Package ‘SIBER’

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**addEllipse**

_addEllipse_ Adds an ellipse around some data to an existing plot

**Description**

This function adds an ellipse based on means and covariance to an existing plot. The ellipse can be scaled so as to represent any prediction interval of the data you wish, or alternatively any confidence interval of the bivariate means.
addEllipse

Usage

addEllipse(
  mu,
  sigma,
  m = NULL,
  n = 100,
  p.interval = NULL,
  ci.mean = FALSE,
  small.sample = FALSE,
  do.plot = TRUE,
  ...
)

Arguments

mu  a vector of length two specifying the bivariate means
sigma a 2x2 covariance matrix for the data
m  is the sample size of the dataset on which the ellipse is to be plotted. This is only informative if calculating the confidence interval of the bivariate mean, which requires a correction of $1/\sqrt{m}$. Defaults to NULL and has no effect.
n  the number of data points to be used to plot the ellipse. More points makes for a smoother ellipse, especially if it has high eccentricity. Defaults to $n = 100$.
p.interval the quantile to be used to construct a prediction ellipse that contains p.interval proportion of the data. By default, p.interval = NULL and the Standard Ellipse is drawn which contains approximately 40% of the data. Setting p.interval = 0.95 will result in an ellipse that contains approximately 95% of the data.

ci.mean a logical that determines whether the ellipse drawn is a prediction ellipse of the entire data, or a confidence interval of the bivariate means. Defaults to FALSE. If set to TRUE, then p.interval can be used to generate an appropriate % confidence interval of the bivariate means.
small.sample a logical that determines whether or not the small sample size correction is to be applied (TRUE) or not (FALSE). Defaults to FALSE. This allows SEAc rather than SEA to be plotted, but the correction can be applied to any percentile ellipse.
do.plot A logical that determines whether plotting should occur (TRUE and default) or not (FALSE). Setting to false is useful if you want to access the coordinates of the ellipse in order to calculate overlap between ellipses for example.

Value

A $n \times 2$ matrix comprising the x and y coordinates of the ellipse.

Examples

#-- NOT RUN --
# data(demo.siber.data)
# my.siber.data <- createSiberObject(demo.siber.data)
# communityMetricsML(my.siber.data)
# -- END --

## allCentroidVectors
Plot the pairwise distances and angles describing the difference between centroids of all groups

**Description**
Plots the posterior densities

**Usage**
```
allCentroidVectors(centroids, upper = TRUE, do.plot = TRUE, ...)
```

**Arguments**
- **centroids**: the list containing distance and angle matrices as returned by `siberCentroids`
- **upper**: a logical determining whether to plot the upper or lower triangle of angles. Defaults to `TRUE` which is the upper triangle and returns the angle from the second ellipse to the first by centering on the first centroid.
- **do.plot**: a logical indicating whether plotting should be done or not. Defaults to `TRUE`.
- **...**: additional arguments to pass onwards, not currently implemented.

**Value**
A nice plot. You can get the corresponding matrices used to generate the plots if you ask for it nicely: `thedata <- plotCentroidVectors(centroids)`

## bayesianLayman
Calculate Layman metrics on Bayesian posterior samples of a community

**Description**
This function loops over the posterior distribution of group means within each community and generates the corresponding Bayesian estimate of the 6 Layman metrics.

**Usage**
```
bayesianLayman(mu.post)
```
Arguments

mu.post  a list of length n.communities, with each list element containing the estimated means of the groups comprising that community. The typical workflow to generate mu.post follows. The Bayesian ellipses are fitted using `siberEllipses`, then the posterior means (centre of mass of each group) is extracted using `extractPosteriorMeans`. See the example below.

Value

A list of length n.communities, with each element containing a matrix of 6 columns, each representing the Bayesian posterior distribution of the 6 Layman metrics for each of the posterior draws recorded by the fitting process (i.e. which determines the number of rows in this matrix).

Description

This function loops over the posterior distribution of the means and covariances matrices of two specified groups.

Usage

```r
bayesianOverlap(
  ellipse1,
  ellipse2,
  ellipses.posterior,
  draws = 10,
  p.interval = 0.95,
  n = 100,
  do.plot = FALSE
)
```

Arguments

`ellipse1`  character code of the form "x.y" where x is an integer indexing the community, and y an integer indexing the group within that community. This specifies the first of two ellipses whose overlap will be compared.

`ellipse2`  same as `ellipse1` specifying a second ellipse.

`ellipses.posterior`  a list of posterior means and covariances fitted using `siberEllipses`.

`draws`  an integer specifying how many of the posterior draws are to be used to estimate the posterior overlap. Defaults to 10 which uses the first 10 draws. In all cases, the selection will be `1:draws` so independence of the posterior draws is assumed. Setting to `NULL` will use all the draws (WARNING - like to be very slow).
communityMetricsML

communityMetricsML (siber)  Calculate the point estimates of the Layman metrics for each community

description

This function loops over each community, determines the centre of mass (centroid) of each of the groups comprising the community using the basic mean function independently on the marginal x and y vectors, and calculates the corresponding 6 Layman metrics based on these points.

usage

communityMetricsML(siber)

Arguments

 siber   a siber object as created by createSiberObject.R

value

A 6 x m matrix of the 6 Layman metrics of dX_range, dY_range, TA, CD, MNND and SDNND in rows, for each community by column

Examples

data(demo.siber.data)
my.siber.data <- createSiberObject(demo.siber.data)
communityMetricsML(my.siber.data)
**createSiberObject**

**Read in SIBER format data and generate the SIBER object**

**Description**

This function takes raw isotope data and creates a SIBER object which contains information in a structured manner that enables other functions to loop over groups and communities, fit Bayesian ellipses, and afterwards, generate various plots, and additional analyses on the posterior distributions.

**Arguments**

- `data.in` Specified In a basic R data.frame or matrix comprising 4 columns. The first two of which are typically isotope tracers, then the third is a column that indicates the group membership, and the fourth column indicates the community membership of an observation. Communities labels should be entered as sequential numbers. As of v2.0.1 group labels can be entered as strings and/or numbers and need not be sequential.

**Value**

A siber list object, that contains data that helps with various model fitting and plotting.

- `original.data` The original data as passed into this function
- `iso.summary` The max, min, mean and median of the isotope data useful for plotting
- `sample.sizes` The number of observations tabulated by group and community
- `raw.data` A list object of length equal to the number of communities

**Examples**

```r
data(demo.siber.data)
my.siber.data <- createSiberObject(demo.siber.data)
names(my.siber.data)
```

**demo.siber.data**

*Simulated d13C and d15N isotope-space data*

**Description**

Data for two communities, created by `generateSiberData` used to generate the vignette and illustrates the main functionality of SIBER.

**Usage**

```r
data(demo.siber.data)
```
Format

An object of class "data.frame" containing four variables. The first and second variables are generic isotopes called iso1 and iso2. The third variable group identifies which group within a community an observation belongs. Group are required to be integers in sequential order starting at 1 and numbering should restart within each community. The fourth variable community identifies which community an observation belongs, and again is required to be an integer in sequential order staring at 1.

Author(s)

Andrew Jackson

demo.siber.data.2  Simulated d13C and d15N isotope-space data

Description

Data for two communities, created by generateSiberData used to generate the vignette and illustrates the main functionality of SIBER.

Usage

data(demo.siber.data.2)

Format

An object of class "data.frame" containing four variables. The first and second variables are generic isotopes called iso1 and iso2. The third variable group identifies which group within a community an observation belongs. Group are required to be integers in sequential order starting at 1 and numbering should restart within each community. The fourth variable community identifies which community an observation belongs, and again is required to be an integer in sequential order staring at 1.

Author(s)

Andrew Jackson
**ellipseBackTransform**  
*Back-transform a z-score siber ellipse to original location and scale.*

**Description**

Back-transforms a bivariate siber ellipse fitted to z-scored data to the original location and scale. Not intended for direct call by users.

**Usage**

```r
ellipseBackTransform(jags.output, siber, idx.community, idx.group)
```

**Arguments**

- **jags.output**: a `mcmc.list` object of posterior samples created by `rjags`. In `siber` this is created typically by `fitEllipse`
- **siber**: a `siber` object as created by `createSiberObject`
- **idx.community**: an integer specifying which community to back-transform.
- **idx.group**: an integer specifying which group to back-transform.

**Value**

A 6 x n matrix representing the back-transformed posterior distributions of the bivariate normal distribution for a specified group within a specified community, where n is the number of posterior draws in the saved sample. The first four columns are the covariance matrix Sigma in vector format. This vector converts to the covariance matrix as `matrix(v[1:4], nrow = 2, ncol = 2)`. The remaining two columns are the back-transformed means.

---

**ellipseInOut**  
*Test whether a set of points are inside or outside a defined circle*

**Description**

Takes a

**Usage**

```r
ellipseInOut(Z, p = 0.95, r = NULL)
```

**Arguments**

- **Z**: the i x d matrix of data points to be tested
- **p**: the percentile of the ellipse to be tested
- **r**: a manually defined radius of the circle to be used. Setting r to be anything other than NULL will override the choice of p.
extractPosteriorMeans

Value
   A logical vector indicating whether the point is inside or outside the circle

ellipsoidTransform  Apply a normalisation transformation to vectors of data onto ellipsoids

Description
   Takes a vector \( x \) and transforms the points onto the same geometry of a normalised ellipse given by the inverse of the covariance matrix \( \text{SigSqrt} \) and the location \( \mu \)

Usage
   ellipsoidTransform(\( x, \text{SigSqrt}, \mu \))

Arguments
   \( x \)  the vector of data points to be transformed
   \( \text{SigSqrt} \)  the inverse of the covariance matrix
   \( \mu \)  the vector of means of the ellipse

Value
   A vector of transformed data points

extractPosteriorMeans  Extract posterior means from call to siberMVN

Description
   This function extracts the posterior means from a call to siberMVN which can then be used to calculate bayesian layman metrics. This function is designed to create an array of posterior means that is more easily interrogated for plotting and summary statistics.

Usage
   extractPosteriorMeans(siber, post)

Arguments
   siber  a siber object as created by createSiberObject
   post  a list containing the posterior estimated parameters fitted to each group within every community. See siberMVN which creates this object for details.
Value
A list of length n.communities with each entry representing a n.draws * 2 * n.groups array of rows equal to the number of posterior samples, 2 columns representing the two means of the multivariate data and n.groups the number of groups within the focal community.

Description
This function contains and defines the jags model script used to fit a bivariate normal distribution to a vector of x and y data. Although not intended for direct calling by users, it presents a quick way to fit a model to a single group of data. Advanced users should be able to manipulate the contained jags model to fit more complex models using different likelihoods, such as multivariate lognormal distributions, multivariate gamma distributions etc...

Usage

fitEllipse(x, y, parms, priors, id = NULL)

Arguments

x a vector of data representing the x-axis
y a vector of data representing the y-axis
parms a list containing four items providing details of the rjags run to be sampled.
  • n.iterThe number of iterations to sample
  • n.burninThe number of iterations to discard as a burnin from the start of sampling.
  • n.thinThe number of samples to thin by.
  • n.chainsThe number of chains to fit.
priors a list of three items specifying the priors to be passed to the jags model.
  • RThe scaling vector for the diagonal of Inverse Wishart distribution prior on the covariance matrix Sigma. Typically set to a 2x2 matrix matrix(c(1, 0, 0, 1), 2, 2).
  • kThe degrees of freedom of the Inverse Wishart distribution for the covariance matrix Sigma. Typically set to the dimensionality of Sigma, which in this bivariate case is 2.
  • tauThe precision on the normal prior on the means mu.
id a character string to prepend to the raw saved jags model output. This is typically passed on from the calling function siberMVN and identifies the community and group with an integer labelling system. Defaults to NULL which will prevent the output object being saved even if parms$save.output is set to TRUE. The file itself will be saved to the user-specified location via parms$save.dir.
Value

A mcmc.list object of posterior samples created by jags.

Examples

```r
x <- stats::rnorm(50)
y <- stats::rnorm(50)
parms <- list()
parms$n.iter <- 2 * 10^3
parms$n.burnin <- 500
parms$n.thin <- 2
parms$n.chains <- 2
priors <- list()
priors$R <- 1 * diag(2)
priors$k <- 2
priors$tau.mu <- 1.0E-3
fitEllipse(x, y, parms, priors)
```

Description

This is a helper function that creates a sequence of points on a circle of radius \( r \) as a resolution determined by \( n \). It is not intended for direct calling, and is used by the ellipse plotting function `addEllipse`. NB not an exported function.

Usage

```r
genCircle(n = 100, r)
```

Arguments

- **n**: the number of points to create around the circle. Defaults to 100.
- **r**: the radius of the circle to create.

Value

A 2 x \( n \) matrix of x and y coordinates of points on a circle.
generateSiberCommunity

A utility function to simulate a single community comprised of groups

Description

This function simulates data for a single community by sampling from a normal distribution with different means for each group within some specified boundaries.

Usage

generateSiberCommunity(
  n.groups = 3,
  community.id = 1,
  n.obs = 30,
  mu.range = c(-1, 1, -1, 1),
  wishSigmaScale = 1
)

Arguments

n.groups the an integer specifying the number of groups to simulate. Defaults to 3.
community.id an integer identifying the community’s ID number. Defaults to 1.
n.obs the number of observations to draw per group.
mu.range a vector of length 4, specifying the min and max x and y values to sample means from. Group means are sampled from a uniform distribution within this range. The first two entries are the min and max of the x-axis, and the second two the min and max of the y-axis. Defaults to c(-1,1,-1,1).
wishSigmaScale is a simple multiplier for the call to rWishart which scales the diagonal sigma matrix using wishSigmaScale * diag(2) that is ultimately passed on to generateSiberGroup.

Value

A data.frame object comprising a column of x and y data, a group identifying column and a community identifying column, all of which are numeric.

generateSiberData

A utility function to simulate isotope data for several communities

Description

This function simulates data for a specified number of communities. It is a wrapper function for generateSiberCommunity.
Usage

generateSiberGroup(
  n.groups = 3,
  n.communities = 2,
  n.obs = 30,
  mu.range = c(-1, 1, -1, 1),
  wishSigmaScale = 1
)

Arguments

n.groups the an integer specifying the number of groups per community to simulate. De-
defaults to 3.
n.communities the number of communities to simulate data for. Defaults to 2.
n.obs the number of observations to draw per group.
mu.range a vector of length 4, specifying the mix and max x and y values to sample means
from. Group means are sampled from a uniform distribution within this range. The first two
entries are the min and max of the x-axis, and the second two the min and max of the y-axis.
Defaults to c(-1,1,-1,1).
wishSigmaScale is a simple multiplier for the call to rWishart which scales the diagonal sigma
matrix using wishSigmaScale * diag(2) that is ultimately passed on to generateSiberGroup.

Value

A data.frame object comprising a column of x and y data, a group identifying column and a com-
munity identifying column, all of which are numeric.

Examples

generateSiberData()
Arguments

**mu.range** a vector of length 4, specifying the min and max x and y values to sample means from. Group means are sampled from a uniform distribution within this range. The first two entries are the min and max of the x-axis, and the second two the min and max of the y-axis.

**n.obs** the number of observations to draw per group. Defaults to 30.

**wishSigmaScale** is a simple multiplier for the call to `rWishart` which scales the diagonal sigma matrix using `wishSigmaScale * diag(2)`.

Value

A data.frame object comprising a column of x and y data, a group identifying column and a community identifying column, all of which are numeric.

Examples

```
# generateSiberGroup()
```

---

**groupMetricsML**  
*Calculate maximum likelihood based measures of dispersion of bivariate data*

Description

This function loops over each group within each community and calculates the convex hull total area, Standard Ellipse Area (SEA) and its corresponding small sample size corrected version SEAc based on the maximum likelihood estimates of the means and covariance matrices of each group.

Usage

```
groupMetricsML(siber)
```

Arguments

**siber** a siber object as created by `createSiberObject`.

Value

A 3 x m matrix of the 6 Layman metrics of dX_range, dY_range, TA, CD, MNND and SDNND in rows, where each column is a different group nested within a community.

Examples

```
data(demo.siber.data)
my.siber.data <- createSiberObject(demo.siber.data)
groupMetricsML(my.siber.data)
```
### hullArea

*Calculate the area of a convex hull given its coordinates*

**Description**

Given the coordinates of a convex hull (i.e. a polygon), this function calculates its area. Not intended for direct use outside of `siberConvexhull`.

**Usage**

```r
hullArea(x, y)
```

**Arguments**

- `x` a vector of x-axis data
- `y` a vector of y-axis data

**Value**

A scalar representing the area of the convex hull in units of \(x \times y\); i.e. most commonly in permille squared for isotope data.

### kapow

*KAPOW!*

**Description**

This function packs a punch and makes a pretty figure.

**Usage**

```r
kapow(cd = 7, ng = 25, n = 50, sc = 10, do.plot = TRUE)
```

**Arguments**

- `cd` sets the random seed to this
- `ng` the number of ellipses to draw
- `n` the number of data points to simulate per group, but never displayed
- `sc` the scaling factor the `rwishart` sigma called by `rWishart`
- `do.plot` a logical indicating whether the plot should be printed (defaults to TRUE).

**Value**

A ggplot object
**laymanMetrics**

Calculates the 6 Layman metrics on a vector of x and y data

### Description

This function takes two x and y vectors, and calculates the corresponding 6 Layman metrics based on these points. Note that for generality, the original metrics of dC_range and dN_range have been renamed dX_range and dY_range respectively. These modified names represent the x and y axes in terms of the order in which the data have been entered, and relate typically to how one plots the data. These x and y vectors could represent the means of the group members comprising a community as is preferred under the SIBER model framework. However, one could use them to calculate the point estimates of the 6 Layman metrics for an entire group of data. In fact, you are free to pass this function any set of x and y data you wish.

### Usage

```r
laymanMetrics(x, y)
```

### Arguments

- **x**
  - a vector of locations in the x-axis direction.
- **y**
  - a vector of locations in the y-axis direction.

### Value

A vector of the 6 Layman metrics of dX_range, dY_range, TA, CD, MNND and SDNND

### Examples

```r
x <- stats::runif(10)
y <- stats::runif(10)
laymanMetrics(x, y)
```

---

**maxLikOverlap**

Calculate the overlap between two ellipses based on the maximum likelihood fitted ellipses.

### Description

This function uses the ML estimated means and covariances matrices of two specified groups to calculate the area of overlap.

---
maxLikOverlap

Usage

maxLikOverlap(
  ellipse1,
  ellipse2,
  siber.object,
  p.interval = 0.95,
  n = 100,
  do.plot = FALSE
)

Arguments

  ellipse1  character code of the form "x.y" where x is an integer indexing the community,
            and y an integer indexing the group within that community. This specifies the
            first of two ellipses whose overlap will be compared.

  ellipse2  same as ellipse1 specifying a second ellipse.

  siber.object  an object created by createSiberObject which contains the ML estimates for
                the means and covariance matrices for each group.

  p.interval  the prediction interval used to scale the ellipse as per addEllipse.

  n  the number of points on the edge of the ellipse used to define it. Defaults to 100
     as per addEllipse.

  do.plot  logical switch to determine whether the corresponding ellipses should be plotted
           or not. A use-case would be in conjunction with a low numbered draws so as to
           visualise a relatively small number of the posterior ellipses. Defaults to FALSE.

Value

A vector comprising three columns: the area of overlap, the area of the first ellipse and the area of
the second ellipse and as many rows as specified by draws.

Examples

  # load in the included demonstration dataset data("demo.siber.data")
  siber.example <- createSiberObject(demo.siber.data)

  # The first ellipse is referenced using a character string representation
  # where in "x.y", "x" is the community, and "y" is the group within that
  # community.
  ellipse1 <- "1.2"

  # Ellipse two is similarly defined: community 1, group3
  ellipse2 <- "1.3"

  # the overlap between the corresponding 95% prediction ellipses is given by:
  ellipse95.overlap <- maxLikOverlap(ellipse1, ellipse2, siber.example,
    p.interval = 0.95, n = 100)
Description

This function loops over each community and plots the convex hull based on the centres of each of the groups that make up the community. See the demonstration scripts for example implementation.

Usage

```r
plotCommunityHulls(
  siber,
  plot.args = list(col = 1, lty = 2),
  iso.order = c(1, 2),
  ...
)
```

Arguments

- `siber`: A siber object as created by `createSiberObject.R`
- `plot.args`: A list of plotting arguments with the following suggested, but non-exhaustive inputs. Additional plotting arguments for passing to the internal call to `plot` can either be specified here, or as additional arguments under the `...` method.
  - `col`: The color of the lines of the convex hull. See `lines` for more details.
  - `lty`: The line type of the convex hull. See `lines` for more details.
  - `lwd`: The line width of the convex hulls. See `lines` for more details.
- `iso.order`: A vector of length 2, either c(1,2) or c(2,1). The order determines which of the columns of raw data are plotted on the x (1) or y (2) axis. N.B. this will be deprecated in a future release, and plotting order will be achieved at point of data-entry.
- `...`: Additional arguments for passing to `plot`.

Value

Convex hulls, drawn as lines on an existing figure.
plotGroupEllipses *Adds ellipses to an existing plot for each of your groups*

**Description**

This function loops over each community and group within, and plots an ellipse around the data. See demonstration scripts for more examples.

**Usage**

```r
plotGroupEllipses(siber, plot.args = list(), iso.order = c(1, 2), ...)
```

**Arguments**

- `siber` a siber object as created by `createSiberObject`
- `plot.args` a list of plotting arguments for passing to `addEllipse`. See `addEllipse` for details of the options, and you can also pass additional arguments such as line widths and styles. See also the demonstration scripts for examples of use.
- `iso.order` a vector of length 2, either `c(1,2)` or `c(2,1)`. The order determines which of the columns of raw data are plotted on the x (1) or y (2) axis. N.B. this will be deprecated in a future release, and plotting order will be achieved at point of data-entry.
- `...` additional arguments to be passed to `addEllipse`.

**Value**

Ellipses, drawn as lines on an existing figure.

plotGroupHulls *Plots illustrative convex hulls for each group within all communities*

**Description**

This function loops over each community and group within, and plots a convex hull around the data. N.B. use of convex hulls to compare isotopic niche width among groups within or between communities is not recommended owing to strong sample size bias. Use of ellipse area is recommended instead. This feature is provided for illustrative purposes only, and because some people have expressed a desire for this feature for figure generation. See demonstration scripts for more examples.

**Usage**

```r
plotGroupHulls(siber, plot.args = NULL, iso.order = c(1, 2), ...)
```
plotSiberObject

Arguments

- `siber`: a siber object as created by `createSiberObject`
- `plot.args`: a list of plotting arguments for passing to `lines`. See `lines` for details of the options. See also the demonstration scripts for examples of use.
- `iso.order`: a vector of length 2, either `c(1, 2)` or `c(2, 1)`. The order determines which of the columns of raw data are plotted on the x (1) or y (2) axis. N.B. this will be deprecated in a future release, and plotting order will be achieved at point of data-entry.
- `...`: additional arguments to be passed to `addEllipse`.

Value

A series of convex hulls added to an existing plot.

plotSiberObject

Creates an isotope biplot and provides a wrapper to ellipse and hull plotting

Description

This function takes a SIBER object as created by `createSiberObject`, and loops over communities and their groups, creating a biplot, and adding ellipses and hulls as desired. Ellipses can be added to groups, while convex hulls can be added at both the group and community level (the former for illustrative purposes only, with no analytical tools in SIBER to fit Bayesian hulls to individual groups. This is not mathematically possible in a Bayesian framework.).

Usage

```r
plotSiberObject(
  siber,
  iso.order = c(1, 2),
  ax.pad = 1,
  hulls = TRUE,
  community.hulls.args = NULL,
  ellipses = TRUE,
  group.ellipses.args = NULL,
  group.hulls = FALSE,
  group.hulls.args = NULL,
  bty = "L",
  xlab = "Isotope 1",
  ylab = "Isotope 2",
  las = 1,
  x.limits = NULL,
  y.limits = NULL,
  points.order = 1:25,
  ...
)
```
plotSiberObject

Arguments

- **siber**: a siber object as created by `createSiberObject`.
- **iso.order**: a vector of length 2, either c(1,2) or c(2,1). The order determines which of the columns of raw data are plotted on the x (1) or y (2) axis. N.B. this will be deprecated in a future release, and plotting order will be achieved at point of data-entry.
- **ax.pad**: a padding amount to apply to the x-axis either side of the extremes of the data. Defaults to 1.
- **hulls**: a logical defaulting to TRUE determining whether or not hulls based on the means of groups within communities should be drawn. That is, a community-level convex hull.
- **community.hulls.args**: a list of plotting arguments to pass to `plotCommunityHulls`. See `plotCommunityHulls` for further details.
- **ellipses**: a logical defaulting to TRUE determining whether or not an ellipse should be drawn around each group within each community.
- **group.ellipses.args**: a list of plotting arguments to pass to `plotGroupEllipses`. See `plotGroupEllipses` for further details.
- **group.hulls**: a logical defaulting to FALSE determining whether or not convex hulls should be drawn around each group within each community.
- **group.hulls.args**: a list of plotting options to pass to `plotGroupHulls`. See `plotGroupHulls` for further details.
- **bty**: a string specifying the box type for the plot. See `par` for details.
- **xlab**: a string for the x-axis label.
- **ylab**: a string for the y-axis label.
- **las**: a scalar determining the rotation of the y-axis labels. Defaults to horizontal with `las = 1`. See `par` for more details.
- **x.limits**: allows you to specify a two-element vector of lower and upper x-axis limits. Specifying this argument over-rides the automatic plotting and ax.pad option. Defaults to NULL.
- **y.limits**: allows you to specify a two-element vector of lower and upper y-axis limits. Specifying this argument over-rides the automatic plotting and ax.pad option. Defaults to NULL.
- **points.order**: a vector of integers specifying the order of point types to use. See `points` for how integers map onto point types. Defaults to the sequence 1:15 as per `points`. It must have at least as many entries as there are communities to plot, else a warning will be issued, and the order will default to the sequence 1:25.
- **...**: additional arguments to be passed to `plot`.

Value

An isotope biplot.
pointsToEllipsoid

*Test whether a set of points are inside or outside a defined ellipse*

**Description**

Takes a $i \times d$ matrix of points where $d$ is the dimension of the space considered, and $i$ is the number of points and returns **TRUE** or **FALSE** for whether each point is inside or outside a $d$-dimensional ellipsoid defined by a covariance matrix $\Sigma$ and vector of means $\mu$.

**Usage**

```
pointsToEllipsoid(X, Sigma, mu)
```

**Arguments**

- **X**
  - the $i \times d$ matrix of data points to be transformed
- **Sigma**
  - the $d \times d$ covariance matrix of the ellipsoid
- **mu**
  - the vector of means of the ellipse of length $d$

**Value**

A matrix of transformed data points corresponding to $X$

**Examples**

```r
X <- matrix(runif(200,-2.5, 2.5), ncol = 2, nrow = 100)
SIG <- matrix(c(1,0,0,1), ncol = 2, nrow = 2)
mu <- c(0, 0)
Z <- pointsToEllipsoid(X, SIG, mu)
```

**posteriorSEA**

*Calculate the SEA based on a posterior distribution of $\Sigma$*

**Description**

This function loops over each posterior draw of a single group’s Bayesian bivariate ellipse and calculates the Standard Ellipse Area (SEA) for each draw, thereby generating a distribution of SEA estimates. Not intended for direct calling outside of `siberEllipses`.

**Usage**

```
posteriorSEA(post)
```
siberCentroids

Arguments

post a matrix of posterior covariance matrices and mean estimates for a bivariate ellipse. In SIBER, this is typically one list element of the object returned by link{siberMVN}.

Value

A vector of posterior Bayesian Standard Ellipse Areas (SEA_B)

SIBER

SIBER: A package for fitting Bayesian Ellipses to Stable Isotope Data

Description

The SIBER package provides tools for fitting multivariate normal distributions to bivariate data using Bayesian Inference. These distributions can then be used to calculate probability distributions of Standard Ellipse Areas for comparing groups of data, or to calculate the 6 Layman metrics for comparing entire communities.

siberCentroids

Calculate the polar form of the vector between pairs of ellipse centroids

Description

This function loops over each group within each community and calculates the vector in polar form between the estimated centroids of each ellipse to each other ellipse.

Usage

siberCentroids(corrected.posteriors)

Arguments

corrected.posteriors

the Bayesian ellipses as returned by siberMVN.

Value

A list containing two arrays, one \( r \) contains the pairwise distances between ellipse centroids in as the first two dimensions, with the third dimension containing the same for each posterior draw defining the ellipse. The second array \( \theta \) has the same structure and contains the angle in radians (from 0 to \( 2\pi \)) between the pairs. A third object \( \text{labels} \) refers to which community/group combination is in each of the first two dimensions of the arrays.
**siberConvexhull**  
*Calculate metrics and plotting information for convex hulls*

**Description**

This function calculates the area of the convex hull describing a set of bivariate points, and returns other information useful for plotting the hull.

**Usage**

`siberConvexhull(x, y)`

**Arguments**

- `x` a vector of x-axis data
- `y` a vector of y-axis data

**Value**

A list of length four comprising:

- `TA` the area of the convex hull.
- `hullX` the x-coordinates of the points describing the convex hull.
- `hullY` the y-coordinates of the points describing the convex hull.
- `ind` the indices of the original data in `x` and `y` that form the boundaries of the convex hull.

**Examples**

```r
x <- stats::rnorm(15)
y <- stats::rnorm(15)
siberConvexhull(x, y)
```

---

**siberDensityPlot**  
*Plot credible intervals as shaded boxplots using hdr.boxplot*

**Description**

This function is essentially `hdr.boxplot` but it more easily works with matrices of data, where each column is a different variable of interest. It has some limitations though....
Usage

siberDensityPlot(
  dat,
  probs = c(95, 75, 50),
  xlab = "Group",
  ylab = "Value",
  xticklabels = NULL,
  yticklabels = NULL,
  clr = matrix(rep(grDevices::gray((9:1)/10), ncol(dat)), nrow = 9, ncol = ncol(dat)),
  scl = 1,
  xspc = 0.5,
  prn = F,
  ct = "mode",
  ylims = NULL,
  lbound = -Inf,
  ubound = Inf,
  main = "",
  ylab.line = 2,
  ...
)

Arguments

dat a matrix of data for which density region boxplots will be constructed and plotted for each column.
probs a vector of credible intervals to represent as box edges. Defaults to c(95, 75, 50).
xlab a string for the x-axis label. Defaults to "Group".
ylab a string of the y-axis label. Defaults to "Value".
xticklabels a vector of strings to override the x-axis tick labels.
yticklabels a vector of strings to override the y-axis tick labels.
clr a matrix of colours to use for shading each of the box regions. Defaults to greyscale
  grDevices::gray((9:1)/10) replicated for as many columns as there are in dat. When specified by the user, rows contain the colours of each of the confidence regions specified in probs and columns represent each of the columns of data in dat. In this way, one could have shades of blue, red and yellow for each of the groups.
scl a scalar multiplier to scale the box widths. Defaults to 1.
xspc a scalar determining the amount of spacing between each box. Defaults to 0.5.
prn a logical value determining whether summary statistics of each column should be printed to screen prn = TRUE or suppressed as per default prn = FALSE.
ct a string of either c("mode", "mean", "median") which determines which measure of central tendency will be plotted as a point in the middle of the boxes. Defaults to "mode".
ylims a vector of length two, specifying the lower and upper limits for the y-axis. Defaults to NULL which inspects the data for appropriate limits.
a lower boundary to specify on the distribution to avoid the density kernel estimating values beyond that which can be expected a priori. Useful for example when plotting dietary proportions which must lie in the interval $0 \leq Y \leq 1$. Defaults to -Inf.

**ubound**

an upper boundary to specify on the distribution to avoid the density kernel estimating values beyond that which can be expected a priori. Useful for example when plotting dietary proportions which must lie in the interval $0 \leq Y \leq 1$. Defaults to +Inf.

**main**

title for the figure. Defaults to blank.

**ylab.line**

a positive scalar indicating the line spacing for rendering the y-axis label. This is included as using the permille symbol has a tendency to push the axis label off the plotting window margins. See the **line** option in **axis** for more details as **ylab.line** passes to this.

... further graphical parameters for passing to **plot**

---

**Value**

A new figure window.

**Warning**

This function will not currently recognise and plot multimodal distributions, unlike **hdr.boxplot**. You should take care, and plot basic histograms of each variable (column in the object you are passing) to **siardensityplot** and check that they are indeed unimodal as expected.

**Examples**

```r
# A basic default greyscale density plot
Y <- matrix(stats::rnorm(1000), 250, 4)
siberDensityPlot(Y)

# A more colourful example
my_clrs <- matrix(c("lightblue", "blue", "darkblue", "red1", "red3", "red4", "yellow1", "yellow3", "yellow4", "turquoise", "turquoise3", "turquoise4"), nrow = 3, ncol = 4)
siberDensityPlot(Y, clr = my_clrs)
```

---

**siberEllipses**

*Calculate the bayesian Standard Ellipse Area for all groups*

**Description**

This function loops over each group within each community and calculates the posterior distribution describing the corresponding Standard Ellipse Area.
Usage

siberEllipses(corrected.posteriors)

Arguments

corrected.posteriors

the Bayesian ellipses as returned by siberMVN.

Value

A matrix of with each column containing the posterior estimates of the SEA.

---

Usage

siberKapow(
  dtf,
  isoNames = c("iso1", "iso2"),
  group = "group",
  pEll = stats::pchisq(1, df = 2)
)

Arguments

dtf a data.frame object comprising bivariate data as a requirement, and possibly other variables too but these are currently ignored.

isoNames a character vector of length 2 providing the names of the variables containing the x and y data respectively.

group a character vector of length 1 providing the name of the grouping variable on which to calculate the KAPOW ellipse.

pEll the probability ellipse to draw for each group. Defaults to the Standard Ellipse with \[ pEll = \text{stats::pchisq}(1, df = 2). \]

Value

an object of class owin containing the numerically calculated ellipses and their union along with the raw ellipse boundaries in both raw and owin format.
siberMVN

Fit Bayesian bivariate normal distributions to each group in each community

Description

This function loops over each community and then loops over each group member, fitting a Bayesian multivariate (bivariate in this case) normal distribution to each group of data. Not intended for direct calling by users.

Usage

siberMVN(siber, parms, priors)

Arguments

siber
   a siber object as created by createSiberObject

parms
   a list containing four items providing details of the rjags run to be sampled.
      • n.iter The number of iterations to sample
      • n.burnin The number of iterations to discard as a burnin from the start of sampling.
      • n.thin The number of samples to thin by.
      • n.chains The number of chains to fit.

priors
   a list of three items specifying the priors to be passed to the jags model.
      • R The scaling vector for the diagonal of Inverse Wishart distribution prior on the covariance matrix Sigma. Typically set to a 2x2 matrix matrix(c(1, 0, 0, 1), 2, 2).
      • k The degrees of freedom of the Inverse Wishart distribution for the covariance matrix Sigma. Typically set to the dimensionality of Sigma, which in this bivariate case is 2.
      • tau The precision on the normal prior on the means mu.

Value

A list of length equal to the total number of groups in all communities. Each entry is named 1.1 1.2... 2.1.. with the first number designating the community, and the second number the group within that community. So, 2.3 would be the third group within the second community. Each list entry is a 6 x n matrix representing the back-transformed posterior distributions of the bivariate normal distribution, where n is the number of posterior draws in the saved sample. The first two columns are the back-transformed means, and the remaining four columns are the covariance matrix Sigma in vector format. This vector converts to the covariance matrix as matrix(v[1:4], nrow = 2, ncol = 2).
sigmaSEA

**Calculate metrics corresponding to the Standard Ellipse based on a covariance matrix**

**Description**

This function takes a covariance 2x2 matrix Sigma and returns various metrics relating to the corresponding Standard Ellipse. The function is limited to the 2-dimensional case, as many of the ancillary summary statistics are not defined for higher dimensions (e.g. eccentricity).

**Usage**

```r
sigmaSEA(sigma)
```

**Arguments**

- `sigma` a 2x2 covariance ellipse.

**Value**

A list comprising the following metrics for summarising the Standard Ellipse:

- SEA the Standard Ellipse Area (not sample size corrected)
- eccentricity a measure of the elongation of the ellipse.
- `a` the length of the semi-major axis
- `b` the length of the semi-minor axis

**Note**

This function is currently based on the eigenvalue and eigenvector approach which is more flexible for higher dimensional problems method for calculating the standard ellipse, and replaces the parametric method used previously in siar and siber.

**Examples**

```r
# A perfect circle
sigma <- matrix(c(1, 0, 0, 1), 2, 2)
sigmaSEA(sigma)
```
specificCentroidVectors

Calculate the pairwise distances and angles describing the difference between centroids of paired groups

---

Description

Plots the posterior densities

Usage

specificCentroidVectors(centroids, do.these, upper = TRUE, do.plot = TRUE, ...)

Arguments

centroids: the list containing distance and angle matrices as returned by `siberCentroids`.
do.these: a character vector of the pattern used to find paired matches in the matrix of all comparisons. Usually the group names within any of the communities.
upper: a logical determining whether to plot the upper or lower triangle of angles. Defaults to TRUE which is the upper triangle and returns the angle from the second ellipse to the first by centering on the first centroid.
do.plot: a logical indicating whether plotting should be done or not. Defaults to TRUE.
...: additional arguments to pass onwards, not currently implemented.

Value

A nice plot. You can get the corresponding matrices used to generate the plots if you ask for it nicely: `thedata <- plotCentroidVectors(centroids)`
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