# Package ‘pavo’

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**Title**  Perceptual Analysis, Visualization and Organization of Spectral Colour Data

**Version**  2.3.0

**Description**  A cohesive framework for parsing, analyzing and organizing colour from spectral data.

**License**  GPL (>= 2)

**URL**  http://rafaelmaia.net/pavo/, https://github.com/rmaia/pavo/

**BugReports**  https://github.com/rmaia/pavo/issues

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adjacent

Run an adjacency and boundary strength analysis

**Description**

Calculate summary variables from the adjacency (Endler 2012) and boundary-strength (Endler et al. 2018) analyses, along with overall pattern contrast (Endler & Mielke 2005).

**Usage**

```r
adjacent(
  classimg,  
  xpts = NULL,  
  xscale = NULL,  
  bkgID = NULL,  
  polygon = NULL,  
  exclude = c("none", "background", "object"),  
  coldists = NULL,  
  hsl = NULL,  
  cores = getOption("mc.cores", 2L)
)
```

**Arguments**

- `classimg` (required) an xyz matrix, or list of matrices, in which x and y correspond to spatial (e.g. pixel) coordinates, and z is a numeric code specifying a colour-class. Preferably the result of `classify()`, or constructed from grid-sampled spectra that have been visually modelled and clustered (as per Endler 2012).

- `xpts` an integer specifying the number of sample points along the x axis, from which the evenly-spaced sampling grid is constructed (if required). Defaults to the smallest dimension of `classimg`, though this should be carefully considered.

- `xscale` (required) an integer specifying the true length of the x-axis, in preferred units. Not required, and ignored, only if image scales have been set via `procimg()`.

- `bkgID` an integer or vector specifying the colour-class ID number(s) of pertaining to the background alone, for relatively homogeneous and uniquely-identified backgrounds (e.g. the matte background of pinned specimens). Examine the attributes of, or call `summary` on, the result of `classify()` to visualise the RGB values corresponding to colour-class ID numbers for classified images. Ignored if the focal object and background has been identified using `procimg()`.

- `polygon` a data.frame of x-y coordinates delineating a closed polygon that separates the focal object from the background. Not required, and ignored, if the focal object outline is specified using `procimg()`.
exclude the portion of the scene to be excluded from the analysis, if any.

- 'none': default
- 'background': exclude everything outside the closed polygon specified using procimg(), or the argument polygon. Alternatively, if the background is relatively homogeneous the colour-class ID(s) uniquely corresponding to the background can be specified via bkgID, and subsequently excluded.
- 'object': exclude everything inside the closed polygon specified using procimg(), or the argument polygon.

coldists a data.frame specifying the visually-modelled chromatic (dS) and/or achromatic (dL) distances between colour-categories. The first two columns should be named 'c1' and 'c2', and specify all possible combinations of numeric colour-class ID’s (viewable by calling summary(image, plot = TRUE) on a colour classified image), with the remaining columns named dS (for chromatic distances) and/or dL (for achromatic distances). See vismodel() and colspace() for visual modelling with spectral data.

hsl data.frame specifying the hue, saturation, and luminance of color patch elements, as might be estimated via vismodel() and colspace(). The first column, named 'patch', should contain numeric color category IDs, with the remaining columns specifying one or more of 'hue' (angle, in radians), 'sat', and/or 'lum'.

cores number of cores to be used in parallel processing. If 1, parallel computing will not be used. Defaults to getOption("mc.cores",2L). Not available on Windows.

Value

a data frame of summary variables:

- 'k': The number of user-specified colour and/or luminance classes.
- 'N': The grand total (sum of diagonal and off-diagonal) transitions.
- 'n_off': The total off-diagonal transitions.
- 'p_i': The overall frequency of colour class i.
- 'q_i_j': The frequency of transitions between all colour classes i and j, such that \( \sum q_{i,j} = 1 \).
- 't_i_j': The frequency of off-diagonal (i.e. class-change transitions) transitions i and j, such that \( \sum t_{i,j} = 1 \).
- 'm': The overall transition density (mean transitions), in units specified in the argument xscale.
- 'm_r': The row-wise transition density (mean row transitions), in user-specified units.
- 'm_c': The column-wise transition density (mean column transitions), in user-specified units.
- 'A': The transition aspect ratio (< 1 = wide, > 1 = tall).
- 'Sc': Simpson colour class diversity, \( Sc = 1/(\sum p_i^2) \). If all colour and luminance classes are equal in relative area, then \( Sc = k \).
• 'St': Simpson transition diversity, \( St = 1/\sum(t_{i,j}^2) \).

• 'Jc': Simpson colour class diversity relative to its achievable maximum. \( Jc = Sc/k \).

• 'Jt': Simpson transition diversity relative to its achievable maximum. \( Jt = St/(k*(k-1)/2) \).

• 'B': The animal/background transition ratio, or the ratio of class-change transitions entirely within the focal object and those involving the object and background, \( B = \sum(O_{a_a}/O_{a_b}) \).

• 'Rt': Ratio of animal-animal and animal-background transition diversities, \( Rt = St_{a_a}/St_{a_b} \).

• 'Rab': Ratio of animal-animal and background-background transition diversities, \( Rt = St_{a_a}/St_{b_b} \).

• 'm_dS', 's_dS', 'cv_dS': weighted mean, sd, and coefficient of variation of the chromatic boundary strength.

• 'm_dL', 's_dL', 'cv_dL': weighted mean, sd, and coefficient of variation of the achromatic boundary strength.

• 'm_hue', 's_hue', 'var_hue': circular mean, sd, and variance of overall pattern hue (in radians).

• 'm_sat', 's_sat', 'cv_sat': weighted mean, sd, and coefficient variation of overall pattern saturation.

• 'm_lum', 's_lum', 'cv_lum': weighted mean, sd, and coefficient variation of overall pattern luminance.

Author(s)

Thomas E. White <thomas.white026@gmail.com>

References


See Also

classify(), summary.rimg(), procimg()

Examples

```r
# Not run:
# Set a seed, for reproducibility
set.seed(153)

# Single image
papilio <- getimg(system.file("testdata/images/papilio.png", package = "pavo"))
papilio_class <- classify(papilio, kcols = 4)
papilio_adj <- adjacent(papilio_class, xscale = 100)
```
# Single image, with (fake) color distances and hsl values
# Fake color distances
distances <- data.frame(
  c1 = c(1, 1, 1, 2, 2, 3),
  c2 = c(2, 3, 4, 3, 4, 4),
  dS = c(5.3, 3.5, 5.7, 2.9, 6.1, 3.2),
  dL = c(5.5, 6.6, 3.3, 2.2, 4.4, 6.6)
)

# Fake hue, saturation, luminance values
hsl_vals <- data.frame(
  patch = seq_len(4),
  hue = c(1.5, 2.2, 1.0, 0.5),
  lum = c(10, 5, 7, 3),
  sat = c(3.5, 1.1, 6.3, 1.3)
)

# Full analysis, including the white background’s ID
papilio_adj <- adjacent(papilio_class,
  xscale = 100, bkgID = 1,
  coldists = distances, hsl = hsl_vals)

# Multiple images
snakes <- getimg(system.file("testdata/images/snakes", package = "pavo"))
snakes_class <- classify(snakes, kcols = 3)
snakes_adj <- adjacent(snakes_class, xpts = 120, xscale = c(50, 55))

## End(Not run)

---

### aggplot

**Plot aggregated reflectance spectra**

#### Description

Combines and plots spectra (by taking the average and the standard deviation, for example) according to an index or a vector of identities.

#### Usage

```r
aggplot(
  rspecdata,
  by = NULL,
  FUN.center = mean,
  FUN.error = sd,
  lcol = NULL,
  shadecol = NULL,
  alpha = 0.2,
)```

aggplot

legend = FALSE,
...
)

Arguments

rspecdata (required) a data frame, possibly of class rspec, which contains a column containing a wavelength range, named 'wl', and spectra data in remaining columns.

by (required) either a single value specifying the range of spectra within the data frame to be combined (for example, by = 3 indicates the function will be applied to groups of 3 consecutive columns in the spectra data frame); a vector containing identifications for the columns in the spectra data frame (in which case the function will be applied to each group of spectra sharing the same identification); or a list of vectors, e.g., by = list(sex,species).

FUN.center the function to be applied to the groups of spectra, calculating a measure of central tendency (defaults to base::mean()).

FUN.error the function to be applied to the groups of spectra, calculating a measure of variation (defaults to stats::sd()).

lcol colour of plotted lines indicating central tendency.

shadecol colour of shaded areas indicating variance measure.

alpha transparency of the shaded areas.

legend automatically add a legend.

... additional graphical parameters to be passed to plot.

Value

Plot containing the lines and shaded areas of the groups of spectra.

Author(s)

Rafael Maia <rm72@zips.uakron.edu>, Chad Eliason <cme16@zips.uakron.edu>

References


Examples

# Load reflectance data
data(sicalis)

# Create grouping variable based on spec names
bysic <- gsub("\d\d\d\d\d\d\", ",", names(sicalis)[-1])

# Plot using various error functions and options
aggplot(sicalis, bysic)
aggplot(sicalis, bysic, FUN.error = function(x) quantile(x, c(0.0275, 0.975)))
aggplot(sicalis, bysic, shadecol = spec2rgb(sicalis), lcol = 1)
aggplot(sicalis, bysic, lcol = 1, FUN.error = function(x) sd(x) / sqrt(length(x)))

---

**aggspec**

*Aggregate reflectance spectra*

**Description**

Combines spectra (by taking the average, for example) according to an index or a vector of identities.

**Usage**

```r
aggspec(rspecdata, by = NULL, FUN = mean, trim = TRUE)
```

**Arguments**

- **rspecdata** (required) a data frame, possibly of class rspec, which contains a column containing a wavelength range, named 'wl', and spectra data in remaining columns.
- **by** (required) either a single value specifying the range of spectra within the data frame to be combined (for example, by = 3 indicates the function will be applied to groups of 3 consecutive columns in the spectra data frame); a vector containing identifications for the columns in the spectra data frame (in which case the function will be applied to each group of spectra sharing the same identification); or a list of vectors, e.g., by = list(sex, species).
- **FUN** the function to be applied to the groups of spectra. (defaults to `mean()`)  
- **trim** logical. if TRUE (default), the function will try to identify and remove numbers at the end of the names of the columns in the new rspec object.

**Value**

A data frame of class rspec containing the spectra after applying the aggregating function.

**Author(s)**

Chad Eliason <cme16@zips.uakron.edu>

**References**

Examples

data(teal)

# Average every two spectra
teal.sset1 <- aggspec(teal, by = 2)
plot(teal.sset1)

# Create factor and average spectra by levels 'a' and 'b'
ind <- rep(c("a", "b"), times = 6)
teal.sset2 <- aggspec(teal, by = ind)
plot(teal.sset2)

as.rimg  

Convert data to an rimg object

Description

Converts an array of RGB values, a cimg object, or a magick-image object, to an rimg object.

Usage

as.rimg(object, name = "img")

## Default S3 method:
as.rimg(object, name = "img")

## S3 method for class 'cimg'
as.rimg(object, name = "img")

is.rimg(object)

Arguments

object  
(required) a three-dimensional array containing RGB values.

name  
the name(s) of the image(s).

Value

an object of class rimg for use in further pavo functions

a logical value indicating whether the object is of class rimg

Author(s)

Thomas E. White <thomas.white026@gmail.com>
Hugo Gruson <hugo.gruson+R@normalesup.org>
# Examples

```r
# Generate some fake image data
defake <- array(c(
    as.matrix(rep(c(0.2, 0.4, 0.6), each = 250)),
    as.matrix(rep(c(0.4, 0.7, 0.8), each = 250)),
    as.matrix(rep(c(0.6, 0.1, 0.2), each = 250))
),
dim = c(750, 750, 3)
)

# Inspect it
head(defake)

# Determine if is rimg object
is.rimg(defake)

# Convert to rimg object and check again
defake2 <- as.rimg(defake)
is.rimg(defake2)
```

---

**as.rspec**

Convert data to an rspec object

## Description
Converting data frames or matrices containing spectral data to rspec object

## Usage

```r
as.rspec(
  object,
  whichwl = NULL,
  interp = TRUE,
  lim = NULL,
  exceed.range = TRUE
)
```

is.rspec(object)

## Arguments

- **object** *(required)* a data frame or matrix containing spectra to process.
- **whichwl** a numeric or character vector specifying which column contains wavelengths. If NULL (default), function searches for column containing equally spaced numbers and sets it as wavelengths "wl". If no wavelengths are found or whichwl is not given, returns arbitrary index values.
- **interp** whether to interpolate wavelengths in 1-nm bins (defaults to TRUE).
axistetra

lim vector specifying wavelength range to interpolate over (e.g. c(300, 700)).

exceed.range logical. Should data be interpolated to the limits specified by lim if lim exceeds the range of the actual data? Useful, and relatively safe, when the data range falls slightly within lim (e.g. 300.1 - 699 nm), but will produce spurious results if lim far exceeds the range of input data. Defaults to TRUE.

Value

an object of class rspec for use in further pavo functions

a logical value indicating whether the object is of class rspec

Author(s)

Chad Eliason <cme16@zips.uakron.edu>

Examples

# Generate some fake reflectance data
fakedat <- data.frame(wl = 300:700, refl1 = rnorm(401), refl2 = rnorm(401))
head(fakedat)

# Determine if is rspec object
is.rspec(fakedat)

# Convert to rspec object
fakedat2 <- as.rspec(fakedat)
is.rspec(fakedat2)
head(fakedat2)

Description

Plots reference x, y and z arrows showing the direction of the axes in a static tetrahedral colourspace plot.

Usage

axistetra(
  x = 0,
  y = 1.3,
  size = 0.1,
  arrowhead = 0.05,
  col = par("fg"),
  lty = par("lty"),
  lwd = par("lwd"),
)
label = TRUE,
adj.label = list(x = c(0.003, 0), y = c(0.003, 0.003), z = c(0, 0.003)),
label.cex = 1,
label.col = NULL
```

Arguments

x, y  position of the legend relative to plot limits (usually a value between 0 and 1, but because of the perspective distortion, values greater than 1 are possible)
size  length of the arrows. Can be either a single value (applied for x, y and z) or a vector of 3 separate values for each axis.
arrows  size of the arrowhead.
col, lty, lwd  graphical parameters for the arrows.
label  logical, include x, y and z labels (defaults to TRUE).
adj.label  position adjustment for the labels. A list of 3 named objects for x, y and z arrows, each with 2 values for x and y adjustment.
label.cex, label.col  graphical parameters for the labels.

Value

axistetra adds reference arrows showing the direction of the 3-dimensional axes in a static tetrahedral colourspace plot.

Author(s)

Rafael Maia <rm72@zips.uakron.edu>

Examples

data(sicalis)
vis.sicalis <- vismodel(sicalis, visual = "avg.uv")
tcs.sicalis <- colspace(vis.sicalis, space = "tcs")
plot(tcs.sicalis)
axistetra()

---

bgandilum  Default background and illuminant data

Description

Default background and illuminant data

Author(s)

Rafael Maia <rm72@zips.uakron.edu>
References


bootcoldist

Bootstrap colour distance confidence intervals

Description

Uses a bootstrap procedure to generate confidence intervals for the mean colour distance between two or more samples of colours

Usage

bootcoldist(
  vismodeldata,
  by,
  boot.n = 1000,
  alpha = 0.95,
  cores = getOption("mc.cores", 2L),
  ...
)

Arguments

vismodeldata (required) quantum catch colour data. Can be the result from vismodel(), or colspace(). Data may also be independently calculated quantum catches, in the form of a data frame with columns representing photoreceptors.

by (required) a numeric or character vector indicating the group to which each row from the object belongs to.

boot.n number of bootstrap replicates (defaults to 1000)

alpha the confidence level for the confidence intervals (defaults to 0.95)

cores number of cores to be used in parallel processing. If 1, parallel computing will not be used. Defaults to getOption("mc.cores", 2L)

... other arguments to be passed to coldist(). Must at minimum include n and weber. See coldist() for details.

Value

a matrix including the empirical mean and bootstrapped confidence limits for dS (and dL if achro = TRUE).

References

## Not run:
data(sicalis)
vm <- vismodel(sicalis, achro = "bt.dc")
gr <- gsub("ind\..", ",", rownames(vm))
bootcoldist(vm, by = gr, n = c(1, 2, 2, 4), weber = 0.1, weber.achro = 0.1, cores = 1)

## End(Not run)

classify  Identify colour classes in an image for adjacency analyses

descriptions
Classify image pixels into discrete colour classes.

Usage

```r
classify(imgdat, 
  method = c("kMeans", "kMedoids"), 
  kcols = NULL, 
  refID = NULL, 
  interactive = FALSE, 
  plotnew = FALSE, 
  col = "red", 
  cores = getOption("mc.cores", 2L), 
  ...
)
```

Arguments

- `imgdat` (required) image data. Either a single image, or a series of images stored in a list. Preferably the result of `getimg()`.
- `method` methods for image segmentation/classification.
  - 'kMeans': k-means clustering (default)
  - 'kMedoids': k-medoids clustering, using the partitioning-around-medoids ('pam') algorithm for large datasets.
- `kcols` the number of discrete colour classes present in the input image(s). Can be a single integer when only a single image is present, or if `kcols` is identical for all images. When passing a list of images, `kcols` can also be a vector the same length as `imgdat`, or a data.frame with two columns specifying image file names and corresponding `kcols`. This argument can optionally be disregarded when `interactive = TRUE`, and `kcols` will be inferred from the number of selections.
classify

refID

either the numeric index or name of a 'reference' image, for use when passing a list of images. Other images will be k-means classified using centres identified in the single reference image, thus helping to ensure that homologous pattern elements will be reliably classified between images, if so desired.

interactive

interactively specify the colour-category 'centers', for k-means clustering. When TRUE, the user is asked to click a number of points (equal to kcols, if specified, otherwise user-determined) that represent the distinct colours of interest. If a reference image is specified, it will be the only image presented.

plotnew

Should plots be opened in a new window when interactive = TRUE? Defaults to FALSE.

col

the color of the marker points, when interactive = TRUE.

cores

number of cores to be used in parallel processing. If 1, parallel computing will not be used. Defaults to getOption("mc.cores",2L). Not available on Windows.

... additional graphical parameters when interactive = TRUE. Also see graphics::par().

Value

A matrix, or list of matrices, of class rimg containing the colour class classifications ID at each pixel location. The RGB values corresponding to cluster centres (i.e. colour classes) are stored as object attributes.

Note

Since the kmeans process draws on random numbers to find initial cluster centres when interactive = FALSE, use set.seed() if reproducible cluster ID's are desired between runs.

Author(s)

Thomas E. White <thomas.white026@gmail.com>

See Also

stats::kmeans

Examples

# Single image
papilio <- getimg(system.file("testdata/images/papilio.png", package = "pavo"))
papilio_class <- classify(papilio, kcols = 4)

# Multiple images, with interactive classification and a reference image
snakes <- getimg(system.file("testdata/images/snakes", package = "pavo"))
# snakes_class <- classify(snakes, refID = "snake_01", interactive = TRUE)
**coldist**

*Colour distances*

**Description**

Calculates colour distances. When data are the result of `vismodel()`, it applies the receptor-noise model of Vorobyev et al. (1998) to calculate colour distances with noise based on relative photoreceptor densities. It also accepts `colspace()` data in which case unweighted Euclidean distances or Manhattan distances (coc model only) are returned.

**Usage**

```r
coldist(
  modeldata, 
  noise = c("neural", "quantum"),
  subset = NULL,
  achromatic = FALSE,
  qcatch = NULL,
  n = c(1, 2, 2, 4),
  weber = 0.1,
  weber.ref = "longest",
  weber.achro = 0.1
)
```

**Arguments**

- `modeldata` (required) quantum catch colour data. Can be the result from `vismodel()` for noise-weighted Euclidean distances, or `colspace()` for unweighted (typically) Euclidean distances. Data may also be independently calculated quantum catches, in the form of a data frame with columns representing photoreceptors.

- `noise` how the noise will be calculated (ignored for `colspace` objects):

  - `neural` (default): noise is proportional to the Weber fraction and is independent of the intensity of the signal received (i.e. assumes bright conditions).
  - `quantum`: noise is the sum of the neural noise and receptor noise, and is thus proportional to the Weber fraction and inversely proportional to the intensity of the signal received (the quantum catches). Note that the quantum option will only work with objects of class `vismodel`.

- `subset` If only some of the comparisons should be returned, a character vector of length 1 or 2 can be provided, indicating which samples are desired. The subset vector must match the labels of the input samples, but partial matching (and regular expressions) are supported.

- `achromatic` Logical. If `TRUE`, last column of the data frame is used to calculate the achromatic contrast, the form of which will depend on the input data and will be indicated by a message during execution. For noise-weighted distances, noise is based on the Weber fraction given by the argument `weber.achro`. 

  - `weber` Weber fraction.
  - `weber.ref` Reference Weber fraction.
  - `weber.achro` Weber fraction for achromatic contrast.
if the object is of class `vismodel` or `colspace`, this argument is ignored. If the object is a data frame of quantal catches from another source, this argument is used to specify what type of quantum catch is being used, so that the noise can be calculated accordingly: * $Q_i$: Quantum catch for each photoreceptor * $f_i$: Quantum catch according to Fechner’s law (the signal of the receptor channel is proportional to the logarithm of the quantum catch)

Quantum catch according to Fechner’s law (the signal of the receptor channel is proportional to the logarithm of the quantum catch)

$\text{n}$

photoreceptor densities for the cones used in visual modeling. must have same length as number of columns (excluding achromatic receptor if used; defaults to the Pekin robin *Leiothrix lutea* densities: `c(1,2,2,4)`). Ignored for `colspace` objects.

The Weber fraction(s) to be used (often also referred to as receptor noise, or $e$). The noise-to-signal ratio $v$ is unknown, and therefore must be calculated based on the empirically estimated Weber fraction of one or (more rarely) all the cone classes. When noise is only known for one receptor, as is typical, $v$ is then applied to estimate the Weber fraction of the other cones. By default, the value of 0.1 is used (the empirically estimated value for the LWS cone from *Leiothrix lutea*). See Olsson et al. 2017 for a review of published values in the literature. Ignored for `colspace` objects.

the cone class used to obtain the empirical estimate of the Weber fraction used for the `weber` argument, if a single value is specified. By default, n4 is used, representing the LWS cone for *Leiothrix lutea*. Ignored for `colspace` objects.

the Weber fraction to be used to calculate achromatic contrast, when `achromatic = TRUE`. Defaults to 0.1. Ignored for `colspace` objects.

Value

A data frame containing up to 4 columns. The first two (patch1, patch2) refer to the two colors being contrasted; $d_S$ is the chromatic contrast (delta S) and $d_L$ is the achromatic contrast (delta L). Units of $d_S$ JND’s in the receptor-noise model, unweighted Euclidean distances in colorspace models, and Manhattan distances in the colour-opponent-coding space. Units of $d_L$ vary, and are either simple contrast, Weber contrast, or Michelson contrast, as indicated by the output message.

Note on previous versions

Generic di- tri- and tetra-chromatic `colspace` objects were previously passed through the receptor-noise limited model to return noise-weighted Euclidean distances. This behaviour has been amended, and generic spaces now return unweighted Euclidean distances. Equivalent results to the former behaviour can be attained by sending the results of `vismodel()` directly to `coldist()`, as previously, which also offers greater flexibility and reliability. Thus `coldist()` now returns unweighted Euclidean distances for `colspace` objects (with the exception of Manhattan distances for the coc space), and noise-weighted Euclidean distances for `vismodel` objects.

Author(s)

Rafael Maia <rm72@zips.uakron.edu>
References


Examples

```r
## Not run:
# Dichromat
data(flowers)
vis.flowers <- vismodel(flowers, visual = "canis", relative = FALSE)
didist.flowers <- coldist(vis.flowers, n = c(1, 2))

# Trichromat
vis.flowers <- vismodel(flowers, visual = "apis", relative = FALSE)
tridist.flowers <- coldist(vis.flowers, n = c(1, 2, 1))

# Trichromat, colour-hexagon model (euclidean distances)
vis.flowers <- vismodel(flowers, visual = "apis", qcatch = "Ei",
relative = FALSE, vonkries = TRUE, achro = "1", bkg = "green"
)
hex.flowers <- colspace(vis.flowers, space = "hexagon")
hexdist.flowers <- coldist(hex.flowers)

# Trichromat, colour-opponent-coding model (manhattan distances)
vis.flowers <- vismodel(flowers, visual = "apis", qcatch = "Ei",
relative = FALSE, vonkries = TRUE)
occ.flowers <- colspace(vis.flowers, space = "coc")
occdist.flowers <- coldist(coc.flowers)

# Tetrachromat
data(sicalis)
vis.sicalis <- vismodel(sicalis, visual = "avg.uv", relative = FALSE)
tetradist.sicalis.n <- coldist(vis.sicalis)

## End(Not run)
```
Model spectra in a colorspace

Description
Models reflectance spectra in a colorspace. For information on plotting arguments and graphical parameters, see `plot.colspace()`.

Usage

colspace(
  vismodeldata,
  space = c("auto", "di", "tri", "tcs", "hexagon", "coc", "categorical", "ciexyz",
            "cielab", "cielch", "segment"),
  qcatch = NULL
)

Arguments

- **vismodeldata** *(required)* quantum catch color data. Can be either the result from `vismodel()` or independently calculated data (in the form of a data frame with columns representing quantum catches).

- **space** Which colorspace/model to use. Options are:
  - auto: if data is a result from `vismodel()`, applies di, tri or tcs if input visual model had two, three or four cones, respectively.
  - di: dichromatic colourspace. See `dispace()` for details. *(plotting arguments)*
  - tri: trichromatic colourspace (i.e. Maxwell triangle). See `trispace()` for details. *(plotting arguments)*
  - tcs: tetrahedral colourspace. See `tcspace()` for details. *(plotting arguments)*
  - hexagon: the trichromatic colour-hexagon of Chittka (1992). See `hexagon()` for details. *(plotting arguments)*
  - categorical: the tetrachromatic categorical fly-model of Troje (1993). See `categorical()` for details. *(plotting arguments)*
  - ciexyz: CIEXYZ space. See `cie()` for details. *(plotting arguments)*
  - cielab: CIELAB space. See `cie()` for details. *(plotting arguments)*
  - cielch: CIELCh space. See `cie()` for details. *(plotting arguments)*

- **qcatch** Which quantal catch metric is being inputted. Only used when input data is NOT an output from `vismodel()`. Must be Qi, fi or Ei.
Author(s)

Rafael Maia <rm72@zips.uakron.edu>
Thomas White <thomas.white026@gmail.com>

References


Examples

data(flowers)

# A dichromat in a segment colourspace
di.flowers <- colspace(vis.flowers, space = "di")

# Honeybee in the colour hexagon
hex.flowers <- colspace(vis.flowers, space = "hexagon")

# A trichromat in a Maxwell triangle
tri.flowers <- colspace(vis.flowers, space = "tri")
plot(tri.flowers)
# A tetrachromat in a tetrahedral colorspace
vis.flowers <- vismodel(flowers, visual = "bluetit")
tcs.flowers <- colspace(vis.flowers, space = "tcs")

# A housefly in the categorical colorspace
vis.flowers <- vismodel(flowers, visual = "musca", achro = "md.r1")
cat.flowers <- colspace(vis.flowers, space = "categorical")

---

### explorespec

**Plot spectral curves**

**Description**

Plots one or multiple spectral curves in the same graph to rapidly compare groups of spectra.

**Usage**

```r
explorespec(
  rspecdata,
  by = NULL,
  scale = c("equal", "free"),
  legpos = "topright",
  ...
)
```

**Arguments**

- `rspecdata` (required) a data frame, possibly of class `rspec`, which contains a column containing a wavelength range, named 'wl', and spectra data in remaining columns.
- `by` number of spectra to include in each graph (defaults to 1)
- `scale` defines how the y-axis should be scaled. "free": panels can vary in the range of the y-axis; "equal": all panels have the y-axis with the same range.
- `legpos` legend position control. Either a vector containing x and y coordinates or a single keyword from the list: "bottomright", "bottom", "bottomleft", "left", "topleft", "top", "topright", "right" and "center".
- `...` additional parameters to be passed to plot

**Value**

Spectral curve plots

**Note**

Number of plots presented per page depends on the number of graphs produced.
Author(s)

Pierre-Paul Bitton <bittonp@uwindsor.ca>

Examples

data(sicalis)
explorespec(sicalis, 3)
explorespec(sicalis, 3, ylim = c(0, 100), legpos = c(500, 80))

flowers

Reflectance spectra from a suite of native Australian flowers, collected around Cairns, Queensland.

Description

dataset containing reflectance measurements from 36 native Australian angiosperm species, indicated by column names.

Author(s)

Thomas White <thomas.white026@gmail.com>

References


getimg

Import image data

Description

Finds and imports PNG, JPEG, and/or BMP images.
Usage

getimg(
  imgpath = getwd(),
  subdir = FALSE,
  subdir.names = FALSE,
  max.size = 1,
  cores
)

Arguments

imgpath (required) either the full file-path or URL to an image (including extension), or
the path to a folder in which multiple image files are located. Mixed file formats
within a folder are accepted.

subdir should subdirectories within the imgpath folder be included in the search? (de-
defaults to FALSE).

subdir.names should subdirectory path be included in the name of the images? (defaults to
FALSE).

max.size maximum size of all images to be allowed in memory, in GB. Defaults to 1.

cores deprecated argument.

Value

a image, or list of images, of class rimg, for use in further pavo functions.

Author(s)

Thomas E. White <thomas.white026@gmail.com>

Examples

# Single image
papilio <- getimg(system.file("testdata/images/papilio.png", package = "pavo"))

# Multiple images
snakes <- getimg(system.file("testdata/images/snakes", package = "pavo"))

Description

Finds and imports spectra files from a folder. Currently works for reflectance files generated in
Ocean Optics SpectraSuite (USB2000, USB4000 and Jaz spectrometers), CRAIC software (after
exporting) and Avantes (before or after exporting).
getspec

Usage

getspec(
  where = getwd(),
  ext = "txt",
  lim = c(300, 700),
  decimal = ".",
  sep = NULL,
  subdir = FALSE,
  subdir.names = FALSE,
  cores = getOption("mc.cores", 2L),
  ignore.case = TRUE
)

Arguments

where (required) folder in which files are located.
ext file extension to be searched for, without the "." (defaults to "txt").
lim a vector with two numbers determining the wavelength limits to be considered (defaults to 300 and 700).
decimal character to be used to identify decimal plates (defaults to ".").
sep column delimiting characters to be considered in addition to the default (which are: tab, space, and ";;").
subdir should subdirectories within the where folder be included in the search? (defaults to FALSE).
subdir.names should subdirectory path be included in the name of the spectra? (defaults to FALSE).
cores Number of cores to be used. If greater than 1, import will use parallel processing (not available in Windows).
ignore.case Logical. Should the extension search be case insensitive? (defaults to "TRUE")

Value

A data frame, of class rspec, containing individual imported spectral files as columns. Reflectance values are interpolated to the nearest wavelength integer.

Author(s)

Rafael Maia <rm72@zips.uakron.edu>
Hugo Gruson <hugo.gruson+R@normalesup.org>

References

Examples
rspecdata <- getspec(system.file("testdata", package = "pavo"), ext = "ttt", lim = c(400, 700))
head(rspecdata)

img_conversion

Convert images between class rimg and cimg or magick-image

Description
Conveniently convert single objects of class rimg to class cimg (from the package imager) or magick-image (from the package magick), both of which contains a suite of useful image-processing capabilities.

Usage
rimg2cimg(image)
rimg2magick(image)

Arguments
image an object of class rimg

Value
an image of the specified class

Note
Attributes (e.g. scales, color-classes) will not be preserved following conversion from class rimg, so it’s best to use early in the analysis workflow.

Author(s)
Thomas E. White <thomas.white026@gmail.com>
Hugo Gruson <hugo.gruson+R@normalesup.org>

Examples
papilio <- getimg(system.file("testdata/images/papilio.png", package = "pavo"))

# From class rimg to cimg
papilio_cimg <- rimg2cimg(papilio)
class(papilio_cimg)

# From class rimg to magick-image
papilio_magick <- rimg2magick(papilio)
class(papilio_magick)
is.colspace

\textit{Test if object is of class 'colspace'}

\begin{description}
\item[Description] Test if object is of class 'colspace'
\item[Usage] \texttt{is.colspace(object)}
\item[Arguments] \texttt{object} an R object
\item[Value] a logical value indicating whether the object is of class colspace
\end{description}

irrad2flux \hspace{1cm} \textit{Converts between irradiance and photon (quantum) flux}

\begin{description}
\item[Description] Some spectrometers will give illuminant values in units of irradiance ($\mu$Watt.cm$^{-2}$), but physiological models require illuminants in units of photon (quantum) flux ($\mu$mol.s$^{-1}$.m$^{-2}$). The functions \texttt{irrad2flux} and \texttt{flux2irrad} allows for easy conversion of rspec objects between these units.
\item[Usage] \texttt{irrad2flux(rspecdata)} \texttt{flux2irrad(rspecdata)}
\item[Arguments] \texttt{rspecdata} (required) a rspec object containing illuminant values.
\item[Value] a converted rspec object.
\item[Author(s)] Rafael Maia <rm72@zips.uakron.edu>
\end{description}
is.vismodel

Test if object is of class 'vismodel'

Description
Test if object is of class 'vismodel'

Usage
is.vismodel(object)

Arguments
object an R object

Value
a logical value indicating whether the object is of class vismodel.

See Also
vismodel()
Arguments

coldistres  (required) the output from a coldist() call.
center    logical indicating if the data should be centered on its centroid (defaults to TRUE).
rotate    logical indicating if the data should be rotated (defaults to TRUE).
rotcenter should the vectors for rotation be centered in the achromatic center ("achro") or the data centroid ("mean", the default)?
ref1      the cone to be used as a the first reference. May be NULL (for no first rotation in the 3-dimensional case) or must match name in the original data that was used for coldist(). Defaults to 'l'.
ref2      the cone to be used as a the second reference. May be NULL (for no first rotation in the 3-dimensional case) or must match name in the original data that was used for coldist(). Defaults to 'u'. (only used if data has 3 dimensions).
axis1     A vector of length 3 composed of 0's and 1's, with 1's representing the axes (x,y,z) to rotate around. Defaults to c(1,1,0), such that the rotation aligns with the xy plane (only used if data has 2 or 3 dimensions). Ignored if ref1 is NULL (in 3-dimensional case only)
axis2     A vector of length 3 composed of 0's and 1's, with 1's representing the axes (x,y,z) to rotate around. Defaults to c(0,0,1), such that the rotation aligns with the z axis (only used if data has 3 dimensions). Ignored if ref2 is NULL (in 3-dimensional case only)

Author(s)

Rafael Maia <rm72@zips.uakron.edu>

References


Examples

data(flowers)
vis.flowers <- vismodel(flowers)
cd.flowers <- coldist(vis.flowers)
jnd2xyz(cd.flowers)
**legendtetra**  
*Add legend to a static tetrahedral colourspace*

**Description**

Adds a legend to a static tetrahedral colourspace plot.

**Usage**

`legendtetra(x = 0.8, y = 1.2, ...)`

**Arguments**

- `x`: position of the legend relative to plot limits (usually a value between 0 and 1, but because of the perspective distortion, values greater than 1 are possible)
- `y`: position of the legend relative to plot limits (usually a value between 0 and 1, but because of the perspective distortion, values greater than 1 are possible)
- `...`: additional arguments passed to `legend()`.

**Value**

`legendtetra()` adds a legend to a static tetrahedral colourspace plot. for additional information on which arguments are necessary and how they are used, see `legend()`.

**Author(s)**

Rafael Maia `<rm72@zips.uakron.edu>`

**Examples**

```r
data(sicalis)
vis_sicalis <- vismodel(sicalis)
tcs_sicalis <- colspace(vis_sicalis)

cols <- c("#1B9E77", "#D95F02", "#7570B3")
plot(tcs_sicalis, col = cols)
legendtetra(
  legend = c("Crown", "Throat", "Breast"),
  col = cols, pch = 16
)
```
merge.rspec

Merge two rspec objects

Description

Merges two rspec or data.frame objects into a single rspec object.

Usage

## S3 method for class 'rspec'
merge(x, y, ...)

Arguments

x, y
(required) rspec objects to merge.

... additional class arguments.

Value

an object of class rspec for use with pavo functions. Will use by = "wl" if unspecified, or automatically append wl to the by argument if one is specified.

Author(s)

Chad Eliason <cme16@zips.uakron.edu>

See Also

as.rspec(), aggspec()

Examples

# Load and split dataset into 2 sections
data(teal)
teal1 <- teal[, c(1, 3:5)]
teal2 <- teal[, c(1, 2, 6:12)]
teal.mer <- merge(teal1, teal2, by = "wl")
head(teal.mer)
par(mfrow = c(1, 2))
plot(teal.mer)
plot(teal)
Description

Calculates height, location and width of peak at the reflectance midpoint (FWHM). Note: bounds should be set wide enough to incorporate all minima in spectra. Smoothing spectra using `procspec()` is also recommended.

Usage

```r
peakshape(
  rspecdata,
  select = NULL,
  lim = NULL,
  plot = TRUE,
  ask = FALSE,
  absolute.min = FALSE,
  ...
)
```

Arguments

- `rspecdata` (required) a data frame, possibly of class `rspec`, which contains a column containing a wavelength range, named `wl`, and spectra data in remaining columns.
- `select` specification of which spectra to plot. Can be a numeric vector or factor (e.g., `sex=="male"`).
- `lim` a vector specifying the wavelength range to analyze.
- `plot` logical. Should plots indicating calculated parameters be returned? (Defaults to `TRUE`).
- `ask` logical, specifies whether user input needed to plot multiple plots when number of spectra to analyze is greater than 1 (defaults to `FALSE`).
- `absolute.min` logical. If `TRUE`, full width at half maximum will be calculated using the absolute minimum reflectance of the spectrum, even if that value falls outside the range specified by `lim`. (defaults to `FALSE`)
- `...` additional arguments to be passed to `plot`.

Value

A data frame containing column names (id); peak height (max value, B3), location (hue, H1) and full width at half maximum (FWHM), as well as half widths on left (HWHM.l) and right side of peak (HWHM.r). Incl.min column indicates whether user-defined bounds incorporate the actual minima of the spectra. Function will return a warning if not.
Author(s)
Chad Eliason <cme16@zips.uakron.edu>
Rafael Maia <rm72@zips.uakron.edu>
Hugo Gruson <hugo.gruson+R@normalesup.org>

See Also
procspec()

Examples
data(teal)

peakshape(teal, select = 3)
peakshape(teal, select = 10)

# Use wavelength bounds to narrow in on peak of interest
peakshape(teal, select = 10, lim = c(400, 550))

plot.colspace  
Plot spectra in a colourspace

Description
Plots reflectance spectra in the appropriate colourspace.

Usage
## S3 method for class 'colspace'
plot(x, ...)

Arguments
x  (required) an object of class colspace.
...
additional graphical options, which vary by modeled space. Refer to their individual documentation:
  • diplot(): dichromat space
  • triplot(): trichromat space
  • tetraplot(): tetrahedral space
  • catplot(): categorical space
  • hexplo(): colour hexagon
  • cocplot(): colour-opponent-coding space
  • cieplot(): cie spaces
  • segplot(): segment analysis space
  • jndplot(): perceptual, ‘noise corrected’ space (for the results of jnd2xyz())
Also see par().
Value

A colourspace plot appropriate to the input data.

Author(s)

Rafael Maia <rm72@zips.uakron.edu>
Thomas White <thomas.white026@gmail.com>
Chad Eliason <cme16@zips.uakron.edu>

References


See Also

plot()

Examples

data(flowers)
data(sicalis)

# Dichromat
vis.flowers <- vismodel(flowers, visual = "canis")
di.flowers <- colspace(vis.flowers, space = "di")
plot(di.flowers)
# Colour hexagon
vis.flowers <- vismodel(flowers,
  visual = "apis", qcatch = "Ei", relative = FALSE,
  vonkries = TRUE, achro = "l", bkg = "green"
) hex.flowers <- colspace(vis.flowers, space = "hexagon") plot(hex.flowers, sectors = "coarse")

# Tetrahedron (static)
vis.sicalis <- vismodel(sicalis, visual = "avg.uv") tcs.sicalis <- colspace(vis.sicalis, space = "tcs") plot(tcs.sicalis)
## Not run:
# Tetrahedron (interactive)
vis.sicalis <- vismodel(sicalis, visual = "avg.uv") tcs.sicalis <- colspace(vis.sicalis, space = "tcs") tcsplot(tcs.sicalis, size = 0.005)

## Add points to interactive tetrahedron
patch <- rep(c("C", "T", "B"), 7) tcs.crown <- subset(tcs.sicalis, "C") tcs.breast <- subset(tcs.sicalis, "B") tcsplot(tcs.crown, col = "blue") tcspoints(tcs.breast, col = "red")

## Plot convex hull in interactive tetrahedron
tcsplot(tcs.sicalis, col = "blue", size = 0.005) tcsvol(tcs.sicalis)
## End(Not run)

---

**plot.rimg**

Plot unprocessed or colour-classified images

### Description

Plot unprocessed or colour-classified image data. If the images are in a list, they will be stepped through one by one.

### Usage

```r
## S3 method for class 'rimg'
plot(x, axes = TRUE, col = NULL, ...)
```

### Arguments

- **x** (required) an image of class rimg, or list thereof.
- **axes** should axes be drawn? (defaults to TRUE)
col

optional vector of colours when plotting colour-classified images. Defaults to the mean RGB values of the k-means centres (i.e. the average ‘original’ colours). additional graphical parameters. Also see `par()`.

Value

an image plot.

Author(s)

Thomas E. White <thomas.white026@gmail.com>

Examples

```r
## Not run:
papilio <- getimg(system.file("testdata/images/papilio.png", package = "pavo"))
plot(papilio)
papilio_class <- classify(papilio, kcols = 4)
plot(papilio_class)

# Multiple images
snakes <- getimg(system.file("testdata/images/snakes", package = "pavo"))
plot(snakes)
snakes_class <- classify(snakes, kcols = 3)
plot(snakes_class)

## End(Not run)
```

Description

Plots reflectance spectra in different arrangements.

Usage

```r
## S3 method for class 'rspec'
plot(
  x,
  select = NULL,
  type = c("overlay", "stack", "heatmap"),
  varying = NULL,
  n = 100,
  labels.stack = NULL,
  wl.guide = TRUE,
  ...
)
```
plotsmooth

Plot loess smoothed curves

Args

x
(required) a data frame, possibly an object of class rspec, with a column with wavelength data, named 'wl', and the remaining column containing spectra to plot.

select
specification of which spectra to plot. Can be a numeric vector or factor (e.g., sex=="male")

type
what type of plot should be drawn. Possibilities are:
  • overlay (default) for plotting multiple spectra in a single panel with a common y-axis.
  • stack for plotting multiple spectra in a vertical arrangement.
  • heatmap for plotting reflectance values by wavelength and a third variable (varying).

varying
a numeric vector giving values for y-axis in type = "heatmap".

n
number of bins with which to interpolate colors and varying for the heatplot.

labels.stack
a vector of labels for the stacked spectra when using type = "stack". Defaults to the numeric column ID’s.

wl.guide
logical determining whether visible light spectrum should be added to the x-axis.

...
additional arguments passed to plot() (or image() for "heatmap").

Author(s)

Chad Eliason <cme16@zips.uakron.edu>
Thomas White <thomas.white026@gmail.com>

See Also

spec2rgb(), image(), plot()

Examples

data(teal)
plot(teal, type = "overlay")
plot(teal, type = "stack")
plot(teal, type = "heatmap")
Usage

```r
plotsmooth(
  rspecdata,
  minsmooth = 0.05,
  maxsmooth = 0.2,
  curves = 5,
  specnum = "ALL",
  ask = TRUE
)
```

Arguments

- `rspecdata` (required) a data frame, possibly of class `rspec`, which contains a column containing a wavelength range, named 'wl', and spectra data in remaining columns.
- `minsmooth` the minimum f value of the loess function to visualize (defaults to 0.05).
- `maxsmooth` the maximum f value of the loess function to visualize (defaults to 0.20).
- `curves` the number of curves to display on the same plot (defaults to 5).
- `specnum` the number of spectral curves, from the data frame, to visualize (defaults to ALL).
- `ask` logical. if TRUE, asks for user input before changing plot pages

Value

Series of plot with curves processed with varying level of loess smoothing

Author(s)

Pierre-Paul Bitton <bittonp@uwindsor.ca>

See Also

`procspec()`

Examples

```r
data(sicalis)
plotsmooth(sicalis, minsmooth = 0.05, maxsmooth = 0.1, curves = 7, specnum = 6)
```

points.colspace  Plot points in a colourspace

Description

Add points to a colourspace plot
### procimg

**Usage**

```r
## S3 method for class 'colspace'
points(x, ...)
```

**Arguments**

- `x` *(required)* an object of class `colspace`.
- `...` additional graphical options. See `par()`.

**Value**

`points.colspace` adds points to a colorspace plot. When `space = 'tcs'`, it creates 3D points in a tetrahedral color space plot using functions of the package `rgl`, based on openGL capabilities.

**Author(s)**

Rafael Maia &lt;rm72@zips.uakron.edu&gt;

Thomas White &lt;thomas.white026@gmail.com&gt;

---

### procimg

**Process images**

**Description**

Specify scales, resize, and/or define focal objects within images.

**Usage**

```r
procimg(
  image,
  resize = NULL,
  rotate = NULL,
  scaledist = NULL,
  outline = FALSE,
  reclass = NULL,
  smooth = FALSE,
  iterations = 1L,
  col = "red",
  plotnew = FALSE,
  ...
)
```
Arguments

- **image** (required) image data. Either a single image array, or a number of images stored in a list. Preferably the result of `getimg()`.
- **resize** an integer specifying a percentage for resizing images, if so desired. E.g. 50 to half the size of an image, 200 to double it.
- **rotate** an integer specifying the angle of image rotation, in degrees. Images are rotated around the centre, and linearly interpolated.
- **scaledist** an integer, or numeric vector equal in length to the number of images, specifying the length of the scale in the image(s). Image(s) will then be presented, and the user asked to select either end of the scale corresponding to the input value.
- **outline** interactively specify the focal object in an image by clicking around its outline. The xy-coordinates of the resulting closed polygon are saved as an attribute, for use in generating a masking layer & separating animals/plants from backgrounds in further analyses. This is particularly useful when backgrounds are complex, such as in natural settings.
- **reclass** interactively specify an area on a colour-classified image that is to be reclassified as the numeric value provided. e.g. when `reclass = 1`, the user will be asked to select a polygon on the image, within which all colour-category values will be changes to 1.
- **smooth** should the polygon specified when `outline = TRUE` be smoothed using Chaikin’s corner-cutting algorithm? Defaults to `FALSE`.
- **iterations** the number of smoothing iterations, when `smooth = TRUE`. Defaults to 1.
- **col** the color of the marker points and/or line, when using interactive options.
- **plotnew** should plots be opened in a new window? Defaults to `FALSE`.
- `...` additional graphical parameters. Also see `par()`.

Value

An image, or list of images, for use in further `pavo` functions.

Author(s)

Thomas E. White <thomas.white026@gmail.com>

References


Examples

```r
## Not run:
# Single image
papilio <- getimg(system.file("testdata/images/papilio.png", package = "pavo"))
papilio <- procimg(papilio, scaledist = 10)
```
# Assign individual scales to each image, after slightly reducing their size.
snakes <- getimg(system.file("testdata/images/snakes", package = "pavo"))
snakes <- procimg(snakes, scaledist = c(10, 14), resize = 90)

## End(Not run)

## procspec

### Process spectra

#### Description
Applies normalization and/or smoothing to spectra for further analysis or plotting.

#### Usage

```r
textprocspec(  
rspecdata,  
opt = c("none", "smooth", "maximum", "minimum", "bin", "sum", "center"),  
fixneg = c("none", "addmin", "zero"),  
span = 0.25,  
bins = 20  
)
```

#### Arguments

- **rspecdata** (required) a data frame, possibly of class `rspec`, which contains a column containing a wavelength range, named 'wl', and spectra data in remaining columns.
- **opt** what type of processing options to apply. User can select multiple options by providing a vector. Possibilities are:
  - "none" does not perform any processing (default).
  - "smooth" applies LOESS smoothing to each spectrum using `loess.smooth()`. Optimal smoothing parameter can be assessed by using `plotsmooth()`.
  - "minimum" subtracts the minimum from each individual spectra.
  - "maximum" divides each spectrum by its maximum value.
  - "sum" divides each spectrum by summed values.
  - "bin" bins each spectrum into the specified number of bins. `bins` argument must be set.
  - "center" centers individual spectra by subtracting mean reflectance from all values.
- **fixneg** how to handle negative values. Possibilities are:
  - "none" does not perform negative value correction (default).
  - "zero" sets all negative values to zero.
  - "addmin" adds the absolute value of the maximally negative values of each spectra to the reflectance at all other wavelengths (setting the minimum value to zero, but scaling other values accordingly).
projplot

span
sets the smoothing parameter used by `loess.smooth()`.

bins
sets the number of equally sized wavelength bins for opt = "bin".

Value

A data frame of class `rspec` with the processed data.

Author(s)

Chad Eliason `<cme16@zips.uakron.edu>`

References


See Also

`loess.smooth()`, `plotsmooth()`

Examples

```r
data(teal)
plot(teal, select = 10)

# Smooth data to remove noise
teal.sm <- procspec(teal, opt = "smooth", span = 0.25)
plot(teal.sm, select = 10)

# Normalize to max of unity
teal.max <- procspec(teal, opt = c("max"))
plot(teal.max, select = 10)
```

projplot

2D projection of a tetrahedral colourspace

Description

Produces a 2D projection plot of points in a tetrahedral colour space

Adds points to a tetrahedral colorspace projection
Usage

projplot(tcsdata, ...)
projpoints(tcsdata, ...)

Arguments

tcsdata (required) tetrahedral color space coordinates, possibly a result from \texttt{colspace()}, containing values for the 'h.theta' and 'h.phi' coordinates as columns (labeled as such).

... additional parameters to be passed to the plotting of data points.

Value

\texttt{projplot()} creates a 2D plot of color points projected from the tetrahedron to its encapsulating sphere, and is ideal to visualize differences in hue.

\texttt{projpoints()} creates points in a projection color space plot produced by \texttt{projplot()}.

Note

\texttt{projplot()} uses the Mollweide projection, and not the Robinson projection, which has been used in the past. Among other advantages, the Mollweide projection preserves area relationships within latitudes without distortion.

Author(s)
Rafael Maia <rm72@zips.uakron.edu>

References


Examples

data(sicalis)
vis.sicalis <- vismodel(sicalis, visual = "avg.uv")
tcs.sicalis <- colspace(vis.sicalis, space = "tcs")
projplot(tcs.sicalis, pch = 16, col = setNames(rep(seq_len(3), 7), rep(c("C", "T", "B"), 7)))
sensdata

Retrieve or plot in-built spectral sensitivity data

Description

Retrieve (as an rspec object) or plot pavo’s in-built spectral sensitivity data.

Usage

sensdata(
  visual = c("none", "all", "avg.uv", "avg.v", "bluetit", "ctenophorus", "star", 
             "pfowl", "apis", "canis", "cie2", "cie10", "musca", "habronattus", "rhinecanthus"),
  achromatic = c("none", "all", "bt.dc", "ch.dc", "st.dc", "md.r1", "ra.dc", "cf.r"),
  illum = c("none", "all", "bluesky", "D65", "forestshade"),
  trans = c("none", "all", "bluetit", "blackbird"),
  bkg = c("none", "all", "green"),
  plot = FALSE,
  ...
)

Arguments

visual visual systems. Options are:
  • "none": no visual sensitivity data.
  • "all": all visual sensitivity data.
  • "apis": Honeybee Apis mellifera visual system.
  • "avg.uv": average avian UV system.
  • "avg.v": average avian V system.
  • "bluetit": Blue tit Cyanistes caeruleus visual system.
  • "canis": Canid Canis familiaris visual system.
  • "cie2": 2-degree colour matching functions for CIE models of human colour vision. Functions are linear transformations of the 2-degree cone fundamentals of Stockman & Sharpe (2000), as ratified by the CIE (2006).
  • "cie10": 10-degree colour matching functions for CIE models of human colour vision. Functions are linear transformations of the 10-degree cone fundamentals of Stockman & Sharpe (2000), as ratified by the CIE (2006).
  • "ctenophorus": Ornate dragon lizard Ctenophorus ornatus.
  • "musca": Housefly Musca domestica visual system.
  • "pfowl": Peafowl Pavo cristatus visual system.
  • "star": Starling Sturnus vulgaris visual system.
  • "habronattus": Jumping spider Habronattus pyrrithrix.
  • "rhinecanthus": Triggerfish Rhinecanthus aculeatus.

achromatic the sensitivity data used to calculate luminance (achromatic) receptor stimulation. Options are:
sensdata

- "none": no achromatic sensitivity data.
- "all": all achromatic sensitivity data.
- "bt.dc": Blue tit *Cyanistes caeruleus* double cone.
- "ch.dc": Chicken *Gallus gallus* double cone.
- "st.dc": Starling *Sturnus vulgaris* double cone.
- "cf.r": Canid *Canis familiaris* rod
- "md.r1": Housefly *Musca domestica* R1-6 photoreceptor.
- "ra.dc": Triggerfish *Rhinocanthus aculeatus* double cone.

**illum**

illuminants. Options are:

- "none": no illuminant data.
- "all": all background spectral data.
- "bluesky": open blue sky.
- "D65": standard daylight.
- "forestshade": forest shade.

**trans**

Ocular transmission data. Options are:

- "none": no transmission data.
- "all": all transmission data.
- "bluetit": blue tit *Cyanistes caeruleus* ocular transmission (from Hart et al. 2000).
- "blackbird": blackbird *Turdus merula* ocular transmission (from Hart et al. 2000).

**bkg**

background spectra. Options are:

- "none": no background spectral data.
- "all": all background spectral data.
- "green": green foliage.

**plot**

should the spectral data be plotted, or returned instead (defaults to FALSE)?

... additional graphical options passed to `plot.rspec()` when plot = TRUE.

**Value**

An object of class `rspec` (when plot = FALSE), containing a wavelength column "wl" and spectral data binned at 1 nm intervals from 300-700 nm.

**Author(s)**

Thomas White <thomas.white026@gmail.com>

Rafael Maia <rm72@zips.uakron.edu>

**Examples**

```r
# Plot the honeybee's receptors
sensdata(visual = "apis", ylab = "Absorbance", plot = TRUE)

# Plot the average UV vs V avian receptors
```
### sensmodel

**Description**

Models spectral sensitivity (with oil droplets; optional) based on peak cone sensitivity according to the models of Govardovskii et al. (2000) and Hart & Vorobyev (2005).

**Usage**

```r
sensmodel(
  peaksens,
  range = c(300, 700),
  lambdacut = NULL,
  Bmid = NULL,
  oiltype = NULL,
  beta = TRUE,
  om = NULL,
  integrate = TRUE
)
```

**Arguments**

- **peaksens** (required) a vector with peak sensitivities for the cones to model.
- **range** a vector of length 2 for the range over which to calculate the spectral sensitivities (defaults to 300nm to 700nm).
- **lambdacut** a vector of same length as peaksens that lists the cut-off wavelength value for oil droplets. Needs either Bmid or oiltype to also be entered. See Hart and Vorobyev (2005).
- **Bmid** a vector of same length as peaksens that lists the gradient of line tangent to the absorbance spectrum of the oil droplets. See Hart and Vorobyev (2005).
- **oiltype** a list of same length as peaksens that lists the oil droplet types (currently accepts only "T", "C", "Y", "R", "P") when Bmid is not known. Calculates Bmid based on the regression equations found in Hart and Vorobyev (2005).
- **beta** logical. If TRUE the sensitivities will include the beta peak See Govardovskii et al.(2000) (defaults to TRUE).
- **om** a vector of same length as range1-range2 that contains ocular media transmission data. If included, cone sensitivity will be corrected for ocular media transmission. Currently accepts "bird" using values from Hart et al. (2005), or user-defined values.
integrate logical. If TRUE, each curve is transformed to have a total area under the curve of 1 (best for visual models; defaults to TRUE). NOTE: integration is applied before any effects of ocular media are considered, for compatibility with visual model procedures.

Value
A data frame of class rspec containing each cone model as a column.

Author(s)
Pierre-Paul Bitton <bittonp@uwindsor.ca>, Chad Eliason <cme16@zips.uakron.edu>

References

Examples
# Blue tit visual system based on Hart et al (2000)
bluesens <- sensmodel(c(371, 448, 502, 563),
    beta = FALSE,
    lambdacut = c(330, 413, 507, 572),
    oiltype = c("T", "C", "Y", "R"), om = TRUE
)
# Danio aequipinnatus based on Govardovskii et al. (2000)
daniosens <- sensmodel(c(357, 411, 477, 569))

sicalis
Spectral curves from three body regions of stripe-tailed yellow finch (Sicalis citrina) males

Description
dataset containing reflectance measurements from 3 body parts ("C": crown, "B": breast, "T": throat) from seven male stripe-tailed yellow finches

Author(s)
Rafael Maia <rm72@zips.uakron.edu>
Description

Calculates rgb values from spectra based on human colour matching functions.

Usage

spec2rgb(rspecdata, alpha = 1)

Arguments

rspecdata (required) a data frame, possibly of class rspec, which contains a column containing a wavelength range, named 'wl', and spectra data in remaining columns.

alpha alpha value to use for colours (defaults to 1, opaque).

Value

A character vector consisting of hexadecimal colour values for passing to further plotting functions.

Author(s)

Hugo Gruson <hugo.gruson+R@normalesup.org>

Chad Eliason <cme16@zips.uakron.edu>

References


Examples

data(teal)
spec2rgb(teal)

# Plot data using estimated perceived colour
plot(teal, col = spec2rgb(teal), type = "overlay")
subset.rspec  Subset rspec, vismodel, and colspace objects

Description

Subsets various object types based on a given vector or grep partial matching of data names.

Usage

## S3 method for class 'rspec'
subset(x, subset, ...)

## S3 method for class 'colspace'
subset(x, subset, ...)

## S3 method for class 'vismodel'
subset(x, subset, ...)

Arguments

x (required) an object of class rspec, vismodel, or colspace, containing spectra, visual model output or colourspace data to subset.
subset a string used for partial matching of observations.
... additional attributes passed to grep. Ignored if subset is logical.

Value

a subsetted object of the same class as the input object.

Note

if more than one value is given to subset, any spectra that matches either condition will be included. It’s a union, not an intersect.

Author(s)

Chad Eliason <cme16@zips.uakron.edu>

Examples

data(sicalis)
vis.sicalis <- vismodel(sicalis)
tcs.sicalis <- colspace(vis.sicalis, space = "tcs")

# Subset all 'crown' patches (C in file names)
head(subset(sicalis, "C"))
head(subset(sicalis, c("B", "C")))
head(subset(sicalis, "T", invert = TRUE))
subset(vis.sicalis, "C")
subset(tcs.sicalis, "C")[, seq_len(5)]

## Colourspace data summary

### Description

Returns the attributes of colspace objects.

### Usage

```r
## S3 method for class 'colspace'
summary(object, by = NULL, ...)
```

### Arguments

- `object` (required) a colspace object.
- `by` when the input is in tcs colourspace, by is either a single value specifying the range of colour points for which summary tetrahedral-colourspace variables should be calculated (for example, by = 3 indicates summary will be calculated for groups of 3 consecutive colour points (rows) in the quantum catch colour data frame) or a vector containing identifications for the rows in the quantum catch colour data frame (in which case summaries will be calculated for each group of points sharing the same identification). If by is left blank, the summary statistics are calculated across all colour points in the data.
- `...` class consistency (ignored).

### Value

returns all attributes of the data as mapped to the selected colourspace, including options specified when calculating the visual model. Also return the default data.frame summary, except when the object is the result of `tcspace()`, in which case the following variables are output instead:

- `centroid.u, .s, .m, .l` the centroids of usml coordinates of points.
- `c.vol` the total volume occupied by the points.
- `rel.c.vol` volume occupied by the points relative to the tetrahedron volume.
- `colspan.m` the mean hue span.
- `colspan.v` the variance in hue span.
- `huedisp.m` the mean hue disparity.
- `huedisp.v` the variance in hue disparity.
- `mean.ra` mean saturation.
- `max.ra` maximum saturation achieved by the group of points.
Author(s)

Rafael Maia <rm72@zips.uakron.edu>

References


Examples

# Colour hexagon
data(flowers)
vis.flowers <- vismodel(flowers, visual = "apis", qcatch = "Ei", relative = FALSE, vonkries = TRUE, bkg = "green")
flowers.hex <- hexagon(vis.flowers)
summary(flowers.hex)

# Tetrahedral model
data(sicalis)
vis.sicalis <- vismodel(sicalis, visual = "avg.uv")
csp.sicalis <- colspace(vis.sicalis)
summary(csp.sicalis, by = rep(c("C", "T", "B"), 7))

summary.rimg

Image summary

Description

Returns the attributes of, and optionally plots, an image.

Usage

## S3 method for class 'rimg'
summary(object, plot = FALSE, axes = TRUE, col = NULL, ...)

Arguments

object (required) an image of class rimg, or list thereof.
plot logical; plot the image and, if the image is color-classified, the colours corresponding to colour class categories side-by-side? Defaults to FALSE.
axes should axes be drawn when plot = TRUE? (defaults to TRUE).
col optional vector of colours when plotting colour-classified images with plot = TRUE. Defaults to the mean RGB values of the k-means centres (i.e. the 'original' colours).
... additional graphical options when plot = TRUE. Also see par().
Value

Either the RGB values of the k-means centres from the colour-classified image, or a plot of both the image and specified colours (when plot = TRUE).

Author(s)

Thomas E. White <thomas.white026@gmail.com>

Examples

```r
## Not run:
papilio <- getimg(system.file("testdata/images/papilio.png", package = "pavo"))
papilio_class <- classify(papilio, kcols = 4)
summary(papilio_class)

# Plot the colour-classified image alongside the colour class palette
summary(papilio_class, plot = TRUE)

# Multiple images
snakes <- getimg(system.file("testdata/images/snakes", package = "pavo"))
snakes_class <- classify(snakes, kcols = 3)
summary(snakes_class, plot = TRUE)

## End(Not run)
```

Description

Calculates all 23 colourimetric variables reviewed in Montgomerie (2006).

Usage

```r
## S3 method for class 'rspec'
summary(object, subset = FALSE, wlmin = NULL, wlmax = NULL, ...)
```

Arguments

- **object**: (required) a data frame, possibly an object of class rspec, with a column with wavelength data, named 'wl', and the remaining column containing spectra to process.
- **subset**: Either FALSE (the default), TRUE, or a character vector. If FALSE, all variables calculated are returned. If TRUE, only a subset of the complete output (composed of B2, S8 and H1; the variables described in Andersson and Prager 2006) are returned. Finally, a user-specified string of variable names can be used in order to filter and show only those variables.
wlmin, wlmax: minimum and maximum used to define the range of wavelengths used in calculations (default is to use entire range in the rspec object).

Value

A data frame containing either 23 or 5 (subset = TRUE) variables described in Montgomerie (2006) with spectra name as row names. The colorimetric variables calculated by this function are described in Montgomerie (2006) with corrections included in the README CLR file from the May 2008 distribution of the CLR software. Authors should reference both this package, Montgomerie (2006), and the original reference(s). Description and notes on the measures:

B1 (Total brightness): Sum of the relative reflectance over the entire spectral range (area under the curve). Frequently used but should be discouraged because values are difficult to compare across studies (B2 is preferred). REF 1-3, 7, 9-11, 13

B2 (Mean brightness): Mean relative reflectance over the entire spectral range. This is preferred to B1 since values are easier to compare across studies. REF 4, 12

B3 (Intensity): Maximum relative reflectance (Reflectance at wavelength of maximum reflectance). Note that may be sensitive to noise near the peak. REF 1, 5, 6

S1 (Chroma): Relative contribution of a spectral range to the total brightness (B1) S1 is arbitrarily divided in 6 measures of chroma based on the wavelength ranges normally associated with specific hues. The values are calculated using the following ranges: S1U (UV, if applicable): lambda min-400nm; S1V (Violet) lambda min-415nm; S1B (Blue) 400nm-510nm; S1G (Green) 510nm-605nm; S1Y (Yellow) 550nm-625nm; S1R (Red) 605nm-lambda max. REF 2, 7, 8, 11-13

S2 (Spectral saturation): Rmax/Rmin This measure is sensitive to spectral noise. Proper interpretation of this value may be difficult for spectra with multiple peaks in the range of interest. REF 1

S3 (Chroma): Reflectance over the Rmax +- 50nm range divided by B1. Values for peaks within 50nm of either the minimum or maximum range of the data will not be comparable since the area under the curve for the area of interest will not always be based on the same wavelength range. Therefore, S3 should be interpreted with caution for peaks in the UV or Red range. REF 11

S4 (Spectral purity): |bmaxneg|, calculated by approximating the derivative of the spectral curve. As such, it is very sensitive to noise and should only be considered when data is adequately smoothed. NAs are returned for curves which do not, at any range of wavelength, decrease in intensity. Therefore, reflectance curves for brown and red surfaces, for example, should not generate a values. REF 1

S5 (Chroma): Similar in design to segment classification measures (see Montgomerie 2006 for details). REF 10

S6 (Contrast): Rmax - Rmin. Because it uses both Rmin and Rmax, this measure may be sensitive to spectral noise. REF 5, 6

S7 (Spectral saturation): Difference between the relative reflectance before and after the wavelength at which reflectance is halfway between its minimum (Rmin) and its maximum (Rmax). Somewhat sensitive to noise and can be misleading when more than one maxima and/or minima are present. REF 3, 9

S8 (Chroma): (Rmax - Rmin)/B2. Because it uses both Rmin and Rmax, this measure may be sensitive to spectral noise. REF 3, 13
S9 (Carotenoid chroma): \( \frac{(R700 - R450)}{R700} \). Should only be used when the colour of the surface is clearly due to carotenoid pigmentation and \( R450 \) is lower than \( R700 \). Could be sensitive to noise. REF 8

S10 (Peaky chroma): \( \frac{(R_{\text{max}} - R_{\text{min}})}{B2 \times |b_{\text{maxneg}}|} \). Should be used with properly smoothed curves. REF 7

H1 (Peak wavelength, hue): Wavelength of maximum reflectance. May be sensitive to noise and may be variable if there is more than one maxima. REF 1, 2, 4, 6, 7, 10-13

H2 (Hue): Wavelength at \( b_{\text{maxneg}} \). Should be calculated using smoothed data. REF 2, 13

H3 (Hue): Wavelength at \( R_{\text{mid}} \). Sensitive to noisy spectra and may be variable if there are more than one maxima and minima. REF 3, 9, 13

H4 (Hue): Similar in design to segment classification measures see Montgomerie (2006) for details. REF 10

H5 (Hue): Wavelength at \( b_{\text{max}} \). Sensitive to noise and may be variable if there is more than one maxima and minima. REF 5

**Note**

If minimum wavelength is over 400, UV chroma is not computed.

Variables which compute \( b_{\text{max}} \) and \( b_{\text{maxneg}} \) should be used with caution, for they rely on smoothed curves to remove noise, which would otherwise result in spurious results. Make sure chosen smoothing parameters are adequate.

Smoothing affects only B3, S2, S4, S6, S10, H2, and H5 calculation. All other variables can be reliably extracted using non-smoothed data.

**Author(s)**

Pierre-Paul Bitton <bittonp@windsor.ca>, Rafael Maia <rm72@zips.uakron.edu>

**References**


References describing variables:


Examples

data(sicalis)
summary(sicalis)
summary(sicalis, subset = TRUE)
summary(sicalis, subset = c("B1", "H4"))

summary.vismodel Visual model summary

Description

Returns the attributes used when calculating a visual model using vismodel().

Usage

## S3 method for class 'vismodel'
summary(object, ...)

Arguments

object (required) Results of vismodel()
... class consistency (ignored)
Value

Returns all attributes chosen when calculating the visual model, as well as the default data.frame summary

Author(s)

Rafael Maia <rm72@zips.uakron.edu>

References


Examples

data(sicalis)
vis.sicalis <- vismodel(sicalis, visual = “avg.uv”)
summary(vis.sicalis)

Description

Produces an interactive 3D plot of a tetrahedral colourspace using OpenGL capabilities.
Plots points in a tetrahedral colour space

Usage

tcsplot(
tcsdata,
size = 0.02,
alpha = 1,
col = “black”,
vertexsize = 0.02,
achro = TRUE,
achrosize = 0.01,
achrocol = “grey”,
lwd = 1,
lcol = "lightgrey",
new = FALSE,
hspin = FALSE,
vspin = FALSE,
floor = TRUE
)

tcspoints(tcsdata, size = 0.02, col = "black", alpha = 1)

tcsvol(
  tcsdata,
  col = "black",
  alpha = 0.2,
  grid.alpha = 1,
  grid = TRUE,
  fill = TRUE,
  lwd = 1
)

Arguments

tcsdata (required) a data frame, possibly a result from the colspace() or tcspace() function, containing values for the 'x', 'y' and 'z' coordinates as columns (labeled as such).

size size of the points in the plot (defaults to 0.02)

alpha transparency of points (or volume fill in tcsvol())

col colour of the points in the plot (defaults to black)

vertexsize size of the points at the vertices

achro should a point be plotted at the origin (defaults to TRUE)?

achrosize size of the point at the origin when achro = TRUE (defaults to 0.8).

achrocol color of the point at the origin achro = TRUE (defaults to 'grey').

lwd, lcol graphical parameters for the edges of the tetrahedron.

new should a new 3D plot be called (defaults to FALSE)?

hspin if TRUE, the graphic will spin horizontally (around the 'z' axis)(defaults to FALSE)

vspin if TRUE, the graphic will spin vertically (around the 'x' axis)(defaults to FALSE)

floor if TRUE, a reference xy plane is plotted under the tetrahedron (defaults to TRUE)

grid.alpha transparency of the volume polygon grid lines

grid if TRUE, connects the polygon outlining the volume occupied by points (defaults to TRUE)

fill if TRUE, fills the volume occupied by points (WARNING: transparency is not saved properly if exported using rgl.postscript)(defaults to TRUE).
Value

tcsplot() creates a 3D plot using functions of the package rgl, based on openGL capabilities. Plot is interactive and can be manipulated with the mouse (left button: rotate along 'z' axis; right button: rotate along 'x' axis; third button: zoom).

tcspoints() adds points to the plot. Points are currently plotted only as spheres to maintain export capabilities.

tcsvol() creates a 3D convex hull within a tcsplot object.

Author(s)

Rafael Maia <rm72@zips.uakron.edu>

References


See Also

rgl::spheres3d(), rgl::rgl.postscript(), rgl::rgl.snapshot(), rgl::rgl.material()

Examples

## Not run:
# For plotting
data(sicalis)
vis.sicalis <- vismodel(sicalis, visual = "avg.uv")
tcs.sicalis <- colspace(vis.sicalis, space = "tcs")
tcsplot(tcs.sicalis, size = 0.005)
rgl::rgl.postscript("testplot.pdf", fmt = "pdf")
rgl::rgl.snapshot("testplot.png")

# For adding points
patch <- rep(c("C", "T", "B"), 7)
tcs.crown <- subset(tcs.sicalis, "C")
tcs.breast <- subset(tcs.sicalis, "B")
tcsplot(tcs.crown, col = "blue")
tcspoints(tcs.breast, col = "red")

# For plotting convex hull
tcsplot(tcs.sicalis, col = "blue", size = 0.005)
tcsvol(tcs.sicalis)

## End(Not run)
teal

*Angle-resolved reflectance data for the iridescent wing patch of a male green-winged teal (Anas carolinensis)*

**Description**

dataset containing reflectance measurements from the wing patch of a single male at different incident angles (15-75 degrees in 5-degree increments).

**Author(s)**

Chad Eliason <cme16@zips.uakron.edu>

**transmissiondata**

*Default ocular transmission data*

**Description**

Default ocular transmission data

**Author(s)**

Rafael Maia <rm72@zips.uakron.edu>

**References**


**ttvertex**

*vertex for the tetrahedral color space*

**Description**

internal data for plotting devices.

**Author(s)**

Rafael Maia <rm72@zips.uakron.edu>

**References**

vismodel

Visual models

Description
Calculates quantum catches at each photoreceptor. Both raw and relative values can be returned, for use in a suite of colourspace and non-colourspace models.

Usage

```R
vismodel(
  rspecdata,
  visual = c("avg.uv", "avg.v", "bluetit", "ctenophorus", "star", "pfowl", "apis", "canis", "cie2", "cie10", "musca", "segment", "habronattus", "rhinecanthus"),
  achromatic = c("none", "bt.dc", "ch.dc", "st.dc", "md.r1", "ra.dc", "cf.r", "ml", "1", "all"),
  illum = c("ideal", "bluesky", "D65", "forestshade"),
  trans = c("ideal", "bluetit", "blackbird"),
  qcatch = c("Qi", "fi", "Ei"),
  bkg = c("ideal", "green"),
  vonkries = FALSE,
  scale = 1,
  relative = TRUE
)
```

Arguments

- `rspecdata` (required) a data frame, possibly of class `rspec`, which contains a column containing a wavelength range, named `wl`, and spectra data in remaining columns.
- `visual` the visual system to be used. Options are:
  - a data frame such as one produced containing by `sensmodel()`, containing user-defined sensitivity data for the receptors involved in colour vision. The data frame must contain a `wl` column with the range of wavelengths included, and the sensitivity for each other cone as a column.
  - 'apis': Honeybee *Apis mellifera*.
  - 'avg.uv': average avian UV system (default).
  - 'avg.v': average avian V system.
  - 'bluetit': Blue tit *Cyanistes caeruleus*.
  - 'canis': Canid *Canis familiaris*.
  - 'cie2': 2-degree colour matching functions for CIE models of human colour vision. Functions are linear transformations of the 2-degree cone fundamentals of Stockman & Sharpe (2000), as ratified by the CIE (2006).
  - 'cie10': 10-degree colour matching functions for CIE models of human colour vision. Functions are linear transformations of the 10-degree cone fundamentals of Stockman & Sharpe (2000), as ratified by the CIE (2006).
• 'ctenophorus': Ornate dragon lizard \textit{Ctenophorus ornatus}.
• 'musca': Housefly \textit{Musca domestica}.
• 'pfo\textbackslash w\textbackslash l': Peafowl \textit{Pavo cristatus}.
• 'star': Starling \textit{Sturnus vulgaris}.
• 'haboronattus': Jumping spider \textit{Habronattus pyrrithrix}.
• 'rhinecanthus': Triggerfish \textit{Rhinecanthus aculeatus}.

\textbf{achromatic} the sensitivity data to be used to calculate luminance (achromatic) receptor stimulation. Either a vector containing the sensitivity for a single receptor, or one of the options:
• 'none': no achromatic stimulation calculated (default).
• 'bt.dc': Blue tit \textit{Cyanistes caeruleus} double cone.
• 'ch.dc': Chicken \textit{Gallus gallus} double cone.
• 'st.dc': Starling \textit{Sturnus vulgaris} double cone.
• 'md.r1': Housefly \textit{Musca domestica} R1-6 photoreceptor.
• 'ra.dc': Triggerfish \textit{Rhinecanthus aculeatus} double cone.
• 'cf.r': Canid \textit{Canis familiaris} cone.
• 'ml': the summed response of the two longest-wavelength photoreceptors.
• 'l': the longest-wavelength photoreceptor.
• 'all': the summed response of all photoreceptors.

\textbf{illum} either a vector containing the illuminant, or one of the options:
• 'ideal': homogeneous illuminance of 1 across wavelengths (default)
• 'bluesky': open blue sky.
• 'D65': standard daylight.
• 'forestshade': forest shade.

\textbf{trans} either a vector containing the ocular or environmental transmission spectra, or one of the options:
• 'ideal': homogeneous transmission of 1 across all wavelengths (default)
• 'bluetit': blue tit \textit{Cyanistes caeruleus} ocular transmission (from Hart et al. 2000).
• 'blackbird': blackbird \textit{Turdus merula} ocular transmission (from Hart et al. 2000).

\textbf{qcatch} Which quantal catch metric to return. Options are:
• 'Qi': Quantum catch for each photoreceptor (default)
• 'fi': Quantum catch according to Fechner's law (the signal of the receptor channel is proportional to the logarithm of the quantum catch)
• 'Ei': Hyperbolic-transformed quantum catch, where \(Ei = Qi / (Qi + 1)\).

\textbf{bkg} background spectrum. Note that this will have no effect when \texttt{vonkries = FALSE}. Either a vector containing the spectral data, or one of the options:
• 'ideal': homogeneous illuminance of 1 across all wavelengths (default).
• 'green': green foliage.
vonkries: logical. Should the von Kries colour correction transformation be applied? (defaults to FALSE).

scale: a value by which the illuminant will be multiplied. Useful for when the illuminant is a relative value (i.e. transformed to a maximum of 1 or to a percentage), and does not correspond to quantum flux units (µmol.s\(^{-1}\).m\(^{-2}\)). Useful values are, for example, 500 (for dim light) and 10000 (for bright illumination). Note that if vonkries = TRUE this transformation has no effect.

relative: should relative quantum catches be returned (i.e. is it a colour space model? Defaults to TRUE).

Value
An object of class vismodel containing the photon catches for each of the photoreceptors considered. Information on the parameters used in the calculation are also stored and can be called using the summary.vismodel() function.

Author(s)
Rafael Maia <rm72@zips.uakron.edu>
Thomas White <thomas.white026@gmail.com>

References


Examples

# Dichromat (dingo)
data(flowers)
vis.dingo <- vismodel(flowers, visual = "canis")
di.dingo <- colspace(vis.dingo, space = "di")

# Trichromat (honeybee)
data(flowers)
vis.bee <- vismodel(flowers, visual = "apis")
tri.bee <- colspace(vis.bee, space = "tri")

# Tetrachromat (blue tit)
data(sicalis)
vis.bluetit <- vismodel(sicalis, visual = "bluetit")
tcs.bluetit <- colspace(vis.bluetit, space = "tcs")

# Tetrachromat (starling), receptor-noise model
data(sicalis)
vis.star <- vismodel(sicalis, visual = "star", achromatic = "bt.dc", relative = FALSE)
dist.star <- coldist(vis.star, achromatic = TRUE)

# Custom trichromatic visual system
custom <- sensmodel(c(330, 440, 550))
names(custom) <- c("wl", "s", "m", "l")
vis.custom <- vismodel(flowers, visual = custom)
tri.custom <- colspace(vis.custom, space = "tri")

vissyst  
Animal visual systems data

Description

Internal data for visual model calculations.

Author(s)

Rafael Maia <rm72@zips.uakron.edu>

References

vol

**Plot a tetrahedral colour space**

Description

Produces a 3D convex hull in tetrahedral colour space when plotting a non-interactive tetrahedral plot.

Usage

```r
vol(tcsdata, alpha = 0.2, grid = TRUE, fill = TRUE, new = FALSE, ...)
```

Arguments

- `tcsdata` (required) a data frame, possibly a result from the `colspace()` or `tcspace()` function, containing values for the 'x', 'y' and 'z' coordinates as columns (labeled as such).
- `alpha` transparency of volume (if `fill = TRUE`).
- `grid` logical. if `TRUE` (default), draws the polygon outline defined by the points.
- `fill` logical. if `TRUE` (default), fills the volume defined by the points.
- `new` logical. Should a new plot be started or draw over an open plot? (defaults to `FALSE`)
- `...` additional graphical options. See `polygon()` and `tetraplot()`.

Value

`vol()` creates a 3D convex hull within a static tetrahedral plot.

Author(s)

Rafael Maia `<rm72@zips.uakron.edu>`

Examples

```r
# For plotting
data(sicalis)
vis.sicalis <- vismodel(sicalis, visual = "avg.uv")
tcs.sicalis <- colspace(vis.sicalis, space = "tcs")
plot(tcs.sicalis)
vol(tcs.sicalis)
```
voloverlap  

**Colour volume overlap**

**Description**

Calculates the overlap between the volumes defined by two sets of points in cartesian space.

**Usage**

```r
voloverlap(
  colsp1, colsp2,
  plot = FALSE,
  interactive = FALSE,
  col = c("blue", "red", "darkgrey"),
  fill = FALSE,
  new = TRUE,
  montecarlo = NULL,
  nsamp = NULL,
  psize = NULL,
  lwd = 1,
  ...
)
```

**Arguments**

- `colsp1, colsp2` (required) data frame, possibly a result from the `colspace()` function, containing values for the `x`, `y` (and possibly `z`) coordinates as columns (labeled as such)
- `plot` logical. Should the volumes and points be plotted? (defaults to `FALSE`). This only works for tetrahedral colourspaces at the moment.
- `interactive` logical. If `TRUE`, uses the rgl engine for interactive plotting; if `FALSE` then a static plot is generated.
- `col` a vector of length 3 with the colours for (in order) the first volume, the second volume, and the overlap.
- `fill` logical. should the two volumes be filled in the plot? (defaults to `FALSE`)
- `new` logical. Should a new plot window be called? If `FALSE`, volumes and their overlap are plotted over the current plot (defaults to `TRUE`).
- `montecarlo` deprecated argument
- `nsamp` deprecated argument
- `psize` deprecated argument
- `lwd` if `plot = TRUE`, sets the line width for volume grids.
- `...` additional arguments passed to the plot. See `vol()`
Value

 Calculates the overlap between the volumes defined by two set of points in colourspace. The volume from the overlap is then given relative to:

- $v_{\text{smallest}}$: the volume of the overlap divided by the smallest of that defined by the the two input sets of colour points. Thus, if one of the volumes is entirely contained within the other, this overlap will be $v_{\text{smallest}} = 1$.
- $v_{\text{both}}$: the volume of the overlap divided by the combined volume of both input sets of colour points.

Note

Stoddard & Stevens (2011) originally obtained the volume overlap through Monte Carlo simulations of points within the range of the volumes, and obtaining the frequency of simulated values that fall inside the volumes defined by both sets of colour points.

Stoddard & Stevens (2011) also return the value of the overlap relative to one of the volumes (in that case, the host species). However, for other applications this value may not be what one expects to obtain if (1) the two volumes differ considerably in size, or (2) one of the volumes is entirely contained within the other. For this reason, we also report the volume relative to the union of the two input volumes, which may be more adequate in most cases.

Author(s)

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Hugo Gruson <hugo.gruson+R@normalesup.org>

References


Examples

data(sicalis)
tcs.sicalis.C <- subset(colspace(vismodel(sicalis)), "C")
tcs.sicalis.T <- subset(colspace(vismodel(sicalis)), "T")
tcs.sicalis.B <- subset(colspace(vismodel(sicalis)), "B")
voloverlap(tcs.sicalis.T, tcs.sicalis.B)
voloverlap(tcs.sicalis.T, tcs.sicalis.C, plot = TRUE)
voloverlap(tcs.sicalis.T, tcs.sicalis.C, plot = TRUE, col = seq_len(3))
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