_type='ies-tm30', out='Rf', wl=None, sampleset=None, ref_type=None,
                                   cieobs=None, avg=None, rf_from_avg_rounded_rfi=None, scale=None,
                                   opt_scale_factor=False, cspace=None, catf=None, cri_specific_pars=None,
                                   rg_pars=None, fit_gamut_ellipse=False)
      Calculates the color rendering fidelity index, Rf, of spectral data.
      Args:
                  St
                        ndarray with spectral data
                        (can be multiple SPDs, first axis are the wavelengths)
                  out
                        'Rf' or str, optional
                        Specifies requested output (e.g. 'Rf,cct,duv')
                  wl
                        None, optional
                        Wavelengths (or [start, end, spacing]) to interpolate the SPDs to.
                        None: default to no interpolation
                  cri_type
                        _CRI_TYPE_DEFAULT or str or dict, optional
                              - 'str: specifies dict with default cri model parameters
                                     (for supported types, see luxpy.cri._CRI_DEFAULTS['cri_types'])
                              - dict: user defined model parameters
                                     (see e.g. luxpy.cri._CRI_DEFAULTS['cierf']
                                     for required structure)
                        Note that any non-None input arguments to the function will
                        override default values in cri_type dict.
                  sampleset
                        None or ndarray or str, optional
                        Specifies set of spectral reflectance samples for cri calculations.
```

- None defaults to standard set for metric in cri\_type.
- ndarray: user defined set of spectral reflectance functions

(.shape = (N+1, number of wavelengths);

first axis are wavelengths)

# ref\_type

None or str or ndarray, optional

Specifies type of reference illuminant type.

- None: defaults to metric\_specific reference illuminant in accordance with cri\_type.
- str: 'BB': Blackbody radiatiors,

'DL': daylightphase,

'ciera': used in CIE CRI-13.3-1995,

'cierf': used in CIE 224-2017,

'iesrf': used in TM30-15, ...

- ndarray: user defined reference SPD

#### cieobs

None or dict, optional

Specifies which CMF sets to use for the calculation of the sample

XYZs and the CCT (for reference illuminant calculation).

None defaults to the one specified in :cri\_type: dict.

- key: 'xyz': str specifying CMF set for calculating xyz of samples and white
- key: 'cct': str specifying CMF set for calculating cct

### cspace

None or dict, optional

Specifies which color space to use.

None defaults to the one specified in :cri\_type: dict.

- key: 'type': str specifying color space used to calculate color differences in.
- key: 'xyzw': None or ndarray with white point of color space
   If None: use xyzw of test / reference (after chromatic adaptation, if specified)
- other keys specify other possible parameters needed for color space calculation,

see lx.cri.\_CRI\_DEFAULTS['iesrf']['cspace'] for details.

# catf

None or dict, optional

Perform explicit CAT before converting to color space coordinates.

- None: don't apply a cat (other than perhaps the one built into the colorspace)
- dict: with CAT parameters:
  - key: 'D': ndarray with degree of adaptation
  - key: 'mcat': ndarray with sensor matrix specification
  - key: 'xyzw': None or ndarray with white point

None: use xyzw of reference otherwise transform both test and ref to xyzw

#### cri\_specific\_pars

None or dict, optional

Specifies other parameters specific to type of cri

(e.g. maxC for CQS calculations)

- None: default to the one specified in :cri\_type: dict.
- dict: user specified parameters.

For its use, see for example:

luxpy.cri.\_CRI\_DEFAULTS['mcri']['cri\_specific\_pars']

## rg\_pars

None or dict, optional

Dict containing specifying parameters for slicing the gamut and calculating hue bin specific indices.

Dict structure:

{'nhbins': None, 'start\_hue': 0, 'normalize\_gamut': False, 'normalized\_chroma\_ref': 100.0}

- key: 'nhbins': int, number of hue bins to slice gamut (None use the one specified in :cri\_type: dict).
- key: 'start\_hue': float (°), hue at which to start slicing
- key: 'normalize\_gamut': True or False: normalize gamut or not before calculating a gamut area index Rg.
- key: 'normalized\_chroma\_ref': 100.0 or float, optional Controls the size (chroma/radius) of the normalization circle/gamut.
- key 'use\_bin\_avg\_DEi': True or False
   Note that following IES-TM30 DEhj from gamut\_slicer()
   is obtained by averaging the DEi per hue bin (True),
   and NOT by averaging the jabt and jabr per hue bin
   and then calculating the DEhj (False).

#### avg

None or fcn handle, optional

Averaging function (handle) for color differences, DEi

(e.g. numpy.mean, .math.rms, .math.geomean)

None use the one specified in :cri\_type: dict.

## rf\_from\_avg\_rounded\_rfi

None, optional

If None: use as specified in the :cri\_type: dict If False: calculate Rf directly from DEa.

If True: round Rfi to integer numbers and average them to Rf (method used in CIE-13.3-1995 Ra calculation)

#### scale

None or dict, optional

Specifies scaling of color differences to obtain CRI.

- None use the one specified in :cri type: dict.
- dict: user specified dict with scaling parameters.
  - key: 'fcn': function handle to type of cri scale,

```
e.g.
             * linear_scale -> (100 - scale_factor*DEi),
            * log_scale -> (cfr. Ohno's CQS),
             * psy_scale (Smet et al.'s cri2012,See: LRT 2013)
- key: 'cfactor': factors used in scaling function,
      If None:
                         Scaling factor value(s) will be optimized to
                         minimize the rms between the Rf's of the
                         requested metric and the target metric specified
                   - key: 'opt_cri_type': str
                                * str: one of the preset _CRI_DEFAULTS
                                * dict: user speciied
                                (dict must contain all keys as normal)
                         Note that if key not in :scale: dict,
                         then 'opt_cri_type' is added with default
                         setting = 'ciera'.
            - key: 'opt_spd_set': ndarray with set of light
                   source spds used to optimize cfactor.
                   Note that if key not in :scale: dict,
                   then default = 'F1-F12'.
```

# opt\_scale\_factor

True or False, optional

True: optimize scaling-factor, else do nothing and use value of scaling-factor in :scale: dict.

## fit\_gamut\_ellipse

fit ellipse to normalized color gamut (extract from function using out; also stored in hue\_bin\_data['gamut\_ellipse\_fit'])

#### **Returns:**

#### returns

float or ndarray with Rf for :out: 'Rf'
Other output is also possible by changing the :out: str value.
E.g. out == 'Rg,data' would output an ndarray with Rf values

and a dictionary :data: with keys:

- 'St, Sr': ndarray of test SPDs and corresponding ref. illuminants.
- 'xyz\_cct': xyz of white point calculate with cieobs defined for cct calculations in cri\_type['cieobs']
- 'cct, duv': CCT and Duv obtained with cieobs in cri\_type['cieobs']['cct']
- 'xyzti, xyzri': ndarray tristimulus values of test and ref. samples (obtained with with cieobs in cri\_type['cieobs']['xyz'])
- 'xyztw, xyzrw': ndarray tristimulus values of test and ref. white points (obtained with with cieobs in cri\_type['cieobs']['xyz'])
- 'DEi, DEa': ndarray with individual sample color differences DEi and average DEa between test and ref.

- 'Rf': ndarray with general color fidelity index values
- 'Rg': ndarray with color gamut area index values
- 'Rfi': ndarray with specific (sample) color fidelity indices
- 'Rfhj': ndarray with local (hue binned) fidelity indices
- 'DEhj': ndarray with local (hue binned) color differences
- 'Rcshj': ndarray with local chroma shifts indices
- 'Rhshj': ndarray with local hue shifts indices
- 'hue\_bin\_data': dict with output from \_get\_hue\_bin\_data() [see its help for more info]
- 'cri\_type': same as input (for reference purposes)

**References:** 1. IES TM30, Method for Evaluating Light Source Color Rendition. New York, NY: The Illuminating Engineering Society of North America.

- 2. A. David, P. T. Fini, K. W. Houser, Y. Ohno, M. P. Royer, K. A. G. Smet, M. Wei, and L. Whitehead, "Development of the IES method for evaluating the color rendition of light sources," Opt. Express, vol. 23, no. 12, pp. 15888–15906, 2015.
- 3. CIE224:2017. CIE 2017 Colour Fidelity Index for accurate scientific use. Vienna, Austria: CIE. (2017).
- 4. Smet, K., Schanda, J., Whitehead, L., & Luo, R. (2013). CRI2012: A proposal for updating the CIE colour rendering index. Lighting Research and Technology, 45, 689–709.
- 5. CIE13.3-1995. Method of Measuring and Specifying Colour Rendering Properties of Light Sources (Vol. CIE13.3-19). Vienna, Austria: CIE. (1995).

Calculate hue bin measures: Rcshj, Rhshj, Rfhj, DEhj.

Reshj: local chroma shift Rhshj: local hue shift

Rfhj: local (hue bin) color fidelity DEhj: local (hue bin) color differences

(See IES TM30)

# Args:

## hue\_bin\_data

Dict with hue bin data obtained with \_get\_hue\_bin\_data().

# use\_bin\_avg\_DEi

True, optional

Note that following IES-TM30 DEhj from gamut\_slicer() is obtained by averaging the DEi per hue bin (True), and NOT by averaging the jabt and jabr per hue bin and then calculating the DEhj (False).

If None: use value in rg\_pars dict in cri\_type dict!

#### scale fcn

function handle to type of cri scale,

e.g.

- \* linear()\_scale -> (100 scale\_factor\*DEi),
- \* log scale -> (cfr. Ohno's CQS),

```
* psy_scale (Smet et al.'s cri2012,See: LRT 2013)
                 scale factor
                       factors used in scaling function
     Returns:
                 returns
                       ndarrays of Rcshj, Rhshj, Rfhj, DEhj
     References: 1. IES TM30, Method for Evaluating Light Source Color Rendition. New York, NY: The Illumi-
            nating Engineering Society of North America.
luxpy.color.cri._hue_bin_data_to_rfi(hue_bin_data=None, cri_type='ies-tm30', scale_factor=None,
                                              scale_fcn=None)
     Get sample color differences DEi and calculate color fidelity values Rfi.
     Rfi: Sample color fidelity
     DEi: Sample color differences
     (See IES TM30)
     Args:
                 hue bin data
                       Dict with hue bin data obtained with get hue bin data().
                 scale fcn
                       function handle to type of cri scale,
                       e.g.
                             * linear()_scale -> (100 - scale_factor*DEi),
                             * log_scale -> (cfr. Ohno's CQS),
                             * psy_scale (Smet et al.'s cri2012,See: LRT 2013)
                 scale_factor
                       factors used in scaling function
     Returns:
                 returns
                       ndarrays of Rfi, DEi
     References: 1. IES TM30, Method for Evaluating Light Source Color Rendition. New York, NY: The Illumi-
            nating Engineering Society of North America.
luxpy.color.cri._hue_bin_data_to_rg(hue_bin_data, max_scale=100, normalize_gamut=False)
     Calculates gamut area index, Rg.
     Args:
                 hue_bin_data
                       Dict with hue bin data obtained with _get_hue_bin_data().
                 max scale
                       100.0, optional
                       Value of Rg when Rf = \max_{s} (i.e. DEavg = 0)
                 normalize_gamut
                       False, optional
                       True normalizes the gamut of test to that of ref.
                       (perfect agreement results in circle).
```

```
out
                        'Rg', optional
                        Specifies which variables to output as ndarray
      Returns:
                  Rg
                        float or ndarray with gamut area indices Rg.
luxpy.color.cri.spd_to_ciera(SPD, out='Rf', wl=None)
      Wrapper function the 'ciera' color rendition (fidelity) metric (CIE 13.3-1995).
      Args:
                  SPD
                        ndarray with spectral data
                        (can be multiple SPDs, first axis are the wavelengths)
                  wl
                        None, optional
                        Wavelengths (or [start, end, spacing]) to interpolate :SPD: to.
                        None: default to no interpolation
                  out
                        'Rf' or str, optional
                        Specifies requested output (e.g. 'Rf,Rfi,cct,duv')
      Returns:
                  returns
                        float or ndarray with CIE13.3 Ra for :out: 'Rf'
                        Other output is also possible by changing the :out: str value.
      References: 1. CIE13.3-1995. Method of Measuring and Specifying Colour Rendering Properties of Light
            Sources (Vol. CIE13.3-19). Vienna, Austria: CIE. (1995).
luxpy.color.cri.spd_to_cierf(SPD, out='Rf', wl=None)
      Wrapper function the 'cierf' color rendition (fidelity) metric (CIE224-2017).
      Args:
                  SPD
                        ndarray with spectral data (can be multiple SPDs,
                        first axis are the wavelengths)
                  wl
                        None, optional
                        Wavelengths (or [start, end, spacing]) to interpolate :SPD: to.
                        None: default to no interpolation
                  out
                        'Rf' or str, optional
                        Specifies requested output (e.g. 'Rf,Rfi,cct,duv')
      Returns:
                  returns
                        float or ndarray with CIE224-2017 Rf for :out: 'Rf'
                        Other output is also possible by changing the :out: str value.
      References: 1. CIE224:2017. CIE 2017 Colour Fidelity Index for accurate scientific use. Vienna, Austria: CIE.
            (2017).
luxpy.color.cri.spd_to_ciera_133_1995(SPD, out='Rf', wl=None)
      Wrapper function the 'ciera' color rendition (fidelity) metric (CIE 13.3-1995).
```

```
Args:
                  SPD
                        ndarray with spectral data
                        (can be multiple SPDs, first axis are the wavelengths)
                  wl
                        None, optional
                        Wavelengths (or [start, end, spacing]) to interpolate :SPD: to.
                        None: default to no interpolation
                  out
                        'Rf' or str, optional
                        Specifies requested output (e.g. 'Rf,Rfi,cct,duv')
      Returns:
                  returns
                        float or ndarray with CIE13.3 Ra for :out: 'Rf'
                        Other output is also possible by changing the :out: str value.
      References: 1. CIE13.3-1995. Method of Measuring and Specifying Colour Rendering Properties of Light
            Sources (Vol. CIE13.3-19). Vienna, Austria: CIE. (1995).
luxpy.color.cri.spd_to_cierf_224_2017(SPD, out='Rf', wl=None)
      Wrapper function the 'cierf' color rendition (fidelity) metric (CIE224-2017).
      Args:
                  SPD
                        ndarray with spectral data (can be multiple SPDs,
                        first axis are the wavelengths)
                  wl
                        None, optional
                        Wavelengths (or [start, end, spacing]) to interpolate :SPD: to.
                        None: default to no interpolation
                  out
                        'Rf' or str, optional
                        Specifies requested output (e.g. 'Rf,Rfi,cct,duv')
      Returns:
                  returns
                        float or ndarray with CIE224-2017 Rf for :out: 'Rf'
                        Other output is also possible by changing the :out: str value.
      References: 1. CIE224:2017. CIE 2017 Colour Fidelity Index for accurate scientific use. Vienna, Austria: CIE.
            (2017).
luxpy.color.cri.spd_to_iesrf(SPD, out='Rf', wl=None, cri_type='iesrf-tm30-20')
      Wrapper function for the 'iesrf' color fidelity index (IES TM30-20 = TM30-18).
      Args:
                  SPD
                        ndarray with spectral data (can be multiple SPDs,
                        first axis are the wavelengths)
                  wl
                        None, optional
                        Wavelengths (or [start, end, spacing]) to interpolate the SPDs to.
                        None: default to no interpolation
```

out

'Rf' or str, optional

Specifies requested output (e.g. 'Rf,Rfi,cct,duv')

#### **Returns:**

#### returns

float or ndarray with IES TM30-20 Rf for :out: 'Rf'

Other output is also possible by changing the :out: str value.

**References:** 1. IES TM30 (99, 4880 spectrally uniform samples)

- 2. A. David, P. T. Fini, K. W. Houser, Y. Ohno, M. P. Royer, K. A. G. Smet, M. Wei, and L. Whitehead, "Development of the IES method for evaluating the color rendition of light sources," Opt. Express, vol. 23, no. 12, pp. 15888–15906, 2015.
- 3. K. A. G. Smet, A. David, and L. Whitehead, "Why color space uniformity and sample set spectral uniformity are essential for color rendering measures," LEUKOS, vol. 12, no. 1–2, pp. 39–50, 2016

luxpy.color.cri.spd\_to\_iesrg(SPD, out='Rg', wl=None, cri\_type='iesrf-tm30-20')

Wrapper function for the 'spd\_to\_rg' color gamut area index (IES TM30-18 = TM30-20). **Args:** 

SPD

ndarray with spectral data (can be multiple SPDs,

first axis are the wavelengths)

wl

None, optional

Wavelengths (or [start, end, spacing]) to interpolate the SPDs to.

None: default to no interpolation

out

'Rg' or str, optional

Specifies requested output (e.g. 'Rg,Rf,Rfi,cct,duv')

### **Returns:**

## returns

float or ndarray with IES TM30-20 Rg for :out: 'Rg'

Other output is also possible by changing the :out: str value.

**References:** 1. IES TM30 (99, 4880 spectrally uniform samples)

- 2. A. David, P. T. Fini, K. W. Houser, Y. Ohno, M. P. Royer, K. A. G. Smet, M. Wei, and L. Whitehead, "Development of the IES method for evaluating the color rendition of light sources," Opt. Express, vol. 23, no. 12, pp. 15888–15906, 2015.
- 3. K. A. G. Smet, A. David, and L. Whitehead, "Why color space uniformity and sample set spectral uniformity are essential for color rendering measures," LEUKOS, vol. 12, no. 1–2, pp. 39–50, 2016

luxpy.color.cri.spd\_to\_iesrf\_tm30(SPD, out='Rf', wl=None, cri\_type='iesrf-tm30-20')

Wrapper function for the 'iesrf' color fidelity index (IES TM30-20 = TM30-18).

**Args:** 

**SPD** 

ndarray with spectral data (can be multiple SPDs,

first axis are the wavelengths)

wl

None, optional

out

'Rf' or str, optional

Specifies requested output (e.g. 'Rf,Rfi,cct,duv')

#### **Returns:**

#### returns

float or ndarray with IES TM30-20 Rf for :out: 'Rf'

Other output is also possible by changing the :out: str value.

**References:** 1. IES TM30 (99, 4880 spectrally uniform samples)

- 2. A. David, P. T. Fini, K. W. Houser, Y. Ohno, M. P. Royer, K. A. G. Smet, M. Wei, and L. Whitehead, "Development of the IES method for evaluating the color rendition of light sources," Opt. Express, vol. 23, no. 12, pp. 15888–15906, 2015.
- 3. K. A. G. Smet, A. David, and L. Whitehead, "Why color space uniformity and sample set spectral uniformity are essential for color rendering measures," LEUKOS, vol. 12, no. 1–2, pp. 39–50, 2016

luxpy.color.cri.spd\_to\_iesrg\_tm30(SPD, out='Rg', wl=None, cri\_type='iesrf-tm30-20')

Wrapper function for the 'spd\_to\_rg' color gamut area index (IES TM30-18 = TM30-20).

**Args:** 

**SPD** 

ndarray with spectral data (can be multiple SPDs,

first axis are the wavelengths)

wl

None, optional

Wavelengths (or [start, end, spacing]) to interpolate the SPDs to.

None: default to no interpolation

out

'Rg' or str, optional

Specifies requested output (e.g. 'Rg,Rf,Rfi,cct,duv')

## **Returns:**

### returns

float or ndarray with IES TM30-20 Rg for :out: 'Rg'

Other output is also possible by changing the :out: str value.

**References:** 1. IES TM30 (99, 4880 spectrally uniform samples)

- 2. A. David, P. T. Fini, K. W. Houser, Y. Ohno, M. P. Royer, K. A. G. Smet, M. Wei, and L. Whitehead, "Development of the IES method for evaluating the color rendition of light sources," Opt. Express, vol. 23, no. 12, pp. 15888–15906, 2015.
- 3. K. A. G. Smet, A. David, and L. Whitehead, "Why color space uniformity and sample set spectral uniformity are essential for color rendering measures," LEUKOS, vol. 12, no. 1–2, pp. 39–50, 2016

luxpy.color.cri.spd\_to\_iesrf\_tm30\_15(SPD, out='Rf', wl=None, cri\_type='iesrf-tm30-15')

Wrapper function for the 'iesrf' color fidelity index (IES TM30-15).

Args:

**SPD** 

ndarray with spectral data (can be multiple SPDs,

first axis are the wavelengths)

wl

None, optional

out

'Rf' or str, optional

Specifies requested output (e.g. 'Rf,Rfi,cct,duv')

#### **Returns:**

## returns

float or ndarray with IES TM30-15 Rf for :out: 'Rf'

Other output is also possible by changing the :out: str value.

**References:** 1. IES TM30 (99, 4880 spectrally uniform samples)

- 2. A. David, P. T. Fini, K. W. Houser, Y. Ohno, M. P. Royer, K. A. G. Smet, M. Wei, and L. Whitehead, "Development of the IES method for evaluating the color rendition of light sources," Opt. Express, vol. 23, no. 12, pp. 15888–15906, 2015.
- 3. K. A. G. Smet, A. David, and L. Whitehead, "Why color space uniformity and sample set spectral uniformity are essential for color rendering measures," LEUKOS, vol. 12, no. 1–2, pp. 39–50, 2016

luxpy.color.cri.spd\_to\_iesrg\_tm30\_15(SPD, out='Rg', wl=None, cri\_type='iesrf-tm30-15')

Wrapper function for the 'spd\_to\_rg' color gamut area index (IES TM30-15).

#### Args:

**SPD** 

ndarray with spectral data (can be multiple SPDs,

first axis are the wavelengths)

wl

None, optional

Wavelengths (or [start, end, spacing]) to interpolate the SPDs to.

None: default to no interpolation

out

'Rg' or str, optional

Specifies requested output (e.g. 'RgRf,Rfi,cct,duv')

## **Returns:**

### returns

float or ndarray with IES TM30-15 Rg for :out: 'Rg'

Other output is also possible by changing the :out: str value.

**References:** 1. IES TM30 (99, 4880 spectrally uniform samples)

- 2. A. David, P. T. Fini, K. W. Houser, Y. Ohno, M. P. Royer, K. A. G. Smet, M. Wei, and L. Whitehead, "Development of the IES method for evaluating the color rendition of light sources," Opt. Express, vol. 23, no. 12, pp. 15888–15906, 2015.
- 3. K. A. G. Smet, A. David, and L. Whitehead, "Why color space uniformity and sample set spectral uniformity are essential for color rendering measures," LEUKOS, vol. 12, no. 1–2, pp. 39–50, 2016

luxpy.color.cri.spd\_to\_iesrf\_tm30\_18(SPD, out='Rf', wl=None, cri\_type='iesrf-tm30-18')

Wrapper function for the 'iesrf' color fidelity index (IES TM30-18).

## Args:

#### SPD

ndarray with spectral data (can be multiple SPDs,

first axis are the wavelengths)

wl

None, optional

out

'Rf' or str, optional

Specifies requested output (e.g. 'Rf,Rfi,cct,duv')

#### **Returns:**

#### returns

float or ndarray with IES TM30-18 Rf for :out: 'Rf'

Other output is also possible by changing the :out: str value.

**References:** 1. IES TM30 (99, 4880 spectrally uniform samples)

- 2. A. David, P. T. Fini, K. W. Houser, Y. Ohno, M. P. Royer, K. A. G. Smet, M. Wei, and L. Whitehead, "Development of the IES method for evaluating the color rendition of light sources," Opt. Express, vol. 23, no. 12, pp. 15888–15906, 2015.
- 3. K. A. G. Smet, A. David, and L. Whitehead, "Why color space uniformity and sample set spectral uniformity are essential for color rendering measures," LEUKOS, vol. 12, no. 1–2, pp. 39–50, 2016

luxpy.color.cri.spd\_to\_iesrg\_tm30\_18(SPD, out='Rg', wl=None, cri\_type='iesrf-tm30-18')

Wrapper function for the 'spd\_to\_rg' color gamut area index (IES TM30-18).

# **Args:**

**SPD** 

ndarray with spectral data (can be multiple SPDs,

first axis are the wavelengths)

wl

None, optional

Wavelengths (or [start, end, spacing]) to interpolate the SPDs to.

None: default to no interpolation

out

'Rg' or str, optional

Specifies requested output (e.g. 'Rg,Rf,Rfi,cct,duv')

## **Returns:**

### returns

float or ndarray with IES TM30-18 Rg for :out: 'Rg'

Other output is also possible by changing the :out: str value.

**References:** 1. IES TM30 (99, 4880 spectrally uniform samples)

- 2. A. David, P. T. Fini, K. W. Houser, Y. Ohno, M. P. Royer, K. A. G. Smet, M. Wei, and L. Whitehead, "Development of the IES method for evaluating the color rendition of light sources," Opt. Express, vol. 23, no. 12, pp. 15888–15906, 2015.
- 3. K. A. G. Smet, A. David, and L. Whitehead, "Why color space uniformity and sample set spectral uniformity are essential for color rendering measures," LEUKOS, vol. 12, no. 1–2, pp. 39–50, 2016

luxpy.color.cri.spd\_to\_iesrf\_tm30\_20(SPD, out='Rf', wl=None, cri\_type='iesrf-tm30-20')

Wrapper function for the 'iesrf' color fidelity index (IES TM30-20 = TM30-18).

Args:

#### **SPD**

ndarray with spectral data (can be multiple SPDs,

first axis are the wavelengths)

wl

None, optional

out

'Rf' or str, optional

Specifies requested output (e.g. 'Rf,Rfi,cct,duv')

#### **Returns:**

#### returns

float or ndarray with IES TM30-20 Rf for :out: 'Rf'

Other output is also possible by changing the :out: str value.

**References:** 1. IES TM30 (99, 4880 spectrally uniform samples)

- 2. A. David, P. T. Fini, K. W. Houser, Y. Ohno, M. P. Royer, K. A. G. Smet, M. Wei, and L. Whitehead, "Development of the IES method for evaluating the color rendition of light sources," Opt. Express, vol. 23, no. 12, pp. 15888–15906, 2015.
- 3. K. A. G. Smet, A. David, and L. Whitehead, "Why color space uniformity and sample set spectral uniformity are essential for color rendering measures," LEUKOS, vol. 12, no. 1–2, pp. 39–50, 2016

luxpy.color.cri.spd\_to\_iesrg\_tm30\_20(SPD, out='Rg', wl=None, cri\_type='iesrf-tm30-20')

Wrapper function for the 'spd\_to\_rg' color gamut area index (IES TM30-18 = TM30-20).

Args:

#### **SPD**

ndarray with spectral data (can be multiple SPDs,

first axis are the wavelengths)

wl

None, optional

Wavelengths (or [start, end, spacing]) to interpolate the SPDs to.

None: default to no interpolation

out

'Rg' or str, optional

Specifies requested output (e.g. 'Rg,Rf,Rfi,cct,duv')

## **Returns:**

### returns

float or ndarray with IES TM30-20 Rg for :out: 'Rg'

Other output is also possible by changing the :out: str value.

**References:** 1. IES TM30 (99, 4880 spectrally uniform samples)

- 2. A. David, P. T. Fini, K. W. Houser, Y. Ohno, M. P. Royer, K. A. G. Smet, M. Wei, and L. Whitehead, "Development of the IES method for evaluating the color rendition of light sources," Opt. Express, vol. 23, no. 12, pp. 15888–15906, 2015.
- 3. K. A. G. Smet, A. David, and L. Whitehead, "Why color space uniformity and sample set spectral uniformity are essential for color rendering measures," LEUKOS, vol. 12, no. 1–2, pp. 39–50, 2016

# luxpy.color.cri.spd\_to\_cri2012(SPD, out='Rf', wl=None)

Wrapper function for the 'cri2012' color rendition (fidelity) metric with the spectally uniform HL17 mathematical sampleset.

**Args:** 

#### **SPD**

ndarray with spectral data (can be multiple SPDs, first axis are the wavelengths)

wl

None, optional

```
Wavelengths (or [start, end, spacing]) to interpolate the SPDs to.
                        None: default to no interpolation
                  out
                        'Rf' or str, optional
                        Specifies requested output (e.g. 'Rf,Rfi,cct,duv')
      Returns:
                  returns
                        float or ndarray with CRI2012 Rf for :out: 'Rf'
                        Other output is also possible by changing the :out: str value.
      References:
            ..[1] Smet, K., Schanda, J., Whitehead, L., & Luo, R. (2013). CRI2012: A proposal for updating the
                  CIE colour rendering index. Lighting Research and Technology, 45, 689–709. Retrieved from http:
                  //lrt.sagepub.com/content/45/6/689
luxpy.color.cri.spd_to_cri2012_hl17(SPD, out='Rf', wl=None)
      Wrapper function for the 'cri2012' color rendition (fidelity) metric with the spectally uniform HL17 mathematical
      sampleset.
      Args:
                  SPD
                        ndarray with spectral data (can be multiple SPDs,
                        first axis are the wavelengths)
                  wl
                        None, optional
                        Wavelengths (or [start, end, spacing]) to interpolate the SPDs to.
                        None: default to no interpolation
                  out
                        'Rf' or str, optional
                        Specifies requested output (e.g. 'Rf,Rfi,cct,duv')
      Returns:
                  returns
                        float or ndarray with CRI2012 Rf for :out: 'Rf'
                        Other output is also possible by changing the :out: str value.
      Reference: 1. Smet, K., Schanda, J., Whitehead, L., & Luo, R. (2013). CRI2012: A proposal for updating the
            CIE colour rendering index. Lighting Research and Technology, 45, 689–709.
luxpy.color.cri.spd_to_cri2012_hl1000(SPD, out='Rf', wl=None)
      Wrapper function for the 'cri2012' color rendition (fidelity) metric with the spectally uniform Hybrid HL1000
      sampleset.
      Args:
                  SPD
                        ndarray with spectral data (can be multiple SPDs,
                        first axis are the wavelengths)
                  wl
                        None, optional
                        Wavelengths (or [start, end, spacing]) to interpolate the SPDs to.
                        None: default to no interpolation
                  out
                        'Rf' or str, optional
```

Specifies requested output (e.g. 'Rf,Rfi,cct,duv')

```
Returns:
                  returns
                        float or ndarray with CRI2012 Rf for :out: 'Rf'
                        Other output is also possible by changing the :out: str value.
      Reference: 1. Smet, K., Schanda, J., Whitehead, L., & Luo, R. (2013). CRI2012: A proposal for updating the
            CIE colour rendering index. Lighting Research and Technology, 45, 689–709.
luxpy.color.cri.spd_to_cri2012_real210(SPD, out='Rf', wl=None)
      Wrapper function the 'cri2012' color rendition (fidelity) metric with the Real-210 sampleset (normally for special
      color rendering indices).
      Args:
                  SPD
                        ndarray with spectral data (can be multiple SPDs,
                        first axis are the wavelengths)
                  wl
                        None, optional
                        Wavelengths (or [start, end, spacing]) to interpolate the SPDs to.
                        None: default to no interpolation
                  out
                        'Rf' or str, optional
                        Specifies requested output (e.g. 'Rf,Rfi,cct,duv')
      Returns:
                  returns
                        float or ndarray with CRI2012 Rf for :out: 'Rf'
                        Other output is also possible by changing the :out: str value.
      Reference: 1. Smet, K., Schanda, J., Whitehead, L., & Luo, R. (2013). CRI2012: A proposal for updating the
            CIE colour rendering index. Lighting Research and Technology, 45, 689–709.
luxpy.color.cri.spd_to_mcri(SPD, D=0.9, E=None, Yb=20.0, out='Rm', wl=None)
      Calculates the MCRI or Memory Color Rendition Index, Rm
      Args:
                  SPD
                        ndarray with spectral data (can be multiple SPDs,
                              first axis are the wavelengths)
                  D
                        0.9, optional
                        Degree of adaptation.
                  \mathbf{E}
                        None, optional
                        Illuminance in lux
                              (used to calculate La = (Yb/100)*(E/pi) to then calculate D
                              following the 'cat02' model).
                        If None: the degree is determined by :D:
                              If (:E: is not None) & (:Yb: is None): :E: is assumed to contain
                              the adapting field luminance La (cd/m<sup>2</sup>).
                  Yb
```

Luminance factor of background. (used when calculating La from E)

20.0, optional

```
If None, E contains La (cd/m^2).
                  out
                        'Rm' or str, optional
                        Specifies requested output (e.g. 'Rm,Rmi,cct,duv')
                  wl
                        None, optional
                        Wavelengths (or [start, end, spacing]) to interpolate the SPDs to.
                        None: default to no interpolation
      Returns:
                  returns
                        float or ndarray with MCRI Rm for :out: 'Rm'
                        Other output is also possible by changing the :out: str value.
      References: 1. K.A.G. Smet, W.R. Ryckaert, M.R. Pointer, G. Deconinck, P. Hanselaer, (2012) "A memory
            colour quality metric for white light sources," Energy Build., vol. 49, no. C, pp. 216–225.
luxpy.color.cri.spd_to_cqs(SPD, version='v9.0', out='Qa', wl=None)
      Calculates CQS Qa (Qai) or Qf (Qfi) or Qp (Qpi) for versions v9.0 or v7.5.
      Args:
                  SPD
                        ndarray with spectral data (can be multiple SPDs,
                        first axis are the wavelengths)
                  version
                        'v9.0' or 'v7.5', optional
                  out
                        'Qa' or str, optional
                        Specifies requested output (e.g. 'Qa,Qai,Qf,cct,duv')
                  wl
                        None, optional
                        Wavelengths (or [start, end, spacing]) to interpolate the SPDs to.
                        None: default to no interpolation
      Returns:
                  returns
                        float or ndarray with CQS Qa for :out: 'Qa'
                        Other output is also possible by changing the :out: str value.
      References: 1. W. Davis and Y. Ohno, "Color quality scale," (2010), Opt. Eng., vol. 49, no. 3, pp.
            33602-33616.
luxpy.color.cri.spd_to_fci(spd, use_cielab=True)
      Calculate Feeling of Contrast Index (FCI).
      Args:
                  spd
                        ndarray with spectral power distribution(s) of the test light source(s).
                  use_cielab
                              True, optional
                        True: use original formulation of FCI, which adopts a CIECAT94
                        chromatic adaptation transform followed by a conversion to
                        CIELAB coordinates before calculating the gamuts.
                        False: use CIECAM02 coordinates and embedded CAT02 transform.
```

```
Returns:
                  fci
                        ndarray with FCI values.
      References: 1. Hashimoto, K., Yano, T., Shimizu, M., & Nayatani, Y. (2007). New method for specifying color-
            rendering properties of light sources based on feeling of contrast. Color Research and Application, 32(5),
            361-371.
luxpy.color.cri.spd_to_thornton_cpi(spd)
      Calculate Thornton's Color Preference Index (CPI).
      Args:
                  spd
                        nd array with spectral power distribution(s) of the test light source(s).
      Returns:
                  cpi
                        ndarray with CPI values.
      Reference: 1. Thornton, W. A. (1974). A Validation of the Color-Preference Index. Journal of the Illuminating
            Engineering Society, 4(1), 48–52.
luxpy.color.cri.plot_hue_bins(hbins=16, start_hue=0.0, scalef=100, plot_axis_labels=False,
                                      bin_labels='#', plot_edge_lines=True, plot_center_lines=False,
                                      plot_bin_colors=True, plot_10_20_circles=False, axtype='polar', ax=None,
                                      force_CVG_layout=False, hbin_color_map=None)
      Makes basis plot for Color Vector Graphic (CVG).
      Args:
                  hbins
                        16 or ndarray with sorted hue bin centers (°), optional
                  start_hue
                        0.0, optional
                  scalef
                        100, optional
                        Scale factor for graphic.
                  plot_axis_labels
                        False, optional
                        Turns axis ticks on/off (True/False).
                  bin labels
                        None or list[str] or '#', optional
                        Plots labels at the bin center hues.
                              - None: don't plot.
                              - list[str]: list with str for each bin.
                                    (len(:bin labels:) = :nhbins:)
                              - '#': plots number.
                  plot_edge_lines
                        True or False, optional
                        Plot grey bin edge lines with '-'.
                  plot_center_lines
                        False or True, optional
                        Plot colored lines at 'center' of hue bin.
                  plot bin colors
```

```
Colorize hue bins.
                  plot_10_20_circles
                        False, optional
                        If True and :axtype: == 'cart': Plot white circles at
                        80%, 90%, 100%, 110% and 120% of :scalef:
                  axtype
                        'polar' or 'cart', optional
                        Make polar or Cartesian plot.
                  ax
                        None or 'new' or 'same', optional
                              - None or 'new' creates new plot
                              - 'same': continue plot on same axes.
                              - axes handle: plot on specified axes.
                 force CVG layout
                        False or True, optional
                        True: Force plot of basis of CVG on first encounter.
                 hbin_color_map
                        ndarray with predefined RGB color map
                        If None or hbin_color_map.shape[0]<nhbins: cmap will be created, else use values in
                        ndarray.
     Returns:
                  returns
                        gcf(), gca(), list with rgb colors for hue bins (for use in other plotting fcns)
luxpy.color.cri.plot_ColorVectorGraphic(jabt, jabr, hbins=16, start_hue=0.0, scalef=100,
                                                  plot axis labels=False, bin labels=None,
                                                  plot_edge_lines=True, plot_center_lines=False,
                                                  plot_bin_colors=True, plot_10_20_circles=True,
                                                  plot_vectors=True, gamut_line_color=None,
                                                  gamut_line_style='-', gamut_line_marker='o',
                                                  gamut_line_label=None, axtype='polar', ax=None,
                                                  force_CVG_layout=False, hbin_color_map=None,
                                                  hvector_color_map=None, jabti=None, jabri=None,
                                                  hbinnr=None)
     Plot Color Vector Graphic (CVG).
     Args:
                 jabt
                        ndarray with jab data under test SPD
                 jabr
                        ndarray with jab data under reference SPD
                 hbins
                        16 or ndarray with sorted hue bin centers (°), optional
                  start hue
                        0.0, optional
                 scalef
                        100, optional
```

True, optional

Scale factor for graphic.

```
plot_axis_labels
      False, optional
      Turns axis ticks on/off (True/False).
bin labels
      None or list[str] or '#', optional
      Plots labels at the bin center hues.
            - None: don't plot.
            - list[str]: list with str for each bin.
                   (len(:bin_labels:) = :nhbins:)
            - '#': plots number.
plot_edge_lines
      True or False, optional
      Plot grey bin edge lines with '-'.
plot_center_lines
      False or True, optional
      Plot colored lines at 'center' of hue bin.
plot_bin_colors
      True, optional
      Colorize hue-bins.
plot_10_20_circles
      True, optional
      If True and :axtype: == 'cart': Plot white circles at
      80%, 90%, 100%, 110% and 120% of :scalef:
plot_vectors
      True, optional
      True: plot vectors from reference to test colors.
gamut_line_color
      'grey', optional
      Color to plot the test color gamut in.
gamut_line_style
      '-', optional
      Line style to plot the test color gamut in.
gamut_line_marker
      'o', optional
      Markers to plot the test color gamut points for each hue bin in
      (only used when plot_vectors = False).
gamut_line_label
      None, optional
      Label for gamut line. (only used when plot_vectors = False).
axtype
      'polar' or 'cart', optional
      Make polar or Cartesian plot.
ax
      None or 'new' or 'same', optional
```

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- None or 'new' creates new plot
- 'same': continue plot on same axes.
- axes handle: plot on specified axes.

# force\_CVG\_layout

False or True, optional

True: Force plot of basis of CVG.

# hbin\_color\_map

ndarray with predefined RGB color map for the hue bins

If None or hbin\_color\_map.shape[0]<nhbins: cmap will be created, else use values in ndarray.

# hvector\_color\_map

in ndarray.

ndarray with predefined RGB color map for the color shift vectors in each hue bin. If None or hvector\_color\_map.shape[0]<hbins: cmap will be created, else use values

## jabti

None, optional

ndarray with jab data of all samples under test SPD (scaled to 'unit' circle)

If not None: plot chromaticity coordinates of test samples relative to

the mean chromaticity of the samples under the reference illuminant.

# jabri

None, optional

ndarray with jab data of all samples under reference SPD (scaled to 'unit' circle)

Must be supplied when jabti is not None!

# hbinnr

None, optional

ndarray with hue bin number of each sample.

Must be supplied when jabti is not None!

#### **Returns:**

#### returns

gcf(), gca(), list with rgb colors for hue bins (for use in other plotting fcns)

```
luxpy.color.cri.spd_to_ies_tm30_metrics(St, cri_type=None, hbins=16, start_hue=0.0, scalef=100, vf_model_type='M6', vf_pcolorshift={'Cref': 40, 'href': array([3.7835, 3.3161, 2.8272, 1.9093, 5.2787, 4.3081, 0.37762, 6.2055, 1.4564, 0.88927]), 'labels': array(['5B', '5BG', '5G', '5GY', '5P', '5PB', '5R', '5RP', '5Y', '5YR'], dtype=object), 'sig': 0.3}, scale_vf_chroma_to_sample_chroma=False)
```

Calculates IES TM30 metrics from spectral data.

## Args:

St

numpy.ndarray with spectral data

#### cri\_type

None, optional

If None: defaults to cri\_type = 'iesrf'.

Not none values of :hbins:, :start\_hue: and :scalef: overwrite

input in cri\_type['rg\_pars']

#### **hbins**

None or numpy.ndarray with sorted hue bin centers (°), optional

#### start\_hue

None, optional

#### scalef

None, optional

Scale factor for reference circle.

#### vf\_pcolorshift

VF PCOLORSHIFT or user defined dict, optional

The polynomial models of degree 5 and 6 can be fully specified or summarized by the model parameters themselved OR by calculating the dCoverC and dH at resp. 5 and 6 hues. :VF\_pcolorshift: specifies these hues and chroma level.

# scale\_vf\_chroma\_to\_sample\_chroma

False, optional

Scale chroma of reference and test vf fields such that average of binned reference chroma equals that of the binned sample chroma before calculating hue bin metrics.

#### **Returns:**

#### data

Dictionary with color rendering data:

- 'St, Sr': ndarray of test SPDs and corresponding ref. illuminants.
- 'xyz\_cct': xyz of white point calculate with cieobs defined for cct calculations in cri\_type['cieobs']
- 'cct, duv': CCT and Duv obtained with cieobs in cri\_type['cieobs']['cct']
- 'xyzti, xyzri': ndarray tristimulus values of test and ref. samples (obtained with with cieobs in cri\_type['cieobs']['xyz'])
- 'xyztw, xyzrw': ndarray tristimulus values of test and ref. white points (obtained with with cieobs in cri\_type['cieobs']['xyz'])
- 'DEi, DEa': ndarray with individual sample color differences DEi and average DEa between test and ref.
- 'Rf': ndarray with general color fidelity index values
- 'Rg': ndarray with color gamut area index values
- 'Rfi': ndarray with specific (sample) color fidelity indices
- 'Rfhj': ndarray with local (hue binned) fidelity indices
- 'DEhj': ndarray with local (hue binned) color differences
- 'Rcshj': ndarray with local chroma shifts indices
- 'Rhshj': ndarray with local hue shifts indices
- 'hue\_bin\_data': dict with output from \_get\_hue\_bin\_data() [see its help for more info]
- 'cri\_type': same as input (for reference purposes)
- 'vf': dictionary with vector field measures and data.

## Keys:

- 'Rt': ndarray with general metameric uncertainty index Rt
- 'Rti': ndarray with specific metameric uncertainty indices Rti

```
- 'Rfhj': ndarray with local (hue binned) fidelity indices
                                           obtained from VF model predictions at color space
                                           pixel coordinates
                                     - 'DEhj': ndarray with local (hue binned) color differences
                                           (same as above)
                                     - 'Rcshj': ndarray with local chroma shifts indices for vectorfield
                                     coordinates
                                           (same as above)
                                     - 'Rhshj': ndarray with local hue shifts indices for vector field coordinates
                                           (same as above)
                                     - 'Rfi': ndarray with sample fidelity indices for vectorfield coordinates
                                           (same as above)
                                     - 'DEi': ndarray with sample color differences for vectorfield coordinates
                                           (same as above)
                                     - 'hue_bin_data': dict with output from _get_hue_bin_data() for
                                     vectorfield coordinates
                                     - 'dataVF': dictionary with output of cri.VFPX.VF_colorshift_model()
luxpy.color.cri._tm30_process_spd(spd, cri_type='ies-tm30', **kwargs)
      Calculate all required parameters for plotting from spd using cri.spd_to_cri()
      Args:
                  spd
                         ndarray or dict
                         If ndarray: single spectral power distribution.
                         If dict: dictionary with pre-computed parameters.
                               required keys:
                                     dict_keys(['St', 'Sr', 'xyztw_cct', 'cct', 'duv',
                                            'xyzti', 'xyztw', 'xyzri', 'xyzrw',
                                           'DEi', 'DEa', 'Rf', 'Rg',
                                           'Rcshj', 'Rhshj', 'Rfhj', 'hue bin data'])
                         see cri.spd_to_cri() for more info on parameters.
                  cri_type
                         _CRI_TYPE_DEFAULT or str or dict, optional
                               - 'str: specifies dict with default cri model parameters
                                     (for supported types, see luxpy.cri._CRI_DEFAULTS['cri_types'])
                               - dict: user defined model parameters
                                     (see e.g. luxpy.cri._CRI_DEFAULTS['cierf']
                                     for required structure)
                         Note that any non-None input arguments (in kwargs)
                         to the function will override default values in cri_type dict.
                  kwargs
                         Additional optional keyword arguments,
                         the same as in cri.spd_to_cri()
      Returns:
                  data
                         dictionary with required parameters for plotting functions.
```

```
luxpy.color.cri.plot_tm30_cvg(spd, cri_type='ies-tm30', gamut_line_color=None, gamut_line_style='-',
                                      gamut_line_marker='o', gamut_line_label=None, plot_vectors=True,
                                      plot index values=True, axh=None, axtype='cart',
                                      show_annexE_priority=True, show_Rcsh1_Rfh1=True, **kwargs)
      Plot TM30 Color Vector Graphic (CVG).
      Args:
                  spd
                        ndarray or dict
                        If ndarray: single spectral power distribution.
                        If dict: dictionary with pre-computed parameters (using _tm30_process_spd()).
                              required keys:
                                    dict_keys(['St', 'Sr', 'xyztw_cct', 'cct', 'duv',
                                           'xyzti', 'xyztw', 'xyzri', 'xyzrw',
                                           'DEi', 'DEa', 'Rf', 'Rg',
                                           'Rcshj', 'Rhshj', 'Rfhj', 'hue_bin_data'])
                        see cri.spd_to_cri() for more info on parameters.
                  cri_type
                        _CRI_TYPE_DEFAULT or str or dict, optional
                              - 'str: specifies dict with default cri model parameters
                                    (for supported types, see luxpy.cri._CRI_DEFAULTS['cri_types'])
                              - dict: user defined model parameters
                                    (see e.g. luxpy.cri._CRI_DEFAULTS['cierf']
                                    for required structure)
                        Note that any non-None input arguments (in kwargs)
                        to the function will override default values in cri_type dict.
                  gamut_line_color
                        'r', optional
                        Plotting line style for the line connecting the
                        average test chromaticity in the hue bins.
                        None defaults to red (240,80,70)/255 (IES-TM30-20 recommended).
                  gamut_line_style
                        '-', optional
                        Plotting color for the line connecting the
                        average test chromaticity in the hue bins.
                  gamut_line_marker
                        '-', optional
                        Markers to plot the test color gamut points for each hue bin in
                        (only used when plot_vectors = False).
                  gamut_line_label
                        None, optional
                        Label for gamut line. (only used when plot_vectors = False).
                  plot_vectors
                        True, optional
                        Plot color shift vectors in CVG (True) or not (False).
                  plot_index_values
                        True, optional
```

```
Print Rf, Rg, CCT and Duv in corners of CVG (True) or not (False).
                        If False: turns of potential prints of Rcsh1, Rfh1
                        and annexE_priority levelels as well. This way this argument can be
                        easily used to turn off all plotting and printing when graphs are
                        to be generated with gamuts of multiple sources.
                  axh
                        None, optional
                        If None: create new figure with single axes, else plot on specified axes.
                  axtype
                         'cart' (or 'polar'), optional
                        Make Cartesian (default) or polar plot.
                  show_annexE_priority
                        True, optional
                        Add Annex E priority levels for source.
                  show Rcsh1 Rfh1
                        True, optional
                        Add the local chroma shift (%) and the local color fidelity index
                        for hue bin 1 at the bottom of the graph.
                  kwargs
                        Additional optional keyword arguments,
                        the same as in cri.spd_to_cri()
      Returns:
                  axh
                        handle to figure axes.
                  data
                        dictionary with required parameters for plotting functions.
luxpy.color.cri.plot_tm30_Rfi(spd, cri_type='ies-tm30', axh=None, font_size=11, **kwargs)
      Plot Sample Color Fidelity values (Rfi).
      Args:
                  spd
                        ndarray or dict
                        If ndarray: single spectral power distribution.
                        If dict: dictionary with pre-computed parameters (using _tm30_process_spd()).
                               required keys:
                                     dict_keys(['St', 'Sr', 'xyztw_cct', 'cct', 'duv',
                                           'xyzti', 'xyztw', 'xyzri', 'xyzrw',
                                           'DEi', 'DEa', 'Rf', 'Rg',
                                           'Rcshj', 'Rhshj', 'Rfhj', 'hue_bin_data'])
                        see cri.spd_to_cri() for more info on parameters.
                  cri_type
                        _CRI_TYPE_DEFAULT or str or dict, optional
                               - 'str: specifies dict with default cri model parameters
                                     (for supported types, see luxpy.cri._CRI_DEFAULTS['cri_types'])
                               - dict: user defined model parameters
                                     (see e.g. luxpy.cri._CRI_DEFAULTS['cierf']
```

```
for required structure)
                         Note that any non-None input arguments (in kwargs)
                         to the function will override default values in cri type dict.
                  axh
                         None, optional
                         If None: create new figure with single axes, else plot on specified axes.
                  font size
                         _TM30_FONT_SIZE, optional
                         Font size of text, axis labels and axis values.
                  kwargs
                         Additional optional keyword arguments,
                         the same as in cri.spd_to_cri()
      Returns:
                  axh
                         handle to figure axes.
                  data
                         dictionary with required parameters for plotting functions.
luxpy.color.cri.plot_tm30_Rxhj (spd, cri_type='ies-tm30', axh=None, figsize=(6, 15), font_size=11,
                                         **kwargs)
      Plot Local Chroma Shifts (Rcshj), Local Hue Shifts (Rhshj) and Local Color Fidelity values (Rfhj) (one for each
      hue-bin).
      Args:
                  spd
                         ndarray or dict
                         If ndarray: single spectral power distribution.
                         If dict: dictionary with pre-computed parameters (using _tm30_process_spd()).
                               required keys:
                                     dict_keys(['St', 'Sr', 'xyztw_cct', 'cct', 'duv',
                                            'xyzti', 'xyztw', 'xyzri', 'xyzrw',
                                           'DEi', 'DEa', 'Rf', 'Rg',
                                            'Rcshj', 'Rhshj', 'Rfhj', 'hue_bin_data'])
                         see cri.spd_to_cri() for more info on parameters.
                  cri_type
                         _CRI_TYPE_DEFAULT or str or dict, optional
                               - 'str: specifies dict with default cri model parameters
                                     (for supported types, see luxpy.cri._CRI_DEFAULTS['cri_types'])
                               - dict: user defined model parameters
                                     (see e.g. luxpy.cri._CRI_DEFAULTS['cierf']
                                     for required structure)
                         Note that any non-None input arguments (in kwargs)
                         to the function will override default values in cri_type dict.
                  axh
                         None, optional
                         If None: create new figure with single axes, else plot on specified axes.
                  figsize
                         (6,15), optional
```

```
Figure size of pyplot figure.
                  font size
                         _TM30_FONT_SIZE, optional
                         Font size of text, axis labels and axis values.
                  kwargs
                         Additional optional keyword arguments,
                         the same as in cri.spd_to_cri()
      Returns:
                  axh
                         handle to figure axes.
                  data
                         dictionary with required parameters for plotting functions.
luxpy.color.cri.plot_tm30_Rcshj(spd, cri_type='ies-tm30', axh=None, xlabel=True, y_offset=0,
                                         font_size=11, **kwargs)
      Plot Local Chroma Shift values (Rcshj) (one for each hue-bin).
      Args:
                  spd
                         ndarray or dict
                         If ndarray: single spectral power distribution.
                         If dict: dictionary with pre-computed parameters (using _tm30_process_spd()).
                               required keys:
                                     dict_keys(['St', 'Sr', 'xyztw_cct', 'cct', 'duv',
                                            'xyzti', 'xyztw', 'xyzri', 'xyzrw',
                                           'DEi', 'DEa', 'Rf', 'Rg',
                                           'Rcshj', 'Rhshj', 'Rfhj', 'hue_bin_data'])
                         see cri.spd_to_cri() for more info on parameters.
                  cri_type
                         _CRI_TYPE_DEFAULT or str or dict, optional
                               - 'str: specifies dict with default cri model parameters
                                     (for supported types, see luxpy.cri._CRI_DEFAULTS['cri_types'])
                               - dict: user defined model parameters
                                     (see e.g. luxpy.cri._CRI_DEFAULTS['cierf']
                                     for required structure)
                         Note that any non-None input arguments (in kwargs)
                         to the function will override default values in cri_type dict.
                  axh
                         None, optional
                         If None: create new figure with single axes, else plot on specified axes.
                  xlabel
                         True, optional
                         If False: don't add label and numbers to x-axis
                         (useful when plotting plotting all 'Local Rfhi, Rcshi, Rshhi'
                               values in 3x1 subplots with 'shared x-axis': saves vertical space)
                  y_offset
                         0, optional
                         text-offset from top of bars in barplot.
```

```
font size
                         _TM30_FONT_SIZE, optional
                         Font size of text, axis labels and axis values.
                  kwargs
                         Additional optional keyword arguments,
                         the same as in cri.spd_to_cri()
      Returns:
                  axh
                         handle to figure axes.
                  data
                         dictionary with required parameters for plotting functions.
luxpy.color.cri.plot_tm30_Rhshj(spd, cri_type='ies-tm30', axh=None, xlabel=True, y_offset=0,
                                         font size=11, **kwargs)
      Plot Local Hue Shift values (Rhshj) (one for each hue-bin).
      Args:
                  spd
                         ndarray or dict
                         If ndarray: single spectral power distribution.
                         If dict: dictionary with pre-computed parameters (using _tm30_process_spd()).
                               required keys:
                                     dict_keys(['St', 'Sr', 'xyztw_cct', 'cct', 'duv',
                                            'xyzti', 'xyztw', 'xyzri', 'xyzrw',
                                           'DEi', 'DEa', 'Rf', 'Rg',
                                            'Rcshj', 'Rhshj', 'Rfhj', 'hue_bin_data'])
                         see cri.spd_to_cri() for more info on parameters.
                  cri_type
                         _CRI_TYPE_DEFAULT or str or dict, optional
                               - 'str: specifies dict with default cri model parameters
                                     (for supported types, see luxpy.cri._CRI_DEFAULTS['cri_types'])
                               - dict: user defined model parameters
                                     (see e.g. luxpy.cri._CRI_DEFAULTS['cierf']
                                     for required structure)
                         Note that any non-None input arguments (in kwargs)
                         to the function will override default values in cri_type dict.
                  axh
                         None, optional
                         If None: create new figure with single axes, else plot on specified axes.
                  xlabel
                         True, optional
                         If False: don't add label and numbers to x-axis
                         (useful when plotting plotting all 'Local Rfhi, Rcshi, Rshhi'
                               values in 3x1 subplots with 'shared x-axis': saves vertical space)
                  y_offset
                         0, optional
                         text-offset from top of bars in barplot.
                  font size
```

```
_TM30_FONT_SIZE, optional
                        Font size of text, axis labels and axis values.
                  kwargs
                        Additional optional keyword arguments,
                        the same as in cri.spd_to_cri()
      Returns:
                  axh
                        handle to figure axes.
                  data
                        dictionary with required parameters for plotting functions.
luxpy.color.cri.plot_tm30_Rfhj(spd, cri_type='ies-tm30', axh=None, xlabel=True, y_offset=0, font_size=11,
                                        **kwargs)
      Plot Local Color Fidelity values (Rfhj) (one for each hue-bin).
      Args:
                  spd
                        ndarray or dict
                        If ndarray: single spectral power distribution.
                        If dict: dictionary with pre-computed parameters (using _tm30_process_spd()).
                               required keys:
                                     dict_keys(['St', 'Sr', 'xyztw_cct', 'cct', 'duv',
                                           'xyzti', 'xyztw', 'xyzri', 'xyzrw',
                                           'DEi', 'DEa', 'Rf', 'Rg',
                                           'Rcshj', 'Rhshj', 'Rfhj', 'hue_bin_data'])
                        see cri.spd_to_cri() for more info on parameters.
                  cri_type
                         _CRI_TYPE_DEFAULT or str or dict, optional
                               -'str: specifies dict with default cri model parameters
                                     (for supported types, see luxpy.cri._CRI_DEFAULTS['cri_types'])
                               - dict: user defined model parameters
                                     (see e.g. luxpy.cri._CRI_DEFAULTS['cierf']
                                     for required structure)
                        Note that any non-None input arguments (in kwargs)
                        to the function will override default values in cri_type dict.
                  axh
                        None, optional
                        If None: create new figure with single axes, else plot on specified axes.
                  xlabel
                        True, optional
                        If False: don't add label and numbers to x-axis
                        (useful when plotting plotting all 'Local Rfhi, Rcshi, Rshhi'
                               values in 3x1 subplots with 'shared x-axis': saves vertical space)
                  y_offset
                        0, optional
                        text-offset from top of bars in barplot.
                  font_size
                         _TM30_FONT_SIZE, optional
```

```
Font size of text, axis labels and axis values.
                  kwargs
                        Additional optional keyword arguments,
                        the same as in cri.spd_to_cri()
      Returns:
                  axh
                        handle to figure axes.
                  data
                        dictionary with required parameters for plotting functions.
luxpy.color.cri.plot_tm30_spd(spd, cri_type='ies-tm30', axh=None, font_size=11, **kwargs)
      Plot test SPD and reference illuminant, both normalized to the same luminous power.
      Args:
                  spd
                        ndarray or dict
                        If ndarray: single spectral power distribution.
                        If dict: dictionary with pre-computed parameters (using _tm30_process_spd()).
                               required keys:
                                     dict_keys(['St', 'Sr', 'xyztw_cct', 'cct', 'duv',
                                           'xyzti', 'xyztw', 'xyzri', 'xyzrw',
                                           'DEi', 'DEa', 'Rf', 'Rg',
                                           'Rcshj', 'Rhshj', 'Rfhj', 'hue_bin_data'])
                        see cri.spd_to_cri() for more info on parameters.
                  cri_type
                        _CRI_TYPE_DEFAULT or str or dict, optional
                               - 'str: specifies dict with default cri model parameters
                                     (for supported types, see luxpy.cri._CRI_DEFAULTS['cri_types'])
                               - dict: user defined model parameters
                                     (see e.g. luxpy.cri._CRI_DEFAULTS['cierf']
                                     for required structure)
                        Note that any non-None input arguments (in kwargs)
                        to the function will override default values in cri_type dict.
                  axh
                        None, optional
                        If None: create new figure with single axes, else plot on specified axes.
                  font size
                        _TM30_FONT_SIZE, optional
                        Font size of text, axis labels and axis values.
                  kwargs
                        Additional optional keyword arguments,
                        the same as in cri.spd_to_cri()
      Returns:
                  axh
                        handle to figure axes.
                  data
                        dictionary with required parameters for plotting functions.
```

```
luxpy.color.cri.plot_tm30_report(spd, cri_type='ies-tm30', report_type='full', source='', manufacturer='',
                                          date=", model=", notes=", max_len_notes_line=40, figsize=None,
                                          save fig name=None, dpi=300, plot report top=True,
                                          plot_report_bottom=True, show_annexE_priority=True,
                                          show Rcsh1 Rfh1=True, suptitle='ANSI/IES TM-30-18 Color Rendition
                                          Report', font size=None, **kwargs)
      Create TM30 Color Rendition Report.
      Args:
                  spd
                        ndarray or dict
                        If ndarray: single spectral power distribution.
                        If dict: dictionary with pre-computed parameters (using _tm30_process_spd()).
                              required keys:
                                    dict_keys(['St', 'Sr', 'xyztw_cct', 'cct', 'duv',
                                           'xyzti', 'xyztw', 'xyzri', 'xyzrw',
                                           'DEi', 'DEa', 'Rf', 'Rg',
                                           'Rcshj', 'Rhshj', 'Rfhj', 'hue_bin_data'])
                        see cri.spd_to_cri() for more info on parameters.
                  cri_type
                        CRI TYPE DEFAULT or str or dict, optional
                              -'str: specifies dict with default cri model parameters
                                    (for supported types, see luxpy.cri._CRI_DEFAULTS['cri_types'])
                              - dict: user defined model parameters
                                    (see e.g. luxpy.cri._CRI_DEFAULTS['cierf']
                                    for required structure)
                        Note that any non-None input arguments (in kwargs)
                        to the function will override default values in cri_type dict.
                  report_type
                        'full', optional
                        Generate a full report as in ANSI/IES-TM30-2020
                        Options:
                              - 'full': full report with spectrum plot, color vector graphic, local indices,
                              sample indices'simple', ...
                              - 'intermediate': color vector graphic + local chroma and hue shifts
                              - 'simple': color vector graphic only
                              - 'spd_cvg': spectrum plot + color vector graphic
                  source
                        string with source name.
                  manufacturer
                        string with source manufacturer.
                  model
                        string with source model.
                  date
                        string with source measurement date.
                  notes
                        string to be split
                  max_len_notes_line
```

```
40, optional
      Maximum length of a single line when splitting the string.
figsize
      None, optional
      Figure size of pyplot figure.
      If None a default depending on the report type is used:
            - 'full': (7,12)
            - 'intermediate': (14,6)
            - 'simple' : (6,6)
            -'spd_cvg': (14,6)
save_fig_name
      None, optional
      Filename (+path) to which the report will be saved as an image (png).
      If None: don't save, just display.
dpi
      300, optional
      Dots-Per-Inch of image file (PNG).
plot_report_top
      execute _plot_tm30_report_top()
plot report bottom
      execute _plot_tm30_report_bottom()
show\_annexE\_priority
      True, optional
      Add Annex E priority levels for source.
show_Rcsh1_Rfh1
      True, optional
      Add the local chroma shift (%) and the local color fidelity index
      for hue bin 1 at the bottom of the graph.
suptitle
      'ANSI/IES TM-30-18 Color Rendition Report' or str, optional
      report title (input for plt.suptitle).
font_size
      None, optional
      Font size of text, axis labels and axis values (adjust when changing figsizes).
      Defaults: ('full': TM30 FONT SIZE FULLREPORT, other options:
      _TM30_FONT_SIZE)
kwargs
      Additional optional keyword arguments,
      the same as in cri.spd_to_cri()
axs
      dictionary with handles to each axes.
data
      dictionary with required parameters for plotting functions.
```

**Returns:** 

```
luxpy.color.cri.spd_to_tm30_report(spd, cri_type='ies-tm30', report_type='full', source='',
                                            manufacturer=", date=", model=", notes=", max_len_notes_line=40,
                                            figsize=None, save fig name=None, dpi=300, plot report top=True,
                                            plot_report_bottom=True, show_annexE_priority=True,
                                            show Rcsh1 Rfh1=True, suptitle='ANSI/IES TM-30-18 Color
                                             Rendition Report', font size=None, **kwargs)
      Create TM30 Color Rendition Report.
      Args:
                  spd
                        ndarray or dict
                        If ndarray: single spectral power distribution.
                        If dict: dictionary with pre-computed parameters (using _tm30_process_spd()).
                              required keys:
                                    dict_keys(['St', 'Sr', 'xyztw_cct', 'cct', 'duv',
                                           'xyzti', 'xyztw', 'xyzri', 'xyzrw',
                                           'DEi', 'DEa', 'Rf', 'Rg',
                                           'Rcshj', 'Rhshj', 'Rfhj', 'hue_bin_data'])
                        see cri.spd_to_cri() for more info on parameters.
                  cri_type
                        CRI TYPE DEFAULT or str or dict, optional
                              -'str: specifies dict with default cri model parameters
                                    (for supported types, see luxpy.cri._CRI_DEFAULTS['cri_types'])
                              - dict: user defined model parameters
                                    (see e.g. luxpy.cri._CRI_DEFAULTS['cierf']
                                    for required structure)
                        Note that any non-None input arguments (in kwargs)
                        to the function will override default values in cri_type dict.
                  report_type
                        'full', optional
                        Generate a full report as in ANSI/IES-TM30-2020
                        Options:
                              - 'full': full report with spectrum plot, color vector graphic, local indices,
                              sample indices'simple', ...
                              - 'intermediate': color vector graphic + local chroma and hue shifts
                              - 'simple': color vector graphic only
                              - 'spd_cvg': spectrum plot + color vector graphic
                  source
                        string with source name.
                  manufacturer
                        string with source manufacturer.
                  model
                        string with source model.
                  date
                        string with source measurement date.
                  notes
                        string to be split
                  max_len_notes_line
```

```
40, optional
      Maximum length of a single line when splitting the string.
figsize
      None, optional
      Figure size of pyplot figure.
      If None a default depending on the report type is used:
            - 'full': (7,12)
            - 'intermediate': (14,6)
            - 'simple' : (6,6)
            -'spd_cvg': (14,6)
save_fig_name
      None, optional
      Filename (+path) to which the report will be saved as an image (png).
      If None: don't save, just display.
dpi
      300, optional
      Dots-Per-Inch of image file (PNG).
plot_report_top
      execute _plot_tm30_report_top()
plot report bottom
      execute _plot_tm30_report_bottom()
show\_annexE\_priority
      True, optional
      Add Annex E priority levels for source.
show_Rcsh1_Rfh1
      True, optional
      Add the local chroma shift (%) and the local color fidelity index
      for hue bin 1 at the bottom of the graph.
suptitle
      'ANSI/IES TM-30-18 Color Rendition Report' or str, optional
      report title (input for plt.suptitle).
font_size
      None, optional
      Font size of text, axis labels and axis values (adjust when changing figsizes).
      Defaults: ('full': TM30 FONT SIZE FULLREPORT, other options:
      _TM30_FONT_SIZE)
kwargs
      Additional optional keyword arguments,
      the same as in cri.spd_to_cri()
axs
      dictionary with handles to each axes.
data
      dictionary with required parameters for plotting functions.
```

**Returns:** 

```
luxpy.color.cri.plot_cri_graphics(data, cri_type=None, hbins=16, start_hue=0.0, scalef=100,
                                            plot_axis_labels=False, bin_labels=None, plot_edge_lines=True,
                                            plot center lines=False, plot bin colors=True, axtype='polar',
                                            ax=None, force_CVG_layout=True, vf_model_type='M6',
                                            vf_pcolorshift={'Cref': 40, 'href': array([3.7835, 3.3161, 2.8272,
                                            1.9093, 5.2787, 4.3081, 0.37762, 6.2055, 1.4564, 0.88927]), 'labels':
                                            array(['5B', '5BG', '5G', '5GY', '5P', '5PB', '5R', '5RP', '5Y', '5YR'],
                                            dtype=object), 'sig': 0.3}, vf_color='k', vf_bin_labels=array(['5B',
                                            '5BG', '5G', '5GY', '5P', '5PB', '5R', '5RP', '5Y', '5YR'], dtype=object),
                                            vf_plot_bin_colors=True, scale_vf_chroma_to_sample_chroma=False,
                                            plot_VF=True, plot_CF=False, plot_SF=False,
                                            plot_test_sample_coord=False)
      Plot graphical information on color rendition properties (custom design).
      Args:
                  data
                         ndarray with spectral data or dict with pre-computed metrics.
                  cri_type
                         None, optional
                         If None: defaults to cri_type = 'iesrf'.
                         :hbins:, :start hue: and :scalef: are ignored if cri type not None
                         and values are replaced by those in cri_type['rg_pars']
                  hbins
                         16 or ndarray with sorted hue bin centers (°), optional
                  start hue
                         0.0, optional
                  scalef
                         100, optional
                         Scale factor for graphic.
                  plot_axis_labels
                         False, optional
                         Turns axis ticks on/off (True/False).
                  bin labels
                         None or list[str] or '#', optional
                         Plots labels at the bin center hues.
                               - None: don't plot.
                               - list[str]: list with str for each bin.
                                     (len(:bin_labels:) = :nhbins:)
                               - '#': plots number.
                  plot_edge_lines
                         True or False, optional
                         Plot grey bin edge lines with '-'.
                  plot_center_lines
                         False or True, optional
                         Plot colored lines at 'center' of hue bin.
                  plot_bin_colors
                         True, optional
```

Colorize hue bins.

#### axtype

'polar' or 'cart', optional

Make polar or Cartesian plot.

ax

None or 'new' or 'same', optional

- None or 'new' creates new plot
- 'same': continue plot on same axes.
- axes handle: plot on specified axes.

## force\_CVG\_layout

True, optional

True: Force plot of basis of CVG.

#### vf\_model\_type

\_VF\_MODEL\_TYPE or 'M6' or 'M5', optional

Type of polynomial vector field model to use for the calculation of base color shift and metameric uncertainty.

### vf\_pcolorshift

\_VF\_PCOLORSHIFT or user defined dict, optional

The polynomial models of degree 5 and 6 can be fully specified or summarized by the model parameters themselved OR by calculating the dCoverC and dH at resp. 5 and 6 hues. :VF\_pcolorshift: specifies these hues and chroma level.

### vf\_color

'k', optional

For plotting the vector fields.

### vf\_plot\_bin\_colors

True, optional

Colorize hue bins of VF graph.

### scale\_vf\_chroma\_to\_sample\_chroma

False, optional

Scale chroma of reference and test vf fields such that average of binned reference chroma equals that of the binned sample chroma before calculating hue bin metrics.

#### vf bin labels

see:bin labels:

Set VF model hue-bin labels.

# plot\_CF

False, optional

Plot circle fields.

# plot\_VF

True, optional

Plot vector fields.

# plot\_SF

True, optional

Plot sample shifts.

# plot\_test\_sample\_coord

Plot the coordinates of the samples under the test illuminant relative to the mean chromaticity under the reference illuminant (in the CVG plot).

#### **Returns:**

#### returns

```
(data, [plt.gcf(),ax_spd, ax_CVG, ax_locC, ax_locH, ax_VF], cmap )
```

:data: is a dictionary with color rendering data with keys:

- 'St, Sr': ndarray of test SPDs and corresponding ref. illuminants.
- 'xyz\_cct': xyz of white point calculate with cieobs defined for cct calculations in cri\_type['cieobs']
- 'cct, duv': CCT and Duv obtained with cieobs in cri\_type['cieobs']['cct']
- 'xyzti, xyzri': ndarray tristimulus values of test and ref. samples (obtained with with cieobs in cri\_type['cieobs']['xyz'])
- 'xyztw, xyzrw': ndarray tristimulus values of test and ref. white points (obtained with with cieobs in cri\_type['cieobs']['xyz'])
- 'DEi, DEa': ndarray with individual sample color differences DEi and average DEa between test and ref.
- 'Rf': ndarray with general color fidelity index values
- 'Rg': ndarray with color gamut area index values
- 'Rfi': ndarray with specific (sample) color fidelity indices
- 'Rfhj': ndarray with local (hue binned) fidelity indices
- 'DEhj': ndarray with local (hue binned) color differences
- 'Rcshj': ndarray with local chroma shifts indices
- 'Rhshj': ndarray with local hue shifts indices
- 'hue\_bin\_data': dict with output from \_get\_hue\_bin\_data() [see its help for more info]
- 'cri\_type': same as input (for reference purposes)
- 'vf': dictionary with vector field measures and data.

## Keys:

- 'Rt': ndarray with general metameric uncertainty index Rt
- 'Rti' : ndarray with specific metameric uncertainty indices Rti
- 'Rfhj': ndarray with local (hue binned) fidelity indices obtained from VF model predictions at color space pixel coordinates
- 'DEhj': ndarray with local (hue binned) color differences (same as above)
- 'Rcshj': ndarray with local chroma shifts indices for vectorfield coordinates

(same as above)

- 'Rhshj': ndarray with local hue shifts indicesfor vectorfield coordinates (same as above)
- 'Rfi': ndarray with sample fidelity indices for vectorfield coordinates (same as above)
- 'DEi': ndarray with sample color differences for vectorfield coordinates (same as above)

- 'hue\_bin\_data': dict with output from \_get\_hue\_bin\_data() for vectorfield coordinates
- 'dataVF': dictionary with output of cri.VFPX.VF\_colorshift\_model()

:[...]: list with handles to current figure and 5 axes.

:cmap: list with rgb colors for hue bins (for use in other plotting fcns)

```
luxpy.color.cri.spd_to_tm30_fast(St)
Calculate tm30 measures from spd.
```

```
luxpy.color.cri.cri_ref_fast(ccts, wl3=array([360, 361, 362, 363, 364, 365, 366, 367, 368, 369, 370, 371,
                                     372, 373, 374, 375, 376, 377, 378, 379, 380, 381, 382, 383, 384, 385, 386, 387,
                                     388, 389, 390, 391, 392, 393, 394, 395, 396, 397, 398, 399, 400, 401, 402, 403,
                                     404, 405, 406, 407, 408, 409, 410, 411, 412, 413, 414, 415, 416, 417, 418, 419,
                                     420, 421, 422, 423, 424, 425, 426, 427, 428, 429, 430, 431, 432, 433, 434, 435,
                                     436, 437, 438, 439, 440, 441, 442, 443, 444, 445, 446, 447, 448, 449, 450, 451,
                                     452, 453, 454, 455, 456, 457, 458, 459, 460, 461, 462, 463, 464, 465, 466, 467,
                                     468, 469, 470, 471, 472, 473, 474, 475, 476, 477, 478, 479, 480, 481, 482, 483,
                                     484, 485, 486, 487, 488, 489, 490, 491, 492, 493, 494, 495, 496, 497, 498, 499,
                                     500, 501, 502, 503, 504, 505, 506, 507, 508, 509, 510, 511, 512, 513, 514, 515,
                                     516, 517, 518, 519, 520, 521, 522, 523, 524, 525, 526, 527, 528, 529, 530, 531,
                                     532, 533, 534, 535, 536, 537, 538, 539, 540, 541, 542, 543, 544, 545, 546, 547,
                                     548, 549, 550, 551, 552, 553, 554, 555, 556, 557, 558, 559, 560, 561, 562, 563,
                                     564, 565, 566, 567, 568, 569, 570, 571, 572, 573, 574, 575, 576, 577, 578, 579,
                                     580, 581, 582, 583, 584, 585, 586, 587, 588, 589, 590, 591, 592, 593, 594, 595,
                                     596, 597, 598, 599, 600, 601, 602, 603, 604, 605, 606, 607, 608, 609, 610, 611,
                                     612, 613, 614, 615, 616, 617, 618, 619, 620, 621, 622, 623, 624, 625, 626, 627,
                                     628, 629, 630, 631, 632, 633, 634, 635, 636, 637, 638, 639, 640, 641, 642, 643,
                                     644, 645, 646, 647, 648, 649, 650, 651, 652, 653, 654, 655, 656, 657, 658, 659,
                                     660, 661, 662, 663, 664, 665, 666, 667, 668, 669, 670, 671, 672, 673, 674, 675,
                                     676, 677, 678, 679, 680, 681, 682, 683, 684, 685, 686, 687, 688, 689, 690, 691,
                                     692, 693, 694, 695, 696, 697, 698, 699, 700, 701, 702, 703, 704, 705, 706, 707,
                                     708, 709, 710, 711, 712, 713, 714, 715, 716, 717, 718, 719, 720, 721, 722, 723,
                                     724, 725, 726, 727, 728, 729, 730, 731, 732, 733, 734, 735, 736, 737, 738, 739,
                                     740, 741, 742, 743, 744, 745, 746, 747, 748, 749, 750, 751, 752, 753, 754, 755,
                                     756, 757, 758, 759, 760, 761, 762, 763, 764, 765, 766, 767, 768, 769, 770, 771,
                                     772, 773, 774, 775, 776, 777, 778, 779, 780, 781, 782, 783, 784, 785, 786, 787,
                                     788, 789, 790, 791, 792, 793, 794, 795, 796, 797, 798, 799, 800, 801, 802, 803,
                                     804, 805, 806, 807, 808, 809, 810, 811, 812, 813, 814, 815, 816, 817, 818, 819,
                                     820, 821, 822, 823, 824, 825, 826, 827, 828, 829, 830]), ref_type='iestm30',
                                     mix_range=[4000, 5000], cieobs='1931_2',
                                     force_daylight_below4000K=False, n=None, daylight_locus=None, wl=[360,
                                     830, 11)
```

Calculates multiple reference illuminant spectra based on ccts for color rendering index calculations.

```
luxpy.color.cri.xyz_to_jab_cam02ucs_fast(xyz, xyzw, ucs=True, conditions=None)
Calculate CAM02-UCS J'a'b' coordinates from xyz tristimulus values of sample and white point.
Args:
```

XYZ

ndarray with sample tristimulus values

xyzw

ndarray with white point tristimulus values conditions

```
None, optional
                       Dictionary with viewing conditions.
                       None results in:
                              {'La':100, 'Yb':20, 'D':1, 'surround':'avg'}
                       For more info see luxpy.cam.ciecam02()?
     Returns:
                 jab
                       ndarray with J'a'b' coordinates.
4.4.9 cri/VFPX/
           рy
                     • __init__.py
                     • VF_PX_models.py
                     · vectorshiftmodel.py

    pixelshiftmodel.py

           namespace luxpy.cri.VFPX
luxpy.color.cri.VFPX.get_poly_model(jabt, jabr, modeltype='M6')
     Setup base color shift model (delta_a, delta_b), determine model parameters and accuracy.
     Calculates a base color shift (delta) from the ref. chromaticity ar, br.
     Args:
                 jabt
                       ndarray with jab color coordinates under the test SPD.
                 jabr
                       ndarray with jab color coordinates under the reference SPD.
                 modeltype
                       _VF_MODEL_TYPE or 'M6' or 'M5', optional
                       Specifies degree 5 or degree 6 polynomial model in ab-coordinates.
                       (see notes below)
     Returns:
                 returns
                       (poly_model,
                             pmodel,
                             dab_model,
                                   dab_res,
                                   dCHoverC_res,
                                   dab std,
                                   dCHoverC_std)
                       :poly_model: function handle to model
                       :pmodel: ndarray with model parameters
                       :dab_model: ndarray with ab model predictions from ar, br.
```

```
:dab_res: ndarray with residuals between 'da,db' of samples and
                                                                  'da,db' predicted by the model.
                                                     :dCHoverC res: ndarray with residuals between 'dCoverC,dH'
                                                                               of samples and 'dCoverC,dH' predicted by the model.
                                                                  Note: dCoverC = (Ct - Cr)/Cr and dH = ht - hr
                                                                               (predicted from model, see notes below)
                                                     :dab_std: ndarray with std of :dab_res:
                                                     :dCHoverC_std: ndarray with std of :dCHoverC_res:
             Notes:
                                 1. Model types:
                                                     poly5\_model = lambda \ a,b,p: \ p[0]*a + p[1]*b + p[2]*(a**2) + p[3]*a*b + p[4]*(b**2)
                                                     poly6\_model = lambda \ a,b,p: \ p[0] + p[1]*a + p[2]*b + p[3]*(a**2) + p[4]*a*b + p[4]
                                                     p[5]*(b**2)
                                 2. Calculation of dCoverC and dH:
                                                     dCoverC = (np.cos(hr)*da + np.sin(hr)*db)/Cr
                                                     dHoverC = (np.cos(hr)*db - np.sin(hr)*da)/Cr
luxpy.color.cri.VFPX.apply_poly_model_at_x(poly_model, pmodel, axr, bxr)
             Applies base color shift model at cartesian coordinates axr, bxr.
             Args:
                                        poly_model
                                                     function handle to model
                                        pmodel
                                                     ndarray with model parameters.
                                       axr
                                                     ndarray with a-coordinates under the reference conditions
                                       bxr
                                                     ndarray with b-coordinates under the reference conditions
             Returns:
                                        returns
                                                     (axt,bxt,Cxt,hxt,
                                                                  axr,bxr,Cxr,hxr)
                                                     ndarrays with ab-coordinates, chroma and hue
                                                     predicted by the model (xt), under the reference (xr).
luxpy.color.cri.VFPX.generate_vector_field(poly model, pmodel, axr=array([-40, -35, -30, -25, -20, -
                                                                                                                        15, -10, -5, 0, 5, 10, 15, 20, 25, 30, 35, 40]), bxr=array([-
                                                                                                                       40, -35, -30, -25, -20, -15, -10, -5, 0, 5, 10, 15, 20, 25, 30,
                                                                                                                       35, 40]), make_grid=True, limit_grid_radius=0, color='k')
             Generates a field of vectors using the base color shift model.
             Has the option to plot vector field.
             Args:
                                        poly_model
                                                     function handle to model
```

```
axr
                        np.arange(-_VF_MAXR,_VF_MAXR+_VF_DELTAR,_VF_DELTAR), optional
                        Ndarray specifying the a-coordinates at which to apply the model.
                  bxr
                        np.arange(-_VF_MAXR,_VF_MAXR+_VF_DELTAR,_VF_DELTAR), optional
                        Ndarray specifying the b-coordinates at which to apply the model.
                  make_grid
                        True, optional
                        True: generate a 2d-grid from :axr:, :bxr:.
                  limit_grid_radius
                        0, optional
                              A value of zeros keeps grid as specified by axr,bxr.
                              A value > 0 only keeps (a,b) coordinates within :limit_grid_radius:
                  color
                        'k', optional
                        For plotting the vector field.
                        If :color: == 0, no plot will be generated.
      Returns:
                  returns
                        If :color: == 0: ndarray of axt,bxt,axr,bxr
                        Else: handle to axes used for plotting.
luxpy.color.cri.VFPX.VF_colorshift_model(S, cri_type='iesrf', model_type='M6', cspace={'Yw': None,
                                                    'conditions': {'D': 1.0, 'Dtype': None, 'La': 100.0, 'Yb': 20.0,
                                                    'surround': 'avg'}, 'mcat': 'cat02', 'type': 'jab_cam02ucs',
                                                    'xyzw': None, 'yellowbluepurplecorrect': None},
                                                    sampleset=None, pool=False, pcolorshift={'Cref': 40, 'href':
                                                    array([0.31416, 0.94248, 1.5708, 2.1991, 2.8274, 3.4558,
                                                    4.0841, 4.7124, 5.3407, 5.969]), 'sig': 0.3}, vfcolor='k',
                                                    verbosity=0)
      Applies full vector field model calculations to spectral data.
      Args:
                  S
                        nump.ndarray with spectral data.
                  cri_type
                        _VF_CRI_DEFAULT or str or dict, optional
                        Specifies type of color fidelity model to use.
                        Controls choice of ref. ill., sample set, averaging, scaling, etc.
                        See luxpy.cri.spd_to_cri for more info.
                  modeltype
                        _VF_MODEL_TYPE or 'M6' or 'M5', optional
                        Specifies degree 5 or degree 6 polynomial model in ab-coordinates.
                  cspace
                        _VF_CSPACE or dict, optional
                        Specifies color space. See _VF_CSPACE_EXAMPLE for example structure.
```

pmodel

ndarray with model parameters.

#### sampleset

None or str or ndarray, optional

Sampleset to be used when calculating vector field model.

## pool

False, optional

If :S: contains multiple spectra, True pools all jab data before modeling the vector field, while False models a different field for each spectrum.

### pcolorshift

default dict (see below) or user defined dict, optional

Dict containing the specification input

for apply\_poly\_model\_at\_hue\_x().

Default dict = { 'href': np.arange(np.pi/10,2\*np.pi,2\*np.pi/10),

```
'Cref': _VF_MAXR, 
'sig': _VF_SIG, 
'labels': '#'}
```

The polynomial models of degree 5 and 6 can be fully specified or summarized by the model parameters themselved OR by calculating the dCoverC and dH at resp. 5 and 6 hues.

### vfcolor

'k', optional

For plotting the vector fields.

# verbosity

0, optional

Report warnings or not.

#### **Returns:**

#### returns

list[dict] (each list element refers to a different test SPD) with the following keys:

- 'Source': dict with ndarrays of the S, cct and duv of source spd.
- 'metrics': dict with ndarrays for:
  - \* Rf (color fidelity: base + metameric shift)
  - \* Rt (metameric uncertainty index)
  - \* Rfi (specific color fidelity indices)
  - \* Rti (specific metameric uncertainty indices)
  - \* cri\_type (str with cri\_type)
- 'Jab': dict with with ndarrays for Jabt, Jabr, DEi
- 'dC/C\_dH\_x\_sig':

```
np.vstack((dCoverC_x,dCoverC_x_sig,dH_x,dH_x_sig)).T
```

See get\_poly\_model() for more info.

- 'fielddata': dict with dicts containing data on the calculated vector-field and circle-fields:

```
- 'modeldata': dict with model info:
                                    {'pmodel': pmodel,
                                    'pcolorshift' : pcolorshift,
                                          'dab_model' : dab_model,
                                          'dab_res': dab_res,
                                          'dab_std': dab_std,
                                          'modeltype': modeltype,
                                          'fmodel' : poly_model,
                                          'Jabtm': Jabtm,
                                          'Jabrm': Jabrm,
                                          'DEim': DEim},
                             - 'vshifts' :dict with various vector shifts:
                                    * 'Jabshiftvector_r_to_t' : ndarray with difference vectors
                                          between jabt and jabr.
                                   * 'vshift ab s': vshift ab s: ab-shift vectors of samples
                                   * 'vshift_ab_s_vf' : vshift_ab_s_vf: ab-shift vectors of
                                          VF model predictions of samples.
                                   * 'vshift_ab_vf' : vshift_ab_vf: ab-shift vectors of VF
                                         model predictions of vector field grid.
luxpy.color.cri.VFPX.initialize_VF_hue_angles(hx=None, Cxr=40, cri_type='iesrf', modeltype='M6',
                                                         determine hue angles=True)
     Initialize the hue angles that will be used to 'summarize' the VF model fitting parameters.
     Args:
                 hx
                        None or ndarray, optional
                        None defaults to Munsell H5 hues.
                  Cxr
                        _VF_MAXR, optional
                 cri_type
                        _VF_CRI_DEFAULT or str or dict, optional,
                        Cri_type parameters for cri and VF model.
                 modeltype
                        _VF_MODEL_TYPE or 'M5' or 'M6', optional
                        Determines the type of polynomial model.
                  determine_hue_angles
                        _DETERMINE_HUE_ANGLES or True or False, optional
                        True: determines the 10 primary / secondary Munsell hues ('5..').
                        Note that for 'M6', an additional
     Returns:
                 pcolorshift
                        {'href': href,
                              'Cref': _VF_MAXR,
                              'sig': VF SIG,
                              'labels' : list[str]}
```

```
luxpy.color.cri.VFPX.generate_grid(jab_ranges=None, out='grid', ax=array([-40, -35, -30, -25, -20, -
                                            15, -10, -5, 0, 5, 10, 15, 20, 25, 30, 35, 40]), bx=array([-40, -35, -
                                            30, -25, -20, -15, -10, -5, 0, 5, 10, 15, 20, 25, 30, 35, 40]), jx=None,
                                            limit_grid_radius=0)
      Generate a grid of color coordinates.
      Args:
                  out
                        'grid' or 'vectors', optional
                              - 'grid': outputs a single 2d numpy.nd-vector with the grid coordinates
                              - 'vector': outputs each dimension seperately.
                 jab_ranges
                        None or ndarray, optional
                        Specifies the pixelization of color space.
                        (ndarray.shape = (3,3), with first axis: J,a,b, and second
                        axis: min, max, delta)
                  ax
                        default ndarray or user defined ndarray, optional
                        default = np.arange(-_VF_MAXR,_VF_MAXR+_VF_DELTAR,_VF_DELTAR)
                  bx
                        default ndarray or user defined ndarray, optional
                        default = np.arange(-_VF_MAXR,_VF_MAXR+_VF_DELTAR,_VF_DELTAR)
                 jх
                        None, optional
                        Note that not-None :jab ranges: override :ax:, :bx: and :jx input.
                  limit_grid_radius
                        0, optional
                        A value of zeros keeps grid as specified by axr,bxr.
                        A value > 0 only keeps (a,b) coordinates within :limit_grid_radius:
      Returns:
                  returns
                        single ndarray with ax,bx [,jx]
                              or
                        seperate ndarrays for each dimension specified.
luxpy.color.cri.VFPX.calculate_shiftvectors(jabt, jabr, average=True, vtype='ab')
      Calculate color shift vectors.
      Args:
                  jabt
                        ndarray with jab coordinates under the test SPD
                  jabr
                        ndarray with jab coordinates under the reference SPD
                  average
                        True, optional
                        If True, take mean of difference vectors along axis = 0.
                  vtype
                        'ab' or 'jab', optional
                        Reduce output ndarray to only a,b coordinates of shift vector(s).
```

### **Returns:**

#### returns

```
ndarray of (mean) shift vector(s).
```

Plots vector or circle fields generated by VFcolorshiftmodel() or PXcolorshiftmodel().

# Args:

#### data

dict generated by VFcolorshiftmodel() or PXcolorshiftmodel()

Must contain 'fielddata'- key, which is a dict with possible keys:

- key: 'vectorfield': ndarray with vector field data
- key: 'circlefield': ndarray with circle field data

### color

'k', optional

Color for plotting the vector-fields.

### axtype

'polar' or 'cart', optional

Make polar or Cartesian plot.

ax

None or 'new' or 'same', optional

- None or 'new' creates new plot
- 'same': continue plot on same axes.
- axes handle: plot on specified axes.

## hbins

16 or ndarray with sorted hue bin centers (°), optional

#### start\_hue

\_VF\_MAXR, optional

Scale factor for graphic.

## plot\_axis\_labels

False, optional

Turns axis ticks on/off (True/False).

#### bin labels

None or list[str] or '#', optional

Plots labels at the bin center hues.

- None: don't plot.
- list[str]: list with str for each bin.

(len(:bin\_labels:) = :nhbins:)

- '#': plots number.

### plot\_edge\_lines

True or False, optional

Plot grey bin edge lines with '-'.

#### plot\_center\_lines

```
False or True, optional
                        Plot colored lines at 'center' of hue bin.
                  plot_bin_colors
                        True, optional
                        Colorize hue-bins.
                  force_CVG_layout
                        False or True, optional
                        True: Force plot of basis of CVG.
      Returns:
                  returns
                        figCVG, hax, cmap
                        :figCVG: handle to CVG figure
                        :hax: handle to CVG axes
                        :cmap: list with rgb colors for hue bins
                              (for use in other plotting fcns)
luxpy.color.cri.VFPX.plotcircle(radii=array([0, 10, 20, 30, 40, 50]), angles=array([0, 10, 20, 30, 40, 50])
                                         60, 70, 80, 90, 100, 110, 120, 130, 140, 150, 160, 170, 180, 190, 200, 210,
                                         220, 230, 240, 250, 260, 270, 280, 290, 300, 310, 320, 330, 340]),
                                         color='k', linestyle='--', out=None)
      Plot one or more concentric circles around (0,0).
      Args:
                        radii
                              np.arange(0,60,10) or ndarray with radii of circle(s), optional
                        angles
                              np.arange(0,350,10) or ndarray with angles (°), optional
                        color
                               'k', optional
                              Color for plotting.
                        linestyle
                               '-', optional
                              Linestyle of circles.
                        out
                              None, optional
                              If None: plot circles, return (x,y) otherwise.
            Returns:
                        x,y
                              ndarrays with circle coordinates (only returned if out is 'x,y')
luxpy.color.cri.VFPX.get_pixel_coordinates(jab, jab_ranges=None, jab_deltas=None,
                                                       limit_grid_radius=0)
      Get pixel coordinates corresponding to array of jab color coordinates.
      Args:
                  jab
                        ndarray of color coordinates
                  jab_ranges
```

None or ndarray, optional

Specifies the pixelization of color space.

(ndarray.shape = (3,3), with first axis: J,a,b, and second axis: min, max, delta)

## jab\_deltas

float or ndarray, optional

Specifies the sampling range.

A float uses jab\_deltas as the maximum Euclidean distance to select samples around each pixel center. A ndarray of 3 deltas, uses

a city block sampling around each pixel center.

### limit\_grid\_radius

0, optional

A value of zeros keeps grid as specified by axr,bxr.

A value > 0 only keeps (a,b) coordinates within :limit\_grid\_radius:

#### **Returns:**

#### returns

gridp, idxp, jabp, samplenrs, samplesIDs

- : gridp: ndarray with coordinates of all pixel centers.
- :idxp: list[int] with pixel index for each non-empty pixel
- : jabp: ndarray with center color coordinates of non-empty pixels
- -:samplenrs: list[list[int]] with sample numbers belong to each non-empty pixel
- :sampleIDs: summarizing list,

with column order: 'idxp, jabp, samplenrs'

# 

Pixelates the color space and calculates the color shifts in each pixel.

## **Args:**

#### Jabt

ndarray with color coordinates under the (single) test SPD.

#### **Jabr**

ndarray with color coordinates under the (single) reference SPD.

## jab\_ranges

None or ndarray, optional

Specifies the pixelization of color space.

(ndarray.shape = (3,3), with first axis: J,a,b, and second

axis: min, max, delta)

## jab\_deltas

float or ndarray, optional

Specifies the sampling range.

A float uses jab\_deltas as the maximum Euclidean distance to select samples around each pixel center. A ndarray of 3 deltas, uses a city block sampling around each pixel center.

### limit\_grid\_radius

0, optional

A value of zeros keeps grid as specified by axr,bxr.

A value > 0 only keeps (a,b) coordinates within :limit\_grid\_radius:

#### **Returns:**

#### returns

```
dict with the following keys:
```

- 'Jab': dict with with ndarrays for:

Jabt, Jabr, DEi, DEi\_ab (only ab-coordinates), DEa (mean) and DEa ab

- 'vshifts': dict with:
  - \* 'vectorshift': ndarray with vector shifts between average Jabt and Jabr for each pixel
  - \* 'vectorshift\_ab': ndarray with vector shifts averaged over J for each pixel
  - \* 'vectorshift\_ab\_J0': ndarray with vector shifts averaged over J for each pixel of J=0 plane.
  - \* 'vectorshift\_len': length of 'vectorshift'
  - \* 'vectorshift\_ab\_len': length of 'vectorshift\_ab'
  - \* 'vectorshift\_ab\_J0\_len': length of 'vectorshift\_ab\_J0'
  - \* 'vectorshift\_len\_DEnormed': length of 'vectorshift' normalized to 'DEa'
  - \* 'vectorshift\_ab\_len\_DEnormed': length of 'vectorshift\_ab' normalized to 'DEa\_ab'
  - \* 'vectorshift\_ab\_J0\_len\_DEnormed': length of

'vectorshift\_ab\_J0'

normalized to 'DEa\_ab'

- 'pixeldata': dict with pixel info:
  - \* 'grid' ndarray with coordinates of all pixel centers.
  - \* 'idx': list[int] with pixel index for each non-empty pixel
  - \* 'Jab': ndarray with center coordinates of non-empty pixels
  - \* 'samplenrs': list[list[int]] with sample numbers belong to each non-empty pixel
  - \* 'IDs: summarizing list,

with column order: 'idxp, jabp, samplenrs'

- 'fielddata' : dict with dicts containing data on the calculated vector-field and circle-fields
  - \* 'vectorfield': dict with ndarrays for the ab-coordinates under the ref. (axr, bxr) and test (axt, bxt) illuminants, centered at the pixel centers corresponding to the ab-coordinates of the reference illuminant.

luxpy.color.cri.VFPX.calculate\_VF\_PX\_models(S, cri\_type='iesrf', sampleset=None, pool=False,

pcolorshift={'Cref': 40, 'href': array([0.31416, 0.94248, 1.5708, 2.1991, 2.8274, 3.4558, 4.0841, 4.7124, 5.3407, 5.969]), 'labels': '#', 'sig': 0.3}, vfcolor='k', verbosity=0)

Calculate Vector Field and Pixel color shift models.

### Args:

#### cri type

\_VF\_CRI\_DEFAULT or str or dict, optional Specifies type of color fidelity model to use. Controls choice of ref. ill., sample set, averaging, scaling, etc. See luxpy.cri.spd\_to\_cri for more info.

## sampleset

None or str or ndarray, optional

Sampleset to be used when calculating vector field model.

#### pool

False, optional

If :S: contains multiple spectra, True pools all jab data before modeling the vector field, while False models a different field for each spectrum.

## pcolorshift

default dict (see below) or user defined dict, optional

Dict containing the specification input

```
for apply_poly_model_at_hue_x().
```

Default dict = {'href': np.arange(np.pi/10,2\*np.pi,2\*np.pi/10),

```
'Cref': _VF_MAXR,
```

'sig': \_VF\_SIG,

'labels': '#'}

The polynomial models of degree 5 and 6 can be fully specified or summarized by the model parameters themselved OR by calculating the dCoverC and dH at resp. 5 and 6 hues.

#### vfcolor

'k', optional

For plotting the vector fields.

#### verbosity

0, optional

Report warnings or not.

#### **Returns:**

#### returns

:dataVF:, :dataPX:

Dicts, for more info, see output description of resp.:

luxpy.cri.VF\_colorshift\_model() and luxpy.cri.PX\_colorshift\_model()

```
luxpy.color.cri.VFPX.subsample_RFL_set(rfl, rflpath=", samplefcn='rand', S=array([[360.0, 361.0, 362.0,
                                                363.0, 364.0, 365.0, 366.0, 367.0, 368.0, 369.0, 370.0, 371.0,
                                                372.0, 373.0, 374.0, 375.0, 376.0, 377.0, 378.0, 379.0, 380.0,
                                                381.0, 382.0, 383.0, 384.0, 385.0, 386.0, 387.0, 388.0, 389.0,
                                                390.0, 391.0, 392.0, 393.0, 394.0, 395.0, 396.0, 397.0, 398.0,
                                                399.0, 400.0, 401.0, 402.0, 403.0, 404.0, 405.0, 406.0, 407.0,
                                                408.0, 409.0, 410.0, 411.0, 412.0, 413.0, 414.0, 415.0, 416.0,
                                                417.0, 418.0, 419.0, 420.0, 421.0, 422.0, 423.0, 424.0, 425.0,
                                                426.0, 427.0, 428.0, 429.0, 430.0, 431.0, 432.0, 433.0, 434.0,
                                                435.0, 436.0, 437.0, 438.0, 439.0, 440.0, 441.0, 442.0, 443.0,
                                                444.0, 445.0, 446.0, 447.0, 448.0, 449.0, 450.0, 451.0, 452.0,
                                                453.0, 454.0, 455.0, 456.0, 457.0, 458.0, 459.0, 460.0, 461.0,
                                                462.0, 463.0, 464.0, 465.0, 466.0, 467.0, 468.0, 469.0, 470.0,
                                                471.0, 472.0, 473.0, 474.0, 475.0, 476.0, 477.0, 478.0, 479.0,
                                                480.0, 481.0, 482.0, 483.0, 484.0, 485.0, 486.0, 487.0, 488.0,
                                                489.0, 490.0, 491.0, 492.0, 493.0, 494.0, 495.0, 496.0, 497.0,
                                                498.0, 499.0, 500.0, 501.0, 502.0, 503.0, 504.0, 505.0, 506.0,
                                                507.0, 508.0, 509.0, 510.0, 511.0, 512.0, 513.0, 514.0, 515.0,
                                                516.0, 517.0, 518.0, 519.0, 520.0, 521.0, 522.0, 523.0, 524.0,
                                                525.0, 526.0, 527.0, 528.0, 529.0, 530.0, 531.0, 532.0, 533.0,
                                                534.0, 535.0, 536.0, 537.0, 538.0, 539.0, 540.0, 541.0, 542.0,
                                                543.0, 544.0, 545.0, 546.0, 547.0, 548.0, 549.0, 550.0, 551.0,
                                                552.0, 553.0, 554.0, 555.0, 556.0, 557.0, 558.0, 559.0, 560.0,
                                                561.0, 562.0, 563.0, 564.0, 565.0, 566.0, 567.0, 568.0, 569.0,
                                                570.0, 571.0, 572.0, 573.0, 574.0, 575.0, 576.0, 577.0, 578.0,
                                                579.0, 580.0, 581.0, 582.0, 583.0, 584.0, 585.0, 586.0, 587.0,
                                                588.0, 589.0, 590.0, 591.0, 592.0, 593.0, 594.0, 595.0, 596.0,
                                                597.0, 598.0, 599.0, 600.0, 601.0, 602.0, 603.0, 604.0, 605.0,
                                                606.0, 607.0, 608.0, 609.0, 610.0, 611.0, 612.0, 613.0, 614.0,
                                                615.0, 616.0, 617.0, 618.0, 619.0, 620.0, 621.0, 622.0, 623.0,
                                                624.0, 625.0, 626.0, 627.0, 628.0, 629.0, 630.0, 631.0, 632.0,
                                                633.0, 634.0, 635.0, 636.0, 637.0, 638.0, 639.0, 640.0, 641.0,
                                                642.0, 643.0, 644.0, 645.0, 646.0, 647.0, 648.0, 649.0, 650.0,
                                                651.0, 652.0, 653.0, 654.0, 655.0, 656.0, 657.0, 658.0, 659.0,
                                                660.0, 661.0, 662.0, 663.0, 664.0, 665.0, 666.0, 667.0, 668.0,
                                                669.0, 670.0, 671.0, 672.0, 673.0, 674.0, 675.0, 676.0, 677.0,
                                                678.0, 679.0, 680.0, 681.0, 682.0, 683.0, 684.0, 685.0, 686.0,
                                                687.0, 688.0, 689.0, 690.0, 691.0, 692.0, 693.0, 694.0, 695.0,
                                                696.0, 697.0, 698.0, 699.0, 700.0, 701.0, 702.0, 703.0, 704.0,
                                                705.0, 706.0, 707.0, 708.0, 709.0, 710.0, 711.0, 712.0, 713.0,
                                                714.0, 715.0, 716.0, 717.0, 718.0, 719.0, 720.0, 721.0, 722.0,
                                                723.0, 724.0, 725.0, 726.0, 727.0, 728.0, 729.0, 730.0, 731.0,
                                                732.0, 733.0, 734.0, 735.0, 736.0, 737.0, 738.0, 739.0, 740.0,
                                                741.0, 742.0, 743.0, 744.0, 745.0, 746.0, 747.0, 748.0, 749.0,
                                                750.0, 751.0, 752.0, 753.0, 754.0, 755.0, 756.0, 757.0, 758.0,
                                                759.0, 760.0, 761.0, 762.0, 763.0, 764.0, 765.0, 766.0, 767.0,
                                                768.0, 769.0, 770.0, 771.0, 772.0, 773.0, 774.0, 775.0, 776.0,
                                                777.0, 778.0, 779.0, 780.0, 781.0, 782.0, 783.0, 784.0, 785.0,
                                                786.0, 787.0, 788.0, 789.0, 790.0, 791.0, 792.0, 793.0, 794.0,
                                                795.0, 796.0, 797.0, 798.0, 799.0, 800.0, 801.0, 802.0, 803.0,
                                                804.0, 805.0, 806.0, 807.0, 808.0, 809.0, 810.0, 811.0, 812.0,
                                                813.0, 814.0, 815.0, 816.0, 817.0, 818.0, 819.0, 820.0, 821.0,
                                                822.0, 823.0, 824.0, 825.0, 826.0, 827.0, 828.0, 829.0, 830.0],
                                                232
                                                1.0, 1.0, 1.0, 1.0, 1.0, 1.Qhapter.4.1.buxpy.packageostructure
```

```
Args:
            rfl
                  ndarray or str
                  Array with of str referring to a set of spectral reflectance
                        functions to be subsampled.
                  If str to file: file must contain data as columns, with first
                        column the wavelengths.
            rflpath
                  " or str, optional
                  Path to folder with rfl-set specified in a str :rfl: filename.
            samplefcn
                  'rand' or 'mean', optional
                        -'rand': selects a random sample from the samples within each pixel
                        -'mean': returns the mean spectral reflectance in each pixel.
            S
                  _CIE_ILLUMINANTS['E'], optional
                  Illuminant used to calculate the color coordinates of the spectral
                        reflectance samples.
            jab_ranges
                  None or ndarray, optional
                  Specifies the pixelization of color space.
                        (ndarray.shape = (3,3), with first axis: J,a,b, and second
                              axis: min, max, delta)
            jab_deltas
                  float or ndarray, optional
                  Specifies the sampling range.
                  A float uses jab_deltas as the maximum Euclidean distance to select
                  samples around each pixel center. A ndarray of 3 deltas, uses
                  a city block sampling around each pixel center.
            cspace
                  _VF_CSPACE or dict, optional
                  Specifies color space. See _VF_CSPACE_EXAMPLE for example structure.
            cieobs
                  VF CIEOBS or str, optional
                  Specifies CMF set used to calculate color coordinates.
            ax
                  default ndarray or user defined ndarray, optional
                  default = np.arange(-_VF_MAXR,_VF_MAXR+_VF_DELTAR,_VF_DELTAR)
            bx
                  default ndarray or user defined ndarray, optional
                  default = np.arange(-_VF_MAXR,_VF_MAXR+_VF_DELTAR,_VF_DELTAR)
            jx
```

Note that not-None :jab\_ranges: override :ax:, :bx: and :jx input.

None, optional

limit\_grid\_radius

```
0, optional
```

A value of zeros keeps grid as specified by axr,bxr.

A value > 0 only keeps (a,b) coordinates within :limit grid radius:

#### **Returns:**

#### returns

rflsampled, jabp

ndarrays with resp. the subsampled set of spectral reflectance

functions and the pixel coordinate centers.

```
luxpy.color.cri.VFPX.plot_VF_PX_models(dataVF=None, dataPX=None, plot_VF=True, plot_PX=True,
                                             axtype='polar', ax='new', plot circle field=True,
                                             plot_sample_shifts=False,
```

plot samples shifts at pixel center=False, jabp\_sampled=None, plot\_VF\_colors=['g'], plot\_PX\_colors=['r'], hbin\_cmap=None, bin\_labels=None,

plot\_bin\_colors=True, force\_CVG\_layout=False)

Plot the VF and PX model color shift vectors.

### Args:

#### dataVF

None or list[dict] with VF\_colorshift\_model() output, optional

None plots nothing related to VF model.

Each list element refers to a different test SPD.

#### dataPX

None or list[dict] with PX colorshift model() output, optional

None plots nothing related to PX model.

Each list element refers to a different test SPD.

## plot\_VF

True, optional

Plot VF model (if :dataVF: is not None).

## plot\_PX

True, optional

Plot PX model (if :dataPX: is not None).

## axtype

'polar' or 'cart', optional

Make polar or Cartesian plot.

ax

None or 'new' or 'same', optional

- None or 'new' creates new plot
- 'same': continue plot on same axes.
- axes handle: plot on specified axes.

## plot\_circle\_field

True or False, optional

Plot lines showing how a series of circles of color coordinates is

distorted by the test SPD.

The width (wider means more) and color (red means more) of the

lines specify the intensity of the hue part of the color shift.

## plot\_sample\_shifts

False or True, optional

Plots the shifts of the individual samples of the rfl-set used to calculated the VF model.

## plot\_samples\_shifts\_at\_pixel\_center

False, optional

Offers the possibility of shifting the vector shifts of subsampled sets from the reference illuminant positions to the pixel centers.

Note that the pixel centers must be supplied in :jabp\_sampled:.

### jabp\_sampled

None, ndarray, optional

Corresponding pixel center for each sample in a subsampled set.

## plot\_VF\_colors

['g'] or list[str], optional

Specifies the plot color the color shift vectors of the VF model.

If len(:plot\_VF\_colors:) == 1: same color for each list element of :dataVF:.

## plot\_VF\_colors

['g'] or list[str], optional

Specifies the plot color the color shift vectors of the VF model.

If len(:plot\_VF\_colors:) == 1: same color for each list element of :dataVF:.

# hbin\_cmap

None or colormap, optional

Color map with RGB entries for each of the hue bins specified by

the hues in \_VF\_PCOLORSHIFT.

If None: cmap will be obtained on first run by

luxpy.cri.plot\_shift\_data() and returned for use in other functions

### plot\_bin\_colors

True, optional

Colorize hue-bins.

## bin labels

None or list[str] or '#', optional

Plots labels at the bin center hues.

- None: don't plot.
- list[str]: list with str for each bin.

(len(:bin\_labels:) = :nhbins:)

- '#': plots number.
- 'VF PCOLORSHIFT': uses the labels in VF PCOLORSHIFT['labels']
- 'pcolorshift': uses the labels in dataVF['modeldata']['pcolorshift']['labels']

## force\_CVG\_layout

False or True, optional

True: Force plot of basis of CVG.

#### **Returns:**

### returns

ax (handle to current axes), cmap (hbin\_cmap)

# 4.4.10 XYZ,LAB classes

```
рy
                      • CDATA.py
           namespace luxpy
class luxpy.color.CDATA.XYZ(value=None, relative=True, cieobs='1931_2', dtype='xyz')
      ctf(dtype='Yuv', **kwargs)
            Convert XYZ tristimulus values to color space coordinates.
            Args:
                        dtype
                              _CSPACE or str, optional
                              Convert to this color space.
                        kwargs
                              additional input arguments required for
                              color space transformation.
                              See specific luxpy function for more info
                                    (e.g. ?luxpy.xyz_to_lab)
            Returns:
                        returns
                              luxpy.LAB with .value field that is a ndarray
                                    with color space coordinates
      plot(ax=None, title=None, **kwargs)
            Plot tristimulus or cone fundamental values.
            Args:
                        ax
                              None or axes handles, optional
                              None: create new figure axes, else use :ax: for plotting.
                        title
                              None or str, optional
                              Give plot a title.
                        kwargs
                              additional arguments for use with
                              matplotlib.pyplot.scatter
            Returns:
                        gca
                              handle to current axes.
      to_Yxy()
            Convert XYZ tristimulus values CIE Yxy chromaticity values.
            Returns:
                        Yxy
                              luxpy.LAB with .value field that is a ndarray
                              with Yxy chromaticity values.
                              (Y value refers to luminance or luminance factor)
      to_Yuv(**kwargs)
            Convert XYZ tristimulus values CIE 1976 Yu'v' chromaticity values.
```

#### **Returns:**

Yuv

luxpy.LAB with .value field that is a ndarray with CIE 1976 Yu'v' chromaticity values.
(Y value refers to luminance or luminance factor)

# to\_Yuv76(\*\*kwargs)

Convert XYZ tristimulus values CIE 1976 Yu'v' chromaticity values.

#### **Returns:**

Yuv

luxpy.LAB with .value field that is a ndarray with CIE 1976 Yu'v' chromaticity values.
(Y value refers to luminance or luminance factor)

## to\_Yuv60(\*\*kwargs)

Convert XYZ tristimulus values CIE 1960 Yuv chromaticity values.

#### **Returns:**

Yuv

luxpy.LAB with .value field that is a ndarray with CIE 1960 Yuv chromaticity values.
(Y value refers to luminance or luminance factor)

### **to\_wuv**(xyzw=array([100.0, 100.0, 100.0]))

Convert XYZ tristimulus values CIE 1964 U\*V\*W\* color space.

### Args:

**xyzw** 

ndarray with tristimulus values of white point, optional Defaults to luxpy.\_COLORTF\_DEFAULT\_WHITE\_POINT

### **Returns:**

wiiv

luxpy.LAB with .value field that is a ndarray with W\*U\*V\* values.

#### to\_lms()

Convert XYZ tristimulus values or LMS cone fundamental responses to LMS cone fundamental responses.

#### **Returns:**

lms

luxpy.XYZ with .value field that is a ndarray with LMS cone fundamental responses.

#### to\_xyz()

Convert XYZ tristimulus values or LMS cone fundamental responses to XYZ tristimulus values.

## **Returns:**

xyz

luxpy.XYZ with .value field that is a ndarray with XYZ tristimulus values.

## to\_lab(xyzw=None, cieobs='1931\_2')

Convert XYZ tristimulus values to CIE 1976 L\*a\*b\* (CIELAB) coordinates.

## Args:

xyzw

```
None or ndarray with xyz values of white point, optional
                       None defaults to xyz of CIE D65 using the :cieobs: observer.
                 cieobs
                       luxpy._CIEOBS, optional
                       CMF set to use when calculating xyzw.
      Returns:
                 lab
                       luxpy.LAB with .value field that is a ndarray
                       with CIE 1976 L*a*b* (CIELAB) color coordinates
to_luv(xyzw=None, cieobs='1931_2')
     Convert XYZ tristimulus values to CIE 1976 L*u*v* (CIELUV) coordinates.
      Args:
                 xyzw
                       None or ndarray with xyz values of white point, optional
                       None defaults to xyz of CIE D65 using the :cieobs: observer.
                 cieobs
                       luxpy._CIEOBS, optional
                       CMF set to use when calculating xyzw.
      Returns:
                 luv
                       luxpy.LAB with .value field that is a ndarray
                       with CIE 1976 L*u*v* (CIELUV) color coordinates
to_Vrb_mb(cieobs='1931_2', scaling=[1, 1], M=None)
      Convert XYZ tristimulus values to V,r,b (Macleod-Boynton) coordinates.
     Macleod Boynton: V = R+G, r = R/V, b = B/V
     Note that R,G,B ~ L,M,S
     Args:
                 cieobs
                       luxpy._CIEOBS, optional
                       CMF set to use when calculating xyzw.
                 scaling
                       list of scaling factors for r and b dimensions.
                 M
                       None, optional
                       Conversion matrix for going from XYZ to RGB (LMS)
                       If None, :cieobs: determines the M (function does inversion)
      Returns:
                 Vrb
                       luxpy.LAB with .value field that is a ndarray
                       ndarray with V,r,b (Macleod-Boynton) color coordinates
to_ipt(cieobs='1931_2', xyzw=None, M=None)
     Convert XYZ tristimulus values to IPT color coordinates.
```

I: Lightness axis, P, red-green axis, T: yellow-blue axis. Args: xyzw None or ndarray with xyz values of white point, optional None defaults to xyz of CIE D65 using the :cieobs: observer. cieobs luxpy.\_CIEOBS, optional CMF set to use when calculating xyzw for rescaling Mxyz2lms (only when not None). M None, optional None defaults to conversion matrix determined by :cieobs: **Returns:** ipt luxpy.LAB with .value field that is a ndarray with IPT color coordinates Note: xyz is assumed to be under D65 viewing conditions!! | If necessary perform chromatic adaptation!! to\_Ydlep(cieobs='1931\_2', xyzw=array([100.0, 100.0, 100.0])) Convert XYZ values to Y, dominant (complementary) wavelength and excitation purity. Args: **xyzw** None or ndarray with xyz values of white point, optional None defaults to xyz of CIE D65 using the :cieobs: observer. cieobs luxpy.\_CIEOBS, optional CMF set to use when calculating spectrum locus coordinates. **Returns: Ydlep** ndarray with Y, dominant (complementary) wavelength and excitation purity to\_srgb(gamma=2.4) Calculates IEC:61966 sRGB values from xyz. Args: xyz ndarray with relative tristimulus values.

gamma

2.4, optional

compression in sRGB

**Returns:** 

rgb

ndarray with R,G,B values (uint8).

to\_jabz(ztype='jabz')

Convert XYZ tristimulus values to Jz,az,bz color coordinates.

Args:

```
xyz
                        ndarray with absolute tristimulus values (Y in cd/m<sup>2</sup>!)
                  ztype
                        'jabz', optional
                        String with requested return:
                        Options: 'jabz', 'iabz'
      Returns:
                  jabz
                        ndarray with Jz,az,bz color coordinates
      Notes:
            1. :xyz: is assumed to be under D65 viewing conditions! If necessary perform chromatic
            adaptation!
            2a. Jz represents the 'lightness' relative to a D65 white with luminance = 10000 \text{ cd/m}^2
                  (note that Jz that not exactly equal 1 for this high value, but rather for 102900 cd/m2)
            2b. az, bz represent respectively a red-green and a yellow-blue opponent axis
                  (but note that a D65 shows a small offset from (0,0))
     Reference: 1. Safdar, M., Cui, G., Kim, Y. J., and Luo, M. R. (2017). Perceptually uniform color space
            for image signals including high dynamic range and wide gamut. Opt. Express, vol. 25, no. 13, pp.
            15131–15151, Jun. 2017.
to_jabM_ciecam02(xyzw=array([[100.0, 100.0, 100.0]]), Yw=100.0, conditions={'D': 1.0, 'Dtype': None,
                     'La': 100.0, 'Yb': 20.0, 'surround': 'avg'}, naka_rushton_parameters=None,
                     unique_hue_data=None, yellowbluepurplecorrect=None, mcat='cat02')
      See ?luxpy.xyz_to_jabM_ciecam02
to_jabC_ciecam02(xyzw=array([[100.0, 100.0, 100.0]]), Yw=100.0, conditions={'D': 1.0, 'Dtype': None,
                     'La': 100.0, 'Yb': 20.0, 'surround': 'avg', naka_rushton_parameters=None,
                     unique_hue_data=None, yellowbluepurplecorrect=None, mcat='cat02')
      See ?luxpy.xyz_to_jabC_ciecam02
to_jab_cam02ucs(xyzw=array([[100.0, 100.0, 100.0]]), Yw=100.0, conditions={'D': 1.0, 'Dtype': None,
                    'La': 100.0, 'Yb': 20.0, 'surround': 'avg'}, naka_rushton_parameters=None,
                    unique hue data=None, yellowbluepurplecorrect=None, mcat='cat02')
     See ?luxpy.xyz_to_jab_cam02ucs
to_jab_cam02lcd(xyzw=array([[100.0, 100.0, 100.0]]), Yw=100.0, conditions={'D': 1.0, 'Dtype': None,
                    'La': 100.0, 'Yb': 20.0, 'surround': 'avg'}, naka_rushton_parameters=None,
                    unique_hue_data=None, yellowbluepurplecorrect=None, mcat='cat02')
      See ?luxpy.xyz_to_jab_cam02lcd
to_jab_cam02scd(xyzw=array([[100.0, 100.0, 100.0]]), Yw=100.0, conditions={'D': 1.0, 'Dtype': None,
                    'La': 100.0, 'Yb': 20.0, 'surround': 'avg'}, naka_rushton_parameters=None,
                    unique_hue_data=None, yellowbluepurplecorrect=None, mcat='cat02')
      See ?luxpy.xyz_to_jab_cam02scd
to_jabM_ciecam16(xyzw=array([[100.0, 100.0, 100.0]]), Yw=100.0, conditions={'D': 1.0, 'Dtype': None,
                     'La': 100.0, 'Yb': 20.0, 'surround': 'avg'}, naka_rushton_parameters=None,
                     unique hue data=None, mcat='cat16')
     See ?luxpy.xyz_to_jabM_ciecam16
```

```
'La': 100.0, 'Yb': 20.0, 'surround': 'avg'}, naka_rushton_parameters=None,
                          unique hue data=None, mcat='cat16')
           See ?luxpy.xyz_to_jabC_ciecam16
     to_jab_cam16ucs(xyzw=array([[100.0, 100.0, 100.0]]), Yw=100.0, conditions={'D': 1.0, 'Dtype': None,
                         'La': 100.0, 'Yb': 20.0, 'surround': 'avg'}, naka rushton parameters=None,
                         unique hue data=None, mcat='cat16')
           See ?luxpy.xyz_to_jab_cam02ucs
     to_jab_cam16lcd(xyzw=array([[100.0, 100.0, 100.0]]), Yw=100.0, conditions=['D': 1.0, 'Dtype': None,
                         'La': 100.0, 'Yb': 20.0, 'surround': 'avg'}, naka_rushton_parameters=None,
                         unique_hue_data=None, mcat='cat16')
           See ?luxpy.xyz_to_jab_cam16lcd
     to_jab_cam16scd(xyzw=array([[100.0, 100.0, 100.0]]), Yw=100.0, conditions={'D': 1.0, 'Dtype': None,
                         'La': 100.0, 'Yb': 20.0, 'surround': 'avg'}, naka_rushton_parameters=None,
                         unique_hue_data=None, mcat='cat16')
           See ?luxpy.xyz_to_jab_cam16scd
     to_jabM_zcam(xyzw=None, conditions={'D': 1.0, 'Dtype': None, 'La': 100.0, 'Yb': 20.0, 'surround': 'avg'},
                     mcat='cat16'
           See ?luxpy.xyz_to_jabM_zcam
     to_jabC_zcam(xyzw=array([[100.0, 100.0, 100.0]]), conditions={'D': 1.0, 'Dtype': None, 'La': 100.0, 'Yb':
                     20.0, 'surround': 'avg'}, mcat='cat16')
           See ?luxpy.xyz_to_jabC_zcam
     to_qabW_cam15u(fov=10.0, parameters=None)
           See ?luxpy.xyz_to_qabW_cam15u
     to_lab_cam_sww_2016(xyzw=None, Yb=20.0, Lw=400.0, relative=True, parameters=None, inputtype='xyz',
                             cieobs='2006 10')
           See ?luxpy.xyz_to_lab_cam_sww_2016
     to_qabS_cam18s1(xyz, xyzb=None, Lb=[100], fov=10.0, parameters=None)
           See ?luxpy.xyz_to_qabS_cam18sl
     to_qabM_cam18s1(xyz, xyzb=None, Lb=[100], fov=10.0, parameters=None)
           See ?luxpy.xyz to gabM cam18sl
class luxpy.color.CDATA.LAB(value=None, relative=True, cieobs='1931 2', dtype='lab', xyzw=None,
                                 M=None, scaling=None, Lw=None, Yw=None, Yb=None, conditions=None,
                                 naka_rushton_parameters=None, unique_hue_data=None,
                                 yellowbluepurplecorrect=None, mcat=None, ucstype=None, fov=None,
                                 parameters=None)
     ctf(**kwargs)
           Convert color space coordinates to XYZ tristimulus values.
           Args:
                       dtype
                             'xyz'
                            Convert to this color space.
                       kwargs
                            additional input arguments required for
                            color space transformation.
                            See specific luxpy function for more info
                                  (e.g. ?luxpy.xyz_to_lab)
```

to\_jabC\_ciecam16(xyzw=array([[100.0, 100.0, 100.0]]), Yw=100.0, conditions={'D': 1.0, 'Dtype': None,

```
Returns:
                  returns
                         luxpy.XYZ with .value field that is a ndarray
                        with tristimulus values
plot(plt_type='3d', ax=None, title=None, **kwargs)
      Plot color coordinates.
      Args:
                  plt_type
                         '3d' or 3 or '2d or 2, optional
                               -'3d' or 3: plot all 3 dimensions (lightness and chromaticity)
                               -'2d' or 2: plot only chromaticity dimensions.
                  ax
                        None or axes handles, optional
                        None: create new figure axes, else use :ax: for plotting.
                  title
                        None or str, optional
                        Give plot a title.
                  kwargs
                         additional arguments for use with
                         matplotlib.pyplot.scatter
      Returns:
                  gca
                        handle to current axes.
to_xyz(**kwargs)
      Convert color space coordinates to XYZ tristimulus values.
```

# 4.5 Toolboxes

# 4.5.1 photbiochem/

рy

- \_\_init\_\_.py
- cie\_tn003\_2015.py
- ASNZS\_1680\_2\_5\_1997\_COI.py
- circadian\_CS\_CLa\_lrc.py

namespace luxpy.photbiochem

## Module for calculating CIE (S026:2018 & TN003:2015) photobiological quantities

Photore-	Photopigment (la-	Spectral ef	fi-	Quantity (-opic irradi-	Q-symbol	Unit sym-
ceptor	bel, )	ciency s()		ance)	(Ee,)	bol
1-cone	photopsin (lc)	erythrolabe		erythropic	Ee,lc	W.m2
m-cone	photopsin (mc)	chlorolabe		chloropic	Ee,mc	W.m2
s-cone	photopsin (sc)	cyanolabe		cyanopic	Ee,sc	W.m2
rod	rhodopsin (r)	rhodopic		rhodopic	Ee,r	W.m2
ipRGC	melanopsin (z)	melanopic		melanopic	Ee,z	W.m2

CIE recommends that the -opic irradiance is determined by convolving the spectral irradiance, Ee,() (Wm2), for each wavelength, with the action spectrum, s(), where s() is normalized to one at its peak:

$$Ee_{s} = Ee_{s}(s) d$$

where the corresponding units are Wm2 in each case.

The equivalent luminance is calculated as:

$$E_{1} = Km \quad Ee_{1}(s) d \quad V(s) d / s(s) d$$

To avoid ambiguity, the weighting function used must be stated, so, for example, cyanopic refers to the cyanopic irradiance weighted using the s-cone or ssc() spectral efficiency function.

ndarray with default CIE-S026:2018 alpha-actinic action spectra. (stored in file:

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'./data/cie\_S026\_2018\_SI\_action\_spectra\_CIEToolBox\_v1.049.dat')

### ACTIONSPECTRA CIES026

ndarray with alpha-actinic action spectra. (stored in file: './data/cie\_S026\_2018\_SI\_action\_spectra\_CIEToolBox\_v1.049.dat')

### \_ACTIONSPECTRA\_CIETN003

ndarray with CIE-TN003:2015 alpha-actinic action spectra. (stored in file: './data/cie\_tn003\_2015\_SI\_action\_spectra.dat')

## spd\_to\_aopicE()

Calculate alpha-opic irradiance (Ee,) and equivalent luminance (E) values for the l-cone, m-cone, s-cone, rod and iprgc () photoreceptor cells following CIE S026:2018 (= default actionspectra) or CIE TN003:2015.

## spd\_to\_aopicEDI()

Calculate alpha-opic equivalent daylight (D65) illuminance (lx) for the l-cone, m-cone, s-cone, rod and iprgc () photoreceptor cells.

## spd\_to\_aopicDER()

Calculate -opic Daylight (D65) Efficacy Ratio for the l-cone, m-cone, s-cone, rod and iprgc () photoreceptor cells.

## spd\_to\_aopicELR()

Calculate -opic Efficacy of Luminous Radiation for the l-cone, m-cone, s-cone, rod and iprgc () photoreceptor cells.

**References:** 1. CIE-S026:E2018 (2018). CIE System for Metrology of Optical Radiation for ipRGC-Influenced Responses to Light (Vienna, Austria). (https://files.cie.co.at/CIE%20S%20026%20alpha-opic%20Toolbox%20User%20Guide.pdf)

2. CIE-TN003:2015 (2015). Report on the first international workshop on circadian and neurophysiological photometry, 2013 (Vienna, Austria). (http://files.cie.co.at/785\_CIE\_TN\_003-2015.pdf)

### Module for calculation of cyanosis index (AS/NZS 1680.2.5:1997)

\_COI\_OBS Default CMF set for calculations
\_COI\_CSPACE Default color space (CIELAB)

\_COI\_RFL\_BLOOD ndarray with reflectance spectra of 100% and 50% oxygenated blood

spd\_to\_COI\_ASNZS1680 Calculate the Cyanosis Observartion Index (COI) [ASNZS
1680.2.5-1995]

Reference: AS/NZS1680.2.5 (1997). INTERIOR LIGHTING PART 2.5: HOSPITAL AND MEDICAL TASKS.

## Module for Blue light hazard calculations

```
_BLH Blue Light Hazard function
```

spd\_to\_blh\_eff() Calculate Blue Light Hazard efficacy (K) or efficiency (eta) of radiation.

#### **References:**

- 1. IEC 62471:2006, 2006, Photobiological safety of lamps and lamp systems.
- 2. IEC TR 62778, 2014, Application of IEC 62471 for the assessment of blue light hazard to light sources and luminaires.

```
luxpy.toolboxes.photbiochem.spd_to_aopicE(sid, Ee=None, E=None, Q=None, cieobs='1931_2', sid_units='W/m2', out='Eeas', actionspectra='CIE-S026')
```

Calculate alpha-opic irradiance (Ee,) values  $(W/m^2)$  for the l-cone, m-cone, s-cone, rod and iprgc () photoreceptor cells following CIE S026:2018.

### Args:

```
sid
```

numpy.ndarray with retinal spectral irradiance in :sid\_units: (if 'uW/cm2', sid will be converted to SI units 'W/m2')

Ee

None, optional

If not None: normalize :sid: to an irradiance of :Ee:

E

None, optional

If not None: normalize :sid: to an illuminance of :E:

Note that E is calculate using a Km factor corrected to standard air.

Q

None, optional

If not None: Normalize :sid: to a quantal energy of :Q:

# cieobs

\_CIEOBS or str, optional

Type of cmf set to use for photometric units.

# $sid\_units$

'W/m2', optional

Other option 'uW/m2', input units of :sid:

out

'Eeas' or str, optional

Determines values to return.

(to get also get equivalent illuminance E set :out: to 'Eeas,Eas')

### actionspectra

'CIES026', optional

Actionspectra to use in calculation

options:

- 'CIE-S026': will use action spectra as defined in CIE S026
- 'CIE-TN003': will use action spectra as defined in CIE TN003

# **Returns:**

#### returns

Eeas a numpy.ndarray with the -opic irradiance

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of all spectra in :sid: in SI-units (W/m<sup>2</sup>).

```
(other choice can be set using :out:)
      References: 1. CIE-S026:E2018 (2018). CIE System for Metrology of Optical Radiation for ipRGC-Influenced
            Responses to Light (Vienna, Austria).
                                                          (https://files.cie.co.at/CIE%20S%20026%20alpha-opic%
            20Toolbox%20User%20Guide.pdf)
            2. CIE-TN003:2015 (2015). Report on the first international workshop on circadian and neurophysiolog-
            ical photometry, 2013 (Vienna, Austria). (http://files.cie.co.at/785_CIE_TN_003-2015.pdf)
luxpy.toolboxes.photbiochem.spd_to_aopicEDI(sid, Ee=None, E=None, Q=None, cieobs='1931_2',
                                                        sid_units='W/m2', actionspectra='CIE-S026', ref='D65',
                                                        out='a edi')
      Calculate alpha-opic equivalent daylight (D65) illuminance (lux) for the l-cone, m-cone, s-cone, rod and iprgc ()
      photoreceptor cells.
      Args:
                  sid
                        numpy.ndarray with retinal spectral irradiance in :sid_units:
                        (if 'uW/cm2', sid will be converted to SI units 'W/m2')
                  Ee
                        None, optional
                        If not None: normalize :sid: to an irradiance of :Ee:
                  Е
                        None, optional
                        If not None: normalize :sid: to an illuminance of :E:
                        Note that E is calculate using a Km factor corrected to standard air.
                  O
                        None, optional
                        If not None: nNormalize :sid: to a quantal energy of :Q:
                  cieobs
                        _CIEOBS or str, optional
                        Type of cmf set to use for photometric units.
                  sid_units
                        'W/m2', optional
                        Other option 'uW/m2', input units of :sid:
                  actionspectra
                        'CIES026', optional
                        Actionspectra to use in calculation
                        - 'CIE-S026': will use action spectra as defined in CIE S026
                        - 'CIE-TN003': will use action spectra as defined in CIE TN003
                  ref
                        'D65', optional
                        Reference (daylight) spectrum to use. ('D65' or 'E' or ndarray)
                  out
                        'Eeas, Eas' or str, optional
                        Determines values to return.
```

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**Returns:** 

```
ndarray with the -opic Equivalent Daylight Illuminance (lux) with the
                        for the 1-cone, m-cone, s-cone, rod and iprgc photoreceptors
                        of all spectra in :sid: in SI-units.
luxpy.toolboxes.photbiochem.spd_to_aopicDER(sid, cieobs='1931_2', sid_units='W/m2',
                                                         actionspectra='CIE-S026', ref='D65')
      Calculate -opic Daylight (D65) Efficacy Ratio (= -opic Daylight (D65) Efficiency) for the l-cone, m-cone, s-cone,
      rod and iprgc () photoreceptor cells.
      Args:
                  sid
                        numpy.ndarray with retinal spectral irradiance in :sid_units:
                        (if 'uW/cm2', sid will be converted to SI units 'W/m2')
                  cieobs
                        _CIEOBS or str, optional
                        Type of cmf set to use for photometric units.
                  sid units
                        'W/m2', optional
                        Other option 'uW/m2', input units of :sid:
                  actionspectra
                        'CIES026', optional
                        Actionspectra to use in calculation
                        options:
                        - 'CIE-S026': will use action spectra as defined in CIE S026
                        - 'CIE-TN003': will use action spectra as defined in CIE TN003
                  ref
                        'D65', optional
                        Reference (daylight) spectrum to use. ('D65' or 'E' or ndarray)
      Returns:
                  returns
                        ndarray with the -opic Daylight Efficacy Ratio with the
                        for the 1-cone, m-cone, s-cone, rod and iprgc photoreceptors
                        of all spectra in :sid: in SI-units.
luxpy.toolboxes.photbiochem.spd_to_aopicELR(sid, cieobs='1931_2', sid_units='W/m2',
                                                         actionspectra='CIE-S026', ref='D65')
      Calculate -opic Efficacy of Luminous Radiation (W/lm) for the l-cone, m-cone, s-cone, rod and iprgc () photore-
      ceptor cells.
      Args:
                  sid
                        numpy.ndarray with retinal spectral irradiance in :sid_units:
                        (if 'uW/cm2', sid will be converted to SI units 'W/m2')
                  cieobs
                         _CIEOBS or str, optional
                        Type of cmf set to use for photometric units.
                  sid units
                        'W/m2', optional
                        Other option 'uW/m2', input units of :sid:
```

returns

4.5. Toolboxes 247

```
actionspectra
                        'CIES026', optional
                        Actionspectra to use in calculation
                        options:
                        - 'CIE-S026': will use action spectra as defined in CIE S026
                        - 'CIE-TN003': will use action spectra as defined in CIE TN003
                  ref
                        'D65', optional
                        Reference (daylight) spectrum to use. ('D65' or 'E' or ndarray)
      Returns:
                  returns
                        ndarray with the -opic Efficacy of Luminous Radiation (W/lm) with the
                        for the 1-cone, m-cone, s-cone, rod and iprgc photoreceptors
                        of all spectra in :sid: in SI-units.
luxpy.toolboxes.photbiochem.spd_to_COI_ASNZS1680(S=None, tf='lab', cieobs='1931 2', out='COI,cct',
                                                               extrapolate_rfl=False)
      Calculate the Cyanosis Observation Index (COI) [ASNZS 1680.2.5-1995].
      Args:
                  \mathbf{S}
                        ndarray with light source spectrum (first column are wavelengths).
                  tf
                        _COI_CSPACE, optional
                        Color space in which to calculate the COI.
                        Default is CIELAB.
                  cieobs
                        COI CIEOBS, optional
                        CMF set to use.
                        Default is '1931 2'.
                  out
                        'COI,cct' or str, optional
                        Determines output.
                  extrapolate_rfl
                        False, optional
                        If False:
                              limit the wavelength range of the source to that of the standard
                              reflectance spectra for the 50% and 100% oxygenated blood.
      Returns:
                  COI
                        ndarray with cyanosis indices for input sources.
                  cct
                        ndarray with correlated color temperatures.
      Note: Clause 7.2 of the ASNZS 1680.2.5-1995. standard mentions the properties demanded of the light source
            used in region where visual conditions suitable to the detection of cyanosis should be provided:
```

1. The correlated color temperature (CCT) of the source should be from 3300 to 5300 K.

2. The cyanosis observation index should not exceed 3.3

```
luxpy.toolboxes.photbiochem.spd_to_CS_CLa_lrc(El=None, version='CLa2.0', E=None,
                                                            sum_sources=False, interpolate_sources=True,
                                                            t CS=1.0, f CS=1.0)
      Calculate Circadian Stimulus (CS) and Circadian Light (CLa, CLa2.0).
      Args:
                  El
                         ndarray, optional
                         Defaults to D65
                         light source spectral irradiance distribution
                   version
                         'CLa2.0', optional
                         CLa version to calculate
                         Options:
                         - 'CLa1.0': Rea et al. 2012
                         - 'CLa2.0': Rea et al. 2021
                  \mathbf{E}
                         None, float or ndarray, optional
                         Illuminance of light sources.
                         If None: El is used as is, otherwise El is renormalized to have
                         an illuminance equal to E.
                   sum_sources
                         False, optional
                               - False: calculate CS (1.0,2.0) and CLa (1.0, 2.0) for all sources in El array.
                               - True: sum sources in El to a single source and perform calc.
                  interpolate_sources
                         True, optional
                               - True: El is interpolated to wavelength range of efficiency
                                     functions (as in LRC calculator).
                               - False: interpolate efficiency functions to source range.
                                     Source interpolation is not recommended due to possible
                                     errors for peaky spectra.
                                     (see CIE15-2018, "Colorimetry").
                  t_CS
                         1.0, optional
                         The duration factor (in hours): a continuous value from 0.5 to 3.0
                  f_CS
                         1.0, optional
                         The spatial distribution factor: a discrete value (2, 1, or 0.5)
                         depending upon the spatial distribution of the light source.
                         Default = 1 (for t = 1 h, CS is equal to the 2012 version).
                         Options:
                         - 2.0: full visual field, as with a Ganzfeld.
                         - 1.0: central visual field, as with a discrete light box on a desk.
                         - 0.5: superior visual field, as from ceiling mounted down-light fixtures.
      Returns:
                   CS
```

ndarray with Circadian stimulus values

CLa

ndarray with Circadian Light values

- Notes on CLa1.0 (2012 version): 1. The original 2012 (E.q. 1) had set the peak wavelength of the melanopsin at 480 nm. Rea et al. later published a corrigendum with updated model parameters for k, a\_{b-y} and a\_rod. The comparison table between showing values calculated for a number of sources with the old and updated parameters were very close (~1 unit voor CLa).
  - 2. In that corrrection paper they did not mention a change in the factor (1622) that multiplies the (sum of) the integral(s) in Eq. 1. HOWEVER, the excel calculator released in 2017 and the online calculator show that factor to have a value of 1547.9. The change in values due to the new factor is much larger than their the updated mentioned in note 1!
  - 3. For reasons of consistency the calculator uses the latest model parameters, as could be read from the excel calculator. They values adopted are: multiplier 1547.9, k = 0.2616,  $a_{b-y} = 0.7$  and  $a_{rod} = 3.3$ .
  - 4. The parameter values to convert CLa to CS were also taken from the 2017 excel calculator.
- Notes on CLa2.0 (2021 version): 1. In the original model, 1000 lux of CIE Illuminant A resulted in a CLA = 1000. In the revised model, a photopic illuminance of 1000 lux from CIE Illuminant A (approximately that of an incandescent lamp operated at 2856 K) results in a CLA 2.0 = 813. The value of 813 CLA 2.0 should be used by those wishing to calibrate instrumentation designed to report CLA 2.0 and CS. CLA 2.0 values can still be used to approximate the photopic illuminance, in lux, from a nonspecific "white" light source. For comparison, CLA 2.0 values should be multiplied by 1.23 to estimate the equivalent photopic illuminance from CIE Illuminant A, or by 0.66 to estimate the equivalent photopic illuminance from CIE Illuminant D65 (an approximation of daylight with a CCT of 6500 K).
  - 2. Nov. 6, 2012: To get a value of CLa2.0 = 813, Eq. 3 from the paper must be adjusted to also divide by the transmision of the macula ('mp' in paper) the S-cone and Vlambda functions prior to calculating the integrals in the denominators of the first factor after the a\_rod\_1 and a\_rod\_2 scalars! Failure to do so results in a CLa2.0 of 800, instead of the reported 813 by the online calculator. Verification of the code on github shows indeed that these denominators are calculated by using the macular transmission divided S-cone and Vlambda functions. Is this an error in the code or in the paper?

### References:

- 1. LRC Online Circadian stimulus calculator
- 2. LRC Excel based Circadian stimulus calculator.
- 3. Rea MS, Figueiro MG, Bierman A, and Hamner R (2012). Modelling the spectral sensitivity of the human circadian system. Light. Res. Technol. 44, 386–396.
- 4. Rea MS, Figueiro MG, Bierman A, and Hamner R (2012). Erratum: Modeling the spectral sensitivity of the human circadian system (Lighting Research and Technology (2012) 44:4 (386-396)). Light. Res. Technol. 44, 516.
- 5. Rea, M. S., Nagare, R., & Figueiro, M. G. (2021). Modeling Circadian Phototransduction: Quantitative Predictions of Psychophysical Data. Frontiers in Neuroscience, 15, 44.
- 6. LRC Online Circadian stimulus calculator (CLa2.0, 2021)

luxpy.toolboxes.photbiochem. ${\tt CLa\_to\_CS}(CLa, t=1, f=1, forward=True)$  Convert Circadian Light to Circadian Stimulus (and back). Args:

CLa

ndarray with Circadian Light values or Circadian Stimulus values (if forward == False)

t

1.0, optional

```
1.0, optional
                         The spatial distribution factor: a discrete value (2, 1, or 0.5)
                         depending upon the spatial distribution of the light source.
                         Default = 1 (for t = 1 h, CS is equal to the 2012 version).
                         Options:
                         - 2.0: full visual field, as with a Ganzfeld.
                         - 1.0: central visual field, as with a discrete light box on a desk.
                         - 0.5: superior visual field, as from ceiling mounted down-light fixtures.
                   forward
                         True, optional
                         If True: convert CLa to CS values.
                         If False: convert CS values to CLa values.
      Returns:
                   CS
                         ndarray with CS values or with CLa values (if forward == False)
      References: 1. Rea MS, Figueiro MG, Bierman A, and Hamner R (2012). Modelling the spectral sensitivity of
            the human circadian system. Light. Res. Technol. 44, 386–396.
            2. Rea MS, Figueiro MG, Bierman A, and Hamner R (2012). Erratum: Modeling the spectral sensitivity
            of the human circadian system (Lighting Research and Technology (2012) 44:4 (386-396)). Light. Res.
            Technol. 44, 516.
            3. Rea, M. S., Nagare, R., & Figueiro, M.G. (2021). Modeling Circadian Phototransduction: Quantitative
            Predictions of Psychophysical Data. Frontiers in Neuroscience, 15, 44.
            4. LRC Online Circadian Stimulus calculator (CLa2.0, 2021)
luxpy.toolboxes.photbiochem.spd_to_blh_eff(spd, efficacy=True, cieobs='1931 2', scr='dict', K=None)
      Calculate Blue Light Hazard efficacy (K) or efficiency (eta) of radiation.
      Args:
                   \mathbf{S}
                         ndarray with spectral data
                   cieobs
                         str, optional
                         Sets the type of Vlambda function to obtain.
                   scr
                         'dict' or array, optional
                         - 'dict': get from ybar from _CMF
                         - 'array': ndarray in :cieobs:
                         Determines whether to load cmfs from file (./data/cmfs/)
                         or from dict defined in .cmf.py
                         Vlambda is obtained by collecting Ybar.
                   K
                         None, optional
                               e.g. K = 683 \text{ lm/W for '} 1931_2' \text{ (relative == False)}
                               or K = 100/\text{sum}(\text{spd*dl}) (relative == True)
      Returns:
                   eff
```

The duration factor (in hours): a continuous value from 0.5 to 3.0

f

ndarray with blue light hazard efficacy or efficiency of radiation values.

#### References:

- 1. IEC 62471:2006, 2006, Photobiological safety of lamps and lamp systems.
- 2. IEC TR 62778, 2014, Application of IEC 62471 for the assessment of blue light hazard to light sources and luminaires.

# 4.5.2 indvcmf/

рy

- \_\_init\_\_.py
- individual\_observer\_cmf\_model.py

namespace luxpy.indvcmf

## Module for Individual Observer Ims-CMFs (Asano, 2016 and CIE TC1-97)

**\_DATA\_PATH** path to data files

\_DATA Dict with required data

\_DSRC\_STD\_DEF default data source for stdev of physiological data ('matlab', 'germany')

\_DSRC\_LMS\_ODENS\_DEF default data source for lms absorbances and optical densities ('asano', 'cietc197')

\_LMS\_TO\_XYZ\_METHOD default method to calculate lms to xyz conversion matrix ('asano', 'cietc197')

WL CRIT critical wavelength above which interpolation of S-cone data fails.

**\_WL** default wavelengths of spectral data in INDVCMF\_DATA.

load\_database() Load a database with parameters and data required by the Asano model.

init() Initialize: load database required for Asano Individual Observer Model into the default \_DATA dict and set some options for rounding, sign. figs and chopping small value to zero; for source data to use for spectral data for LMS absorp. and optical densities, ...

query\_state() print current settings for global variables.

compute\_cmfs() Generate Individual Observer CMFs (cone fundamentals) based on CIE2006 cone fundamentals and published literature on observer variability in color matching and in physiological parameters (Use of Asano optical data and model; or of CIE TC1-91 data and 'variability'-extended model possible).

cie2006cmfsEx() Generate Individual Observer CMFs (cone fundamentals) based on CIE2006 cone fundamentals and published literature on observer variability in color matching and in physiological parameters. (Use of Asano optical data and model; or of CIE TC1-91 data and 'variability'-extended model possible)

getMonteCarloParam() Get dict with normally-distributed physiological factors for a population of observers.

getUSCensusAgeDist() Get US Census Age Distribution

**genMonteCarloObs()** Monte-Carlo generation of individual observer color matching functions (cone fundamentals) for a certain age and field size.

getCatObs() Generate cone fundamentals for categorical observers.

**get\_lms\_to\_xyz\_matrix()** Calculate lms to xyz conversion matrix for a specific field size determined as a weighted combination of the 2° and 10° matrices.

**lmsb\_to\_xyzb()** Convert from LMS cone fundamentals to XYZ CMFs using conversion matrix determined as a weighted combination of the 2° and 10° matrices.

add\_to\_cmf\_dict() Add set of cmfs to \_CMF dict.

plot\_cmfs() Plot cmf set.

### References

- 1. Asano Y, Fairchild MD, and Blondé L (2016). Individual Colorimetric Observer Model. PLoS One 11, 1–19.
- 2. Asano Y, Fairchild MD, Blondé L, and Morvan P (2016). Color matching experiment for highlighting interobserver variability. Color Res. Appl. 41, 530–539.
- 3. CIE TC1-36 (2006). Fundamental Chromaticity Diagram with Physiological Axes Part I (Vienna: CIE).
- 4. Asano's Individual Colorimetric Observer Model
- 5. CIE TC1-97 cmf functions python code developed by Ivar Farup and Jan Hendrik Wold.

### **Notes**

1. Port of Matlab code from: https://www.rit.edu/cos/colorscience/re\_AsanoObserverFunctions.php (Accessed April 20, 2018) 2. Adjusted/extended following CIE TC1-97 Python code (and data): github.com/ifarup/ciefunctions (Copyright (C) 2012-2017 Ivar Farup and Jan Henrik Wold) (Accessed Dec 18, 2019)

luxpy.toolboxes.indvcmf.load\_database(wl=None, dsrc\_std=None, dsrc\_lms\_odens=None, path=None) Load database required for Asano Individual Observer Model.

## **Args:**

wl

None, optional

Wavelength range to interpolate data to.

None defaults to the wavelength range associated with data in :dsrc\_lms\_odens:

## path

None, optional

Path where data files are stored (If None: look in ./data/ folder under toolbox path)

## dsrc\_std

None, optional

Data source ('matlab' code, or 'germany') for stdev data on physiological factors.

None defaults to string in \_DSRC\_STD\_DEF

## dsrc\_lms\_odens

None, optional

Data source ('asano', 'cietc197') for LMS absorbance and optical density data.

None defaults to string in \_DSRC\_LMS\_ODENS\_DEF

## **Returns:**

data

dict with data for:

- 'LMSa': LMS absorbances
- 'rmd': relative macular pigment density
- 'docul': ocular media optical density
- 'USCensus2010population': data (age and numbers) on a 2010 US Census
- 'CatObsPfctr': dict with iteratively derived Categorical Observer physiological stdevs.
- 'M2d': Asano 2° lms to xyz conversion matrix
- 'M10d': Asano 10° lms to xyz conversion matrix
- standard deviations on physiological parameters: 'od\_lens', 'od\_macula', 'od\_L', 'od\_M', 'od\_S', 'shft\_L', 'shft\_M', 'shft\_S'

Initialize: load database required for Asano Individual Observer Model into the default \_DATA dict and set some options for rounding, sign. figs and chopping small value to zero; for source data to use for spectral data for LMS absorp. and optical desnities, ...

## Args:

### wl

None, optional

Wavelength range to interpolate data to.

None defaults to the wavelength range associated with data in :dsrc lms odens:

### dsrc std

None, optional

Data source ('matlab' code, or 'germany') for stdev data on physiological factors.

None defaults to string in \_DSRC\_STD\_DEF

# dsrc\_lms\_odens

None, optional

Data source ('asano', 'cietc197') for LMS absorbance and optical density data.

None defaults to string in DSRC LMS ODENS DEF

## lms\_to\_xyz\_method

None, optional

Method to use to determine lms-to-xyz conversion matrix (options: 'asano', 'cietc197')

## use\_my\_round

True, optional

If True: use my\_rounding() conform CIE TC1-91 Python code 'ciefunctions'. (slows down code)

by setting \_USE\_MY\_ROUND.

## use\_sign\_figs

True, optional

If True: use sign\_figs() conform CIE TC1-91 Python code 'ciefunctions'. (slows down code)

by setting \_USE\_SIGN\_FIGS.

## use\_chop

True, optional

```
If True: use chop() conform CIE TC1-91 Python code 'ciefunctions'. (slows down
                        code)
                        by setting _USE_CHOP.
                  path
                        None, optional
                        Path where data files are stored (If None: look in ./data/ folder under toolbox path)
                  out
                        None, optional
                        If None: only set global variables, do not output _DATA.copy()
                  verbosity
                        1, optional
                        Print new state of global settings.
      Returns:
                  data
                        if out is not None: return a dict with dict with data for:
                        - 'LMSa': LMS absorbances
                        - 'rmd': relative macular pigment density
                        - 'docul': ocular media optical density
                        - 'USCensus2010population': data (age and numbers) on a 2010 US Census
                        - 'CatObsPfctr': dict with iteratively derived Categorical Observer physiological
                        stdevs.
                        - 'M2d': Asano 2° lms to xyz conversion matrix
                        - 'M10d': Asano 10° lms to xyz conversion matrix
                        - standard deviations on physiological parameters: 'od_lens', 'od_macula', 'od_L',
                        'od_M', 'od_S', 'shft_L', 'shft_M', 'shft_S'
luxpy.toolboxes.indvcmf.query_state()
      Print current settings for 'global variables'.
luxpy.toolboxes.indvcmf.cie2006cmfsEx(age=32, fieldsize=10, wl=None, var od lens=0,
                                                var_od_macula=0, var_od_L=0, var_od_M=0, var_od_S=0,
                                                var shft L=0, var shft M=0, var shft S=0, norm type=None,
                                                out='lms', base=False, strategy_2=True, odata0=None,
                                                lms_to_xyz_method=None, allow_negative_values=False,
                                                normalize_lms_to_xyz_matrix=False)
      Generate Individual Observer CMFs (cone fundamentals) based on CIE2006 cone fundamentals and published
      literature on observer variability in color matching and in physiological parameters.
      Args:
                  age
                        32 or float or int, optional
                        Observer age
                  fieldsize
                        Field size of stimulus in degrees (between 2^{\circ} and 10^{\circ}).
                  wl
                        None, optional
                        Interpolation/extraplation of :LMS: output to specified wavelengths.
                        None: output original _WL
                  var_od_lens
```

```
0, optional
      Std Dev. in peak optical density [%] of lens.
var_od_macula
      0, optional
      Std Dev. in peak optical density [%] of macula.
var_od_L
      0, optional
      Std Dev. in peak optical density [%] of L-cone.
var_od_M
      0, optional
      Std Dev. in peak optical density [%] of M-cone.
var_od_S
      0, optional
      Std Dev. in peak optical density [%] of S-cone.
var_shft_L
      0, optional
      Std Dev. in peak wavelength shift [nm] of L-cone.
var\_shft\_L
      0, optional
      Std Dev. in peak wavelength shift [nm] of M-cone.
var_shft_S
      0, optional
      Std Dev. in peak wavelength shift [nm] of S-cone.
norm_type
      None, optional
      - 'max': normalize LMSq functions to max = 1
      - 'area': normalize to area
      - 'power': normalize to power
out
      'lms' or 'xyz', optional
      Determines output.
base
      False, boolean, optional
      The returned energy-based LMS cone fundamentals given to the
      precision of 9 sign. figs. if 'True', and to the precision of
      6 sign. figs. if 'False'.
strategy_2
      True, bool, optional
      Use strategy 2 in github.com/ifarup/ciefunctions issue #121 for
      computing the weighting factor. If false, strategy 3 is applied.
odata0
      None, optional
      Dict with uncorrected ocular media and macula density functions and LMS
      absorptance functions
```

None defaults to the ones stored in \_DATA

### lms\_to\_xyz\_method

None, optional

Method to use to determine lms-to-xyz conversion matrix (options: 'asano', 'cietc197')

### allow\_negative\_values

False, optional

Cone fundamentals or color matching functions should not have negative values.

If False: X[X<0] = 0.

## normalize\_lms\_to\_xyz\_matrix

False, optional

Normalize that EEW is always at [100,100,100] in XYZ and LMS system.

### **Returns:**

#### returns

- 'LMS' [or 'XYZ']: ndarray with individual observer equal area-normalized cone fundamentals. Wavelength have been added.

[- 'M': lms to xyz conversion matrix

- 'trans\_lens': ndarray with lens transmission (no interpolation)
- 'trans\_macula': ndarray with macula transmission (no interpolation)
- 'sens\_photopig' : ndarray with photopigment sens.

(no interpolation)]

**References:** 1. Asano Y, Fairchild MD, and Blondé L, (2016), Individual Colorimetric Observer Model. PLoS One 11, 1–19.

- 2. Asano Y, Fairchild MD, Blondé L, and Morvan P (2016). Color matching experiment for highlighting interobserver variability. Color Res. Appl. 41, 530–539.
- 3. CIE TC1-36, (2006), Fundamental Chromaticity Diagram with Physiological Axes Part I (Vienna: CIE).
- 4. Asano's Individual Colorimetric Observer Model
- **5.** CIE TC1-97 Python code for cone fundamentals and XYZ cmf calculations (by Ivar Farup and Jan Henrik Wold, (c) 2012-2017)

```
luxpy.toolboxes.indvcmf.getMonteCarloParam(n_obs=1, stdDevAllParam={'dsrc': 'matlab', 'od_L': 17.9, 'od_M': 17.9, 'od_S': 14.7, 'od_lens': 19.1, 'od_macula': 37.2, 'shft_L': 4.0, 'shft_M': 3.0, 'shft_S': 2.5})
```

Get dict with normally-distributed physiological factors for a population of observers. **Args:** 

## n\_obs

1, optional

Number of individual observers in population.

### stdDevAllParam

### **Returns:**

### returns

dict with n\_obs randomly drawn parameters.

 $\label{loss} \verb|luxpy.toolboxes.indvcmf.genMonteCarloObs|| ($n\_obs=1$, fieldsize=10$, list\_Age=[32]$, $wl=None$, $norm\_type=None$, out='lms'$, base=False$, strategy\_2=True$, $norm\_type=None$, out='lms'$, base=False$, $norm\_type=None$, $norm\_type=None$$ 

odata0=None, lms\_to\_xyz\_method=None, allow\_negative\_values=False)

Monte-Carlo generation of individual observer cone fundamentals.

## Args:

## n\_obs

1, optional

Number of observer CMFs to generate.

## list\_Age

list of observer ages or str, optional

Defaults to 32 (cfr. CIE2006 CMFs)

If 'us\_census': use US population census of 2010

to generate list\_Age.

# fieldsize

fieldsize in degrees (between 2° and 10°), optional

Defaults to 10°.

### wl

None, optional

Interpolation/extraplation of :LMS: output to specified wavelengths.

None: output original \_WL

## norm\_type

None, optional

- 'max': normalize LMSq functions to max = 1
- 'area': normalize to area
- 'power': normalize to power

### out

'lms' or 'xyz', optional

Determines output.

## base

False, boolean, optional

The returned energy-based LMS cone fundamentals given to the precision of 9 sign. figs. if 'True', and to the precision of 6 sign. figs. if 'False'.

## strategy\_2

True, bool, optional

Use strategy 2 in github.com/ifarup/ciefunctions issue #121 for computing the weighting factor. If false, strategy 3 is applied.

### odata0

None, optional

Dict with uncorrected ocular media and macula density functions and LMS absorptance functions

None defaults to the ones stored in \_DATA

```
lms_to_xyz_method
                        None, optional
                        Method to use to determine lms-to-xyz conversion matrix (options: 'asano',
                        'cietc197')
                  allow_negative_values
                        False, optional
                        Cone fundamentals or color matching functions should not have negative values.
                             If False: X[X<0] = 0.
     Returns:
                  returns
                        LMS [,var age, vAll]
                             - LMS: ndarray with population LMS functions.
                             - var_age: ndarray with population observer ages.
                             - vAll: dict with population physiological factors (see .keys())
     References: 1. Asano Y., Fairchild M.D., and Blondé L., (2016), Individual Colorimetric Observer Model.
           PLoS One 11, 1-19.
           2. Asano Y, Fairchild MD, Blondé L, and Morvan P (2016). Color matching experiment for highlighting
           interobserver variability. Color Res. Appl. 41, 530–539.
           3. CIE TC1-36, (2006), Fundamental Chromaticity Diagram with Physiological Axes - Part I. (Vienna:
           CIE).
           4. Asano's Individual Colorimetric Observer Model
luxpy.toolboxes.indvcmf.getCatObs(n cat=10, fieldsize=2, wl=None, norm type=None, out='lms',
                                          base=False, strategy_2=True, odata0=None,
                                          lms to xyz method=None, allow negative values=False)
     Generate cone fundamentals for categorical observers.
     Args:
                  n cat
                        10, optional
                        Number of observer CMFs to generate.
                  fieldsize
                        fieldsize in degrees (between 2° and 10°), optional
                        Defaults to 10°.
                  out
                        'LMS' or str, optional
                        Determines output.
                  wl
                        None, optional
                        Interpolation/extraplation of :LMS: output to specified wavelengths.
                             None: output original _WL
                 norm_type
                        None, optional
                        - 'max': normalize LMSq functions to max = 1
                        - 'area': normalize to area
                        - 'power': normalize to power
                  out
```

'lms' or 'xyz', optional Determines output.

### base

False, boolean, optional

The returned energy-based LMS cone fundamentals given to the precision of 9 sign. figs. if 'True', and to the precision of 6 sign. figs. if 'False'.

## strategy\_2

True, bool, optional

Use strategy 2 in github.com/ifarup/ciefunctions issue #121 for computing the weighting factor. If false, strategy 3 is applied.

#### odata0

None, optional

Dict with uncorrected ocular media and macula density functions and LMS absorptance functions

None defaults to the ones stored in \_DATA

# lms\_to\_xyz\_method

None, optional

Method to use to determine lms-to-xyz conversion matrix (options: 'asano', 'cietc197')

## allow\_negative\_values

False, optional

Cone fundamentals or color matching functions should not have negative values.

If False: X[X<0] = 0.

### **Returns:**

### returns

LMS [,var\_age, vAll]

- LMS: ndarray with population LMS functions.
- var\_age: ndarray with population observer ages.
- vAll: dict with population physiological factors (see .keys())

Notes: 1. Categorical observers are observer functions that would represent color-normal populations. They are finite and discrete as opposed to observer functions generated from the individual colorimetric observer model. Thus, they would offer more convenient and practical approaches for the personalized color imaging workflow and color matching analyses. Categorical observers were derived in two steps. At the first step, 10000 observer functions were generated from the individual colorimetric observer model using Monte Carlo simulation. At the second step, the cluster analysis, a modified k-medoids algorithm, was applied to the 10000 observers minimizing the squared Euclidean distance in cone fundamentals space, and categorical observers were derived iteratively. Since the proposed categorical observers are defined by their physiological parameters and ages, their CMFs can be derived for any target field size. 2. Categorical observers were ordered by the importance; the first categorical observer vas the average observer equivalent to CIEPO06 with 38 year-old for a given field size, followed by the second most important categorical observer, the third, and so on.

3. see: https://www.rit.edu/cos/colorscience/re AsanoObserverFunctions.php

```
luxpy.toolboxes.indvcmf.compute_cmfs(fieldsize=10, age=32, wl=None, var_od_lens=0, var_od_macula=0,
                                               var_shft_LMS=[0, 0, 0], var_od_LMS=[0, 0, 0], norm_type=None,
                                               out='lms', base=False, strategy 2=True, odata0=None,
                                               lms_to_xyz_method=None, allow_negative_values=False,
                                               normalize_lms_to_xyz_matrix=False)
      Generate Individual Observer CMFs (cone fundamentals) based on CIE2006 cone fundamentals and published
      literature on observer variability in color matching and in physiological parameters.
      Args:
                  age
                        32 or float or int, optional
                        Observer age
                  fieldsize
                        10, optional
                        Field size of stimulus in degrees (between 2° and 10°).
                  wl
                        None, optional
                        Interpolation/extraplation of :LMS: output to specified wavelengths.
                        None: output original _WL
                  var_od_lens
                        0, optional
                        Variation of optical density of lens.
                  var_od_macula
                        0, optional
                        Variation of optical density of macula.
                  var_shft_LMS
                        [0, 0, 0] optional
                        Variation (shift) of LMS peak absorptance.
                  var_od_LMS
                        [0, 0, 0] optional
                        Variation of LMS optical densities.
                  norm_type
                        None, optional
                        - 'max': normalize LMSq functions to max = 1
                        - 'area': normalize to area
                        - 'power': normalize to power
                  out
                        'lms' or 'xyz', optional
                        Determines output.
                  base
                        False, boolean, optional
                        The returned energy-based LMS cone fundamentals given to the
                        precision of 9 sign. figs. if 'True', and to the precision of
                        6 sign. figs. if 'False'.
                  strategy_2
                        True, bool, optional
                        Use strategy 2 in github.com/ifarup/ciefunctions issue #121 for
```

```
computing the weighting factor. If false, strategy 3 is applied.
```

### odata0

None, optional

Dict with uncorrected ocular media and macula density functions and LMS absorptance functions

None defaults to the ones stored in \_DATA

# lms\_to\_xyz\_method

None, optional

Method to use to determine lms-to-xyz conversion matrix (options: 'asano', 'cietc197')

## allow\_negative\_values

False, optional

Cone fundamentals or color matching functions should not have negative values.

If False: X[X<0] = 0.

# $normalize\_lms\_to\_xyz\_matrix$

False, optional

Normalize that EEW is always at [100,100,100] in XYZ and LMS system.

### **Returns:**

### returns

- 'LMS' [or 'XYZ']: ndarray with individual observer equal area-normalized cone fundamentals. Wavelength have been added.

[- 'M': lms to xyz conversion matrix

- 'trans\_lens': ndarray with lens transmission

(no interpolation)

- 'trans\_macula': ndarray with macula transmission

(no interpolation)

- 'sens\_photopig' : ndarray with photopigment sens.

(no interpolation)]

**References:** 1. Asano Y, Fairchild MD, and Blondé L, (2016), Individual Colorimetric Observer Model. PLoS One 11, 1–19.

- 2. Asano Y, Fairchild MD, Blondé L, and Morvan P (2016). Color matching experiment for highlighting interobserver variability. Color Res. Appl. 41, 530–539.
- 3. CIE, TC1-36, (2006). Fundamental Chromaticity Diagram with Physiological Axes Part I (Vienna: CIE).
- 4. Asano's Individual Colorimetric Observer Model
- 5. CIE TC1-97 Python code for cone fundamentals and XYZ cmf calculations (by Ivar Farup and Jan Henrik Wold, (c) 2012-2017)

luxpy.toolboxes.indvcmf.add\_to\_cmf\_dict(bar=None, cieobs='indv', K=683, M=array([[1.0, 0.0, 0.0], [0.0, 1.0, 0.0], [0.0, 0.0, 1.0]]))

Add set of cmfs to \_CMF dict.

### Args:

## bar

None, optional

Set of CMFs. None: initializes to empty ndarray.

cieobs

```
'indv' or str, optional
Name of CMF set.

K

683 (lm/W), optional
Conversion factor from radiometric to photometric quantity.

M

np.eye, optional
Matrix for lms to xyz conversion.

luxpy.toolboxes.indvcmf.plot_cmfs(cmf, axh=None, **kwargs)
Plot cmf set.
```

# 4.5.3 spdbuild/

рy

- \_\_init\_\_.py
- spdbuilder.py
- · spdbuilder2020.py
- · spdoptimzer2020.py

namespace luxpy.spdbuild/

# Module for building and optimizing SPDs

spdbuilder.py

# **Functions**

gaussian\_spd() Generate Gaussian spectrum.

butterworth\_spd() Generate Butterworth based spectrum.

**lorentzian2\_spd()** Generate 2nd order Lorentzian based spectrum.

roundedtriangle\_spd() Generate a rounded triangle based spectrum.

**mono\_led\_spd()** Generate monochromatic LED spectrum based on a Gaussian or butterworth profile or according to Ohno (Opt. Eng. 2005).

spd\_builder() Build spectrum based on Gaussians, monochromatic and/or phophor LED spectra.

**color3mixer()** Calculate fluxes required to obtain a target chromaticity when (additively) mixing 3 light sources.

**colormixer**() Calculate fluxes required to obtain a target chromaticity when (additively) mixing N light sources.

**colormixer\_pinv()** Additive color mixer of N primaries using using Moore-Penrose pseudo-inverse matrix.

spd\_builder() Build spectrum based on Gaussians, monochromatic and/or phophor LEDtype spectra.

- **get\_w\_summed\_spd()** Calculate weighted sum of spds.
- **fitnessfcn()** Fitness function that calculates closeness of solution x to target values for specified objective functions.
- spd\_constructor\_2() Construct spd from spectral model parameters using pairs of intermediate sources.
- spd\_constructor\_3() Construct spd from spectral model parameters using trio's of intermediate sources.
- spd\_optimizer\_2\_3() Optimizes the weights (fluxes) of a set of component spectra by combining pairs (2) or trio's (3) of components to intermediate sources until only 3 remain. Color3mixer can then be called to calculate required fluxes to obtain target chromaticity and fluxes are then back-calculated.
- get\_optim\_pars\_dict() Setup dict with optimization parameters.
- initialize\_spd\_model\_pars() Initialize spd\_model\_pars (for spd\_constructor) based on type
   of component\_data.
- **initialize\_spd\_optim\_pars**() Initialize spd\_optim\_pars (x0, lb, ub for use with math.minimizebnd) based on type of component\_data.
- **spd\_optimizer**() Generate a spectrum with specified white point and optimized for certain objective functions from a set of component spectra or component spectrum model parameters.

## Module for building and optimizing SPDs (2)

This module implements a class based spectral optimizer. It differs from the spdoptimizer function in spdbuild.py, in that it can use several different minimization algorithms, as well as a user defined method. It is also written such that the user can easily write his own primary constructor function. It supports the '3mixer' algorithm (but no '2mixer') and a 'no-mixer' algorithm (chromaticity as part of the list of objectives) for calculating the mixing contributions of the primaries.

### **Functions**

gaussian\_prim\_constructor() constructs a gaussian based primary set.

- \_setup\_wlr() Initialize the wavelength range for use with PrimConstructor.
- **\_extract\_prim\_optimization\_parameters**() Extract the primary parameters from the optimization vector x and the pdefs dict for use with PrimConstructor.
- \_stack\_wlr\_spd() Stack the wavelength range 'on top' of the spd values for use with Prim-Constructor.

**PrimConstructor** class for primary (spectral) construction

**Minimizer** class for minimization of fitness of each of the objective functions

**ObjFcns** class to specify one or more objective functions for minimization

SpectralOptimizer class for spectral optimization (initialization and run)

**spd\_optimizer2**() Generate a spectrum with specified white point and optimized for certain objective functions from a set of component spectra or component spectrum model parameters (functional wrapper around SpectralOptimizer class).

## **Notes**

```
1. See examples below (in spdoptimizer2020.'__main__') for use.
luxpy.toolboxes.spdbuild.gaussian_spd(peakwl=530, fwhm=20, wl=[360.0, 830.0, 1.0], with_wl=True)
     Generate Gaussian spectrum.
     Args:
                 peakw
                       int or float or list or ndarray, optional
                       Peak wavelength
                 fwhm
                       int or float or list or ndarray, optional
                       Full-Width-Half-Maximum of gaussian.
                 wl
                       _WL3, optional
                       Wavelength range.
                 with_wl
                       True, optional
                       True outputs a ndarray with first row wavelengths.
     Returns:
                 returns
                       ndarray with spectra.
     Note:
           Gaussian:
                 g = \exp(-0.5*((wl - peakwl)/sig)**2)
           with sig = fwhm/(2*(2*np.log(2))**0.5)
luxpy.toolboxes.spdbuild.butterworth_spd(peakwl=530, fwhm=20, bw\_order=1, wl=[360.0, 830.0, 1.0],
                                                   with wl=True)
     Generate Butterworth based spectrum.
     Args:
                 peakw
                       int or float or list or ndarray, optional
                       Peak wavelength
                 fwhm
                       int or float or list or ndarray, optional
                       Full-Width-Half-Maximum of butterworth.
                 bw order
                       1, optional
                       Order of the butterworth function.
                 wl
                        _WL3, optional
                       Wavelength range.
                 with_wl
                       True, optional
                       True outputs a ndarray with first row wavelengths.
     Returns:
```

```
returns
                       ndarray with spectra.
     Note:
           Butterworth:
                 bw = 1 / (1 + ((2*(wl - peakwl)/fwhm)**2))
luxpy.toolboxes.spdbuild.lorentzian2_spd(peakwl=530, fwhm=20, wl=[360.0, 830.0, 1.0],
                                                   with_wl=True)
     Generate 2nd order Lorentzian spectrum.
     Args:
                 peakw
                       int or float or list or ndarray, optional
                       Peak wavelength
                 fwhm
                       int or float or list or ndarray, optional
                       Full-Width-Half-Maximum of lorentzian.
                 wl
                       _WL3, optional
                       Wavelength range.
                 with_wl
                       True, optional
                       True outputs a ndarray with first row wavelengths.
     Returns:
                 returns
                       ndarray with spectra.
     Note:
           Lorentzian (2nd order):
                 lz = (1 + ((n*(wl - peakwl)/fwhm)**2))**(-2)
                       with n = 2*(2**0.5-1)**0.5
luxpy.toolboxes.spdbuild.roundedtriangle_spd(peakwl=530,fwhm=100,rounding=0.5,wl=[360.0,
                                                        830.0, 1.0], with_wl=True, min_v=0.0, max_v=1.0,
                                                        fw=100, rw=100)
     Generate rounded triangle spectrum.
     Args:
                 peakw
                       int or float or list or ndarray, optional
                       Peak wavelength
                 fwhm
                       int or float or list or ndarray, optional
                       Full-Width-Half-Maximum of rounded triangle.
                 rounding
                       int or float or list or ndarray, optional
                       Amount of rounding of triangle corners (top, bottom-left, bottom-right)
                 wl
                       _WL3, optional
                       Wavelength range.
```

```
with_wl
                        True, optional
                        True outputs a ndarray with first row wavelengths.
                  min_v, max_v
                        0.0, 1.0, optional
                        Minimum and maximum of spd.
                  fw
                        100, optional
                        front width of triangle.
                        Only used when fwhm is set to None.
                  rw
                        100, optional
                        rear width of triangle.
                        Only used when fwhm is set to None.
      Returns:
                  returns
                        ndarray with spectra.
luxpy.toolboxes.spdbuild.mono_led_spd(peakwl=530, fwhm=20, wl=[360.0, 830.0, 1.0], with_wl=True,
                                                strength_shoulder=2, bw_order=- 1)
      Generate monochromatic LED spectrum based on a Gaussian or or Lorentzian or butterworth profile or according
      to Ohno (Opt. Eng. 2005).
      Args:
                  peakw
                        int or float or list or ndarray, optional
                        Peak wavelength
                  fwhm
                        int or float or list or ndarray, optional
                        Full-Width-Half-Maximum of gaussian used to simulate led.
                  wl
                        _WL3, optional
                              Wavelength range.
                  with_wl
                        True, optional
                        True outputs a ndarray with first row wavelengths.
                  strength_shoulder
                        2, optional
                        Determines the strength of the spectrum shoulders of the mono led.
                        A value of 0 reduces to a pure Gaussian model (if bw_order >= -1).
                  bw order
                        -1, optional
                        Order of Butterworth function.
                        If -1 or 0: spd profile is Ohno's gaussian based
                              (to obtain pure Gaussian: set strength_shoulder = 0).
                        If -2: spd profile is Lorentzian,
                        else (>0): Butterworth.
```

```
Returns:
                                        returns
                                                      ndarray with spectra.
             Note:
                          Gaussian:
                                         g = \exp(-0.5*((wl - peakwl)/sig)**2)
                          with sig = fwhm/(2*(2*np.log(2))**0.5)
                          Lorentzian (2nd order):
                                        lz = (1 + ((n*(wl - peakwl)/fwhm)**2))**(-2)
                                                      with n = 2*(2**0.5-1)**0.5
                           Butterworth:
                                        bw = 1 / (1 + ((2*(wl - peakwl)/fwhm)**2))
                          Ohno's model:
                                        ohno = (g + strength shoulder*g**5)/(1+strength shoulder)
                                        mono\_led\_spd = ohno*((bw\_order >= -1) & (bw\_order <= 0)).T + bw*(bw\_order > 0).T + bw*
                                        1z*((bw order >=-2) & (bw order < -1)).T
             Reference: 1. Ohno Y (2005). Spectral design considerations for white LED color rendering. Opt. Eng. 44,
                           111302.
luxpy.toolboxes.spdbuild.phosphor_led_spd(peakwl=450, fwhm=20, wl=[360.0, 830.0, 1.0], bw_order=-
                                                                                                                        1, with_wl=True, strength_shoulder=2, strength_ph=0,
                                                                                                                       peakwl ph1=530, fwhm ph1=80, strength ph1=1,
                                                                                                                       peakwl_ph2=560, fwhm_ph2=80, strength_ph2=None,
                                                                                                                       use piecewise fcn=False, verbosity=0, out='spd')
             Generate phosphor LED spectrum with up to 2 phosphors based on Smet (Opt. Expr. 2011).
             Model:
                           1) If strength_ph2 is not None:
                                                      phosphor_spd = (strength_ph1*mono_led_spd(peakwl_ph1, ..., strength_shoulder = 1)
                                                                   + strength_ph2)*mono_led_spd(peakwl_ph2, ..., strength_shoulder = 1))
                                                                                 /(strength ph1 + strength ph2)
                                        else:
                                                      phosphor_spd = (strength_ph1*mono_led_spd(peakwl_ph1, ..., strength_shoulder = 1)
                                                                   + (1-strength_ph1)*mono_led_spd(peakwl_ph2, ..., strength_shoulder = 1))
                          2) S = (mono\_led\_spd() + strength\_ph*(phosphor\_spd/phosphor\_spd.max()))/(1 + strength\_ph)
                          3) piecewise_fcn = S for wl < peakwl and 1 for wl >= peakwl
                          4) phosphor_led_spd = S*piecewise_fcn
             Args:
                                        peakw
```

```
int or float or list or ndarray, optional
      Peak wavelengths of the monochromatic led.
fwhm
      int or float or list or ndarray, optional
      Full-Width-Half-Maximum of mono_led spectrum.
wl
      _WL3, optional
      Wavelength range.
bw_order
      -1, optional
      Order of Butterworth function.
      If -1 or 0: spd profile is Ohno's gaussian based
            (to obtain pure Gaussian: set strength_shoulder = 0).
      If -2: spd profile is Lorentzian,
      else (>0): Butterworth.
      Note that this only applies to the monochromatic led spds and not
      the phosphors spds (these are always gaussian based).
with_wl
      True, optional
      True outputs a ndarray with first row wavelengths.
strength_shoulder
      2, optiona l
      Determines the strength of the spectrum shoulders of the mono led.
strength_ph
      0, optional
      Total contribution of phosphors in mixture.
peakwl_ph1
      int or float or list or ndarray, optional
      Peak wavelength of the first phosphor.
fwhm_ph1
      int or float or list or ndarray, optional
      Full-Width-Half-Maximum of gaussian used to simulate first phosphor.
strength_ph1
      1, optional
      Strength of first phosphor in phosphor mixture.
      If :strength_ph2: is None: value should be in the [0,1] range.
peakwl_ph2
      int or float or list or ndarray, optional
      Peak wavelength of the second phosphor.
fwhm_ph2
```

Full-Width-Half-Maximum of gaussian used to simulate second phosphor.

int or float or list or ndarray, optional

Strength of second phosphor in phosphor mixture.

strength\_ph2

None, optional

```
If None: strength is calculated as (1-:strength_ph1:)
                                   :target: np2d([100,1/3,1/3]), optional
                             ndarray with Yxy chromaticity of target.
                  verbosity
                       0, optional
                       If > 0: plots spectrum components (mono led, ph1, ph2, ...)
                  out
                        'spd', optional
                       Specifies output.
                  use piecewise fcn
                       False, optional
                       True: uses piece-wise function as in Smet et al. 2011. Can give
                       non smooth spectra optimized from components to which it is applied.
     Returns:
                  returns
                       spd, component_spds
                       ndarrays with spectra (and component spds used to build the
                       final spectra)
     References: 1. Ohno Y (2005). Spectral design considerations for white LED color rendering. Opt. Eng. 44,
            111302.
           2. Smet K, Ryckaert WR, Pointer MR, Deconinck G, and Hanselaer P (2011). Optimal colour quality of
           LED clusters based on memory colours. Opt. Express 19, 6903–6912.
luxpy.toolboxes.spdbuild.spd_builder(flux=None, component spds=None, peakwl=450, fwhm=20,
                                              bw order=-1, pair strengths=None, wl=[360.0, 830.0, 1.0],
                                              with_wl=True, strength_shoulder=2, strength_ph=0,
                                              peakwl\_ph1=530, fwhm\_ph1=80, strength\_ph1=1,
                                              peakwl_ph2=560, fwhm_ph2=80, strength_ph2=None,
                                              target=None, tar_type='Yuv', cspace_bwtf={}, cieobs='1931_2',
                                              use_piecewise_fcn=False, verbosity=0, out='spd', **kwargs)
     Build spectrum based on Gaussian, monochromatic and/or phophor type spectra.
     Args:
                  flux
                       None, optional
                       Fluxes of each of the component spectra.
                       None outputs the individual component spectra.
                  component spds
                       None or ndarray, optional
                       If None: calculate component spds from input args.
                  peakw
                       int or float or list or ndarray, optional
                       Peak wavelengths of the monochromatic led.
                  fwhm
                       int or float or list or ndarray, optional (but must be same shape as peakw!)
                       Full-Width-Half-Maximum of gaussian.
                  wl
                        _WL3, optional
```

```
Wavelength range.
bw_order
      -1, optional
      Order of Butterworth function.
      If -1 or 0: spd profile is Ohno's gaussian based
            (to obtain pure Gaussian: set strength shoulder = 0).
      If -2: spd profile is Lorentzian,
      else (>0): Butterworth.
      Note that this only applies to the monochromatic led spds and not
      the phosphors spds (these are always gaussian based).
pair_strengths
      ndarray with pair_strengths of mono_led spds, optional
      If None: will be randomly selected, possibly resulting in
      unphysical (out-of-gamut) solution.
with wl
      True, optional
      True outputs a ndarray with first row wavelengths.
strength_shoulder
      2, optiona 1
      Determines the strength of the spectrum shoulders of the mono led.
strength_ph
      0, optional
      Total contribution of phosphors in mixture.
peakwl_ph1
      int or float or list or ndarray, optional
      Peak wavelength of the first phosphor.
fwhm_ph1
      int or float or list or ndarray, optional
      Full-Width-Half-Maximum of gaussian used to simulate first phosphor.
strength_ph1
      1, optional
      Strength of first phosphor in phosphor mixture.
      If :strength_ph2: is None: value should be in the [0,1] range.
peakwl_ph2
      int or float or list or ndarray, optional
      Peak wavelength of the second phosphor.
fwhm_ph2
      int or float or list or ndarray, optional
      Full-Width-Half-Maximum of gaussian used to simulate second phosphor.
strength_ph2
```

:target: np2d([100,1/3,1/3]), optional

Strength of second phosphor in phosphor mixture. If None: strength is calculated as (1-:strength\_ph1:)

ndarray with Yxy chromaticity of target.

None, optional

```
verbosity
                        0, optional
                        If > 0: plots spectrum components (mono_led, ph1, ph2, ...)
                  out
                        'spd', optional
                        Specifies output.
                  use_piecewise_fcn
                        False, optional
                        True: uses piece-wise function as in Smet et al. 2011. Can give
                        non smooth spectra optimized from components to which it is
                        applied.
                  target
                        None, optional
                        ndarray with Yxy chromaticity of target.
                        If None: don't override phosphor strengths, else calculate strength
                              to obtain :target: using color3mixer().
                        If not None AND strength_ph is None or 0: components are
                        monochromatic and colormixer is used to optimize fluxes to
                        obtain target chromaticity (N can be > 3 components)
                  tar_type
                        'Yxy' or str, optional
                        Specifies the input type in :target: (e.g. 'Yxy' or 'cct')
                  cieobs
                        _CIEOBS, optional
                        CIE CMF set used to calculate chromaticity values.
                  cspace_bwtf
                        {}, optional
                        Backward (..._to_xyz) transform parameters
                        (see colortf()) to go from :tar_type: to 'Yxy')
      Returns:
                  returns
                        ndarray with spectra.
      Note: 1. Target-optimization is only for phophor leds with three components (blue pump, ph1 and ph2) span-
            ning a sufficiently large gamut.
      References: 1. Ohno Y (2005). Spectral design considerations for white LED color rendering. Opt. Eng. 44,
            111302.
            2. Smet K, Ryckaert WR, Pointer MR, Deconinck G, and Hanselaer P (2011). Optimal colour quality of
           LED clusters based on memory colours. Opt. Express 19, 6903–6912.
luxpy.toolboxes.spdbuild.get_w_summed_spd(w, spds)
      Calculate weighted sum of spds.
      Args:
                  W
                        ndarray with weigths (e.g. fluxes)
                  spds
                        ndarray with component spds.
      Returns:
```

### returns

ndarray with weighted sum.

 $\label{limits} \begin{tabular}{ll} luxpy.toolboxes.spdbuild. {\bf fitnessfcn}(x, spd\_constructor, spd\_constructor\_pars=None, F\_rss=True, \\ decimals=[3], obj\_fcn=[None], obj\_fcn\_pars=[\{\}], \\ \end{tabular}$ 

obj\_fcn\_weights=[1], obj\_tar\_vals=[0], verbosity=0, out='F')

Fitness function that calculates closeness of solution x to target values for specified objective functions. **Args:** 

X

ndarray with parameter values

# spd\_constructor

function handle to a function that constructs the spd from parameter values in :x:.

## spd\_constructor\_pars

None, optional,

Parameters required by :spd\_constructor:

### F rss

True, optional

Take Root-Sum-of-Squares of 'closeness' values between target and objective function values.

## decimals

3, optional

List of rounding decimals of objective function values.

## obj\_fcn

[None] or list, optional

List of function handles to objective function.

# obj\_fcn\_weights

[1] or list, optional.

List of weigths for each obj. fcn

## obj\_fcn\_pars

[None] or list, optional

List of parameter dicts for each obj. fcn.

## obj\_tar\_vals

[0] or list, optional

List of target values for each objective function.

# verbosity

0, optional

If > 0: print intermediate results.

### out

'F', optional

Determines output.

## **Returns:**

F

float or ndarray with fitness value for current solution :x:.

luxpy.toolboxes.spdbuild.spd\_constructor\_2(x, constructor\_pars={}, \*\*kwargs)

Construct spd from model parameters using pairs of intermediate sources.

```
Pairs (odd, even) of components are selected and combined using
            'pair_strength'. This process is continued until only 3 intermediate
            (combined) sources remain. Color3mixer is then used to calculate the
            fluxes for the remaining 3 sources, after which the fluxes of all
            components are back-calculated.
      Args:
                  X
                        vector of optimization parameters.
                  constructor pars
                        dict with model parameters.
                        Key 'list' determines which parameters are in :x: and key 'len'
                        (Specifies the number of variables representing each parameter).
      Returns:
                  returns
                        spd, M, spds
                        ndarrays with spectrum corresponding to x, M the fluxes of
                        the spectral components of spd and spds the spectral components
                        themselves.
luxpy.toolboxes.spdbuild.color3mixer(Yxyt, Yxy1, Yxy2, Yxy3)
      Calculate fluxes required to obtain a target chromaticity when (additively) mixing 3 light sources.
      Args:
                  Yxvt
                        ndarray with target Yxy chromaticities.
                  Yxy1
                        ndarray with Yxy chromaticities of light sources 1.
                  Yxy2
                        ndarray with Yxy chromaticities of light sources 2.
                  Yxy3
                        ndarray with Yxy chromaticities of light sources 3.
      Returns:
                  M
                        ndarray with fluxes.
      Note: Yxyt, Yxy1, ... can contain multiple rows, referring to single mixture.
luxpy.toolboxes.spdbuild.colormixer(Yxyt=None, Yxyi=None, n=4, pair_strengths=None,
                                              source_order=None)
      Calculate fluxes required to obtain a target chromaticity when (additively) mixing N light sources.
      Args:
                  Yxyt
                        ndarray with target Yxy chromaticities.
                        Defaults to equi-energy white.
                  Yxyi
                        ndarray with Yxy chromaticities of light sources i = 1 to n.
                  n
                        4 or int, optional
```

Number of source components to randomly generate when Yxyi is None.

### pair strengths

ndarray with light source pair strengths.

## source\_order

ndarray with order of source components.

If None: use np.arange(n)

### **Returns:**

M

ndarray with fluxes.

### Note:

## Algorithm

- Loop over all source components and create intermediate sources from all (even,odd)-pairs using the relative strengths of the pair (specified in pair\_strengths).
- 2. Collect any remaining sources.
- 3. Combine with new intermediate source components
- 4. Repeat 1-3 until there are only 3 source components left.
- 5. Use color3mixer to calculate the required fluxes of the 3 final intermediate components to obtain the target chromaticity.
- 6. Backward calculate the fluxes of all original source components from the 3 final intermediate fluxes.

luxpy.toolboxes.spdbuild.colormixer\_pinv(xyzt, xyzi, input fmt='xyz')

Additive color mixer of N primaries using using Moore-Penrose pseudo-inverse matrix.

## Args:

xyzt

ndarray with target XYZ tristimulus values or Yxy chromaticity coordinates.

xvzi

ndarray with XYZ tristimulus values or Yxy chromaticity coordinates of light sources i = 1 to n.

### input\_fmt

'xyz', optional

Format specifier of :xyzt: and :xyzi: input arguments.

- options: 'xyz', 'Yxy'

## **Returns:**

w

ndarray with fluxes (weights) of each of the primaries in the mixture.

luxpy.toolboxes.spdbuild.spd\_constructor\_3(x, constructor\_pars={}, \*\*kwargs)

Construct spd from model parameters using trio's of intermediate sources.

The triangle/trio method creates for all possible combinations of 3 primary component spectra a spectrum that results in the target chromaticity using color3mixer() and then optimizes the weights of each of the latter spectra such that adding them (additive mixing) results in obj\_vals as close as possible to the target values.

### Args:

 $\mathbf{X}$ 

vector of optimization parameters.

## constructor\_pars

dict with model parameters.

Key 'list' determines which parameters are in :x: and key 'len' (specifies the number of variables representing each parameter).

### **Returns:**

### returns

spd, M, spds ndarrays with spectrum corresponding to x, M the fluxes of the spectral components of spd and spds the spectral components themselves.

luxpy.toolboxes.spdbuild.spd\_optimizer\_2\_3(optimizer\_type='2mixer', spd\_constructor=None, spd\_model\_pars=None, component\_data=4, N\_components=None, wl=[360.0, 830.0, 1.0], allow\_nongaussianbased\_mono\_spds=False, Yxy\_target=array([[100.0, 0.33333, 0.33333]]), cieobs='1931\_2', obj\_fcn=[None], obj\_fcn\_pars=[{}], obj\_fcn\_weights=[1], obj\_tar\_vals=[0], decimals=[5], minimize\_method='Nelder-Mead', minimize\_opts=None, F\_rss=True, verbosity=0, \*\*kwargs)

Optimizes the weights (fluxes) of a set of component spectra by combining pairs (2) or trio's (3) of components to intermediate sources until only 3 remain. Color3mixer can then be called to calculate required fluxes to obtain target chromaticity and fluxes are then back-calculated.

# Args:

## optimizer\_type

'2mixer' or '3mixer' or 'user', optional Specifies whether to optimize spectral model parameters by combining pairs or trio's of comonponents.

# $spd\_constructor$

None, optional

Function handle to user defined spd\_constructor function.

Input: fcn(x, constructor\_pars = { }, kwargs)
Output: spd,M,spds

nd array with:

spd: spectrum resulting from xM: fluxes of all component spds

- spds: component spds (in [N+1,wl] format)

(See e.g. spd\_constructor\_2 or spd\_constructor\_3)

# $spd\_model\_pars$

dict with model parameters required by spd\_constructor and with optimization parameters required by minimize (x0, lb, ub).

```
Only used when :optimizer_type: == 'user'.
component_data
      4, optional
      Component spectra data:
      If int: specifies number of components used in optimization
            (peakwl, fwhm and pair strengths will be optimized).
      If dict: generate components based on parameters (peakwl, fwhm,
                  pair_strengths, etc.) in dict.
            (keys with None values will be optimized)
      If ndarray: optimize pair_strengths of component spectra.
N_components
      None, optional
      Specifies number of components used in optimization. (only used
      when :component_data: is dict and user wants to override dict.
      Note that shape of parameters arrays must match N_components).
allow_nongaussianbased_mono_spds
      False, optional
      False: use pure Gaussian based monochrom. spds.
wl
      _WL3, optional
      Wavelengths used in optimization when :component_data: is not
      ndarray with spectral data.
Yxy_target
      np2d([100,1/3,1/3]), optional
      ndarray with Yxy chromaticity of target.
cieobs
      _CIEOBS, optional
      CIE CMF set used to calculate chromaticity values if not provided
      in:Yxyi:.
F_rss
      True, optional
      Take Root-Sum-of-Squares of 'closeness' values between target and
      objective function values.
decimals
      5, optional
      Rounding decimals of objective function values.
obj_fcn
      [None] or list, optional
      Function handles to objective function.
obj_fcn_weights
      [1] or list, optional.
```

Weigths for each obj. fcn

[None] or list, optional

Parameter dicts for each obj. fcn.

obj\_fcn\_pars

obj\_tar\_vals

```
[0] or list, optional
                       Target values for each objective function.
                 minimize method
                       'Nelder-Mead', optional
                       Optimization method used by minimize function.
                 minimize_opts
                       None, optional
                       Dict with minimization options.
                       None defaults to: {'xtol': 1e-5, 'disp': True, 'maxiter': 1000*Nc,
                             'maxfev': 1000*Nc,'fatol': 0.01}
                 verbosity
                       0, optional
                       If > 0: print intermediate results.
     Returns:
                 returns
                       M, spd_opt, obj_vals
                             - 'M': ndarray with fluxes for each component spectrum.
                             - 'spd opt': optimized spectrum.
                             - 'obj_vals': values of the obj. fcns for the optimized spectrum.
luxpy.toolboxes.spdbuild.get_optim_pars_dict(target=array([[100.0, 0.33333, 0.33333]]),
                                                      tar_type='Yxy', cieobs='1931_2',
                                                      optimizer_type='2mixer', spd_constructor=None,
                                                      spd_model_pars=None, cspace='Yuv', cspace_bwtf={},
                                                      cspace_fwtf={}, component_spds=None,
                                                      N_{components}=None, obj_fcn=[None],
                                                      obj_fcn_pars=[{}], obj_fcn_weights=[1],
                                                       obj tar vals=[0], decimals=[5],
                                                      minimize_method='Nelder-Mead', minimize_opts=None,
                                                       F rss=True, peakwl=[450, 530, 610], fwhm=[20, 20, 20],
                                                      allow_nongaussianbased_mono_spds=False,
                                                      bw order=[-1], wl=[360.0, 830.0, 1.0], with wl=True,
                                                      strength\_shoulder=2, strength\_ph=[0],
                                                      use_piecewise_fcn=False, peakwl_ph1=[530],
                                                      fwhm_ph1=[80], strength_ph1=[1], peakwl_ph2=[560],
                                                      fwhm_ph2=[80], strength_ph2=None, verbosity=0,
                                                      pair_strengths=None, triangle_strengths=None,
                                                      peakwl_min=[400], peakwl_max=[700], fwhm_min=[5],
                                                      fwhm_max=[600], bw_order_min=[-2],
                                                       bw\_order\_max=[100])
     Setup dict with optimization parameters.
     Args: See ?spd_optimizer for more info.
     Returns:
                 opts
                       dict with keys and values of the function's keywords and values.
luxpy.toolboxes.spdbuild.initialize_spd_model_pars(component_data, N_components=None,
```

Initialize spd\_model\_pars dict (for spd\_constructor) based on type of component\_data.

allow\_nongaussianbased\_mono\_spds=False, optimizer\_type='2mixer', wl=[360.0, 830.0, 1.0])

### Args:

## component\_data

None, optional

Component spectra data:

If int: specifies number of components used in optimization

(peakwl, fwhm and pair strengths will be optimized).

If dict: generate components based on parameters (peakwl, fwhm,

pair strengths, etc.) in dict.

(keys with None values will be optimized)

If ndarray: optimize pair\_strengths of component spectra.

## N\_components

None, optional

Specifies number of components used in optimization. (only used

when :component\_data: is dict and user wants to override dict.

Note that shape of parameters arrays must match N\_components).

## allow nongaussianbased mono spds

False, optional

- False: use Gaussian based monochrom. spds.
- True: also allow butterworth and lorentzian type monochrom. spds while optimizing.

# optimizer\_type

'2mixer', optional

Type of spectral optimization routine.

(other options: '3mixer', 'search')

wl

WL3, optional

Wavelengths used in optimization when :component\_data: is not an ndarray with spectral data.

## **Returns:**

# spd\_model\_pars

dict with spectrum-model parameters

luxpy.toolboxes.spdbuild.initialize\_spd\_optim\_pars(component\_data, N\_components=None,

allow\_nongaussianbased\_mono\_spds=False, optimizer\_type='2mixer', wl=[360.0, 830.0, 1.0], spd\_model\_pars=None)

Initialize spd optim pars dict based on type of component data.

### Args:

### component data

None, optional

Component spectra data:

If int: specifies number of components used in optimization

(peakwl, fwhm and pair\_strengths will be optimized).

If dict: generate components based on parameters (peakwl, fwhm,

pair strengths, etc.) in dict.

(keys with None values will be optimized)

If ndarray: optimize pair\_strengths of component spectra.

```
N_components
                        None, optional
                        Specifies number of components used in optimization. (only used
                        when :component_data: is dict and user wants to override dict.
                        Note that shape of parameters arrays must match N_components).
                  allow\_nongaus sian based\_mono\_spds
                        False, optional
                        False: use Gaussian based monochrom. spds.
                  optimizer_type
                        '2mixer', optional
                        Type of spectral optimization routine.
                        (other options: '3mixer', 'search')
                  wl
                        _WL3, optional
                        Wavelengths used in optimization when :component data: is not an
                        ndarray with spectral data.
                  spd_model_pars
                        None, optional
                        If None, initialize based on type of component_data.
                        else: initialize on pre-defined spd model pars dict.
      Returns:
                  spd_optim_pars
                        dict with optimization parameters (x0, ub, lb)
luxpy.toolboxes.spdbuild.get_primary_fluxratios(res, primaries, Ytarget=1, ptype='pu',
                                                             cieobs='1931_2', out='M,Sopt')
      Get flux ratios of primaries.
      Args:
                  res
                        dict or ndarray with optimized fluxes for component spds normalized to \max = 1.
                        (output of spd_optimizer)
                  primaries
                        ndarray with primary spectra.
                  Ytarget
                        1, optional
                        M will be scaled to result in a photo-/radio-metric power of Ytarget
                  ptype
                        'pu' or 'ru', optional
                        Type of power:
                        -'pu': photometric units
                        -'ru': radiometric units
                  cieobs
                        _CIEOBS, optional
                        CMF set/Vlambda to use in calculation of power.
      Returns:
                  M
```

ndarray with flux ratios.

### Sopt

ndarray with optimized scaled spectrum.

```
luxpy.toolboxes.spdbuild.spd_optimizer(target=array([[100.0, 0.33333, 0.33333]]), tar_type='Yxy',
                                              cieobs='1931 2', optimizer type='2mixer',
                                              spd constructor=None, spd model pars=None, cspace='Yuv',
                                              cspace_bwtf={}, cspace_fwtf={}, component_spds=None,
                                              N_components=None, obj_fcn=[None], obj_fcn_pars=[{}],
                                              obj_fcn_weights=[1], obj_tar_vals=[0], decimals=[5],
                                              minimize_method='Nelder-Mead', minimize_opts=None,
                                              F_rss=True, peakwl=[450, 530, 610], fwhm=[20, 20, 20],
                                              allow_nongaussianbased_mono_spds=False, bw_order=[-1],
                                              wl = [360.0, 830.0, 1.0], with_wl = True, strength_shoulder = 2,
                                              strength_ph=[0], use_piecewise_fcn=False, peakwl_ph1=[530],
                                              fwhm_ph1=[80], strength_ph1=[1], peakwl_ph2=[560],
                                              fwhm_ph2=[80], strength_ph2=None, verbosity=0,
                                              pair strengths=None, peakwl min=[400], peakwl max=[700],
                                              fwhm_min=[5], fwhm_max=[600], bw_order_min=-2,
                                              bw order max=100, out='spds,M')
```

Generate a spectrum with specified white point and optimized for certain objective functions from a set of component spectra or component spectrum model parameters.

## Args:

```
target
      np2d([100,1/3,1/3]), optional
      ndarray with Yxy chromaticity of target.
tar_type
      'Yxy' or str, optional
      Specifies the input type in :target: (e.g. 'Yxy' or 'cct')
cieobs
      _CIEOBS, optional
      CIE CMF set used to calculate chromaticity values, if not provided
      in:Yxyi:.
optimizer_type
      '2mixer', optional
      Specifies type of chromaticity optimization
      ('3mixer' or '2mixer' or 'search')
      For help on '2mixer' and '3mixer' algorithms, see notes below.
spd_constructor
      None, optional
      Function handle to user defined spd_constructor function.
            Input: fcn(x, constructor_pars = {}, kwargs)
            Output: spd,M,spds
                  nd array with:
                        - spd: spectrum resulting from x
```

```
- M: fluxes of all component spds
                        - spds: component spds (in [N+1,wl] format)
      (See e.g. spd_constructor_2 or spd_constructor_3)
spd model pars
      dict with model parameters required by spd_constructor
      and with optimization parameters required by minimize (x0, lb, ub). .
      Only used when :optimizer_type: == 'user'.
cspace
      'Yuv', optional
      Color space for 'search'-type optimization.
cspace_bwtf
      {}, optional
      Backward (cspace_to_xyz) transform parameters
      (see colortf()) to go from :tar_type: to 'Yxy').
cspace_fwtf
      {}, optional
      Forward (xyz_to_cspace) transform parameters
      (see colortf()) to go from xyz to :cspace:).
component_spds
      ndarray of component spectra.
      If None: they are built from input args.
N_components
      None, optional
      Specifies number of components used in optimization. (only used
      when :component_data: is dict and user wants to override dict value
      Note that shape of parameters arrays must match N_components).
allow_nongaussianbased_mono_spds
      False, optional
      False: use Ohno monochromatic led spectra based on Gaussian spds.
      True: also use Butterworth and Lorentzian spds.
wl
      WL3, optional
      Wavelengths used in optimization when :component_data: is not an
      ndarray with spectral data.
F_rss
      True, optional
      Take Root-Sum-of-Squares of 'closeness' values between target and
      objective function values.
decimals
      5, optional
      Rounding decimals of objective function values.
obj_fcn
      [None] or list, optional
      Function handles to objective function.
obj_fcn_weights
```

```
[1] or list, optional.
                        Weigths for each obj. fcn
                  obj_fcn_pars
                        [None] or list, optional
                        Parameter dicts for each obj. fcn.
                  obj_tar_vals
                        [0] or list, optional
                        Target values for each objective function.
                  minimize method
                        'Nelder-Mead', optional
                        Optimization method used by minimize function.
                  minimize_opts
                        None, optional
                        Dict with minimization options.
                              None defaults to: {'xtol': 1e-5, 'disp': True, 'maxiter': 1000*Nc,
                                     'maxfev': 1000*Nc,'fatol': 0.01}
                  verbosity
                        0, optional
                        If > 0: print intermediate results.
                  out
                        'spds,M', optional
                        Determines output of function.
      Note: peakwl:, :fwhm:, . . . : see ?spd_builder for more info.
      Returns:
                  returns
                        spds, M
                              - 'spds': optimized spectrum.
                               - 'M': ndarray with fluxes for each component spectrum.
      Notes:
                  Optimization algorithms
            1. '2mixer': Pairs (odd, even) of components are selected and combined using 'pair_strength'. This process
            is continued until only 3 (combined) intermediate sources remain. Color3mixer is then used to calculate
            the fluxes for the remaining 3 sources, after which the fluxes of all components are back-calculated.
            2. '3mixer': The triangle/trio method creates for all possible combinations of 3 primary component spectra
            a spectrum that results in the target chromaticity using color3mixer() and then optimizes the weights of
            each of the latter spectra such that adding them (additive mixing) results in obj_vals as close as possible
            to the target values.
class luxpy.toolboxes.spdbuild.PrimConstructor(f=<function gaussian_prim_constructor>,
                                                            ptypes=['peakwl', 'fwhm'], pdefs={})
      get_spd(nprim=None, wlr=[360, 830, 1])
            Get ndarray with spds for prims.
            Args:
                        nprim
                              None, optional
                              If not None: generate nprim random prims (based fixed pars and bounds in
                              pdefs)
```

else: values for all pars should be defined in pdefs!

```
(nprims is determined by number of elements in pdefs[ptypes[0]])
class luxpy.toolboxes.spdbuild.Minimizer(method='Nelder-Mead', opts={|}, x0=None, pareto=False,
                                                  display=True)
     _set_defopts_and_pareto(pareto=None, x0=None, display=None)
           Set default options if not provided, as well as pareto (False: output Root-Sum-Squares of Fi in _fitnessfcn).
     apply(fitness fcn, npars, fitness args dict, bounds, verbosity=1)
           Run minimizer on fitness function with specified fitness_args_dict input arguments and bounds.
class luxpy.toolboxes.spdbuild.0bjFcns(f=None, fp=[{}], fw=[1], ft=[0], ft_tol=[0],
                                               f_requires_solution_info=[False], decimals=[5])
     _equalize_sizes(x)
           Equalize structure of x to that of self.f for ease of looping of the objective functions in the fitness function
     _calculate_fj(spdi, j=0, solution_info={})
           Calculate objective function j for input spd.
     _get_normalization_factors()
           Set normalization factor for F-calculation
     _get_fj_output_str(j, obj_vals_ij, F_ij=nan, verbosity=1)
           get output string for objective function fj
class luxpy.toolboxes.spdbuild.SpectralOptimizer(target=array([[1.0000e+02, 3.3333e-01,
                                                            3.3338e-01]]), tar_type='Yxy', cspace_bwtf={},
                                                            nprim=4, wlr=[360, 830, 1], cieobs='1931 2',
                                                            out='spds,primss,Ms,results',
                                                            optimizer type='3mixer',
                                                            triangle_strengths_bnds=None,
                                                            prim_constructor=<luxpy.toolboxes.spdbuild.spdoptimizer2020.PrimC
                                                            object>, prims=None,
                                                            obj_fcn=<luxpy.toolboxes.spdbuild.spdoptimizer2020.ObjFcns
                                                            object>, mini-
                                                            mizer=<luxpy.toolboxes.spdbuild.spdoptimizer2020.Minimizer
                                                            object>, verbosity=1)
     _update_nprim_prims(nprim=None, prims=None)
           Update prims (and nprim).
     _update_target(target=None, tar_type=None, cspace_bwtf=None)
           Update target chromaticity.
     _update_prim_pars_bnds(nprim=None, **kwargs)
           Get and set fixed and free parameters, as well as bnds on latter for an nprim primary mixture.
     _update_triangle_strengths_bnds(nprim=None, triangle_strengths_bnds=None)
           Update bounds of triangle_strengths for for an nprim primary mixture.
     _update_bnds(nprim=None, triangle_strengths_bnds=None, **prim_kwargs)
           Update all bounds (triangle_strengths and those of free parameters of primary constructor) for an nprim
           primary mixture..
     update(nprim=None, prims=None, cieobs=None, target=None, tar_type=None, cspace_bwtf=None,
              triangle_strengths_bnds=None, **prim_kwargs)
           Updates all that is needed when one of the input arguments is changed.
     _spd_constructor_tri(x)
           Construct a mixture spectrum composed of n primaries using the 3mixer algorithm.
           Args:
                       \mathbf{X}
```

optimization parameters, first n!/(n-3)!\*3! are the strengths of the triangles in the '3mixer' algorithm.

#### **Returns:**

# spd, prims, M

- spd: spectrum resulting from x
- spds: primary spds
- M: fluxes of all primaries

**Notes:** 1. '3mixer' - optimization algorithm: The triangle/trio method creates for all possible combinations of 3 primary component spectra a spectrum that results in the target chromaticity using color3mixer() and then optimizes the weights of each of the latter spectra such that adding them (additive mixing) results in obj\_vals as close as possible to the target values.

#### \_spd\_constructor\_nomixer(x)

Construct a mixture spectrum composed of n primaries using no mixer algorithm (just simple weighted sum of primaries).

#### Args:

X

optimization parameters, first n are the strengths of individual primaries.

#### **Returns:**

#### spd, prims, M

- spd: spectrum resulting from x
- spds: primary spds
- M: fluxes of all primaries

#### **Notes:**

1. 'no-mixer' - simple weighted sum of primaries.

# $_{\mathbf{fitness\_fcn}}(x, out = 'F')$

Fitness function that calculates closeness of solution x to target values for specified objective functions.

#### start(verbosity=None, out=None)

Start optimization of \_fitnessfcn for n primaries using the initialized minimizer and the selected optimizer\_type.

Returns variables specified in :out:

# luxpy.toolboxes.spdbuild.\_extract\_prim\_optimization\_parameters(x, nprims,

prim\_constructor\_parameter\_types,
prim\_constructor\_parameter\_defs)

Extract the primary parameters from the optimization vector x and the prim\_constructor\_parameter\_defs dict, for use with PrimConstructor..

# luxpy.toolboxes.spdbuild.\_stack\_wlr\_spd(wlr, spd)

Stack the wavelength range on top of the spd values for use with PrimConstructor.

# luxpy.toolboxes.spdbuild.\_setup\_wlr(wlr)

Setup the wavelength range for use with PrimConstructor.

```
luxpy.toolboxes.spdbuild.spd_optimizer2(target=array([[1.0000e+0.2, 3.3333e-0.1, 3.3333e-0.1]]), tar_type='Yxy', cspace_bwtf={}, n=4, wlr=[360, 830, 1], prims=None, cieobs='1931_2', out='spds,primss,Ms,results', optimizer_type='3mixer', prim_constructor=<function gaussian_prim_constructor>, prim_constructor_parameter_types=['peakwl', 'fwhm'], prim_constructor_parameter_defs={}, obj_fcn=None, obj_fcn_pars=[{}], obj_fcn_weights=[1], obj_tar_vals=[0], obj_tar_tols=[0], decimals=[5], triangle_strengths_bnds=None, minimize_method='Nelder-Mead', minimize_opts={}, x0=None, pareto=False, display=False, verbosity=1)
```

Generate a spectrum with specified white point and optimized for certain objective functions from a set of primary spectra or primary spectrum model parameters.

```
Args:
            target
                  np2d([100,1/3,1/3]), optional
                  ndarray with Yxy chromaticity of target.
            tar_type
                  'Yxy' or str, optional
                  Specifies the input type in :target: (e.g. 'Yxy' or 'cct')
            cspace bwtf
                  {}, optional
                  Backward (cspace to xyz) transform parameters
                  (see colortf()) to go from :tar_type: to 'Yxy').
            n
                  4, optional
                  Number of primaries in light mixture.
            wl
                  [360,830,1], optional
                  Wavelengths used in optimization when :prims: is not an ndarray with spectral data.
            cieobs
                  _CIEOBS, optional
                  CIE CMF set used to calculate chromaticity values, if not provided
                  in:Yxyi:.
            optimizer_type
                  '3mixer', optional
                  Specifies type of chromaticity optimization
                  For help on '3mixer' algorithm, see notes below.
            prims
                  ndarray of predefined primary spectra.
                  If None: they are built from optimization parameters using the
                  function in :prim_constructor:
            prim_constructor
```

function that constructs the primaries from the optimization parameters

```
Should have the form:
```

prim\_constructor(x, n, wl, prim\_constructor\_parameter\_types,
\*\*prim\_constructor\_parameter\_defs)

# $prim\_constructor\_parameter\_types$

gaussian\_prim\_parameter\_types ['peakwl', 'fwhm'], optional List with strings of the parameters used by prim\_constructor() to calculate the primary spd. All parameters listed and that do not have default values (one for each prim!!!) in prim\_constructor\_parameters\_defs will be optimized.

#### prim\_constructor\_parameters\_defs

{}, optional

Dict with constructor parameters required by prim\_constructor and/or default values for parameters that are not being optimized.

For example: {'fwhm': 30} will keep fwhm fixed and not optimize it.

## obj\_fcn

[None] or list, optional

Function handles to objective function.

#### obj\_fcn\_weights

[1] or list, optional.

Weigths for each obj. fcn

## obj\_fcn\_pars

[{}] or list, optional

Parameter dicts for each obj. fcn.

#### obj tar vals

[0] or list, optional

Target values for each objective function.

# obj\_tar\_tols

[0] or list, optional

Tolerance of objective function values with target values.

#### decimals

[5], optional

Rounding decimals of objective function values.

# minimize\_method

'Nelder-Mead', optional

Optimization method used by minimize function.

options:

- 'Nelder-Mead': Nelder-Mead simplex local optimization using the luxpy.math.minimizebnd wrapper with method set to 'Nelder-Mead'.
- 'demo' : Differential Evolutionary Multiobjective Optimizatizer (using math.DEMO.demo\_opt)
- 'particleswarm': Pseudo-global optimizer using particle swarms (from pyswarm wrapper module luxpy.math.pyswarms\_particleswarm)
- 'nsga\_ii': Pareto multiobjective optimizer using the NSGA-II genetic algorithm

(from pymoo wrapper module luxpy.math.pymoo\_nsga\_ii)

- A user-defined minimization function (see \_start\_optimization\_tri? for info on the requirements of this function)

#### minimize\_opts

None, optional

Dict with minimization options.

None defaults to the options depending on choice of minimize method

```
'Nelder-Mead': {'xtol': 1e-5, 'disp': True, 'maxiter': 1000*Nc, 'maxfev': 1000*Nc, 'fatol': 0.01}
'demo': {'F': 0.5, 'CR': 0.3, 'kmax': 300, 'mu': 100, 'display': True}
'particleswarm': {'iters': 100, 'n_particles': 10, 'ftol': -np.inf, 'ps_opts': {'c1': 0.5, 'c2': 0.3, 'w':0.9}}
'nsga_ii': {'n_gen': 40, 'n_pop': 400, 'n_offsprings': None, 'termination': ('n_gen', 40), 'seed': 1, 'ga_opts': {'sampling': ("real_random",{}), 'crossover': ("real_sbx", {'prob': 0.9, 'eta': 15}), 'mutation': ("real_pm", {'eta': 20})}}
```

- dict with options for user-defined minimization method.

# triangle\_strength\_bnds

(None, None)

Specifies lower- and upper-bounds for the strengths of each of the primary combinations that will be made during the optimization using '3mixer'.

 $\mathbf{x0}$ 

None, optional

If None: a random starting value will be generated for the Nelder-Mead minimization algorithm, else the user defined starting value will be used. Note that it should only contain a value for each peakwl and/or fwhm that is set to be optimized. The triangle\_strengths are added automatically.

#### pareto

False, optional

Specifies whether the output of the fitnessfcn should be the Root-Sum-of-Squares of all weighted objective function values or not. Individual function values are required by true multi-objective optimizers (i.e. pareto == True).

#### display

True, optional

Turn native display options of minimizers on (True) or off (False).

#### verbosity

0, optional

If > 0: print intermediate results.

#### out

'spds,primss,Ms,results', optional

Determines output of function (see :returns:).

#### **Returns:**

#### returns

spds, primss, Ms, results

- 'spds': optimized spectrum (or spectra: for demo, particleswarm and nsga\_ii minimization methods)

- 'primss': primary spectra of each optimized spectrum
- 'Ms': ndarrays with fluxes of each primary
- 'results': dict with optimization results

Notes on the optimization algorithms:

- 1. '3mixer': The triangle/trio method creates for all possible combinations of 3 primary component spectra a spectrum that results in the target chromaticity using color3mixer() and then optimizes the weights of each of the latter spectra such that adding them (additive mixing) results in obj\_vals as close as possible to the target values.
- 2. '2mixer': APRIL 2020, NOT YET IMPLEMENTED!! Pairs (odd,even) of components are selected and combined using 'pair\_strength'. This process is continued until only 3 (combined) intermediate sources remain. Color3mixer is then used to calculate the fluxes for the remaining 3 sources, after which the fluxes of all components are back-calculated.

luxpy.toolboxes.spdbuild.gaussian\_prim\_constructor(x, nprims, wlr, ptypes, \*\*pdefs)

Construct a set of nprim gaussian primaries with wavelengths wlr using the input in x and in kwargs.

Args:

X

ndarray (M x nprim) with optimization parameters.

#### nprim

number of primaries

wlr

wavelength range for which to construct a spectrum

#### prim\_constructor

function that constructs the primaries from the optimization parameters Should have the form:

prim constructor(x, n, wl, ptypes, pdefs)

#### ptypes

gaussian\_prim\_parameter\_types ['peakwl', 'fwhm'], optional List with strings of the parameters used by PrimConstructor()) to calculate the primary spd. All parameters listed and that do not have default values (one for each prim!!!) in pdefs will be optimized.

#### pdefs

Dict with constructor parameters required by PrimConstructor and/or default values for parameters that are not being optimized.

For example: {'fwhm': [30]} will keep fwhm fixed and not optimize it.

#### **Returns:**

spd

ndarray with spectrum of nprim primaries (1st row = wavelengths)

#### **Example on how to create constructor:**

```
`def gaussian_prim_constructor(x, nprims, wlr, ptypes, **pdefs):`
` # Extract the primary parameters from x and pdefs:`
` pars = _extract_prim_optimization_parameters(x, nprims, ptypes, pdefs)`
` # Setup wavelengths:`
` wlr = _setup_wlr(wlr)`
```

```
# Conversion factor for FWHM to sigma of Gaussian:
             fwhm_to_sig = 1/(2*(2*np.log(2))**0.5)
            # Create spectral profile function: `
             spd = np.exp(-0.5*((pars['peakwl']-wlr)/(pars['fwhm']*fwhm_to_sig))**2)`
             # Stack wlr and spd together:
             return _stack_wlr_spd(wlr,spd)`
luxpy.toolboxes.spdbuild._triangle_mixer(Yxy_target, Yxyi, triangle_strengths)
     Calculates the fluxes of each of the primaries to realize the target chromaticity Yxy_target given the trian-
     gle_strengths.
luxpy.toolboxes.spdbuild._color3mixer(Yxyt, Yxy1, Yxy2, Yxy3)
     Calculate fluxes required to obtain a target chromaticity when (additively) mixing 3 light sources.
     Args:
                Yxyt
                      ndarray with target Yxy chromaticities.
                Yxy1
                      ndarray with Yxy chromaticities of light sources 1.
                Yxy2
                      ndarray with Yxy chromaticities of light sources 2.
                Yxy3
                      ndarray with Yxy chromaticities of light sources 3.
     Returns:
                M
                      ndarray with fluxes.
     Note: Yxyt, Yxy1, ... can contain multiple rows, referring to single mixture.
4.5.4 hypspcim/
           рy
```

- \_\_init\_\_.py
- · hyperspectral\_img\_simulator.py

namespace luxpy.hypspcim

#### Module for hyper spectral image simulation

```
_HYPSPCIM_PATH path to module
_HYPSPCIM_DEFAULT_IMAGE path + filename to default image
xyz_to_rfl() approximate spectral reflectance of xyz based on k nearest neighbour interpola-
      tion of samples from a standard reflectance set.
render_image() Render image under specified light source spd.
```

```
luxpy.toolboxes.hypspcim.render_image(img=None, spd=None, rfl=None, out='img_hyp', refspd=None,
                                                D=None, cieobs='1931_2', cspace='xyz', cspace_tf={},
                                                CSF=None, interp_type='nd', k_neighbours=4, show=True,
                                                verbosity=0, show_ref_img=True, stack_test_ref=12,
                                                write_to_file=None, csf_based_rgb_rounding=6)
      Render image under specified light source spd.
      Args:
                  img
                        None or str or ndarray with float (max = 1) rgb image.
                        None load a default image.
                  spd
                        ndarray, optional
                        Light source spectrum for rendering
                        If None: use CIE illuminant F4
                  rfl
                        ndarray, optional
                        Reflectance set for color coordinate to rfl mapping.
                  out
                        'img_hyp' or str, optional
                              (other option: 'img_ren': rendered image under :spd:)
                  refspd
                        None, optional
                        Reference spectrum for color coordinate to rfl mapping.
                        None defaults to D65 (srgb has a D65 white point)
                  D
                        None, optional
                        Degree of (von Kries) adaptation from spd to refspd.
                  cieobs
                        _CIEOBS, optional
                        CMF set for calculation of xyz from spectral data.
                  cspace
                        'xyz', optional
                        Color space for color coordinate to rfl mapping.
                        Tip: Use linear space (e.g. 'xyz', 'Yuv',...) for (interp_type == 'nd'),
                              and perceptually uniform space (e.g. 'ipt') for (interp_type == 'nearest')
                  cspace_tf
                        {}, optional
                        Dict with parameters for xyz_to_cspace and cspace_to_xyz transform.
                  CSF
                        None, optional
                        RGB camera response functions.
                        If None: input :xyz: contains raw rgb values. Override :cspace:
                        argument and perform estimation directly in raw rgb space!!!
                  interp_type
                        'nd', optional
                        Options:
```

- 'nd': perform n-dimensional linear interpolation using Delaunay triangulation.
- 'nearest': perform nearest neighbour interpolation.

#### k\_neighbours

4 or int, optional

Number of nearest neighbours for reflectance spectrum interpolation.

Neighbours are found using scipy.spatial.cKDTree

#### show

True, optional

Show images.

#### verbosity

0, optional

If > 0: make a plot of the color coordinates of original and rendered image pixels.

#### show\_ref\_img

True, optional

True: shows rendered image under reference spd. False: shows original image.

#### write\_to\_file

None, optional

None: do nothing, else: write to filename(+path) in :write\_to\_file:

#### stack\_test\_ref

12, optional

- 12: left (test), right (ref) format for show and imwrite
- 21: top (test), bottom (ref)
- 1: only show/write test
- 2: only show/write ref
- 0: show both, write test

#### csf\_based\_rgb\_rounding

\_ROUNDING, optional

Int representing the number of decimals to round the RGB values (obtained from not-None CSF input) to before applying the search algorithm.

Smaller values increase the search speed, but could cause fatal error that causes python kernel to die. If this happens increase the rounding int value.

#### **Returns:**

#### returns

img\_hyp, img\_ren,

ndarrays with float hyperspectral image and rendered images

```
luxpy.toolboxes.hypspcim.xyz_to_rfl(xyz, CSF=None, rfl=None, out='rfl_est', refspd=None, D=None, cieobs='1931_2', cspace='xyz', cspace_tf={}, interp_type='nd', k_neighbours=4, verbosity=0, csf_based_rgb_rounding=6}
```

Approximate spectral reflectance of xyz values based on nd-dimensional linear interpolation or k nearest neighbour interpolation of samples from a standard reflectance set.

### Args:

xyz

ndarray with xyz values of target points.

**CSF** 

None, optional

```
RGB camera response functions.
      If None: input :xyz: contains raw rgb (float) values. Override :cspace:
      argument and perform estimation directly in raw rgb space!!!
rfl
      ndarray, optional
      Reflectance set for color coordinate to rfl mapping.
out
      'rfl_est' or str, optional
refspd
      None, optional
      Refer ence spectrum for color coordinate to rfl mapping.
      None defaults to D65.
cieobs
      _CIEOBS, optional
      CMF set used for calculation of xyz from spectral data.
cspace
      'xyz', optional
      Color space for color coordinate to rfl mapping.
      Tip: Use linear space (e.g. 'xyz', 'Yuv',...) for (interp_type == 'nd'),
            and perceptually uniform space (e.g. 'ipt') for (interp_type == 'nearest')
cspace_tf
      {}, optional
      Dict with parameters for xyz_to_cspace and cspace_to_xyz transform.
interp_type
      'nd', optional
      Options:
      - 'nd': perform n-dimensional linear interpolation using Delaunay triangulation.
      - 'nearest': perform nearest neighbour interpolation.
k_neighbours
      4 or int, optional
      Number of nearest neighbours for reflectance spectrum interpolation.
      Neighbours are found using scipy.spatial.cKDTree
verbosity
      0, optional
      If > 0: make a plot of the color coordinates of original and
      rendered image pixels.
csf_based_rgb_rounding
      _ROUNDING, optional
      Int representing the number of decimals to round the RGB values (obtained from
      not-None CSF input) to before applying the search algorithm.
      Smaller values increase the search speed, but could cause fatal error that causes
      python kernel to die. If this happens increase the rounding int value.
returns
      :rfl est:
```

4.5. Toolboxes 293

**Returns:** 

ndarrays with estimated reflectance spectra.

```
luxpy.toolboxes.hypspcim.get_superresolution_hsi(lrhsi, hrci, CSF, wl=[380, 780, 1], csf_based_rgb_rounding=6, interp_type='nd',
```

 $k_neighbours=4, verbosity=0)$ 

Get a HighResolution HyperSpectral Image (super-resolution HSI) based on a LowResolution HSI and a High-Resolution Color Image.

### Args:

#### lrhsi

ndarray with float (max = 1) LowResolution HSI [m,m,L].

hrci

ndarray with float (max = 1) HighResolution HSI [M,N,3].

**CSF** 

None, optional

ndarray with camera sensitivity functions

If None: use Nikon D700

wl

[380,780,1], optional

Wavelength range and spacing or ndarray with wavelengths of HSI image.

#### interp\_type

'nd', optional

Options:

- 'nd': perform n-dimensional linear interpolation using Delaunay triangulation.
- 'nearest': perform nearest neighbour interpolation.

### k\_neighbours

4 or int, optional

Number of nearest neighbours for reflectance spectrum interpolation.

Neighbours are found using scipy.spatial.cKDTree

#### verbosity

0, optional

Verbosity level for sub-call to render\_image().

If > 0: make a plot of the color coordinates of original and

rendered image pixels.

#### csf\_based\_rgb\_rounding

\_ROUNDING, optional

Int representing the number of decimals to round the RGB values (obtained from not-None CSF input) to before applying the search algorithm.

Smaller values increase the search speed, but could cause fatal error that causes python kernel to die. If this happens increase the rounding int value.

#### **Returns:**

#### hrhsi

ndarray with HighResolution HSI [M,N,L].

### **Procedure:**

Call render\_image(hrci, rfl = lrhsi\_2, CSF = ...) to estimate a hyperspectral image

from the high-resolution color image hrci with the reflectance spectra

in the low-resolution hyper-spectral image as database for the estimation.

Estimation is done in raw RGB space with the lrhsi converted using the

camera sensitivity functions in CSF. luxpy.toolboxes.hypspcim.hsi\_to\_rgb(hsi, spd=None, cieobs='1931\_2', srgb=False, linear\_rgb=False, CSF=None, normalize to white=True, wl=[380, 780, 1]) Convert HyperSpectral Image to rgb. Args: hsi ndarray with hyperspectral image [M,N,L] spd None, optional ndarray with illumination spectrum cieobs \_CIEOBS, optional CMF set to convert spectral data to xyz tristimulus values. srgb False, optional If False: Use xyz\_to\_srgb(spd\_to\_xyz(...)) to convert to srgb values If True: use camera sensitivity functions. linear\_rgb False, optional If False: use gamma = 2.4 in xyz\_to\_srgb, if False: use gamma = 1 and set :use\_linear\_part: to False. **CSF** None, optional ndarray with camera sensitivity functions If None: use Nikon D700 normalize\_to\_white True, optional If True & CSF is not None: white-balance output rgb to a perfect white diffuser. wl [380,780,1], optional Wavelength range and spacing or ndarray with wavelengths of HSI image. **Returns:** rgb ndarray with rgb image [M,N,3] luxpy.toolboxes.hypspcim.**rfl\_to\_rgb**(rfl, spd=None, CSF=None, wl=None, normalize\_to\_white=True) Convert spectral reflectance functions (illuminated by spd) to Camera Sensitivity Functions. **Args:** rfl ndarray with spectral reflectance functions (1st row is wavelengths if wl is None). spd None, optional ndarray with illumination spectrum **CSF** None, optional ndarray with camera sensitivity functions

If None: use Nikon D700

```
normalize_to_white
                        True, optional
                        If True: white-balance output rgb to a perfect white diffuser.
     Returns:
                  rgb
                        ndarray with rgb values for each spectral reflectance functions
4.5.5 dispcal/
           py
                      • __init__.py

    displaycalibration.py

           namespace luxpy.dispcal
Module for display characterization
           _PATH_DATA path to package data folder
            _RGB set of RGB values that work quite well for display characterization
            _XYZ example set of measured XYZ values corresponding to the RGB values in _RGB
           calibrate() Calculate TR parameters/lut and conversion matrices
           calibration performance() Check calibration performance (cfr. individual and average
                  color differences for each stimulus).
           rgb_to_xyz() Convert input rgb to xyz
           xyz to rgb() Convert input xyz to rgb
           DisplayCalibration() Calculate TR parameters/lut and conversion matrices and store in ob-
                 ject.
luxpy.toolboxes.dispcal.calibrate(rgbcal, xyzcal, L_type='lms', tr_type='lut', cieobs='1931_2', nbit=8,
                                          cspace='lab', avg=<function <lambda>>,
                                          ensure_increasing_lut_at_low_rgb=0.2, verbosity=1, sep=', ',
                                          header=None)
     Calculate TR parameters/lut and conversion matrices.
     Args:
                  rgbcal
                        ndarray [Nx3] or string with filename of RGB values
                        rgcal must contain at least the following type of settings:
                        - pure R,G,B: e.g. for pure R: (R != 0) & (G==0) & (B==0)
                        - white(s): R = G = B = 2**nbit-1
                        - gray(s): R = G = B
                        - black(s): R = G = B = 0
                        - binary colors: cyan (G = B, R = 0), yellow (G = R, B = 0), magenta (R = B, G = 0)
                 xyzcal
                        ndarray [Nx3] or string with filename of measured XYZ values for
                        the RGB settings in rgbcal.
```

```
L_type
      'lms', optional
      Type of response to use in the derivation of the Tone-Response curves.
      options:
            - 'lms': use cone fundamental responses: L vs R, M vs G and S vs B
                  (reduces noise and generally leads to more accurate characterization)
            - 'Y': use the luminance signal: Y vs R, Y vs G, Y vs B
tr_type
      'lut', optional
      options:
            - 'lut': Derive/specify Tone-Response as a look-up-table
            - 'gog': Derive/specify Tone-Response as a gain-offset-gamma function
cieobs
      '1931 2', optional
      CIE CMF set used to determine the XYZ tristimulus values
      (needed when L_{type} == 'lms': determines the conversion matrix to
      convert xyz to lms values)
nbit
      8, optional
      RGB values in nbit format (e.g. 8, 16, ...)
cspace
      color space or chromaticity diagram to calculate color differences in
      when optimizing the xyz_to_rgb and rgb_to_xyz conversion matrices.
avg
      lambda x: ((x**2).mean()**0.5), optional
      Function used to average the color differences of the individual RGB settings
      in the optimization of the xyz_to_rgb and rgb_to_xyz conversion matrices.
ensure_increasing_lut_at_low_rgb
      0.2 or float (max = 1.0) or None, optional
      Ensure an increasing lut by setting all values below the RGB with the maximum
      zero-crossing of np.diff(lut) and RGB/RGB.max() values of
      :ensure_increasing_lut_at_low_rgb:
      (values of 0.2 are a good rule of thumb value)
      Non-strictly increasing lut values can be caused at low RGB values due
      to noise and low measurement signal.
      If None: don't force lut, but keep as is.
verbosity
      1, optional
      > 0: print and plot optimization results
sep
      ',', optional
      separator in files with rgbcal and xyzcal data
header
      None, optional
      header specifier for files with rgbcal and xyzcal data
```

```
(see pandas.read_csv)
      Returns:
                  \mathbf{M}
                        linear rgb to xyz conversion matrix
                  \mathbf{N}
                        xyz to linear rgb conversion matrix
                  tr
                        Tone Response function parameters or lut
                  xyz_black
                        ndarray with XYZ tristimulus values of black
                  xyz_white
                        ndarray with tristimlus values of white
luxpy.toolboxes.dispcal.calibration_performance(rgb, xyztarget, M, N, tr, xyz_black, xyz_white,
                                                              tr_type='lut', cspace='lab', avg=<function
                                                              <lambda>>, rgb_is_xyz=False,
                                                              is_verification_data=False, nbit=8, verbosity=1,
                                                              sep=', ', header=None)
      Check calibration performance. Calculate DE for each stimulus.
      Args:
                  rgb
                        ndarray [Nx3] or string with filename of RGB values
                        (or xyz values if argument rgb_to_xyz == True!)
                  xyztarget
                        ndarray [Nx3] or string with filename of target XYZ values corresponding
                        to the RGB settings (or the measured XYZ values, if argument rgb_to_xyz == True).
                  M
                        linear rgb to xyz conversion matrix
                  N
                        xyz to linear rgb conversion matrix
                  tr
                        Tone Response function parameters or lut
                  xyz black
                        ndarray with XYZ tristimulus values of black
                  xyz_white
                        ndarray with tristimlus values of white
                  tr_type
                        'lut', optional
                        options:
                              - 'lut': Derive/specify Tone-Response as a look-up-table
                              - 'gog': Derive/specify Tone-Response as a gain-offset-gamma function
                  cspace
                        color space or chromaticity diagram to calculate color differences in.
                  avg
                        lambda x: ((x**2).mean()**0.5), optional
                        Function used to average the color differences of the individual RGB settings
```

```
rgb_is_xyz
                        False, optional
                        If True: the data in argument rgb are actually measured XYZ tristimulus values
                               and are directly compared to the target xyz.
                  is_verification_data
                        False, optional
                        If False: the data is assumed to be corresponding to RGB value settings used
                               in the calibration (i.e. containing whites, blacks, grays, pure and binary
                               mixtures)
                        If True: no assumptions on content of rgb, so use this settings when
                               checking the performance for a set of measured and target xyz data
                               different than the ones used in the actual calibration measurements.
                  nbit
                        8, optional
                        RGB values in nbit format (e.g. 8, 16, ...)
                  verbosity
                        1, optional
                        > 0: print and plot optimization results
                  sep
                         ',', optional
                        separator in files with rgbcal and xyzcal data
                  header
                        None, optional
                        header specifier for files with rgbcal and xyzcal data
                        (see pandas.read_csv)
      Returns:
                  M
                        linear rgb to xyz conversion matrix
                  Ν
                        xyz to linear rgb conversion matrix
                  tr
                        Tone Response function parameters or lut
                  xyz_black
                        ndarray with XYZ tristimulus values of black
                  xyz_white
                        ndarray with tristimlus values of white
luxpy.toolboxes.dispcal.rgb_to_xyz(rgb, M, tr, xyz_black, tr_type='lut')
      Convert input rgb to xyz.
      Args:
                  rgb
                        ndarray [Nx3] with RGB values
                  M
                        linear rgb to xyz conversion matrix
                  tr
```

in the optimization of the xyz\_to\_rgb and rgb\_to\_xyz conversion matrices.

```
Tone Response function parameters or lut
                 xyz_black
                        ndarray with XYZ tristimulus values of black
                  tr_type
                        'lut', optional
                        Type of Tone Response in tr input argument
                        options:
                              - 'lut': Tone-Response as a look-up-table
                              - 'gog': Tone-Response as a gain-offset-gamma function
     Returns:
                 XYZ
                        ndarray [Nx3] of XYZ tristimulus values
luxpy.toolboxes.dispcal.xyz_to_rgb(xyz, N, tr, xyz_black, tr_type='lut')
     Convert xyz to input rgb.
     Args:
                 xyz
                        ndarray [Nx3] with XYZ tristimulus values
                  N
                        xyz to linear rgb conversion matrix
                  tr
                        Tone Response function parameters or lut
                 xyz_black
                        ndarray with XYZ tristimulus values of black
                 tr_type
                        'lut', optional
                        Type of Tone Response in tr input argument
                        options:
                              - 'lut': Tone-Response as a look-up-table
                              - 'gog': Tone-Response as a gain-offset-gamma function
     Returns:
                  rgb
                        ndarray [Nx3] of display RGB values
class luxpy.toolboxes.dispcal.DisplayCalibration(rgbcal, xyzcal=None, L type='lms',
                                                              cieobs='1931_2', tr_type='lut', nbit=8, cspace='lab',
                                                              avg=<function DisplayCalibration.<lambda>>,
                                                              ensure_increasing_lut_at_low_rgb=0.2,
                                                              verbosity=1, sep=', ', header=None)
     Class for display calibration.
     Args:
                  rgbcal
                        ndarray [Nx3] or string with filename of RGB values
                        rgcal must contain at least the following type of settings:
                        - pure R,G,B: e.g. for pure R: (R != 0) & (G==0) & (B == 0)
                        - white(s): R = G = B = 2**nbit-1
                        - gray(s): R = G = B
                        - black(s): R = G = B = 0
```

```
- binary colors: cyan (G = B, R = 0), yellow (G = R, B = 0), magenta (R = B, G = 0)
xyzcal
      None, optional
      ndarray [Nx3] or string with filename of measured XYZ values for
      the RGB settings in rgbcal.
      if None: rgbcal is [Nx6] ndarray containing rgb (columns 0-2) and xyz data (columns
      3-5)
L_type
      'lms', optional
      Type of response to use in the derivation of the Tone-Response curves.
      options:
            - 'lms': use cone fundamental responses: L vs R, M vs G and S vs B
                  (reduces noise and generally leads to more accurate characterization)
            - 'Y': use the luminance signal: Y vs R, Y vs G, Y vs B
tr_type
      'lut', optional
      options:
            - 'lut': Derive/specify Tone-Response as a look-up-table
            - 'gog': Derive/specify Tone-Response as a gain-offset-gamma function
cieobs
      '1931_2', optional
      CIE CMF set used to determine the XYZ tristimulus values
      (needed when L_{type} == 'lms': determines the conversion matrix to
      convert xyz to lms values)
nbit
      8, optional
      RGB values in nbit format (e.g. 8, 16, ...)
cspace
      color space or chromaticity diagram to calculate color differences in
      when optimizing the xyz_to_rgb and rgb_to_xyz conversion matrices.
avg
      lambda x: ((x**2).mean()**0.5), optional
      Function used to average the color differences of the individual RGB settings
      in the optimization of the xyz_to_rgb and rgb_to_xyz conversion matrices.
verbosity
      1, optional
      > 0: print and plot optimization results
sep
      ',', optional
      separator in files with rgbcal and xyzcal data
header
      None, optional
      header specifier for files with rgbcal and xyzcal data
      (see pandas.read_csv)
calobject
```

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Return:

```
attributes are:
                        - M: linear rgb to xyz conversion matrix
                         - N: xyz to linear rgb conversion matrix
                         - TR: Tone Response function parameters or lut
                         - xyz_black: ndarray with XYZ tristimulus values of black
                         - xyz_white: ndarray with tristimlus values of white
                  as well as:
                         - rgbcal, xyzcal, cieobs, avg, tr_type, nbit, cspace, verbosity
                         - performance: dictionary with various color differences set to np.nan
                         - (run calobject.performance() to fill it with actual values)
check_performance(rgb=None, xyz=None, verbosity=None, sep=',', header=None, rgb_is_xyz=False,
                       is_verification_data=True)
      Check calibration performance (if rgbcal is None: use calibration data).
      Args:
                  rgb
                         None, optional
                         ndarray [Nx3] or string with filename of RGB values
                         (or xyz values if argument rgb_to_xyz == True!)
                         If None: use self.rgbcal
                  xyz
                         None, optional
                         ndarray [Nx3] or string with filename of target XYZ values corresponding
                         to the RGB settings (or the measured XYZ values, if argument rgb_to_xyz ==
                         True).
                         If None: use self.xyzcal
                  verbosity
                         None, optional
                         if None: use self.verbosity
                         if > 0: print and plot optimization results
                  sep
                         ',', optional
                         separator in files with rgb and xyz data
                  header
                         None, optional
                         header specifier for files with rgb and xyz data
                         (see pandas.read csv)
                  rgb_is_xyz
                         False, optional
                         If True: the data in argument rgb are actually measured XYZ tristimulus values
                               and are directly compared to the target xyz.
                  is_verification_data
                         False, optional
                         If False: the data is assumed to be corresponding to RGB value settings used
```

in the calibration (i.e. containing whites, blacks, grays, pure and binary

Performance results are stored in self.performance.

mixtures)

If True: no assumptions on content of rgb, so use this settings when checking the performance for a set of measured and target xyz data different than the ones used in the actual calibration measurements.

#### **Return:**

# performance

dictionary with various color differences.

```
to_xyz(rgb)
```

Convert display rgb to xyz.

to\_rgb(xyz)

Convert xyz to display rgb.

# 4.5.6 rgb2spec/

рy

- \_\_init\_\_.py
- · smits\_mitsuba.py

namespace luxpy.rgb2spec

Module for RGB to spectrum conversions

**\_BASESPEC\_SMITS** Default dict with base spectra for white, cyan, magenta, yellow, blue, green and red for each intent ('rfl' or 'spd')

rgb\_to\_spec\_smits() Convert an array of RGB values to a spectrum using a smits like conversion as implemented in mitsuba (July 10, 2019)

convert() Convert an array of RGB values to a spectrum (wrapper around rgb\_to\_spec\_smits(), future: implement other methods)

luxpy.toolboxes.rgb2spec.rgb\_to\_spec\_smits(rgb, intent='rfl', bitdepth=8, wlr=[360.0, 830.0, 1.0], rgb2spec=None)

Convert an array of RGB values to a spectrum using a Smits like conversion as implemented in Mitsuba. **Args:** 

rgb

ndarray of list of rgb values

intent

'rfl' (or 'spd'), optional

type of requested spectrum conversion.

bitdepth

8, optional

bit depth of rgb values

wlr

\_WL3, optional

desired wavelength (nm) range of spectrum.

rgb2spec

None, optional

Dict with base spectra for white, cyan, magenta, yellow, blue, green and red for each intent.

If None: use BASESPEC SMITS.

```
Returns:
                  spec
                        ndarray with spectrum or spectra (one for each rgb value, first row are the wavelengths)
luxpy.toolboxes.rgb2spec.convert(rgb, method='smits_mtsb', intent='rfl', bitdepth=8, wlr=[360.0, 830.0,
                                          1.0], rgb2spec=None)
      Convert an array of RGB values to a spectrum.
      Args:
                  rgb
                        ndarray of list of rgb values
                  method
                        'smits_mtsb', optional
                        Method to use for conversion:
                              - 'smits_mtsb': use a smits like conversion as implemented in mitsuba.
                  intent
                        'rfl' (or 'spd'), optional
                        type of requested spectrum conversion.
                  bitdepth
                        8, optional
                        bit depth of rgb values
                  wlr
                        _WL3, optional
                        desired wavelength (nm) range of spectrum.
                  rgb2spec
                        None, optional
                        Dict with base spectra for white, cyan, magenta, yellow, blue, green and red for each
                        If None: use _BASESPEC_SMITS.
      Returns:
                  spec
                        ndarray with spectrum or spectra (one for each rgb value, first row are the wavelengths)
4.5.7 iolidfiles/
```

рy

- \_\_init\_\_.py
- · io\_lid\_files.py

namespace luxpy.iolidfiles

# Module for reading and writing IES and LDT files.

read\_lamp\_data Read in light intensity distribution and other lamp data from LDT or IES
files.

**Notes:** 1.Only basic support. Writing is not yet implemented. 2.Reading IES files is based on Blender's ies2cycles.py 3.This was implemented to build some uvtexture maps for rendering and only tested for a few files. 4. Use at own risk. No warranties.

Read in light intensity distribution and other lamp data from LDT or IES files.

#### Args:

#### datasource

Filename of LID file or StringIO object or string with LID data.

#### multiplier

1.0, optional

Scaler for candela values.

#### verbosity

0, optional

Display messages while reading file.

#### normalize

'I0', optional

If 'I0': normalize LID to intensity at (theta,phi) = (0,0)

If 'max': normalize to max = 1.

#### only common keys

False, optional

If True, output only common dict keys related to angles, values and such of LID.

read\_lid\_lamp\_data(?) for print of common keys and return

empty dict with common keys.

#### **Returns:**

#### **Notes:**

- 1. if only\_common\_keys: output is dictionary with keys: ['datasource', 'version', 'intensity', 'theta', 'phi', 'values', 'map', 'Iv0', 'candela\_values', 'candela\_2d']
- 2. 'theta','phi', 'values' (='candela\_2d') contain the original theta angles, phi angles and normalized candelas as specified in file.
- 3. 'map' contains a dicionary with keys 'thetas', 'phis', 'values'. This data has been complete to full angle ranges thetas: [0,180]; phis: [0,360]

```
4. LDT map completion only supported for Isymm == 4 (since 31/10/2018), and Isymm == 1 (since, 02/10/2021), Map will be filled with original 'theta', 'phi' and normalized 'candela_2d' values!
5. LIDtype is checked by looking for the presence of 'TILT=' in datasource content (if True->'IES' else 'LDT')
```

 $\label{linear} \verb|luxpy.toolboxes.iolidfiles.get_uv_texture| (theta, phi=None, values=None, input_types=('array', 'array'), \\ method='linear', theta_min=0, angle_res=1, \\ close\_phi=False, deg=True, r=1, show=True, \\ out='values\_map') \\$ 

Create a uv-texture map. | with specified angular resolution (°) and with positive z-axis as normal. | u corresponds to phi  $[0^{\circ} - 360^{\circ}]$  | v corresponds to theta  $[0^{\circ} - 180^{\circ}]$ , (or  $[-90^{\circ} - 90^{\circ}]$ )

# Args:

#### theta

Float, int or ndarray

Angle with positive z-axis.

Values corresponding to 0 and 180° must be specified!

#### phi

None, optional

Float, int or ndarray

Angle around positive z-axis starting from x-axis.

If not None: values corresponding to 0 and 360° must be specified!

#### values

None

ndarray or mesh of values at (theta, phi) locations.

#### input\_types

('array', 'array'), optional

Specification of type of input of (angles, values)

#### method

'linear', optional

Interpolation method.

 $(supported\ scipy. interpolate. griddata\ methods:$ 

'nearest', 'linear', 'cubic')

# theta\_min

0, optional

If 0: [0, 180]; If -90: theta range = [-90,90]

# close\_phi

False, optional

Make phi angles array closed (full circle).

#### angle\_res

1, optional

Resolution in degrees.

#### deg

True, optional

Type of angle input (True: degrees, False: radians).

r

1, optional

Float, int or ndarray

```
radius
                  show
                        True, optional
                        Plot results.
                  out
                        'values_map', optional
                        Specifies output: "return eval(out)"
      Returns:
                  returns as specified by :out:.
luxpy.toolboxes.iolidfiles.save_texture(filename, tex, bits=16, transpose=True)
      Save 16 bit grayscale PNG image of uv-texture.
      Args:
                  filename
                        Filename of output image.
                  tex
                        ndarray float uv-texture.
                  transpose
                        True, optional
                        If True: transpose tex (u,v) to set u as columns and v as rows
                        in texture image.
      Returns:
                  None
      Note:
           Texture is rescaled to max = 1 and saved as uint 16.
            -> Before using uv_map: rescale back to set 'normal' to 1.
luxpy.toolboxes.iolidfiles.draw_lid(LID, grid_interp_method='linear', theta_min=0, angle_res=1,
                                              ax=None, projection='2d', polar_plot_Cx_planes=[0, 90],
                                              use_scatter_plot=False, plot_colorbar=True, legend_on=True,
                                              plot_luminaire_position=True, plot_diagram_top=0.001, out='ax',
                                              **plottingkwargs)
      Draw the light intensity distribution.
      Args:
                  LID
                        dict with IES or LDT file data.
                        (obtained with iolidfiles.read_lamp_data())
                  grid_interp_method
                        'linear', optional
                        Interpolation method for (theta,phi)-grid of normalized luminous intensity values.
                        (supported scipy.interpolate.griddata methods:
                              'nearest', 'linear', 'cubic')
                  theta_min
                        0, optional
                        If 0: [0, 180]; If -90: theta range = [-90,90]
                  angle_res
                        1, optional
                        Resolution in degrees.
```

```
ax
      None, optional
      If None: create new 3D-axes for plotting.
projection
      '2d', optional
      If '3d' make 3 plot
      If '2d': make polar plot(s). [not yet implemented (25/03/2021)]
polar\_plot\_Cx\_planes
      [0,90], optional
      Plot (Cx)-(Cx+180) planes; eg. [0,90] will plot C0-C180 and C90-C270 planes in 2D
      polar plot.
use_scatter_plot
      False, optional
      If True: use plt.scatter for plotting intensity values in 3D plot.
      If False: use plt.plot_surface for plotting in 3D plot.
plot_colorbar
      True, optional
      Plot colorbar representing the normalized luminous intensity values in the LID 3D
      plot.
legend_on
      True, optional
      If True: plot legend on polar plot (no legend for 3D plot!).
plot_luminaire_position
      True, optional
      Plot the position of the luminaire (0,0,0) in the 3D graph as a red diamond.
plot_diagram_top
      1e-3, optional
      Plot the top of the polar diagram (True).
      If None: automatic detection of non-zero intensity values in top part.
      If float: automatic detection of intensity values larger than max_intensity*float in top
      part.
            (if smaller: don't plot top.)
out
      'ax', optional
      string with variable to return
      default: ax handle to plot.
returns
```

Whatever requested as determined by the string in :out:

**Returns:** 

```
luxpy.toolboxes.iolidfiles.render_lid(LID=',/data/luxpy test lid file.ies', sensor resolution=100,
                                                 sensor\_position=[0, -1, 0.8], sensor\_n=[0, 1, -0.2], fov=(90, 90),
                                                 Fd=2, luminaire position=[0, 1.3, 2], luminaire n=[0, 0, -1],
                                                 wall_center=[0, 2, 1], wall_n=[0, -1, 0], wall_width=4,
                                                 wall\_height=2, wall\_rho=1, floor\_center=[0, 1, 0], floor\_n=[0, 0, 0]
                                                 1], floor width=4, floor height=2, floor rho=1,
                                                 grid interp method='linear', angle res=5, theta min=0,
                                                ax3D=None, ax2D=None, join_axes=True, legend_on=True,
                                                plot luminaire position=True, plot lumiaire rays=False,
                                                 plot_luminaire_lid=True, plot_sensor_position=True,
                                                plot_sensor_pixels=True, plot_sensor_rays=False,
                                                 plot_wall_edges=True, plot_wall_luminance=True,
                                                plot_wall_intersections=False, plot_floor_edges=True,
                                                 plot_floor_luminance=True, plot_floor_intersections=False,
                                                 out='Lv2D')
      Render a light intensity distribution.
      Args:
                  LID
                        dict with IES or LDT file data or string with path/filename;
                        or String or StringIO object with IES or LDT data.
                        (dict should be obtained with iolidfiles.read lamp data())
                  sensor_resolution
                        100, optional
                        Number of sensor 'pixels' along each dimension.
                  sensor\_position
                        [0,-1,0.8], optional
                        x,y,z position of the sensor 'focal' point (is located Fd meters behind actual sensor
                        plane)
                  sensor_n
                        [0,1,-0.2], optional
                        Sensor plane surface normal
                  fov
                        (90,90), optional
                        Field of view of sensor image in degrees.
                  Fd
                        'Focal' distance in meter. Sensor center is located Fd meter away from
                        :sensor_position:
                  luminaire_position
                        [0,1.3,2], optional
                        x,y,z position of the photometric equivalent point source
                  luminaire_n
                        [0,0,-1], optional
                        Orientation of lumaire LID (default points downward along z-axis away from source)
                  wall_center
                        [0,2,1], optiona
                        x,y,z position of the back wall
                  wall_n
```

```
[0,-1,0], optional
      surface normal of wall
wall_width
      4, optional
      width of wall (m)
wall_height
      2, optional
      height of wall (m)
wall_rho
      1, optional
      Diffuse (Lambertian) reflectance of wall.
floor center
      [0,1,0], optiona
      x,y,z position of the floor
floor_n
      [0,0,1], optional
      surface normal of floor
floor_width
      4, optional
      width of floor (m)
floor_height
      2, optional
      height of floor (m)
floor_rho
      1, optional
      Diffuse (Lambertian) reflectance of floor.
grid_interp_method
      'linear', optional
      Interpolation method for (theta,phi)-grid of normalized luminous intensity values.
      (supported scipy.interpolate.griddata methods:
            'nearest', 'linear', 'cubic')
theta min
      0, optional
      If 0: [0, 180]; If -90: theta range = [-90,90]
      Only used when generating a plot of the LID in the 3D graphs.
angle_res
      1, optional
      Angle resolution in degrees of LID sampling.
      Only used when generating a plot of the LID in the 3D graphs.
ax3D,ax2D
      None, optional
      If None: create new 3D- or 2D- axes for plotting.
      If join_axes == True: try and combine two axes on same figure.
      If False: don't plot..
legend_on
```

```
False, optional plot legend.
```

# plot\_luminaire\_position

True, optional

Plot the position of the luminaire (0,0,0) in the graph as a red diamond.

# plot\_X...

VArious options to customize plotting. Mainly allows for plotting of additional info such as plane-ray intersection points, sensor pixels, sensor-to-plane rays, plane-to-luminaire rays, 3D plot of LID, etc.

out

'Lv2D', optional string with variable to return default: variable storing an grayscale image of the rendered LID.

#### **Returns:**

#### returns

Whatever requested as determined by the string in :out:

# 4.5.8 spectro/

рy

- \_\_init\_\_.py
- · spectro.py

namespace luxpy.spectro

#### Package for spectral measurements

# Supported devices:

- JETI: specbos 1211, etc.
- OceanOptics: QEPro, QE65Pro, QE65000, USB2000, USB650,etc.

**get\_spd()** wrapper function to measure a spectral power distribution using a spectrometer of one of the supported manufacturers.

#### **Notes**

- 1. For info on the input arguments of get\_spd(), see help for each identically named function in each of the sub-packages.
- 2. The use of jeti spectrometers requires access to some dll files (delivered with this package).
- 3. The use of oceanoptics spectrometers requires the manual installation of pyseabreeze, as well as some other 'manual' settings. See help for oceanoptics sub-package.

#### luxpy.toolboxes.spectro.init(manufacturer)

Import module for specified manufacturer. Make sure everything (drivers, external packages, ...) required is installed!

```
luxpy.toolboxes.spectro.get_spd(manufacturer='jeti', dvc=0, Tint=0, autoTint_max=None,
                                         close device=True, out='spd', **kwargs)
      Measure a spectral power distribution using a spectrometer of one of the supported manufacturers.
      Args:
                  manufacturer
                        'jeti' or 'oceanoptics', optional
                        Manufacturer of spectrometer (ensures the correct module is loaded).
                  dvc
                        0 or int or spectrometer handle, optional
                        If int: function will try to initialize the spectrometer to
                              obtain a handle. The int represents the device
                              number in a list of all detected devices of the manufacturer.
                  Tint
                        0 or Float, optional
                        Integration time in seconds. (if 0: find best integration time, but < autoTint_max).
                  autoTint max
                        Limit Tint to this value when Tint = 0.
                  close device
                        True, optional
                        Close spectrometer after measurement.
                        If 'dvc' not in out.split(','): always close!!!
                  out
                        "spd" or e.g. "spd,dvc,Errors", optional
                        Requested return.
                  kwargs
                        For info on additional input (keyword) arguments of get_spd(),
                        see help for each identically named function in each of the subpackages.
      Returns:
                  spd
                        ndarray with spectrum. (row 0: wavelengths, row1: values)
                  dvc
                        Device handle, if successfull open (_ERROR: failure, nan: closed)
                  Errors
                        Dict with error messages.
4.5.9 sherbrooke spectral indices/
            рy

    init .py
```

• sherbrooke\_spectral\_indices\_2013.py

namespace luxpy.sherbrooke\_spectral\_indices

spd\_to\_msi() calculate Melatonin Suppression Index from spectrum.

```
spd_to_sli() calculate Star Light Index from spectrum.
References: 1. Aubé M, Roby J, Kocifaj M (2013) Evaluating Potential Spectral Impacts of Various Artificial Lights
     on Melatonin Suppression, Photosynthesis, and Star Visibility. PLoS ONE 8(7): e67798 https://journals.plos.
     org/plosone/article?id=10.1371/journal.pone.0067798
Created on Fri Jun 11 13:46:33 2021
@author: ksmet1977 [at] gmail dot com
luxpy.toolboxes.sherbrooke_spectral_indices.spd_to_msi(spd, force_5nm_interval=True)
     Calculate Melatonin Suppression Index from spectrum.
     Args:
                 spd
                       ndarray with spectral data (first row are wavelengths)
                 force_5nm_interval
                       True, optional
                       If True: interpolate spd to 5nm wavelengths intervals, else: keep as in spd.
     Returns:
                 msi
                       ndarray with Melatonin Suppression Index values for each input spectrum.
luxpy.toolboxes.sherbrooke_spectral_indices.spd_to_ipi(spd, force_5nm_interval=True)
     Calculate Induced Photosynthesis Index from spectrum.
     Args:
                 spd
                       ndarray with spectral data (first row are wavelengths)
                 force_5nm_interval
                       True, optional
                       If True: interpolate spd to 5nm wavelengths intervals, else: keep as in spd.
     Returns:
                 msi
                       ndarray with Induced Photosynthesis Index values for each input spectrum.
luxpy.toolboxes.sherbrooke_spectral_indices.spd_to_sli(spd, force_5nm_interval=True)
     Calculate Star Light Index from spectrum.
     Args:
                 spd
                       ndarray with spectral data (first row are wavelengths)
                 force_5nm_interval
                       True, optional
                       If True: interpolate spd to 5nm wavelengths intervals, else: keep as in spd.
     Returns:
                 msi
                       ndarray with Star Light Index values for each input spectrum.
```

**spd\_to\_ipi()** calculate Induced Photosynthesis Index from spectrum.

# 4.5.10 spectral mismatch and uncertainty/

рy

- \_\_init\_\_.py
- detector\_spectral\_mismatch.py

**namespace** luxpy.spectral\_mismatch\_and\_uncertainty

# Toolbox for spectral mismatch and measurement uncertainty calculations

spectral\_mismatch\_and\_uncertainty/detector\_spectral\_mismatch.py

**f1prime**() Determine the f1prime spectral mismatch index.

get\_spectral\_mismatch\_correct\_factors() Determine the spectral mismatch factors.

#### Reference

1. Krüger, U. et al. GENERAL V() MISMATCH - INDEX HISTORY, CURRENT STATE, NEW IDEAS (TechnoTeam)

```
Created on Tue Aug 31 10:46:02 2021

@author: ksmet1977 [at] gmail.com

luxpy.toolboxes.spectral_mismatch_and_uncertainty.flprime(s_detector, S_C='A', cieobs='1931_2', s_target_index=2, wlr=None, interp_kind='linear', out='flp')
```

Determine the f1prime spectral mismatch index.

Args:

#### s\_detector

ndarray with detector spectral responsivity (first row = wavelengths)

 $S_C$ 

'A', optional

Standard 'calibration' illuminant.

string specifying the illuminant to use from the luxpy.\_CIE\_ILLUMINANTS dict or ndarray with standard illuminant spectral data.

### cieobs

```
'1931_2', optional string with CIE standard observer color matching functions to use (from luxpy._CMF) or ndarray with CMFs (s_target_index > 0) or target spectral responsivity (s_target_index == 0) (first row contains the wavelengths).
```

### s\_target\_index

2, optional

if > 0: index into CMF set (1->'xbar', 2->'ybar'='Vlambda', 3->'zbar')

if == 0: cieobs is expected to contain an ndarray with the target spectral responsivity.

wlr

s\_detector,  $S_C='A'$ cieobs='1931 2 s target index= wlr=None.

terp\_kind='lined out='F')

in-

```
If None: the wavelengths of the detector are used throughout.
                  interp_kind
                         'linear', optional
                         Interpolation type to use when interpolating function to specified wavelength range.
                  out
                         'f1p', optional
                         Specify requested output of function,
                               e.g. 'flp,s_rel' also outputs the normalized target spectral responsitivity.
      Returns:
                  f1p
                         ndarray (vector) with f1prime values for each of the spectral responsivities in
                         s detector.
luxpy.toolboxes.spectral_mismatch_and_uncertainty.get_spectral_mismatch_correction_factors(S_Z,
      Determine the spectral mismatch factors.
      Args:
                  \mathbf{S} \mathbf{Z}
                         ndarray with spectral power distribution of measured light source (first row =
                         wavelengths).
                  s detector
                         ndarray with detector spectral responsivity (first row = wavelengths)
                  S C
                         'A', optional
                         Standard 'calibration' illuminant.
                         string specifying the illuminant to use from the luxpy._CIE_ILLUMINANTS dict
                         or ndarray with standard illuminant spectral data.
                  cieobs
                         '1931_2', optional
                         string with CIE standard observer color matching functions to use (from luxpy._CMF)
                         or ndarray with CMFs (s_target_index > 0)
                         or target spectral responsivity (s_target_index == 0)
                         (first row contains the wavelengths).
                  s\_target\_index
                         2, optional
                         if > 0: index into CMF set (1->'xbar', 2->'ybar'='Vlambda', 3->'zbar')
                         if == 0: cieobs is expected to contain an indurray with the target spectral responsivity.
                  wlr
                         None, optional
```

None, optional

Wavelength range (None, ndarray or [start, stop, spacing]).

Wavelength range (ndarray or [start, stop, spacing]).

If None: use the wavelength range of S\_Z.

# interp\_kind

'linear', optional

Interpolation type to use when interpolating function to specified wavelength range.

out

'F', optional

Specify requested output of function,

e.g. 'F,f1p' also outputs the f1prime spectral mismatch index.

# **Returns:**

 $\mathbf{F}$ 

ndarray with correction factors for each of the mesured spectra (rows) and spectral responsivities in s\_detector (columns).

# **CHAPTER**

# **FIVE**

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