Package 'gdi'

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Title Volumetric Analysis using Graphic Double Integration
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Description Tools implementing an automated version of the graphic double integration technique first employed by Jerison (1973) <isbn:9780323141086> and Hurlburt (1999) <doi:10.1080 02724634.1999.10011145="">. Graphic double integration is primarily used for volume or mass estimation of (extinct) animals, and the package 'gdi' aims to make this technique as convenient and versatile as possible. The main functions of 'gdi' provide utilities for automatically measuring diameters from digital silhouettes provided as image files, and for calculating volume via graphic double integration with a simple elliptical, superelliptical (following Motani 2001 <doi:10.1666 0094-8373(2001)027%3c0735:ebmfst%3e2.0.co;2="">) or complex cross-sectional model.</doi:10.1666></doi:10.1080></isbn:9780323141086>
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R topics documented:
cscorr fdetect gdi hCOM imghist

2 cscorr

plot.sil	 																		
sellipse																			
sellipse.coo .	 																		
vCOM	 																		

cscorr

Index

Measure and analyze cross-sectional geometry

13

Description

Measure and analyze cross-sectional geometry

Usage

```
cscorr(
  image_file,
  threshold = 0.5,
  channel = 4,
  method = "greater",
  return = "area_corr",
  k = 2,
  scale = 1
)
```

Arguments

channel

method

return

image_file Image to be read. Images can be jpeg or png files, or a previously read image saved as an object in R.

threshold Reference value for colour criterium after which pixels that are part of the cross-section are differentiated from the background.

Colour channel to which to apply the threshold criterium. Default is 4 (alpha channel of rgba image). Channel setting needs to be adjusted depending on the colour mode of the image used (e.g. there are two channels to choose from in a greyscale image, and 3 in an rgb image).

Method for determining which pixels to count. Default "greater" counts pixels with value greater than threshold (e.g. higher opacity, in the case of an alpha channel). "less" counts pixels with a value less than the threshold. "not" counts all pixels not precisely matching threshold. Any other character string results in only pixels exactly matching the value given as threshold being counted.

What value to return. Possible values are "area_corr" (Default, returns ratio between measured area and area of ellipse with same horizontal and vertical diameters), "aspect_ratio" (returns aspect ratio), "diameters" (returns diameters) and "area" (returns area). Any other value for this parameter will prompt the function to return a vector containing all of these.

fdetect 3

k	optional superellipse exponent for the (super)ellipse to which the measurements
	should be compared (for the "area_corr" setting for the parameter return).
scale	Optional scale of the image (for raw area measurements).

Value

Either a numeric of length 1 (depending on the input of the return parameter), defaulting to the area correction factor (if return=="area_corr"), or (if return is left empty or does not match any of the predefined settings) a numeric vector of length 5 containing all the possible outputs (x and y diameters, aspect ratio, area and area correction factor).

Examples

```
fdir <- system.file(package="gdi")
correction_factor <- cscorr(file.path(fdir,"exdata","cross_section.png"))

fdetect

Tool to help determine which threshold value and method to use with
measuresil() or cscorr(). The function analyzes all pixels along the
edges of the image to determine the background colour, to help with
deciding on appropriate settings and avoid errors introduced by inap-
propriate settings
```

Description

Tool to help determine which threshold value and method to use with measuresil() or cscorr(). The function analyzes all pixels along the edges of the image to determine the background colour, to help with deciding on appropriate settings and avoid errors introduced by inappropriate settings

Usage

```
fdetect(image_file, threshold = 0.5, channel = 4, plot = FALSE)
```

Arguments

image_file	Image to be read. Images can be jpeg or png files, or a previously read image saved as an object in R.
threshold	Reference value for colour criterium after which pixels that are part of the silhouette should be differentiated from the background.
channel	Colour channel to which to apply the threshold criterium. Default is 4 (alpha channel of rgba image). Channel setting needs to be adjusted depending on the colour mode of the image used (e.g. there are two channels to choose from in a greyscale image, and 3 in an rgb image).
plot	Whether to plot a histogram with the detected colour values (if TRUE) or not (if FALSE, default).

4 gdi

Value

A list()-object containing: \$edgetable (a table of the different colour values detected and their respective frequencies), \$histogram (a histogram-object of the colour values), \$most_common (the most common colour value found), \$foreground (a character string, indicating whether the foreground colour value is likely "greater" or "less" than the specified threshold), \$result (a character string giving a summary of the results)

Examples

```
fdir <- system.file(package="gdi")
fdetect(file.path(fdir,"exdata","lat.png"))</pre>
```

gdi

Estimate volume using Graphic Double Integration.

Description

Estimate volume using Graphic Double Integration.

Usage

```
gdi(
  lat,
  dors,
  indices = NULL,
  scale = 10,
  sliceL = 1/scale,
  method = "raw",
  k = 2,
  corr = 1,
  smooth.ends = FALSE,
  return = "total"
)
```

Arguments

lat	Measurements of diameter in lateral view/first of two orthogonal views to be used with the gdi. Can be either a numeric vector, a data.frame (output of measuresil(,return="all") with a collumn named "diameter", or a text file with diameter measurements to be scanned.
dors	Measurements of diameter in dorsal view/second of two orthogonal views to be used with the gdi. Can be either a numeric vector, a data.frame (output of measuresil(,return="all") with a collumn named "diameter", or a text file with diameter measurements to be scanned. Must be the same length as lat.
indices	Optional indices specifying a subset of the silhouette measurement vectors to be

analyzed. Useful if separate segment calculations are desired.

hCOM 5

scale	Scale of the data in terms of how many units of the input data are in one side of the desired unit of output volume. Defaults to 10.
sliceL	Length of individual segments to be used in the GDI. Defaults to 1/scale.
method	Method to be used for the GDI. Default "raw" setting calculates each segment as an elliptical cylinder with volume = Area $*$ SliceL. Any other string will result in volume being calculated as an elliptical frustum with base areas based on the measurements of segments i and $i+1$.
k	Superellipse exponent to be used for the cross-sectional area. Defaults to 2.0 (normal ellipse).
corr	Correction factor for area of cross-sections, calculated as the ratio between the actual cross-sectional area and that of a (super)ellipse (depending on the specified exponent k) with the same diameters. This setting enables the function to account for complex, non-elliptical cross-sections. Default value is 1, i.e. no correction. Can be either a single number, or a numeric vector of the same length as lat and dors (in the case of a changing cross-sectional geometry along the length of the body).
smooth.ends	If method != "raw", specify whether first and last segments should be left raw, or taper to 0 (i.e. be approximated as cones). Only applies if there are no leading or following zeros in the measurement vectors.
return	Determines whether to report the estimated total volume (if default/"total"), or a data.frame() with segment radii, areas and volumes (if left empty of any other character string.

Value

Either a single number representing the total volume estimated (with names indicating the horizontal length of the silhouette in the unit determined by scale), or (if return!="total") a data.frame() containing columns with the radii in both dimensions, the estimated elliptical or superelliptical areas, and the segment volumes.

Examples

```
lateral <- rep(2,4) #generate example data
dorsal <- rep(2,4)
gdi(lat=lateral, dors=dorsal, scale=10, method="raw", k=2.0)
gdi(lat=lateral, dors=lateral/2, scale=10, method="smooth", k=2.3)</pre>
```

hCOM

Finds the horizontal (x axis, i.e. the axis vertical to the cross-sections) position of the center of mass (COM) of the volume. Experimental; only valid for "raw" gdi results with segment volumes approximated as elliptical prisms, or for manually supplied segment COMs. COM is calculated as a weighted mean of all segment COMs, with the segment mass as the weighting factor.

6 hCOM

Description

Finds the horizontal (x axis, i.e. the axis vertical to the cross-sections) position of the center of mass (COM) of the volume. Experimental; only valid for "raw" gdi results with segment volumes approximated as elliptical prisms, or for manually supplied segment COMs. COM is calculated as a weighted mean of all segment COMs, with the segment mass as the weighting factor.

Usage

```
hCOM(
   x,
   volumes = NULL,
   align = "h",
   subtract = NULL,
   densities = NULL,
   scale = 1
)
```

Arguments

X	Either a data frame that is the output of gdi(, return="all"), or a numeric vector of horizontal segment COM positions.
volumes	An optional separate vector of volumes, required if x is not a data.frame containing volumes.
align	alignment of the silhouette, if "h" (default) the silhouette is assumed to be horizontally aligned, if any other value (e.g. "v") then the silhouette is assumed to be vertically aligned.
subtract	An optional separate vector of volumes, with length equal to the length or nrow() of x , to be subtracted from the volumes for the COM calculation.
densities	An optional vector of segment densities, with length equal to the length or $nrow()$ of x , to be multiplied with the volumes for the COM calculation. If both subtract and densities are supplied, the density is applied only to the "residual" volume that is left after subtraction.
scale	Optional scale value (number of pixels to chosen unit of measurement)

Value

An object of class numeric() containing the x coordinate of the center of mass of the shape, in pixels (or chosen units, if manually calculated)

```
fdir <- system.file(package="gdi")
measuresil(file.path(fdir,"exdata","lat.png"), return="all")->lat_
measuresil(file.path(fdir,"exdata","dors.png"), return="all")->dors_
gdi(lat_, dors_, return="all")->gdiresults
hCOM(gdiresults)
```

imghist 7

imghist	Simple histogram analysis for all colour values in an input image. Can
	be used to help assess whether a chosen threshold value is appropriate
	for differentiating the silhouette from the background, or for general.

Description

Simple histogram analysis for all colour values in an input image. Can be used to help assess whether a chosen threshold value is appropriate for differentiating the silhouette from the background, or for general.

Usage

```
imghist(
  image_file,
  threshold = 0.5,
  channel = 4,
  breaks = seq(0, 1, 0.05),
  plot = TRUE,
  unique = FALSE
)
```

Arguments

image_file	Image to be read. Images can be jpeg or png files, or a previously read image saved as an object in R.
threshold	Reference value for colour criterium after which pixels that are part of the silhouette should be differentiated from the background.
channel	Colour channel to which to apply the threshold criterium. Default is 4 (alpha channel of rgba image). Channel setting needs to be adjusted depending on the colour mode of the image used (e.g. there are two channels to choose from in a greyscale image, and 3 in an rgb image).
breaks	A vector of breaks for the histogram, defaults to a bin width of 0.05 between colour values of 0 and 1 .
plot	Whether to plot a histogram, defaults to TRUE
unique	Whether to return counts for unique colour values, defaults to FALSE.

Value

A plotted histogram (unless plot==FALSE), and a matrix containing the counts from the histogram (default) or the counts for unique colour values (if unique==TRUE).

```
fdir <- system.file(package="gdi")
imghist(file.path(fdir,"exdata","lat.png"))</pre>
```

8 measuresil

measuresil	Take pixel-by-pixel measurements of a silhouette in jpeg or png format for use with the gdi function.

Description

Take pixel-by-pixel measurements of a silhouette in jpeg or png format for use with the gdi function.

Usage

```
measuresil(
  image_file,
  threshold = 0.5,
  channel = 4,
  method = "greater",
  align = "h",
  return = "diameters"
)
```

Arguments

image_file	Image to be read. Images can be jpeg or png files, or a previously read image saved as an object in R.
threshold	Reference value for colour criterium after which pixels that are part of the silhouette are differentiated from the background.
channel	Colour channel to which to apply the threshold criterium. Default is 4 (alpha channel of rgba image). Channel setting needs to be adjusted depending on the colour mode of the image used (e.g. there are two channels to choose from in a greyscale image, and 3 in an rgb image).
method	Method for determining which pixels to count. Default "greater" counts pixels with value greater than threshold (e.g. higher opacity, in the case of an alpha channel). "less" counts pixels with a value less than the threshold. "not" counts all pixels not precisely matching threshold. Any other character string results in only pixels exactly matching the value given as threshold being counted.
align	Indicate whether the silhouette long axis is aligned horizontally (setting "h", default), or vertically (any other parameter setting).
return	Setting for what to return, default setting ("diameters") returns a single vector containing the diameters, any other setting returns a data frame containing centers and diameters.

Value

A numeric vector giving the measurements of the silhouette

plot.sil 9

Examples

```
fdir <- system.file(package="gdi")
lat <- measuresil(file.path(fdir,"exdata","lat.png"))</pre>
```

plot.sil

Plots a silhouette read by measuresil()

Description

Plots a silhouette read by measuresil()

Usage

```
## S3 method for class 'sil'
plot(sil, flip = FALSE, add = FALSE, ...)
```

Arguments

sil	A data frame that is the output of measuresil(, return="all"), containing the center and the diameter of the silhouette at each value for x.
flip	Whether to flip axes (useful if measuresil() was performed using align=" v ", defaults to FALSE.
add	Whether to add to an existing plot
	Other parameters to pass on to plot() or lines()

Value

A plotted silhuette

```
fdir <- system.file(package="gdi")
measuresil(file.path(fdir,"exdata","lat.png"), return="all")->lat_
plot.sil(lat_)
```

10 sellipse.coo

sellipse

Estimate area of a superellipse. Assistant function for gdi.

Description

Estimate area of a superellipse. Assistant function for gdi.

Usage

```
sellipse(a, b, k)
```

Arguments

a First radius of the superellipse.
 b Second radius of the superellipse.
 k superellipse exponent.

Value

A single number giving the area of the superellipse

Examples

```
major_radius<-2
minor_radius<-3
exponent<-2.3
sellipse(major_radius, minor_radius, exponent)</pre>
```

sellipse.coo

calculate coordinates for plotting a superellipse for visualizing body cross-sections

Description

calculate coordinates for plotting a superellipse for visualizing body cross-sections

Usage

```
sellipse.coo(k, res = 100)
```

Arguments

k superellipse exponent.
res the desired resolution

vCOM 11

Value

a data frame containing

Examples

```
sellipse.coo(2.0)->df #get coordinates for normal ellipse (exponent k=2) plot(df$x,df$y,col="black", type="l") #plot normal ellipse sellipse.coo(2.3)->df2 # get coordinates for superellipse with exponent 2.3 lines(df$x,df$y, col="blue") #plot superellipse
```

vCOM

Finds the vertical (y axis, i.e. the axis parallel to the cross-section diameter) position of the center of mass (COM) of the volume. Experimental; only valid for "raw" gdi results with segment volumes approximated as elliptical prisms, or for manually supplied segment COMs. COM is calculated as a weighted mean of all segment COMs, with the segment mass as the weighting factor. Estimates have lower accuracy compared to hCOM, because cross-sectional geometry and variation in density throughout the cross-section is not taken into account.

Description

Finds the vertical (y axis, i.e. the axis parallel to the cross-section diameter) position of the center of mass (COM) of the volume. Experimental; only valid for "raw" gdi results with segment volumes approximated as elliptical prisms, or for manually supplied segment COMs. COM is calculated as a weighted mean of all segment COMs, with the segment mass as the weighting factor. Estimates have lower accuracy compared to hCOM, because cross-sectional geometry and variation in density throughout the cross-section is not taken into account.

Usage

```
vCOM(
   y,
   volumes = NULL,
   subtract = NULL,
   densities = NULL,
   scale = 1,
   from_top = FALSE
)
```

Arguments

A data.frame that is the output of gdi(..., return="all"), or a numeric vector containing vertical COM positions for segments

volumes An optional separate vector or data.frame (output of gdi(...,return="all") or vec-

tor of volumes.

vCOM

An optional separate vector of volumes, with length equal to the length or nrow() subtract of x, to be subtracted from the volumes for the COM calculation. densities An optional vector of segment densities, with length equal to the length or nrow() of x, to be multiplied with the volumes for the COM calculation. If both subtract and densities are supplied, the density is applied only to the "residual" volume that is left after subtraction. Optional scale value (number of pixels to chosen unit of measurement) scale from_top Whether the output coordinate should be measured from the top of the image (standard for image processing software), if TRUE, or from the bottom (standard for plotting in R (if FALSE, default). If TRUE, an attribute to y, containing the vertical dimension relative to which the measurement should be taken is required.

Value

An object of class numeric() containing the y coordinate of the center of mass of the shape, in pixels (or chosen units, if manually calculated)

```
fdir <- system.file(package="gdi")
measuresil(file.path(fdir,"exdata","lat.png"), return="all")->lat_
measuresil(file.path(fdir,"exdata","dors.png"), return="all")->dors_
gdi(lat_, dors_, return="all")->gdiresults
vCOM(gdiresults)
```

Index

```
cscorr, 2
fdetect, 3
gdi, 4
hCOM, 5
imghist, 7
measuresil, 8
plot.sil, 9
sellipse, 10
sellipse.coo, 10
vCOM, 11
```