Package 'Morpho'

December 6, 2023

Type Package

Title Calculations and Visualisations Related to Geometric Morphometrics

Version 2.12

Date 2023-12-04

Description A toolset for Geometric Morphometrics and mesh processing. This includes (among other stuff) mesh deformations based on reference points, permutation tests, detection of outliers, processing of sliding semi-landmarks and semi-automated surface landmark placement.

Suggests car, lattice, shapes, testthat

Depends R (>= 3.2.0)

Imports Rvcg (>= 0.7), rgl (>= 0.100.18), foreach (>= 1.4.0), Matrix (>= 1.0-1), MASS, parallel, doParallel (>= 1.0.6), colorRamps, Rcpp, graphics, grDevices, methods, stats, utils, jsonlite, sf, bezier

LinkingTo Rcpp, RcppArmadillo (>= 0.4)

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BugReports https://github.com/zarquon42b/Morpho/issues

LazyLoad yes

URL https://github.com/zarquon42b/Morpho

Encoding UTF-8 **RoxygenNote** 7.2.3

NeedsCompilation yes

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Repository CRAN

Date/Publication 2023-12-06 15:20:07 UTC

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Description

A toolbox for Morphometric calculations. Including sliding operations for Semilandmarks, importing, exporting and manipulating of 3D-surface meshes and semi-automated placement of surface landmarks.

Details

Package: Morpho
Type: Package
Version: 2.12
Date: 2023-12-04
License: GPL
LazyLoad: yes

Note

The pdf-version of Morpho-help can be obtained from CRAN on https://cran.r-project.org/package=Morpho

For more advanced operations on triangular surface meshes, check out my package Rvcg: https://cran.r-project.org/package=Rvcg or the code repository on github https://github.com/zarquon42b/Rvcg

Author(s)

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Maintainer: Stefan Schlager <zarquon42@gmail.com>

6 align2procSym

References

Schlager S. 2013. Soft-tissue reconstruction of the human nose: population differences and sexual dimorphism. PhD thesis, Universitätsbibliothek Freiburg. URL: http://www.freidok.uni-freiburg.de/volltexte/9181/.

align2procSym

align new data to an existing Procrustes registration

Description

align new data to an existing Procrustes registration

Usage

```
align2procSym(x, newdata, orp = TRUE)
```

Arguments

x result of a procSym call

newdata matrix or array of with landmarks corresponding to the data aligned in x

orp logical: allows to skip orthogonal projection, even if it was used in the procSym

call.

Value

an array with data aligned to the mean shape in x (and projected into tangent space)

Note

this will never yield the same result as a pooled Procrustes analysis because the sample mean is iteratively updated and new data would change the mean.

```
require(Morpho)
data(boneData)
# run procSym on entire data set
proc <- procSym(boneLM)
# this is the training data
array1 <- boneLM[,,1:60]
newdata <- boneLM[,,61:80]
proc1 <- procSym(array1)
newalign <- align2procSym(proc1,newdata)
## compare alignment for one specimen to Proc. registration using all data
## Not run:
deformGrid3d(newalign[,,1],proc$orpdata[,,61])
## End(Not run)</pre>
```

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angle.calc

calculate angle between two vectors

Description

calculates unsigned angle between two vectors

Usage

```
angle.calc(x, y)
```

Arguments

x numeric vector (or matrix to be interpreted as vector)

y numeric vector (or matrix to be interpreted as vector) of same length as x

Value

angle between x and y in radians.

Examples

```
#calculate angle between two centered and
# superimposed landmark configuration
data(boneData)
opa <- rotonto(boneLM[,,1],boneLM[,,2])
angle.calc(opa$X, opa$Y)</pre>
```

 ${\tt angleTest}$

Test whether the direction of two vectors is similar

Description

Test whether the direction of two vectors is similar

Usage

```
angleTest(x, y)
```

Arguments

x vector y vector 8 anonymize

Details

Under the assumption of all (normalized) n-vectors being represented by an n-dimensional hypersphere, the probability of the angle between two vectors is <= the measured values can be estimated as the area of a cap defined by that angle and divided by the hypersphere's complete surface area.

Value

a list with

angle angle between vectors

p.value p-value for the probability that the angle between two random vectors is smaller or equal to the one calculatted from x and y

References

S. Li, 2011. Concise Formulas for the Area and Volume of a Hyperspherical Cap. Asian Journal of Mathematics & Statistics, 4: 66-70.

Examples

```
x \leftarrow c(1,0); y \leftarrow c(1,1) # for a circle this should give us p = 0.25 as the angle between vectors ## is pi/4 and for any vector the segment +-pi/4 covers a quarter of the circle angleTest(x,y)
```

anonymize

Replace ID-strings of data and associated files.

Description

Replace ID-strings with for digits - e.g. for blind observer error testing.

```
anonymize(
  data,
  remove,
  path = NULL,
  dest.path = NULL,
  ext = ".ply",
  split = "_",
  levels = TRUE,
  prefix = NULL,
  suffix = NULL,
  sample = TRUE
)
```

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Arguments

data	Named array, matrix or vector containing data.
remove	integer: which entry (separated by split) of the name is to be removed
path	Path of associated files to be copied to renamed versions.
dest.path	where to put renamed files.
ext	file extension of files to be renamed.
split	character: by which to split specimen-ID
levels	logical: if a removed entry is to be treated as a factor. E.g. if one specimen has a double entry, the anonymized versions will be named accordingly.
prefix	character: prefix before the alias string.
suffix	character: suffix after the alias ID-string.
sample	logical: whether to randomize alias ID-string.

Value

data with names replaced

anonymkey map of original name and replaced name

Examples

```
anonymize(iris,remove=1)
```

applyTransform

apply affine transformation to data

Description

apply affine transformation to data

```
applyTransform(x, trafo, ...)
## S3 method for class 'matrix'
applyTransform(x, trafo, inverse = FALSE, threads = 1, ...)
## S3 method for class 'mesh3d'
applyTransform(x, trafo, inverse = FALSE, threads = 1, ...)
## Default S3 method:
applyTransform(x, trafo, inverse = FALSE, threads = 1, ...)
```

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Arguments

x matrix or mesh3d

trafo 4x4 transformation matrix or an object of class "tpsCoeff"

... additional arguments, currently not used.

inverse logical: if TRUE, the inverse of the transformation is applied (for TPS coeffi-

cients have to be recomputed)

threads threads to be used for parallel execution in tps deformation.

Value

the transformed object

See Also

```
rotonto, link{rotmesh.onto}, tps3d, computeTransform
```

Examples

```
data(boneData)
rot <- rotonto(boneLM[,,1],boneLM[,,2])
trafo <- getTrafo4x4(rot)
boneLM2trafo <- applyTransform(boneLM[,,2],trafo)</pre>
```

areaSphere

compute the area of an n-dimensional hypersphere

Description

compute the area of an n-dimensional hypersphere

Usage

```
areaSphere(n, r = 1)
```

Arguments

n dimensionality of space the hypersphere is embedded in (e.g.3 for a 3D-sphere)

r radius of the sphere

Value

returns the area

```
areaSphere(2) #gives us the circumference of a circle of radius 1
```

areaSpherePart 11

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compute the area of an n-dimensional hypersphere cap

Description

compute the area of an n-dimensional hypersphere cap

Usage

```
areaSpherePart(n, phi, r = 1)
```

Arguments

n dimensionality of space the hypersphere is embedded in (e.g.3 for a 3D-sphere)

phi angle between vectors defining the cone

r radius of the sphere

Value

returns the area of the hypersphere cap

Examples

```
areaSpherePart(2,pi/2) # covers half the area of a circle
```

armaGinv

calculate Pseudo-inverse of a Matrix using RcppArmadillo

Description

a simple wrapper to call Armadillo's pinv function

Usage

```
armaGinv(x, tol = NULL)
```

Arguments

x numeric matrix

tol numeric: maximum singular value to be considered

Value

Pseudo-inverse

12 arrMean3

Examples

```
mat <- matrix(rnorm(12),3,4)
pinvmat <- armaGinv(mat)</pre>
```

array2list

reverts list2array, converting an array to a list of matrices

Description

reverts list2array, converting an array to a list of matrices

Usage

```
array2list(x)
```

Arguments

Х

array

Value

returns a list containing the matrices

arrMean3

calculate mean of an array

Description

calculate mean of a 3D-array (e.g. containing landmarks) (fast) using the Armadillo C++ Backend

Usage

```
arrMean3(arr)
```

Arguments

arr

k x m x n dimensional numeric array

Value

matrix of dimensions k x m.

Note

this is the same as apply(arr, 1:2, mean), only faster for large configurations.

asymPermute 13

Examples

```
data(boneData)
proc <- ProcGPA(boneLM, silent = TRUE)
mshape <- arrMean3(proc$rotated)</pre>
```

asymPermute Assess differences in amount and direction of asymmetric variation (only object symmetry)

Description

Assess differences in amount and direction of asymmetric variation (only object symmetry)

Usage

```
asymPermute(x, groups, rounds = 1000, which = NULL)
```

Arguments

X	object of class symproc result from calling procSym with pairedLM specified
groups	factors determining grouping.
rounds	number of permutations
which	select which factorlevels to use, if NULL, all pairwise differences will be assessed after shuffling pooled data.

Value

dist	difference between vector lengths of group means
angle	angle (in radians) between vectors of group specific asymmetric deviation
means	actual group averages
p.dist	p-value obtained by comparing the actual distance to randomly acquired distances
p.angle	p-value obtained by comparing the actual angle to randomly acquired angles
permudist	vector containing differences between random group means' vector lenghts
permuangle	vector containing angles between random group means' vectors
groupmeans	array with asymmetric displacement per group
levels	character vector containing the factors used

Note

This test is only sensible if between-group differences concerning directional asymmetry have been established (e.g. by applying a MANOVA on the "asymmetric" PCscores (see also procSym) and one wants to test whether these can be attributed to differences in amount and/or direction of asymmetric displacement. Careful interpretation for very small amounts of directional asymmetry is advised. The Null-Hypothesis is that we have the same directional asymmetry in both groups. If you want to test whether the angle between groups is similar, please use angleTest.

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See Also

procSym

barycenter

calculates the barycenters for all faces of a triangular mesh

Description

calculates the barycenters for all faces of a triangular mesh

Usage

```
barycenter(mesh)
```

Arguments

mesh

triangular mesh of class 'mesh3d'

Value

k x 3 matrix of barycenters for all k faces of input mesh.

See Also

closemeshKD

```
data(nose)
bary <- barycenter(shortnose.mesh)
## Not run:
require(rgl)
##visualize mesh
wire3d(shortnose.mesh)
# visualize barycenters
points3d(bary, col=2)
## now each triangle is equipped with a point in its barycenter
## End(Not run)</pre>
```

bindArr 15

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concatenate multiple arrays/matrices

Description

concatenate multiple 3-dimensional arrays and/or 2-dimensional matrices to one big array

Usage

```
bindArr(..., along = 1, collapse = FALSE)
```

Arguments

... matrices and/or arrays with appropriate dimensionality to combine to one array,

or a single list containing suitable matrices, or arrays).

along dimension along which to concatenate.

collapse logical: if the resulting array is shallow (only 1 dimension deep), it is converted

to a matrix.

Details

dimnames, if present and if differing between entries, will be concatenated, separated by a "_".

Value

returns array of combined matrices/arrays

See Also

```
cbind, rbind, array
```

```
A <- matrix(rnorm(18),6,3)
B <- matrix(rnorm(18),6,3)
C <- matrix(rnorm(18),6,3)

#combine to 3D-array
newArr <- bindArr(A,B,C,along=3)
#combine along first dimension
newArr2 <- bindArr(newArr,newArr,along=1)</pre>
```

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boneData	Landmarks and a triangular mesh	

Description

Landmarks on the osseous human nose and a triangular mesh representing this structure.

Format

boneLM: A 10x3x80 array containing 80 sets of 3D-landmarks placed on the human osseous nose. skull_0144_ch_fe.mesh: The mesh representing the area of the first individual of boneLM

CAC

calculate common allometric component

Description

calculate common allometric component

Usage

```
CAC(x, size, groups = NULL, log = FALSE)
```

Arguments

X	datamatrix (e.g.	with PC-scores)	or 3D-array with	landmark coordinates
---	------------------	-----------------	------------------	----------------------

size vector with Centroid sizes

groups grouping variable

log logical: use log(size)

Value

CACscores	common allometric component scores

CAC common allometric component

x (group-) centered data

sc CAC reprojected into original space by applying CAC ** x

RSCscores residual shape component scores

RSC residual shape components

gmeans groupmeans

CS the centroid sizes (log transformed if log = TRUE)

cExtract 17

References

Mitteroecker P, Gunz P, Bernhard M, Schaefer K, Bookstein FL. 2004. Comparison of cranial ontogenetic trajectories among great apes and humans. Journal of Human Evolution 46(6):679-97.

Examples

```
data(boneData)
proc <- procSym(boneLM)
pop.sex <- name2factor(boneLM,which=3:4)
cac <- CAC(proc$rotated,proc$size,pop.sex)
plot(cac$CACscores,cac$size)#plot scores against Centroid size
cor.test(cac$CACscores,cac$size)#check for correlation
#visualize differences between large and small on the sample's consensus
## Not run:
large <- restoreShapes(max(cac$CACscores),cac$CAC,proc$mshape)
small <- restoreShapes(min(cac$CACscores),cac$CAC,proc$mshape)
deformGrid3d(small,large,ngrid=0)

## End(Not run)</pre>
cExtract

extract information about fixed landmarks, curves and patches from
and atlas generated by "landmark"
```

Description

After exporting the pts file of the atlas from "landmark" and importing it into R via "read.pts" cExtract gets information which rows of the landmark datasets belong to curves or patches.

Usage

```
cExtract(pts.file)
```

Arguments

pts.file

either a character naming the path to a pts.file or the name of an object imported via read.pts.

Value

returns a list containing the vectors with the indices of matrix rows belonging to the in "landmark" defined curves, patches and fix landmarks and a matrix containing landmark coordinates.

Author(s)

Stefan Schlager

See Also

```
read.lmdta,read.pts
```

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checkLM	Visually browse through a sample rendering its landmarks and corresponding surfaces.
	sponding surjuces.

Description

Browse through a sample rendering its landmarks and corresponding surfaces. This is handy e.g. to check if the landmark projection using placePatch was successful, and to mark specific specimen.

Usage

```
checkLM(
  dat.array,
 path = NULL,
 prefix = "",
 suffix = ".ply",
  col = "white",
 pt.size = NULL,
  alpha = 1,
 begin = 1,
 render = c("w", "s"),
 point = c("s", "p"),
 add = FALSE,
 meshlist = NULL,
 Rdata = FALSE,
 atlas = NULL,
  text.lm = FALSE
)
```

Arguments

dat.array	array or list containing landmark coordinates.
path	optional character: path to files where surface meshes are stored locally. If not specified only landmarks are displayed.
prefix	<pre>prefix to attach to the filenames extracted from dimnames(dat.array)[[3]] (in case of an array), or names(dat.array) (in case of a list)</pre>
suffix	suffix to attach to the filenames extracted from dimnames(dat.array)[[3]] (in case of an array), or names(dat.array) (in case of a list)
col	mesh color
pt.size	size of plotted points/spheres. If point="s". pt.size defines the radius of the spheres. If point="p" it sets the variable size used in point3d.
alpha	value between 0 and 1. Sets transparency of mesh 1=opaque 0= fully transparent.
begin	integer: select a specimen to start with.
render	if render="w", a wireframe will be drawn, else the meshes will be shaded.

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how to render landmarks. "s"=spheres, "p"=points. point add logical: add to existing rgl window. meshlist list holding meshes in the same order as dat.array (Overrides path). Rdata logical: if the meshes are previously stored as Rdata-files by calling save(), these are simply loaded and rendered. Otherwise it is assumed that the meshes are stored in standard file formats such as PLY, STL or OBJ, that are then imported with the function file2mesh. provide object generated by createAtlas to specify coloring of surface patches, atlas curves and landmarks text.lm logical: number landmarks. Only applicable when atlas=NULL.

Value

returns an invisible vector of indices of marked specimen.

See Also

```
placePatch, createAtlas, plotAtlas, file2mesh
```

```
data(nose)
###create mesh for longnose
longnose.mesh <- tps3d(shortnose.mesh,shortnose.lm,longnose.lm,threads=1)</pre>
### write meshes to disk
save(shortnose.mesh, file="shortnose")
save(longnose.mesh, file="longnose")
## create landmark array
data <- bindArr(shortnose.lm, longnose.lm, along=3)</pre>
dimnames(data)[[3]] <- c("shortnose", "longnose")</pre>
## Not run:
checkLM(data, path="./",Rdata=TRUE, suffix="")
## End(Not run)
## now visualize by using an atlas:
atlas <- createAtlas(shortnose.mesh, landmarks =</pre>
           shortnose.lm[c(1:5,20:21),],
patch=shortnose.lm[-c(1:5,20:21),])
if (interactive()){
checkLM(data, path="./",Rdata=TRUE, suffix="", atlas=atlas)
## remove data from disk
unlink("shortnose")
unlink("longnose")
```

20 classify

checkNA

check for NA values in a matrix (of landmarks)

Description

check for NA values in a matrix (of landmarks)

Usage

checkNA(x)

Arguments

Х

matrix containing landmarks

Value

returns a vector with missin landmarks and a vector of length=0 if none are missing

classify

classify specimen based on between-group PCA or CVA or typprob-Class

Description

classify specimen based on between-group PCA, CVA or typprobClass

```
classify(x, cv = TRUE, ...)
## S3 method for class 'bgPCA'
classify(x, cv = TRUE, newdata = NULL, ...)
## S3 method for class 'CVA'
classify(x, cv = T, newdata = NULL, prior = NULL, ...)
## S3 method for class 'typprob'
classify(x, cv = TRUE, ...)
```

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Arguments

X	result of groupPCA, CVA or typprobClass
cv	logical: use cross-validated scores if available
	currently not used
newdata	use new data to predict scores and evaluate group affinity
prior	specify prior probability for CVA evaluation if NULL prior from CVA will be used. Be m your number of groups then to set the prior equally for all groups set prior=rep(1,m)/m.

Value

class classification result

groups original grouping variable, only available if newdata=NULL

posterior only for object of CVA and typprob, also the posterior probabilities are returned

See Also

CVA, groupPCA, typprobClass

closemeshKD Project coordinates onto a target triangular surface mesh.

Description

For a set of 3D-coordinates the closest matches on a target surface are determined and normals at as well as distances to that point are calculated.

```
closemeshKD(
   x,
   mesh,
   k = 50,
   sign = FALSE,
   barycoords = FALSE,
   cores = 1,
   method = 0,
   ...
)
```

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Arguments

x k x 3 matrix containing 3D-coordinates or object of class mesh3d.

mesh triangular surface mesh stored as object of class mesh3d.

k neighbourhood of kd-tree to search - the larger, the slower - but the more likely

the absolutely closest point is hit.

sign logical: if TRUE, signed distances are returned.

barycoords logical: if TRUE, barycentric coordinates of the hit points are returned.

cores integer: how many cores to use for the search algorithm.

method integer: either 0 or 1, if 0 ordinary Euclidean distance is used, if 1, the distance

suggested by Moshfeghi(1994) is calculated.

... additional arguments. currently unavailable.

Details

The search for the clostest point is designed as follows: Calculate the barycenter of each target face. For each coordinate of x, determine the k closest barycenters and calculate the distances to the closest point on these faces.

Value

returns an object of class mesh3d. with:

vb 4xn matrix containing n vertices as homolougous coordinates

normals 4xn matrix containing vertex normals

quality vector: containing distances to target. In case of method=1, this is not the Eu-

clidean distance but the distance of the reference point to the faceplane (orthogonally projected) plus the distance to the closest point on one of the face's edges

(the target point). See the literature cited below for details.

it 4xm matrix containing vertex indices forming triangular faces. Only available,

when x is a mesh

Author(s)

Stefan Schlager

References

Baerentzen, Jakob Andreas. & Aanaes, H., 2002. Generating Signed Distance Fields From Triangle Meshes. Informatics and Mathematical Modelling.

Moshfeghi M, Ranganath S, Nawyn K. 1994. Three-dimensional elastic matching of volumes IEEE Transactions on Image Processing: A Publication of the IEEE Signal Processing Society 3:128-138.

See Also

ply2mesh

colors 23

Examples

```
data(nose)
out <- closemeshKD(longnose.lm,shortnose.mesh,sign=TRUE)
### show distances - they are very small because
###longnose.lm is scaled to unit centroid size.
hist(out$quality)</pre>
```

colors

predefined colors for bone and skin

Description

predefined colors for bone and skin

Details

available colors are:

bone1

bone2

bone3

skin1

skin2

skin3

skin4

computeArea

Compute area enclosed within an irregular polygon

Description

Compute area enclosed within an irregular polygon - i.e. defined by curves

Usage

```
computeArea(x)
```

Arguments

Х

k x 2 or k x 3 matrix containing ordered coordinates forming the boundary of the area. For 3D-cases, the area should be closed to a 2D surface (see details below).

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Details

For 3D coordinates, a PCA is computed and only the first two PCs are used to compute the area. This is a projection of the coordinates onto a 2D plane spanned by those PCs.

Value

returns a list containing

area size of the enclosed area

xpro2D projected coordinates of x in the 2D plane.

poly object of class sp as defined by the sp package.

xpro3D For 3D-cases, this contains the projected coordinates of x rotated back into the

original coordinate system

Note

in case custom planes are preferred, the data can first be projected onto such a custom defined plane via points2plane first.

Examples

```
require(shapes)
require(sf)
myarea <- computeArea(gorf.dat[c(1,6:8,2:5),,1])
myarea$area
plot(myarea$poly)

## 3D example
data(boneData)
myarea3D <- computeArea(boneLM[c(4,2,3,7,5,6,8),,1])
plot(myarea3D$poly)
cent <- colMeans(myarea3D$xpro2D)
text(cent[1],cent[2],labels=paste0("Area=",round(myarea3D$area,digits=2)))</pre>
```

computeTransform

calculate an affine transformation matrix

Description

calculate an affine transformation matrix

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Usage

```
computeTransform(
    x,
    y,
    type = c("rigid", "similarity", "affine", "tps"),
    reflection = FALSE,
    lambda = 1e-08,
    weights = NULL,
    centerweight = FALSE,
    threads = 1
)
```

Arguments

x fix landmarks. Can be a k x m matrix or mesh3d.

y moving landmarks. Can be a k x m matrix or mesh3d.

type set type of affine transformation: options are "rigid", "similarity" (rigid + scale)

and "affine",

reflection logical: if TRUE "rigid" and "similarity" allow reflections.

lambda numeric: regularisation parameter of the TPS.

weights vector of length k, containing weights for each landmark (only used in type="rigid"

or "similarity").

centerweight logical or vector of weights: if weights are defined and centerweigths=TRUE,

the matrix will be centered according to these weights instead of the barycenter. If centerweight is a vector of length nrow(x), the barycenter will be weighted

accordingly.

threads number of threads to use in TPS interpolation.

Details

x and y can also be a pair of meshes with corresponding vertices.

Value

returns a 4x4 (3x3 in 2D case) transformation matrix or an object of class "tpsCoeff" in case of type="tps".

Note

all lines containing NA, or NaN are ignored in computing the transformation.

See Also

```
rotonto, link{rotmesh.onto}, tps3d
```

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Examples

```
data(boneData)
trafo <- computeTransform(boneLM[,,1],boneLM[,,2])
transLM <- applyTransform(boneLM[,,2],trafo)</pre>
```

covDist

calculates distances and PC-coordinates of covariance matrices

Description

calculates PC-coordinates of covariance matrices by using the Riemannian metric in their respective space.

Usage

```
covDist(s1, s2)

covPCA(
   data,
   groups,
   rounds = 1000,
   bootrounds = 0,
   lower.bound = 0.05,
   upper.bound = 0.95
)
```

Arguments

s1	m x m covariance matrix
s2	m x m covariance matrix
data	matrix containing data with one row per observation
groups	factor: group assignment for each specimen
rounds	integer: rounds to run permutation of distances by randomly assigning group membership
bootrounds	integer: perform bootstrapping to generate confidence intervals (lower boundary, median and upper boundary) for PC-scores.
lower.bound	numeric: set probability (quantile) for lower boundary estimate from bootstrapping.
upper.bound	numeric: set probability (quantile) for upper boundary estimate from bootstrapping.

covDist 27

Details

covDist calculates the Distance between covariance matrices while covPCA uses a MDS (multidimensional scaling) approach to obtain PC-coordinates from a distance matrix derived from multiple groups. P-values for pairwise distances can be computed by permuting group membership and comparing actual distances to those obtained from random resampling. To calculate confidence intervals for PC-scores, within-group bootstrapping can be performed.

Value

covDist returns the distance between s1 and s2

covPCA returns a list containing:

if scores = TRUE

PCscores PCscores

eigen decomposition of the centered inner product

if rounds > 0

distance matrix

p.matrix p-values for pairwise distances from permutation testing

if bootrounds > 0

bootstrap list containing the lower and upper bound of the confidence intervals of PC-

scores as well as the median of bootstrapped values.

boot.data array containing all results generated from bootstrapping.

Author(s)

Stefan Schlager

References

Mitteroecker P, Bookstein F. 2009. The ontogenetic trajectory of the phenotypic covariance matrix, with examples from craniofacial shape in rats and humans. Evolution 63:727-737.

Hastie T, Tibshirani R, Friedman JJH. 2013. The elements of statistical learning. Springer New York.

See Also

prcomp

```
cpca <- covPCA(iris[,1:4],iris[,5])
cpca$p.matrix #show pairwise p-values for equal covariance matrices
## Not run:</pre>
```

28 covW

```
require(car)
sp(cpca$PCscores[,1],cpca$PCscores[,2],groups=levels(iris[,5]),
   smooth=FALSE,xlim=range(cpca$PCscores),ylim=range(cpca$PCscores))
data(boneData)
proc <- procSym(boneLM)</pre>
pop <- name2factor(boneLM, which=3)</pre>
## compare covariance matrices for PCscores of Procrustes fitted data
cpca1 <- covPCA(proc$PCscores, groups=pop, rounds = 1000)</pre>
## view p-values:
cpca1$p.matrix # differences between covariance matrices
# are significant
## visualize covariance ellipses of first 5 PCs of shape
spm(proc$PCscores[,1:5], groups=pop, smooth=FALSE,ellipse=TRUE, by.groups=TRUE)
## covariance seems to differ between 1st and 5th PC
## for demonstration purposes, try only first 4 PCs
cpca2 <- covPCA(proc$PCscores[,1:4], groups=pop, rounds = 1000)</pre>
## view p-values:
cpca2$p.matrix # significance is gone
## End(Not run)
#do some bootstrapping 1000 rounds
cpca <- covPCA(iris[,1:4],iris[,5],rounds=0, bootrounds=1000)</pre>
#plot bootstrapped data of PC1 and PC2 for first group
plot(t(cpca$boot.data[1,1:2,]),xlim=range(cpca$boot.data[,1,]),
                                ylim=range(cpca$boot.data[,2,]))
points(t(cpca$PCscores[1,]),col="white",pch=8,cex=1.5)##plot actual values
for (i in 2:3) {
  points(t(cpca$boot.data[i,1:2,]),col=i)##plot other groups
  points(t(cpca$PCscores[i,]),col=1,pch=8,cex=1.5)##plot actual values
}
```

covW

calculate the pooled within groups covariance matrix

Description

calculate the pooled within groups covariance matrix

```
covW(data, groups, robust = c("classical", "mve", "mcd"), ...)
```

createAtlas 29

Arguments

data a matrix containing data
groups grouping variables

robust character: determines covariance estimation methods in case sep=TRUE, when covariance matrices and group means can be estimated robustly using MASS::cov.rob.

Default is the standard product-moment covariance matrix.

additional parameters passed to MASS::cov.rob for robust covariance and mean estimations.

Value

Returns the pooled within group covariance matrix. The attributes contain the entry means, containing the respective group means.

Author(s)

Stefan Schlager

See Also

```
cov, typprobClass
```

Examples

```
data(iris)
poolCov <- covW(iris[,1:4],iris[,5])</pre>
```

createAtlas

Create an atlas needed in placePatch

Description

Create an atlas needed in placePatch

```
createAtlas(
  mesh,
  landmarks,
  patch,
  corrCurves = NULL,
  patchCurves = NULL,
  keep.fix = NULL
)
```

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Arguments

mesh triangular mesh representing the atlas' surface landmarks matrix containing landmarks defined on the atlas, as well as on each specimen in the corresponding sample. matrix containing semi-landmarks to be projected onto each specimen in the patch corresponding sample. a vector or a list containing vectors specifying the rowindices of landmarks corrCurves to be curves that are defined on the atlas AND each specimen. e.g. if landmarks 2:4 and 5:10 are two distinct curves, one would specify corrCurves = list(c(2:4), c(5:10)). a vector or a list containing vectors specifiyng the rowindices of landmarks to patchCurves be curves that are defined ONLY on the atlas. E.g. if coordinates 5:10 and 20:40 on the patch are two distinct curves, one would specify patchCurves = list(c(5:10),c(20:40)).keep.fix in case corrCurves are set, specify explicitly which landmarks are not allowed

Value

Returns a list of class "atlas". Its content is corresponding to argument names.

to slide during projection (with placePatch)

Note

This is a helper function of placePatch.

See Also

```
placePatch, plotAtlas
```

Examples

CreateL

Create Matrices necessary for Thin-Plate Spline

Description

Create (Bending Engergy) Matrices necessary for Thin-Plate Spline, and sliding of Semilandmarks

CreateL 31

Usage

```
CreateL(
  matrix,
  lambda = 1e-08,
  output = c("K", "L", "Linv", "Lsubk", "Lsubk3"),
  threads = 1
)
```

Arguments

matrix k x 3 or k x 2 matrix containing landmark coordinates.

lambda numeric: regularization factor

output character vector: select which matrices to create. Can be a vector containing

any combination of the strings: "K", "L", "Linv", "Lsubk", "Lsubk3".

threads to be used for parallel execution calculating K. sliding of semilandmarks.

Value

depending on the choices in output:

Matrix K as specified in Bookstein (1989)

L Matrix L as specified in Bookstein (1989)

Linv Inverse of matrix L as specified in Bookstein (1989)

Lsubk uper left k x k submatrix of Linv

Lsubk3 Matrix used for sliding in slider3d and relaxLM

.

Note

This function is not intended to be called directly - except for playing around to grasp the mechansims of the Thin-Plate Spline.

References

Gunz, P., P. Mitteroecker, and F. L. Bookstein. 2005. Semilandmarks in Three Dimensions, in Modern Morphometrics in Physical Anthropology. Edited by D. E. Slice, pp. 73-98. New York: Kluwer Academic/Plenum Publishers.

Bookstein FL. 1989. Principal Warps: Thin-plate splines and the decomposition of deformations. IEEE Transactions on pattern analysis and machine intelligence 11(6).

See Also

tps3d

32 createMissingList

Examples

```
data(boneData)
L <- CreateL(boneLM[,,1])
## calculate Bending energy between first and second specimen:
be <- t(boneLM[,,2])%*%L$Lsubk%*%boneLM[,,2]
## calculate Frobenius norm
sqrt(sum(be^2))
## the amount is dependant on on the squared scaling factor
# scale landmarks by factor 5 and compute bending energy matrix
be2 <- t(boneLM[,,2]*5)%*%L$Lsubk%*%(boneLM[,,2]*5)
sqrt(sum(be2^2)) # exactly 25 times the result from above
## also this value is not symmetric:
L2 <- CreateL(boneLM[,,2])
be3 <- t(boneLM[,,1])%*%L2$Lsubk%*%boneLM[,,1]
sqrt(sum(be3^2))</pre>
```

createMissingList

create a list with empty entries to be used as missingList in slider3d

Description

create a list with empty entries to be used as missingList in slider3d

Usage

```
createMissingList(x)
```

Arguments

Χ

length of the list to be created

Value

returns a list of length x filled with numerics of length zero.

See Also

```
fixLMtps, fixLMmirror, slider3d
```

```
## Assume in a sample of 10, the 9th individual has (semi-)landmarks 10:50
# hanging in thin air (e.g. estimated using fixLMtps)
# while the others are complete.
## create empty list
missingList <- createMissingList(10)
missingList[[9]] <- 10:50</pre>
```

crossProduct 33

crossProduct

calculate the orthogonal complement of a 3D-vector

Description

calculate the orthogonal complement of a 3D-vector

Usage

```
crossProduct(x, y, normalize = TRUE)
tangentPlane(x)
```

Arguments

x vector of length 3. y vector of length 3.

normalize logical: if TRUE, the resulting vector is normalized

Details

calculate the orthogonal complement of a 3D-vector or the 3D-crossproduct, finding an orthogonal vector to a plane in 3D.

Value

```
tangentPlane:
```

crossProduct: returns a vector of length 3.

y vector orthogonal to x

z vector orthogonal to x and y

Author(s)

Stefan Schlager

```
require(rgl)

x <- c(1,0,0)
y <- c(0,1,0)

#example tangentPlane
z <- tangentPlane(x)
#visualize result
## Not run:</pre>
```

cSize cSize

```
lines3d(rbind(0, x), col=2, lwd=2)
## show complement
lines3d(rbind(z$y, 0, z$z), col=3, lwd=2)
## End(Not run)
# example crossProduct
z <- crossProduct(x, y)
# show x and y
## Not run:
lines3d(rbind(x, 0, y), col=2, lwd=2)
# show z
lines3d(rbind(0, z), col=3, lwd=2)
## End(Not run)</pre>
```

cSize

calculate Centroid Size for a landmark configuration

Description

calculate Centroid Size for a landmark configuration

Usage

cSize(x)

Arguments

Х

k x 3 matrix containing landmark coordinates or mesh of class "mesh3d"

Value

returns Centroid size

```
data(boneData)
cSize(boneLM[,,1])
```

cutMeshPlane 35

cutMeshPlane	cut a mesh by a hyperplane and remove parts above/below that plane

Description

cut a mesh by a hyperplane and remove parts above/below that plane

Usage

```
cutMeshPlane(mesh, v1, v2 = NULL, v3 = NULL, normal = NULL, keep.upper = TRUE)
```

Arguments

mesh	triangular mesh of class "mesh3d"
v1	numeric vector of length=3 specifying a point on the separating plane
v2	numeric vector of length=3 specifying a point on the separating plane
v3	numeric vector of length=3 specifying a point on the separating plane
normal	plane normal (overrides specification by v2 and v3)
keep.upper	logical specify whether the points above or below the plane are should be kept

Details

see cutSpace for more details.

Value

mesh with part above/below hyperplane removed

Description

```
separate a 3D-pointcloud by a hyperplane
```

```
cutSpace(pointcloud, v1, v2 = NULL, v3 = NULL, normal = NULL, upper = TRUE)
```

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Arguments

pointcloud	numeric n x 3 matrix
v1	numeric vector of length=3 specifying a point on the separating plane
v2	numeric vector of length=3 specifying a point on the separating plane
v3	numeric vector of length=3 specifying a point on the separating plane
normal	plane normal (overrides specification by v2 and v3)
upper	logical specify whether the points above or below the plane are to be reported as TRUE.

Details

As above and below are specified by the normal calculated from $(v2-v1)\times(v3-v1)$, where \times denotes the vector crossproduct. This means the normal points "upward" when viewed from the positon where v1, v2 and v3 are arranged counter-clockwise. Thus, which side is "up" depends on the ordering of v1, v2 and v3.

Value

logical vector of length n. Reporting for each point if it is above or below the hyperplane

Examples

```
data(nose)
v1 <- shortnose.lm[1,]
v2 <- shortnose.lm[2,]</pre>
v3 <- shortnose.lm[3,]</pre>
pointcloud <- vert2points(shortnose.mesh)</pre>
upper <- cutSpace(pointcloud, v1, v2, v3)</pre>
## Not run:
require(rgl)
normal <- crossProduct(v2-v1,v3-v1)</pre>
zeroPro <- points2plane(rep(0,3),v1,normal)</pre>
## get sign of normal displacement from zero
sig <- sign(crossprod(-zeroPro,normal))</pre>
d <- sig*norm(zeroPro,"2")</pre>
planes3d(normal[1],normal[2],normal[3],d=d)
points3d(pointcloud[upper,])
## End(Not run)
```

 CVA

Canonical Variate Analysis

Description

performs a Canonical Variate Analysis.

CVA 37

Usage

```
CVA(
   dataarray,
   groups,
   weighting = TRUE,
   tolinv = 1e-10,
   plot = TRUE,
   rounds = 0,
   cv = FALSE,
   p.adjust.method = "none",
   robust = c("classical", "mve", "mcd"),
   prior = NULL,
   ...
)
```

Arguments

dataarray Either a k x m x n real array, where k is the number of points, m is the number

of dimensions, and n is the sample size. Or alternatively a n x m Matrix where n is the number of observations and m the number of variables (this can be PC

scores for example)

groups a character/factor vector containing grouping variable.

weighting Logical: Determines whether the between group covariance matrix and Grand-

mean is to be weighted according to group size.

tolinv Threshold for the eigenvalues of the pooled within-group-covariance matrix to

be taken as zero - for calculating the general inverse of the pooled withing groups

covariance matrix.

plot Logical: determins whether in the two-sample case a histogramm ist to be plot-

ted.

rounds integer: number of permutations if a permutation test of the Mahalanobis dis-

tances (from the pooled within-group covariance matrix) and Euclidean distance

between group means is requested If rounds = 0, no test is performed.

cv logical: requests a Jackknife Crossvalidation.

p.adjust.method

method to adjust p-values for multiple comparisons see p.adjust.methods for

options.

robust character: determines covariance estimation methods, allowing for robust esti-

mations using MASS::cov.rob

prior vector assigning each group a prior probability.

.. additional parameters passed to MASS::cov.rob for robust covariance and mean

estimations

Value

CV A matrix containing the Canonical Variates

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CVscores A matrix containing the individual Canonical Variate scores

Grandm a vector or a matrix containing the Grand Mean (depending if the input is an

array or a matrix)

groupmeans a matrix or an array containing the group means (depending if the input is an

array or a matrix)

Variance explained by the Canonical Variates

CVvis Canonical Variates projected back into the original space - to be used for visual-

ization purposes, for details see example below

Dist Mahalanobis Distances between group means - if requested tested by permuta-

tion test if the input is an array it is assumed to be superimposed Landmark Data

and Procrustes Distance will be calculated

CVcv A matrix containing crossvalidated CV scores

groups factor containing the grouping variable

class classification results based on posteriror probabilities. If cv=TRUE, this will be

done by a leaving-one-out procedure

posterior posterior probabilities
prior prior probabilities

Author(s)

Stefan Schlager

References

Cambell, N. A. & Atchley, W. R.. 1981 The Geometry of Canonical Variate Analysis: Syst. Zool., 30(3), 268-280.

Klingenberg, C. P. & Monteiro, L. R. 2005 Distances and directions in multidimensional shape spaces: implications for morphometric applications. Systematic Biology 54, 678-688.

See Also

groupPCA

Examples

```
## all examples are kindly provided by Marta Rufino
if (require(shapes)) {
# perform procrustes fit on raw data
alldat<-procSym(abind(gorf.dat,gorm.dat))
# create factors
groups<-as.factor(c(rep("female",30),rep("male",29)))
# perform CVA and test Mahalanobis distance
# between groups with permutation test by 100 rounds)
cvall<-CVA(alldat$orpdata,groups,rounds=10000)
## visualize a shape change from score -5 to 5:</pre>
```

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```
cvvis5 <- 5*matrix(cvall$CVvis[,1],nrow(cvall$Grandm),ncol(cvall$Grandm))+cvall$Grandm</pre>
cvvisNeg5 <- -5*matrix(cvall$CVvis[,1],nrow(cvall$Grandm),ncol(cvall$Grandm))+cvall$Grandm</pre>
plot(cvvis5,asp=1)
points(cvvisNeg5,col=2)
for (i in 1:nrow(cvvisNeg5))
 lines(rbind(cvvis5[i,],cvvisNeg5[i,]))
### Morpho CVA
data(iris)
vari <- iris[,1:4]</pre>
facto <- iris[,5]</pre>
cva.1=CVA(vari, groups=facto)
## get the typicality probabilities and resulting classifications - tagging
## all specimens with a probability of < 0.01 as outliers (assigned to no class)
typprobs <- typprobClass(cva.1$CVscores,groups=facto)</pre>
print(typprobs)
## visualize the CV scores by their groups estimated from (cross-validated)
## typicality probabilities:
if (require(car)) {
scatterplot(cva.1$CVscores[,1],cva.1$CVscores[,2],groups=typprobs$groupaffinCV,
                  smooth=FALSE,reg.line=FALSE)
}
# plot the CVA
plot(cva.1$CVscores, col=facto, pch=as.numeric(facto), typ="n",asp=1,
  xlab=paste("1st canonical axis", paste(round(cva.1$Var[1,2],1),"%")),
  ylab=paste("2nd canonical axis", paste(round(cva.1$Var[2,2],1),"%")))
  text(cva.1$CVscores, as.character(facto), col=as.numeric(facto), cex=.7)
 # add chull (merge groups)
 for(jj in 1:length(levels(facto))){
        ii=levels(facto)[jj]
   kk=chull(cva.1$CVscores[facto==ii,1:2])
   lines(cva.1$CVscores[facto==ii,1][c(kk, kk[1])],
   cva.1$CVscores[facto==ii,2][c(kk, kk[1])], col=jj)
    }
 # add 80% ellipses
 if (require(car)) {
  for(ii in 1:length(levels(facto))){
    dataEllipse(cva.1$CVscores[facto==levels(facto)[ii],1],
    cva.1$CVscores[facto==levels(facto)[ii],2],
                    add=TRUE,levels=.80, col=c(1:7)[ii])}
 # histogram per group
 if (require(lattice)) {
 histogram(~cva.1$CVscores[,1]|facto,
 layout=c(1,length(levels(facto))),
          xlab=paste("1st canonical axis", paste(round(cva.1$Var[1,2],1),"%")))
 histogram(~cva.1$CVscores[,2]|facto, layout=c(1,length(levels(facto))),
          xlab=paste("2nd canonical axis", paste(round(cva.1$Var[2,2],1),"%")))
 }
```

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```
# plot Mahalahobis
     dendroS=hclust(cva.1$Dist$GroupdistMaha)
     dendroS$labels=levels(facto)
     par(mar=c(4,4.5,1,1))
     dendroS=as.dendrogram(dendroS)
     plot(dendroS, main='', sub='', xlab="Geographic areas",
                               ylab='Mahalahobis distance')
         # Variance explained by the canonical roots:
         cva.1$Var
         # or plot it:
         barplot(cva.1$Var[,2])
# another landmark based example in 3D:
data(boneData)
groups <- name2factor(boneLM,which=3:4)</pre>
proc <- procSym(boneLM)</pre>
cvall<-CVA(proc$orpdata,groups)</pre>
#' ## visualize a shape change from score -5 to 5:
cvvis5 <- 5*matrix(cvall$CVvis[,1],nrow(cvall$Grandm),ncol(cvall$Grandm))+cvall$Grandm</pre>
\verb|cvvisNeg5| < -5*| \texttt{matrix}(cvall\$CVvis[,1],nrow(cvall\$Grandm),ncol(cvall\$Grandm)) + cvall\$Grandm| | \texttt{model}(cvall\$Grandm) | \texttt{model}(cvall) | \texttt{model}(cv
## Not run:
#visualize it
deformGrid3d(cvvis5,cvvisNeg5,ngrid = 0)
## End(Not run)
#for using (e.g. the first 5) PCscores, one will do:
cvall <- CVA(proc$PCscores[,1:5],groups)</pre>
#' ## visualize a shape change from score -5 to 5:
cvvis5 <- 5*cvall$CVvis[,1]+cvall$Grandm</pre>
cvvisNeg5 <- -5*cvall$CVvis[,1]+cvall$Grandm</pre>
cvvis5 <- restoreShapes(cvvis5,proc$PCs[,1:5],proc$mshape)</pre>
cvvisNeg5 <- restoreShapes(cvvisNeg5,proc$PCs[,1:5],proc$mshape)</pre>
## Not run:
#visualize it
deformGrid3d(cvvis5,cvvisNeg5,ngrid = 0)
## End(Not run)
```

data2platonic

creates 3D shapes from data to be saved as triangular meshes

Description

creates 3D shapes from 3-dimensional data that can be saved as triangular meshes

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Usage

```
data2platonic(
  datamatrix,
  shape = Rvcg::vcgSphere(),
  col = "red",
  scale = FALSE,
  scalefactor = 1
)
```

Arguments

datamatrix k x 3 data matrix
shape a 3D shape
col color value
scale logical: whether to scale the data to unit sd.
scalefactor scale the resulting shapes.

Value

returns all shapes merged into a single mesh

Examples

deformGrid2d

visualise differences between two superimposed sets of 2D landmarks

Description

visualise differences between two superimposed sets of 2D landmarks by deforming a square grid based on a thin-plate spline interpolation

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Usage

```
deformGrid2d(
 matrix,
  tarmatrix,
 ngrid = 0,
 lwd = 1,
  show = c(1:2),
 lines = TRUE,
 lcol = 1,
 lty = 2,
 col1 = 2,
 col2 = 3,
 pcaxis = FALSE,
 add = FALSE,
 wireframe = NULL,
 margin = 0.2,
 gridcol = "grey",
 gridlty = 1,
 cex1 = 1,
 cex2 = 1,
)
```

Arguments

matrix	reference matrix containing 2D landmark coordinates or mesh of class "mesh3d"
tarmatrix	target matrix containing 2D landmark coordinates or mesh of class "mesh3d"
ngrid	number of grid lines to be plotted; ngrid=0 suppresses grid creation.
lwd	width of lines connecting landmarks.
show	integer (vector): if $c(1:2)$ both configs will be plotted, show = 1 only plots the reference and show = 2 the target. plotted. Options are combinations of 1,2 and 3.
lines	logical: if TRUE, lines between landmarks will be plotted.
lcol	color of lines
lty	line type
col1	color of "matrix"
col2	color of "tarmat"
pcaxis	logical: align grid by shape's principal axes.
add	logical: if TRUE, output will be drawn on existing plot.
wireframe	list/vector containing row indices to be plotted as wireframe (see lineplot.)
margin	margin around the bounding box to draw the grid
gridcol	color of the grid
gridlty	linetype for grid

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```
cex1 control size of points belonging to matrix
cex2 control size of points belonging to tarmatrix
... additional parameters passed to plot
```

Value

if ngrid > 1 the coordinates of the displaced grid knots are returned.

Author(s)

Stefan Schlager

See Also

tps3d

Examples

```
if (require(shapes)) {
proc <- procSym(gorf.dat)
deformGrid2d(proc$mshape,proc$rotated[,,1],ngrid=5,pch=19)
}</pre>
```

deformGrid3d

visualise differences between two superimposed sets of 3D landmarks

Description

visualise differences between two superimposed sets of 3D landmarks by deforming a cubic grid based on a thin-plate spline interpolation

Usage

```
deformGrid3d(
  matrix,
  tarmatrix,
  ngrid = 0,
  align = FALSE,
  lwd = 1,
  showaxis = c(1, 2),
  show = c(1, 2),
  lines = TRUE,
  lcol = 1,
  add = FALSE,
  col1 = 2,
  col2 = 3,
  type = c("s", "p"),
```

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```
size = NULL,
pcaxis = FALSE,
ask = TRUE,
margin = 0.2,
createMesh = FALSE,
slice1 = NULL,
slice2 = NULL,
slice3 = NULL,
gridcol = 1,
gridwidth = 1,
...
)
```

Arguments

matrix	reference matrix containing 3D landmark coordinates or mesh of class "mesh3d"
tarmatrix	target matrix containing 3D landmark coordinates or mesh of class "mesh3d"
ngrid	number of grid lines to be plotted; ngrid=0 suppresses grid creation.
align	logical: if TRUE, tarmatrix will be aligned rigidly to matrix
lwd	width of lines connecting landmarks.
showaxis	integer (vector): which dimensions of the grid to be plotted. Options are combinations of 1,2 and 3.
show	integer (vector): if $c(1:2)$ both configs will be plotted, show = 1 only plots the reference and show = 2 the target
lines	logical: if TRUE, lines between landmarks will be plotted.
lcol	color of lines
add	logical: add to existing rgl window.
col1	color of "matrix"
col2	color of "tarmat"
type	"s" renders landmarks as spheres; "p" as points - much faster for very large pointclouds.
size	control size/radius of points/spheres
pcaxis	logical: align grid by shape's principal axes.
ask	logical: if TRUE for > 1000 coordinates the user will be asked to prefer points over spheres.
margin	margin around the bounding box to draw the grid
createMesh	logical: if TRUE, a triangular mesh of spheres and displacement vectors (can take some time depending on number of reference points and grid density).
slice1	integer or vector of integers: select slice(s) for the dimensions
slice2	integer or vector of integers: select slice(s) for the dimensions
slice3	integer or vector of integers: select slice(s) for the dimensions
gridcol	define color of grid
gridwidth	integer: define linewidth of grid
	additional parameters passed to rotonto in case align=TRUE

equidistantCurve 45

Value

if createMesh=TRUE, a mesh containing spheres of reference and target as well as the displacement vectors is returned. Otherwise the knots of the displaced grid is returned.

Author(s)

Stefan Schlager

See Also

tps3d

Examples

```
if (interactive()){
data(nose)
deformGrid3d(shortnose.lm,longnose.lm,ngrid=10)

## select some slices
deformGrid3d(shortnose.lm,longnose.lm,showaxis=1:3,ngrid=10,slice1=2,slice2=5,slice3=7)
}
```

equidistantCurve

make a curve equidistant (optionally up/downsampling)

Description

make a curve equidistant (optionally up/downsampling)

Usage

```
equidistantCurve(
    X,
    n = NULL,
    open = TRUE,
    subsample = 0,
    increment = 2,
    smoothit = 0,
    mesh = NULL,
    iterations = 1
)
```

Arguments

x k x m matrix containing the 2D or 3D coordinates

n integer: number of coordinates to sample. If NULL, the existing curve will be made equidistant.

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open logical: specifies whether the curve is open or closed.

subsample integer: number of subsamples to draw from curve for interpolation. For curves

with < 1000 points, no subsampling is required.

increment integer: if > 1, the curve is estimated iteratively by incrementing the original

points by this factor. The closer this value to 1, the smoother the line but possibly

farther away from the control points.

smoothit integer: smoothing iterations after each step

mesh specify mesh to project point to

iterations integer: how many iterations to run equidistancing.

Details

Equidistancy is reached by iteratively deforming (using TPS) a straight line with equidistantly placed points to the target using control points with the same spacing as the actual curve. To avoid singularity, the straight line containes a small amount of noise, which can (optionally) be accounted for by smoothing the line by its neighbours.

Value

matrix containing equidistantly placed points

Note

if n » number of original points, the resulting curves can show unwanted distortions.

Examples

```
## Not run:
data(nose)
x <- shortnose.lm[c(304:323),]
xsample <- equidistantCurve(x,n=50,iterations=10,increment=2)
require(rgl)
points3d(xsample,size=5)
spheres3d(x,col=2,radius=0.3,alpha=0.5)
## End(Not run)</pre>
```

exVar

calculate variance of a distribution stemming from prediction models

Description

calculates a quotient of the overall varriance within a predicted distribution to that from the original one. This function calculates a naive extension of the univariate R^2-value by dividing the variance in the predicted dat by the variance of the original data. No additional adjustments are made!!

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Usage

```
exVar(model, ...)
## S3 method for class 'lm'
exVar(model, ...)
## S3 method for class 'mvr'
exVar(model, ncomp, val = FALSE, ...)
```

Arguments

```
model a model of classes "lm" or "mvr" (from the package "pls")
... currently unused additional arguments.

ncomp How many latent variables to use (only for mvr models)
val use cross-vaildated predictions (only for mvr models)
```

Value

returns the quotient.

Note

The result is only!! a rough estimate of the variance explained by a multivariate model. And the result can be misleading - especially when there are many predictor variables involved. If one is interested in the value each factor/covariate explains, we recommend a 50-50 MANOVA perfomed by the R-package "ffmanova", which reports this value factor-wise.

Author(s)

Stefan Schlager

References

Langsrud O, Juergensen K, Ofstad R, Naes T. 2007. Analyzing Designed Experiments with Multiple Responses Journal of Applied Statistics 34:1275-1296.

Examples

```
lm1 <- lm(as.matrix(iris[,1:4]) ~ iris[,5])
exVar(lm1)</pre>
```

48 fastKmeans

fastKmeans

fast kmeans clustering for 2D or 3D point clouds

Description

fast kmeans clustering for 2D or 3D point clouds - with the primary purpose to get a spatially equally distributed samples

Usage

```
fastKmeans(x, k, iter.max = 10, project = TRUE, threads = 0)
```

Arguments

x matrix containing coordinates or mesh3d

k number of clusters

iter.max maximum number of iterations

project logical: if x is a triangular mesh, the centers will be projected onto the surface.

threads integer number of threads to use

Value

returns a list containing

selected coordinates closest to the final centers

centers cluster center

class vector with cluster association for each coordinate

Examples

```
require(Rvcg)
data(humface)
set.seed(42)
clust <- fastKmeans(humface,k=1000,threads=1)
## Not run:
require(rg1)
## plot the cluster centers
spheres3d(clust$centers)
## now look at the vertices closest to the centers
wire3d(humface)
spheres3d(vert2points(humface)[clust$selected,],col=2)
## End(Not run)</pre>
```

file2mesh 49

file2mesh

Import 3D surface mesh files

Description

Import 3D surface mesh files

Usage

```
file2mesh(filename, clean = TRUE, readcol = FALSE)

obj2mesh(filename, adnormals = TRUE)

ply2mesh(
  filename,
  adnormals = TRUE,
  readnormals = FALSE,
  readcol = FALSE,
  silent = FALSE
)
```

Arguments

filename character: path to file

clean Logical: Delete dumpfiles.

readcol Logical: Import vertex colors (if available).

adnormals Logical: If the file does not contain normal information, they will be calculated

in R: Can take some time.

readnormals Logical: Import vertex normals (if available), although no face information is

present.

silent logical: suppress messages.

Details

imports 3D mesh files and store them as an R .object of class mesh3d

Value

mesh

list of class mesh3d - see rgl manual for further details, or a matrix containing vertex information or a list containing vertex and normal information

50 find.outliers

Examples

```
data(nose)
mesh2ply(shortnose.mesh)
mesh <- ply2mesh("shortnose.mesh.ply")

mesh2obj(shortnose.mesh)
mesh2 <- obj2mesh("shortnose.mesh.obj")
## cleanup
unlink(c("shortnose.mesh.obj", "shortnose.mesh.ply"))</pre>
```

find.outliers

Graphical interface to find outliers and/or to switch mislabeld landmarks

Description

Graphical interface to find outliers and/or to switch mislabeld landmarks

Usage

```
find.outliers(
   A,
   color = 4,
   lwd = 1,
   lcol = 2,
   mahalanobis = FALSE,
   PCuse = NULL,
   text = TRUE,
   reflection = FALSE
)
```

Arguments

Α	Input k x m x n real array, where k is the number of points, m is the number of
	dimensions, and n is the sample size.

1 07 1 1 1 1 1 1 1 1

color color of Landmarks points to be plotted

lwd linewidth visualizing distances of the individual landmarks from mean.lcol color of lines visualizing distances of the individual landmarks from mean.

mahalanobis logical: use mahalanobis distance to find outliers.

PCuse integer: Restrict mahalanobis distance to the first n Principal components.

text logical: if TRUE, landmark labels (rownumbers) are displayed

reflection logical: specify whether reflections are allowed for superimpositioning.

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Details

This function performs a procrustes fit and sorts all specimen according to their distances (either Procrustes or Mahalanobis-distance) to the sample's consensus. It provides visual help for rearranging landmarks and/or excluding outliers.

Value

data.cleaned	array (in original coordinate system) containing the changes applied and outliers eliminated
outlier	vector with integers indicating the positions in the original array that have been marked as outliers
dist.sort	table showing the distance to mean for each observation - decreasing by distance
type	what kind of distance was used

Author(s)

Stefan Schlager

See Also

typprobClass

Examples

```
data(boneData)
## look for outliers using the mahalanobis distance based on the first
# 10 PCscores
# to perform the example below, you need,of course, uncomment the answers
if (interactive()){
  outliers <- find.outliers(boneLM, mahalanobis= TRUE, PCuse=10)
# n # everything is fine
# n # proceed to next
# s # let's switch some landmarks (3 and 4)
# 3
# 4
# n # we are done
# y # yes, because now it is an outlier
# s #enough for now
}</pre>
```

52 fixLMmirror

fixLMmirror

estimate missing landmarks from their bilateral counterparts

Description

estimate missing landmarks from their bilateral counterparts

Usage

```
fixLMmirror(x, pairedLM, ...)
## S3 method for class 'array'
fixLMmirror(x, pairedLM, ...)
## S3 method for class 'matrix'
fixLMmirror(x, pairedLM, ...)
```

Arguments

x a matrix or an array containing landmarks (3D or 2D)

pairedLM a k x 2 matrix containing the indices (rownumbers) of the paired LM. E.g. the

left column contains the lefthand landmarks, while the right side contains the

corresponding right hand landmarks.

... additional arguments

Details

the configurations are mirrored and the relabled version is matched onto the original using a thinplate spline deformation. The missing landmark is now estimated using its bilateral counterpart. If one side is completely missing, the landmarks will be mirrored and aligned by the unilateral landmarks.

Value

a matrix or array with fixed missing bilateral landmarks.

Note

in case both landmarks of a bilateral pair are missing a message will be issued. As well if there are missing landmarks on the midsaggital plane are detected.

Examples

```
data(boneData)
left <- c(4,6,8)
## determine corresponding Landmarks on the right side:
# important: keep same order
right <- c(3,5,7)</pre>
```

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```
pairedLM <- cbind(left, right)
exampmat <- boneLM[,,1]
exampmat[4,] <- NA #set 4th landmark to be NA
fixed <- fixLMmirror(exampmat, pairedLM=pairedLM)
## Not run:
deformGrid3d(fixed, boneLM[,,1],ngrid=0)
## result is a bit off due to actual asymmetry

## End(Not run)
## example with one side completely missing
oneside <- boneLM[,,1]
oneside[pairedLM[,1],] <- NA
onesidefixed <- fixLMmirror(oneside,pairedLM)
## Not run:
deformGrid3d(onesidefixed, boneLM[,,1],ngrid=0)
## result is a bit off due to actual asymmetry

## End(Not run)</pre>
```

fixLMtps

estimate missing landmarks

Description

Missing landmarks are estimated by deforming a sample average or a weighted estimate of the configurations most similar onto the deficient configuration. The deformation is performed by a Thin-plate-spline interpolation calculated by the available landmarks.

Usage

```
fixLMtps(data, comp = 3, weight = TRUE, weightfun = NULL)
```

Arguments

data array containing landmark data

comp integer: select how many of the closest observations are to be taken to calculate

an initial estimate.

weight logical: requests the calculation of an estimate based on the procrustes distance.

Otherwise the sample's consensus is used as reference.

weightfun custom function that operates on a vector of distances (see examples) and gen-

erates weights accordingly.

Details

This function tries to estimate missing landmark data by mapping weighted averages from complete datasets onto the missing specimen. The weights are the inverted Procrustes (see proc.weight) distances between the 'comp' closest specimen (using the available landmark configuration).

54 fixLMtps

Value

out array containing all data, including fixed configurations - same order as input

mshape meanshape - calculated from complete datasets

checklist list containing information about missing landmarks

check vector containing position of observations in data where at least one missing

coordinate was found

Note

Be aware that these estimates might be grossly wrong when the missing landmark is quite far off the rest of the landmarks (due to the radial basis function used in the Thin-plate spline interpolation.

Author(s)

Stefan Schlager

References

Bookstein FL. 1989. Principal Warps: Thin-plate splines and the decomposition of deformations IEEE Transactions on pattern analysis and machine intelligence 11.

See Also

```
proc.weight, tps3d
```

Examples

```
if (require(shapes)) {
data <- gorf.dat
### set first landmark of first specimen to NA
data[1,,1] <- NA
repair <- fixLMtps(data,comp=5)</pre>
### view difference between estimated and actual landmark
plot(repair$out[,,1],asp=1,pch=21,cex=0.7,col=2)#estimated landmark
points(gorf.dat[,,1],col=3,pch=20)#actual landmark
## 3D-example:
data(boneData)
data <- boneLM
### set first and 5th landmark of first specimen to NA
data[c(1,5),,1] <- NA
repair <- fixLMtps(data,comp=10)</pre>
## view difference between estimated and actual landmark
deformGrid3d(repair$out[,,1], boneLM[,,1],ngrid=0)
## End(Not run)
## Now use a gaussian kernel to compute the weights and use all other configs
gaussWeight <- function(r,sigma=0.05) {</pre>
```

getFaces 55

```
sigma <- 2*sigma^2
  return(exp(-r^2/ sigma))
}
repair <- fixLMtps(data,comp=79,weightfun=gaussWeight)</pre>
```

getFaces

find indices of faces that contain specified vertices

Description

find indices of faces that contain specified vertices

Usage

```
getFaces(mesh, index)
```

Arguments

mesh triangular mesh of class "mesh3d" index vector containing indices of vertices

Value

vector of face indices

 ${\tt getMeaningfulPCs}$

get number of meaningful Principal components

Description

get number of meaningful Principal components

Usage

```
getMeaningfulPCs(values, n, expect = 2, sdev = FALSE)
```

Arguments

values eigenvalues from a PCA

n sample size

expect expectation value for chi-square distribution of df=2

sdev logical: if TRUE, it is assumed that the values are square roots of eigenvalues.

56 getOuterViewpoints

Details

This implements the method suggested by Bookstein (2014, pp. 324), to determine whether a PC is entitled to interpretation. I.e. a PC is regarded meaningful (its direction) if the ratio of this PC and its successor is above a threshold based on a log-likelihood ratio (and dependent on sample size).

Value

tol threshold of ratio specific for n

good integer vector specifying the meaningful Principal Components

References

Bookstein, F. L. Measuring and reasoning: numerical inference in the sciences. Cambridge University Press, 2014

See Also

```
getPCtol
```

Examples

```
data(boneData)
proc <- procSym(boneLM)
getMeaningfulPCs(proc$eigenvalues,n=nrow(proc$PCscores))
## the first 3 PCs are reported as meaningful
## show barplot that seem to fit the bill
barplot(proc$eigenvalues)</pre>
```

getOuterViewpoints

Get viewpoints on a sphere around a 3D mesh

Description

Get viewpoints on a sphere around a 3D mesh to be used with virtualMeshScan

Usage

```
getOuterViewpoints(
    x,
    n,
    inflate = 1.5,
    radius = NULL,
    subdivision = 3,
    PCA = FALSE
)
```

getPCscores 57

Arguments

x triangular mesh of class 'mesh3d' n number of viewpoint to generate

inflate factor for the size of the sphere: inflate=1 means that the sphere around the

object just touches the point farthest away from the mesh's centroid.

radius defines a fix radius for the sphere (overrides arg inflate).

subdivision parameter passed to vcgSphere

PCA logical: if TRUE, the sphere will be deformed to match the principle axes of the

mesh. NOTE: this may result in the sphere not necessarily completely enclosing

the mesh.

Value

a list containing

viewpoints n x 3 matrix containing viewpoints.

sphere sphere from which the points are sampled

radius radius of the sphere

Examples

```
## Not run:
data(boneData)
vp <- getOuterViewpoints(skull_0144_ch_fe.mesh,n=100)

require(rgl)
shade3d(skull_0144_ch_fe.mesh,col="white")
spheres3d(vp$viewpoints)
wire3d(vp$sphere)

### Fit to principal axes
vppca <- getOuterViewpoints(skull_0144_ch_fe.mesh,n=100,PCA=TRUE,inflate=1.5)

require(rgl)
shade3d(skull_0144_ch_fe.mesh,col="white")
spheres3d(vppca$viewpoints)
wire3d(vppca$sphere)

## End(Not run)</pre>
```

getPCscores

Obtain PC-scores for new landmark data

Description

Obtain PC-scores for new landmark data

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Usage

```
getPCscores(x, PC, mshape)
```

Arguments

x landmarks aligned (e.g. using align2procSym to the meanshape of data the PCs

are derived from.

PC Principal components (eigenvectors of the covariance matrix)

mshape matrix containing the meanshape's landmarks (used to center the data)

Value

returns a matrix containing the PC scores

See Also

```
restoreShapes
```

Examples

```
## Not run:
data(boneData)
proc <- procSym(boneLM[,,-c(1:2)])
newdata <- boneLM[,,c(1:2)]
newdataAlign <- align2procSym(proc,newdata)
scores <- getPCscores(newdataAlign,proc$PCs,proc$mshape)
## End(Not run)</pre>
```

getPCtol

determine the minimum ratio for two subsequent eigenvalues to be considered different

Description

determine the minimum ratio for two subsequent eigenvalues to be considered different

Usage

```
getPCtol(n, expect = 2)
```

Arguments

n sample size

expect expectation value for chi-square distribution of df=2

Value

returns the minimum ratio between two subsequent subsequent eigenvalues to be considered different

References

Bookstein, F. L. Measuring and reasoning: numerical inference in the sciences. Cambridge University Press, 2014

See Also

getMeaningfulPCs

Examples

```
## reproduce the graph from Bookstein (2014, p. 324)
## and then compare it to ratios for values to be considered
## statistically significant
myseq \leftarrow seq(from=10, to = 50, by = 2)
myseq \leftarrow c(myseq, seq(from=50, to=1000, by =20))
ratios <- getPCtol(myseq)</pre>
plot(log(myseq), ratios, cex=0, xaxt = "n", ylim=c(1,5.2))
ticks <- c(10,20,50,100,200,300,400,500,600,700,800,900,1000)
axis(1,at=log(ticks),labels=ticks)
lines(log(myseq), ratios)
abline(v=log(ticks), col="lightgray", lty="dotted")
abline(h=seq(from=1.2,to=5, by = 0.2), col="lightgray", lty="dotted")
## now we raise the bar and compute the ratios for values
## to be beyond the 95th percentile of
## the corresponding chi-square distribution:
ratiosSig <- getPCtol(myseq,expect=qchisq(0.95,df=2))</pre>
lines(log(myseq),ratiosSig,col=2)
```

getPLSCommonShape

Get the linear combinations associated with the common shape change in each latent dimension of a pls2B

Description

Get the linear combinations associated with the common shape change in each latent dimension of a pls2B

Usage

```
getPLSCommonShape(pls)
```

60 getPLSfromScores

Arguments

pls object of class "pls2B"

Value

returns a list containing

shapevectors matrix with each containing the shapevectors (in column-major format) of com-

mon shape change associated with each latent dimension

XscoresScaled Xscores scaled according to shapevectors
YscoresScaled Yscores scaled according to shapevectors

commoncenter Vector containing the common mean

lmdim dimension of landmarks

References

Mitteroecker P, Bookstein F. 2007. The conceptual and statistical relationship between modularity and morphological integration. Systematic Biology 56(5):818-836.

See Also

plsCoVarCommonShape

Examples

```
data(boneData)
proc <- procSym(boneLM)
pls <- pls2B(proc$orpdata[1:4,,],proc$orpdata[5:10,,])
commShape <- getPLSCommonShape(pls)
## get common shape for first latent dimension at +-2 sd of the scores
## (you can do this much more convenient using \code{\link{plsCoVarCommonShape}}
scores <- c(-2,2) * sd(c(commShape$XscoresScaled[,1],commShape$YscoresScaled[,1]))
pred <- restoreShapes(scores,commShape$shapevectors[,1],matrix(commShape$commoncenter,10,3))
## Not run:
deformGrid3d(pred[,,1],pred[,,2])
## End(Not run)</pre>
```

getPLSfromScores

compute changes associated with 2-Block PLS-scores

Description

compute changes associated with 2-Block PLS-scores

Usage

```
getPLSfromScores(pls, x, y)
```

getPLSscores 61

Arguments

pls	output of pls2B
X	scores associated with dataset x in original pls2B
у	scores associated with dataset y in original pls2B

Details

other than predictPLSfromScores, providing Xscores will not compute predictions of y, but the changes in the original data x that is associated with the specific scores

Value

returns data in the original space associated with the specified values.

getPLSscores	compute 2-Block PLS scores for new data
--------------	---

Description

compute 2-Block PLS scores for new data from an existing pls2B

Usage

```
getPLSscores(pls, x, y)
```

Arguments

pls	output of pls2B
X	matrix or vector representing new dataset(s) - same kind as in original pls2B $$
у	matrix or vector representing new dataset(s) - same kind as in original pls2B

Value

returns a vector of pls-scores

Note

either x or y must be missing

See Also

```
pls2B, predictPLSfromScores, predictPLSfromData
```

62 getSides

getPointAlongOutline Get a point along a line with a given distance from the start of the line

Description

Get a point along a line with a given distance from the start of the line

Usage

```
getPointAlongOutline(mat, dist = 15, reverse = FALSE)
```

Arguments

mat matrix with rows containing sequential coordinates
dist numeric: distance from the first point on the line.
reverse logical: if TRUE start from the end of the line

Value

returns a vector containing the resulting coordinate

getSides

try to identify bilateral landmarks and sort them by side

Description

try to identify bilateral landmarks and sort them by side

Usage

```
getSides(x, tol = 3, pcAlign = TRUE, icpiter = 100, ...)
```

Arguments

X	matrix containing landmarks (see details)
tol	maximal distance allowed between original and mirrored set.
pcAlign	logical: if TRUE orginal and mirrored landmarks will be initally aligned by their PC-axes
icpiter	integer: number of iterations in ICP alignment.

... more arguments passed to mirror.

getTrafo4x4 63

Details

This function mirrors the landmark set and aligns it to the original. Then it tries to find pairs. If you have a sample, run a Procrustes registration first (without scaling to unit centroid size, or you later have to adapt tol - see examples) and then use the mean as it is usually more symmetrical.

Value

returns a list containing

side1 integer vector containing indices of landmarks on one side side2 integer vector containing indices of landmarks on the other side unilat integer vector containing indices unilateral landmarks

Examples

```
data(boneData)
proc <- procSym(boneLM,CSinit=FALSE)
mysides <- getSides(proc$mshape)
if (interactive()){
#visualize bilateral landmarks
deformGrid3d(boneLM[mysides$side1,,1],boneLM[mysides$side2,,1])
## visualize unilateral landmarks
rgl::spheres3d(boneLM[mysides$unilat,,1],radius=0.5)
}</pre>
```

getTrafo4x4

get 4x4 Transformation matrix

Description

```
get 4x4 Transformation matrix
```

Usage

```
getTrafo4x4(x)
## S3 method for class 'rotonto'
getTrafo4x4(x)
```

Arguments

x object of class "rotonto"

Value

returns a 4x4 transformation matrix

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Examples

```
data(boneData)
rot <- rotonto(boneLM[,,1],boneLM[,,2])
trafo <- getTrafo4x4(rot)</pre>
```

getTrafoRotaxis

compute a 4x4 Transformation matrix for rotation around an arbitrary

axis

Description

compute a 4x4 Transformation matrix for rotation around an arbitrary axis

Usage

```
getTrafoRotaxis(pt1, pt2, theta)
```

Arguments

pt1 numeric vector of length 3, defining first point on the rotation axis. pt2 numeric vector of length 3, defining second point on the rotation axis.

theta angle to rotate in radians. With pt1 being the viewpoint, the rotation is counter-

clockwise.

Note

the resulting matrix can be used in applyTransform

getVisibleVertices

find vertices visible from a given viewpoints

Description

find vertices visible from a given viewpoints

Usage

```
getVisibleVertices(mesh, viewpoints, offset = 0.001, cores = 1)
```

Arguments

mesh triangular mesh of class 'mesh3d'

viewpoints vector or k x 3 matrix containing a set of viewpoints

offset value to generate an offset at the meshes surface (see notes) cores integer: number of cores to use (not working on windows)

groupPCA 65

Value

a vector with (1-based) indices of points visible from at least one of the viewpoints

Note

The function tries to filter out all vertices where the line connecting each vertex with the viewpoints intersects with the mesh itself. As, technically speaking this always occurs at a distance of value=0, a mesh with a tiny offset is generated to avoid these false hits.

Examples

```
SCP1 <- file2mesh(system.file("extdata","SCP1.ply",package="Morpho"))
viewpoints <- read.fcsv(system.file("extdata","SCP1_Endo.fcsv",package="Morpho"))
visivert <- getVisibleVertices(SCP1,viewpoints)</pre>
```

groupPCA

Perform PCA based of the group means' covariance matrix

Description

Calculate covariance matrix of the groupmeans and project all observations into the eigenspace of this covariance matrix. This displays a low dimensional between group structure of a high dimensional problem.

Usage

```
groupPCA(
  dataarray,
  groups,
  rounds = 10000,
  tol = 1e-10,
  cv = TRUE,
  mc.cores = parallel::detectCores(),
  weighting = TRUE
)
```

Arguments

dataarray	of dimensions, and n is the sample size. Or alternatively a n x m Matrix where n is the number of observations and m the number of variables (this can be PC scores for example)
groups	a character/factor vector containgin grouping variable.
rounds	integer: number of permutations if a permutation test of the euclidean distance between group means is requested. If rounds = 0, no test is performed.
tol	threshold to ignore eigenvalues of the covariance matrix.

66 groupPCA

cv logical: requests leaving-one-out crossvalidation

mc.cores integer: how many cores of the Computer are allowed to be used. Default is

use autodetection by using detectCores() from the parallel package. Parallel

processing is disabled on Windows due to occasional errors.

weighting logical: weight between groups covariance matrix according to group sizes.

Value

eigenvalues Non-zero eigenvalues of the groupmean covariance matrix

groupPCs PC-axes - i.e. eigenvectors of the groupmean covariance matrix

Variance table displaying the between-group variance explained by each between group

PC - this only reflects the variability of the group means and NOT the variability

of the data projected into that space

Scores Scores of all observation in the PC-space

probs p-values of pairwise groupdifferences - based on permuation testing

groupdists Euclidean distances between groups' averages

groupmeans matrix with rows containing the Groupmeans, or a k x m x groupsize array if the

input is a k x m x n landmark array

Grandmean vector containing the Grand mean, or a matrix if the input is a k x m x n landmark

array

CV Cross-validated scores

groups grouping Variable

resPCs PCs orthogonal to the between-group PCs

resPCscores Scores of the residualPCs

resVar table displaying the residual variance explained by each residual PC.

Author(s)

Stefan Schlager

References

Mitteroecker P, Bookstein F 2011. Linear Discrimination, Ordination, and the Visualization of Selection Gradients in Modern Morphometrics. Evolutionary Biology 38:100-114.

Boulesteix, A. L. 2005: A note on between-group PCA, International Journal of Pure and Applied Mathematics 19, 359-366.

See Also

CVA

histGroup 67

Examples

```
data(iris)
vari <- iris[,1:4]</pre>
facto <- iris[,5]</pre>
pca.1 <-groupPCA(vari,groups=facto,rounds=100,mc.cores=1)</pre>
### plot scores
if (require(car)) {
scatterplotMatrix(pca.1$Scores,groups=facto, ellipse=TRUE,
        by.groups=TRUE,var.labels=c("PC1","PC2","PC3"))
## example with shape data
data(boneData)
proc <- procSym(boneLM)</pre>
pop_sex <- name2factor(boneLM, which=3:4)</pre>
gpca <- groupPCA(proc$orpdata, groups=pop_sex, rounds=0, mc.cores=2)</pre>
## Not run:
## visualize shape associated with first between group PC
dims <- dim(proc$mshape)</pre>
## calculate matrix containing landmarks of grandmean
grandmean <-gpca$Grandmean</pre>
## calculate landmarks from first between-group PC
                     (+2 and -2 standard deviations)
gpcavis2sd \leftarrow restoreShapes(c(-2,2)*sd(gpca$Scores[,1]), gpca$groupPCs[,1], grandmean)
deformGrid3d(gpcavis2sd[,,1], gpcavis2sd[,,2], ngrid = 0,size=0.01)
require(rgl)
## visualize grandmean mesh
grandm.mesh <- tps3d(skull_0144_ch_fe.mesh, boneLM[,,1],grandmean,threads=1)</pre>
wire3d(grandm.mesh, col="white")
spheres3d(grandmean, radius=0.01)
## End(Not run)
```

histGroup

plot histogram for multiple groups.

Description

plot a histogram for multiple groups, each group colored individually

Usage

```
histGroup(
  data,
  groups,
```

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```
main = paste("Histogram of", dataname),
xlab = dataname,
ylab,
col = NULL,
alpha = 0.5,
breaks = "Sturges",
legend = TRUE,
legend.x = 80,
legend.y = 80,
legend.pch = 15,
freq = TRUE
)
```

Arguments

data vector containing data.
groups grouping factors
main, xlab, ylab

these arguments to title have useful defaults here.

col vector containing color for each group. If NULL, the function "rainbow" is

called.

alpha numeric between 0 and 1. Sets the transparency of the colors

breaks one of:

• a vector giving the breakpoints between histogram cells,

• a single number giving the number of cells for the histogram,

• a character string naming an algorithm to compute the number of cells (see 'Details'),

• a function to compute the number of cells.

In the last three cases the number is a suggestion only.

legend logical: if TRUE, a legend is plotted

legend.x x position of the legend from the upper left corner legend.y y position of the legend from the upper left corners

legend.pch integer: define the symbol to visualise group colors (points)

freq logical: if TRUE, the histogram graphic is a representation of frequencies, the

counts component of the result; if FALSE, probability densities are plotted for

each group.

Details

Just a wrapper for the function hist from the "graphics" package

Author(s)

Stefan Schlager

icpmat 69

See Also

hist

Examples

```
data(iris)
histGroup(iris$Petal.Length,iris$Species)
```

icpmat

match two landmark configurations using iteratively closest point search

Description

match two landmark configurations using iteratively closest point search

Usage

```
icpmat(
    x,
    y,
    iterations,
    mindist = 1e+15,
    subsample = NULL,
    type = c("rigid", "similarity", "affine"),
    weights = NULL,
    threads = 1,
    centerweight = FALSE
)
```

Arguments

x moving landmarksy target landmarks

iterations integer: number of iterations

mindist restrict valid points to be within this distance

subsample use a subsample determined by kmean clusters to speed up computation

type character: select the transform to be applied, can be "rigid", "similarity" or "affine"

weights vector of length nrow(x) containing weights for each row in x

threads integer: number of threads to use.

centerweight logical: if weights are defined and centerweigths=TRUE, the matrix will be

centered according to these weights instead of the barycenter.

70 invertFaces

Value

returns the rotated landmarks

Examples

```
data(nose)
icp <- icpmat(shortnose.lm,longnose.lm,iterations=10)</pre>
## example using weights
## we want to assign high weights to the first three cordinates
icpw <- icpmat(shortnose.lm,longnose.lm,iterations=10,</pre>
               weights=c(rep(100,3),rep(1,620)),centerweight = TRUE)
## the RMSE between those four points and the target is now smaller:
require(Rvcg)
RMSE <- sqrt(sum(vcgKDtree(longnose.lm,icp[1:3,],k=1)$distance^2))</pre>
RMSEW<- sqrt(sum(vcgKDtree(longnose.lm,icpw[1:3,],k=1)$distance^2))
barplot(c(RMSE,RMSEW),names.arg=c("RMSE weighted","RMSE unweighted"))
## Not run:
## plot the differences between unweighted and weighted icp
deformGrid3d(icp,icpw)
## plot the first four coordinates from the icps:
spheres3d(icp[1:3,],col="red",radius = 0.5)
spheres3d(icpw[1:3,],col="green",radius = 0.5)
## plot the target
spheres3d(longnose.lm,col="yellow",radius = 0.2)
## End(Not run)
##2D example using icpmat to determine point correspondences
if (require(shapes)) {
## we scramble rows to show that this is independent of point order
moving <- gorf.dat[sample(1:8),,1]</pre>
plot(moving,asp=1) ## starting config
icpgorf <- icpmat(moving,gorf.dat[,,2],iterations = 20)</pre>
points(icpgorf,asp = 1,col=2)
points(gorf.dat[,,2],col=3)## target
## get correspondences using nearest neighbour search
index <- mcNNindex(icpgorf,gorf.dat[,,2],k=1,cores=1)</pre>
icpsort <- icpgorf[index,]</pre>
for (i in 1:8)
lines(rbind(icpsort[i,],gorf.dat[i,,2]))
```

invertFaces

invert faces' orientation of triangular mesh

Description

inverts faces' orientation of triangular mesh and recomputes vertex normals

kendalldist 71

Usage

```
invertFaces(mesh)
```

Arguments

mesh

triangular mesh of class mesh3d

Value

returns resulting mesh

Author(s)

Stefan Schlager

See Also

updateNormals

Examples

```
data(nose)
## Not run:
rgl::shade3d(shortnose.mesh,col=3)

## End(Not run)
noseinvert <- invertFaces(shortnose.mesh)
## show normals
## Not run:
plotNormals(noseinvert,long=0.01)

## End(Not run)</pre>
```

kendalldist

Calculates the Riemannian distance between two superimposed landmark configs.

Description

Calculates the Riemannian distance between two superimposed landmark configs.

Usage

```
kendalldist(x, y)
```

72 line2plane

Arguments

x Matrix containing landmark coordinates.y Matrix containing landmark coordinates.

Value

returns Riemannian distance

Examples

```
if(require(shapes)) {
OPA <- rotonto(gorf.dat[,,1],gorf.dat[,,2])
kendalldist(OPA$X,OPA$Y)
}</pre>
```

line2plane

get intersection between a line and a plane

Description

get intersection between a line and a plane

Usage

```
line2plane(ptLine, ptDir, planePt, planeNorm)
```

Arguments

ptLine vector of length 3: point on line

ptDir vector of length 3: direction vector of line

planePt vector of length 3: point on plane

planeNorm vector of length 3: plane normal vector

Value

hit point

Note

in case you only have three points on a plane (named pt1, pt2, pt3 you can get the plane's normal by calling crossProduct(pt1-pt2,pt1-pt3).

lineplot 73

lineplot

plot lines between landmarks

Description

add lines connecting landmarks to visualise a sort of wireframe

Usage

```
lineplot(
    x,
    point,
    col = 1,
    lwd = 1,
    line_antialias = FALSE,
    lty = 1,
    add = TRUE
)
```

Arguments

x matrix containing 2D or 3D landmarks

point vector or list of vectors containing rowindices of x, determining which land-

marks to connect.

col color of lines

line_antialias logical: smooth lines

lty line type (only for 2D case) add logical: add to existing plot

Note

works with 2D and 3D configurations

Author(s)

Stefan Schlager

See Also

pcaplot3d

74 LPS2RAS

Examples

```
if (require(shapes)) {
##2D example
plot(gorf.dat[,,1],asp=1)
lineplot(gorf.dat[,,1],point=c(1,5:2,8:6,1),col=2)
}
##3D example
## Not run:
require(rg1)
data(nose)
points3d(shortnose.lm[1:9,])
lineplot(shortnose.lm[1:9,],point=list(c(1,3,2),c(3,4,5),c(8,6,5,7,9)),col=2)
## End(Not run)
```

list2array

converts a list of matrices to an array

Description

converts a list of matrices to an array

Usage

```
list2array(x)
```

Arguments

Χ

a list containing matrices of the same dimensionality

Value

returns an array concatenating all matrices

LPS2RAS

convert data from LPS to RAS space and back

Description

convert data from LPS to RAS space and back

Usage

```
LPS2RAS(x)
```

mcNNindex 75

Arguments

Χ

mesh of class mesh3d or a matrix with 3D Landmarks

Details

As e.g. the Slicer versions >= 4.11 are using LPS space, it might be needed to transform data like fiducials and surface models from and back to that space.

Value

returns the rotated data

mcNNindex

find nearest neighbours for 2D and 3D point clouds

Description

find nearest neighbours for point clouds using a kd-tree search. This is just a wrapper of the function vcgKDtree from package Rvcg. Wwraps the function vcgKDtree from package 'Rvcg' (for backward compatibility)

Usage

```
mcNNindex(target, query, cores = parallel::detectCores(), k = k, ...)
```

Arguments

target k x m matrix containing data which to search.

query 1 x m matrix containing data for which to search.

cores integer: amount of CPU-cores to be used. Only available on systems with OpenMP support.

k integer: how many closest points are sought.

additional arguments - currently unused.

Value

1 x k matrix containing indices of closest points.

See Also

closemeshKD

76 mergeMeshes

Examples

```
require(rgl)
data(nose)
# find closest vertex on surface for each landmark
clost <- mcNNindex(vert2points(shortnose.mesh), shortnose.lm, k=1,
mc.cores=1)
## Not run:
spheres3d(vert2points(shortnose.mesh)[clost,],col=2,radius=0.3)
spheres3d(shortnose.lm,radius=0.3)
wire3d(shortnose.mesh)
## End(Not run)</pre>
```

mergeMeshes

merge multiple triangular meshes into a single one

Description

merge multiple triangular meshes into a single one, preserving color and vertex normals.

Usage

```
mergeMeshes(...)
```

Arguments

... triangular meshes of class 'mesh3d' to merge or a list of triangular meshes.

Value

returns the meshes merged into a single one.

See Also

```
mesh2ply, file2mesh, ply2mesh
```

```
## Not run:
require(rgl)
data(boneData)
data(nose)
mergedMesh <- mergeMeshes(shortnose.mesh, skull_0144_ch_fe.mesh)
require(rgl)
shade3d(mergedMesh, col=3)
## End(Not run)</pre>
```

mesh2grey 77

mesh2grey

convert a colored mesh to greyscale.

Description

convert the colors of a colored mesh to greyscale values

Usage

```
mesh2grey(mesh)
```

Arguments

mesh

Object of class mesh3d

Value

returns a mesh with material\$color replaced by greyscale rgb values.

Author(s)

Stefan Schlager

See Also

```
ply2mesh,file2mesh
```

mesh2obj

export mesh objects to disk

Description

export mesh objects to disk.

Usage

```
mesh2obj(x, filename = dataname, writeNormals = TRUE)
mesh2ply(x, filename = dataname, col = NULL, writeNormals = FALSE)
```

78 mesh2obj

Arguments

X	object of class mesh3d - see rgl documentation for further details or a matrix containing vertices, this can either be a k x 3 or a 3 x k matrix, with rows or columns containing vertex coordinates.
filename	character: Path/name of the requested output - extension will be added atuomatically. If not specified, the file will be named as the exported object.
writeNormals	logical: if TRUE, existing normals of a mesh are written to file - can slow things down for very large meshes.
col	Writes color information to ply file. Can be either a single color value or a vector containing a color value for each vertex of the mesh.

Details

export an object of class mesh3d or a set of coordinates to a common mesh file.

Note

meshes containing quadrangular faces will be converted to triangular meshes by splitting the faces. Additionally, mesh2obj is now simply a wrapper of Rvcg::vcgObjWrite.

Author(s)

Stefan Schlager

See Also

```
ply2mesh, quad2trimesh
```

```
require(rg1)
vb <- c(-1.8,-1.8,-1.8,1.0,1.8,-1.8,-1.8,1.0,-1.8,1.0,-1.8,1.0,1.8,
1.8,-1.8,1.0,-1.8,-1.8,1.8,1.0,1.8,
-1.8,1.8,1.0,-1.8,1.8,1.0,1.8,1.8,1.0,0)
it <- c(2,1,3,3,4,2,3,1,5,5,7,3,5,1,2,2,6,5,6,8,7,7,5,6,7,8,4,4,3,7,4,8,6,6,2,4)
vb <- matrix(vb,4,8) ##create vertex matrix
it <- matrix(it,3,12) ## create face matrix
cube<-list(vb=vb,it=it)
class(cube) <- "mesh3d"
## Not run:
shade3d(cube,col=3) ## view the green cube

## End(Not run)
mesh2ply(cube,filename="cube") # write cube to a file called cube.ply
unlink("cube.ply")</pre>
```

meshcube 79

meshcube

calculate the corners of a mesh's bouning box

Description

calculate the corners of a mesh's bouning box

Usage

```
meshcube(x)
```

Arguments

Х

object of class 'mesh3d'

Value

returns a 8 x 3 matrix with the coordinates of the corners of the bounding box.

Examples

```
require(rg1)
data(boneData)
mc <- meshcube(skull_0144_ch_fe.mesh)
## Not run:
spheres3d(mc)
wire3d(skull_0144_ch_fe.mesh)
## End(Not run)</pre>
```

meshDist

calculates and visualises distances between surface meshes or 3D coordinates and a surface mesh.

Description

calculates and visualises distances between surface meshes or 3D coordinates and a surface mesh.

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Usage

```
meshDist(x, ...)
## S3 method for class 'mesh3d'
meshDist(
 Х,
 mesh2 = NULL,
 distvec = NULL,
  from = NULL,
  to = NULL,
  steps = 20,
  ceiling = FALSE,
  rampcolors = colorRamps::blue2green2red(steps - 1),
 NAcol = "white",
  file = "default",
  imagedim = "100x800",
  uprange = 1,
  ray = FALSE,
  raytol = 50,
  raystrict = FALSE,
  save = FALSE,
  plot = TRUE,
  sign = TRUE,
  tol = NULL,
  tolcol = "green",
  displace = FALSE,
  shade = TRUE,
 method = c("vcglib", "morpho"),
  add = FALSE,
  scaleramp = TRUE,
  threads = 1,
  titleplot = "Distance in mm",
)
## S3 method for class 'matrix'
meshDist(
 х,
 mesh2 = NULL,
  distvec = NULL,
  from = NULL,
  to = NULL,
  steps = 20,
  ceiling = FALSE,
  rampcolors = colorRamps::blue2green2red(steps - 1),
 NAcol = "white",
  uprange = 1,
  plot = TRUE,
```

meshDist 81

```
sign = TRUE,
tol = NULL,
tolcol = "green",
type = c("s", "p"),
radius = NULL,
displace = FALSE,
add = FALSE,
scaleramp = FALSE,
titleplot = "Distance in mm",
...
)
```

Arguments

x reference mesh; object of class "mesh3d" or a n x 3 matrix containing 3D coor-

dinates.

... additional arguments passed to shade3d. See material3d for details.

mesh2 target mesh: either object of class "mesh3d" or a character pointing to a surface

mesh (ply, obj or stl file)

distvec vector: optional, a vector containing distances for each vertex/coordinate of x,

if distvec != NULL, mesh2 will be ignored.

from numeric: minimum distance to be colorised; default is set to 0 mm

to numeric: maximum distance to be colorised; default is set to the maximum

distance

steps integer: determines break points for color ramp: n steps will produce n-1 colors.

ceiling logical: if TRUE, the next larger integer of "to" is used

rampcolors character vector: specify the colors which are used to create a colorramp.

NAcol character: specify color for values outside the range defined by from and to.

file character: filename for mesh and image files produced. E.g. "mydist" will

produce the files mydist.ply and mydist.png

imagedim character of type 100x200 where 100 determines the width and 200 the height

of the image.

uprange numeric between 0 and 1: restricts "to" to a quantile of "to", if to is NULL.

ray logical: if TRUE, the search is along vertex normals.

raytol maximum distance to follow a normal.

raystrict logical: if TRUE, only outward along normals will be sought for closest points.

save logical: save a colored mesh.

plot logical: visualise result as 3D-plot and distance charts

sign logical: request signed distances. Only meaningful, if mesh2 is specified or

distvec contains signed distances.

tol numeric: threshold to color distances within this threshold green.

tolcol a custom color to color vertices below a threshold defined by tol. Default is

green.

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displace logical: if TRUE, displacement vectors between original and closest points are

drawn colored according to the distance.

shade logical: if FALSE, the rendering of the colored surface will be supressed.

method accepts: "vcglib" and "morpho" (and any abbreviation), vcglib uses a command

line tool using vcglib headers, morpho uses fortran routines based on a kd-tree

search for closest triangles.

add logical: if TRUE, visualization will be added to the rgl window currently in

focus

scaleramp logical: if TRUE, the colorramp will be symmetrical for signed distances: span-

ning from -max(from, to) to max(from, to).

threads integer: number of threads to use. 0 = let system decide.

titleplot character: axis description of heatmap.

type character: "s" shows coordinates as spheres, while "p" shows 3D dots.

radius determines size of spheres; if not specified, optimal radius size will be estimated

by centroid size of the configuration.

Details

calculates the distances from a mesh or a set of 3D coordinates to another at each vertex; either closest point or along the normals

Value

Returns an object of class "meshDist" if the input is a surface mesh and one of class "matrixDist" if input is a matrix containing 3D coordinates.

colMesh object of mesh3d with colors added

dists vector with distances
cols vector with color values
params list of parameters used

Author(s)

Stefan Schlager

References

Detection of inside/outside uses the algorithm proposed in:

Baerentzen, Jakob Andreas. & Aanaes, H., 2002. Generating Signed Distance Fields From Triangle Meshes. Informatics and Mathematical Modelling, .

See Also

render.meshDist,, export.meshDist, shade3d

meshPlaneIntersect 83

Examples

```
data(nose)##load data
##warp a mesh onto another landmark configuration:
longnose.mesh <- tps3d(shortnose.mesh, shortnose.lm, longnose.lm,threads=1)
## Not run:
mD <- meshDist(longnose.mesh, shortnose.mesh)
##now change the color ramp
render(mD,rampcolors = c("white","red"))
## End(Not run)
#use unsigned distances and a ramp from blue to red
#color distances < 0.01 green:
## Not run:
meshDist(longnose.mesh, shortnose.mesh, rampcolors = c("blue", "red"),sign=FALSE, tol=0.5)
## End(Not run)</pre>
```

meshPlaneIntersect

get intersections between mesh and a plane

Description

get intersections between mesh and a plane

Usage

```
meshPlaneIntersect(mesh, v1, v2 = NULL, v3 = NULL, normal = NULL)
```

Arguments

mesh	triangular mesh of class "mesh3d"
v1	numeric vector of length=3 specifying a point on the separating plane
v2	numeric vector of length=3 specifying a point on the separating plane
v3	numeric vector of length=3 specifying a point on the separating plane
normal	plane normal (overrides specification by v2 and v3)

Value

returns the intersections of edges and the plane

84 meshres

Examples

```
data(nose)
v1 <- shortnose.lm[1,]
v2 <- shortnose.lm[2,]
v3 <- shortnose.lm[3,]
intersect <- meshPlaneIntersect(shortnose.mesh,v1,v2,v3)
## Not run:
require(rg1)
wire3d(shortnose.mesh)
spheres3d(shortnose.lm[1:3,],col=2)#the plane
spheres3d(intersect,col=3,radius = 0.2)#intersections
## End(Not run)</pre>
```

meshres

calculate average edge length of a triangular mesh

Description

calculate average edge length of a triangular mesh, by iterating over all faces.

Usage

```
meshres(mesh)
```

Arguments

mesh

triangular mesh stored as object of class "mesh3d"

Value

```
returns average edge length (a.k.a. mesh resolution)
```

Author(s)

Stefan Schlager

```
data(boneData)
mres <- meshres(skull_0144_ch_fe.mesh)</pre>
```

mirror 85

mirror

mirror landmarks or triangular mesh in place

Description

mirror landmarks or triangular mesh in place

Usage

```
mirror(
  Х,
  icpiter = 50,
  subsample = NULL,
 pcAlign = FALSE,
 mirroraxis = 1,
  initPC = TRUE,
  initCenter = TRUE,
  v1 = NULL,
  v2 = NULL,
  v3 = NULL,
 normal = NULL,
 mc.cores = 2
## S3 method for class 'matrix'
mirror(
 Х,
  icpiter = 50,
  subsample = NULL,
 pcAlign = FALSE,
 mirroraxis = 1,
  initPC = TRUE,
  initCenter = TRUE,
  v1 = NULL,
  v2 = NULL,
  v3 = NULL,
 normal = NULL,
 mc.cores = 2
)
## S3 method for class 'mesh3d'
mirror(
  Х,
  icpiter = 50,
  subsample = NULL,
  pcAlign = FALSE,
 mirroraxis = 1,
```

86 mirror

```
initPC = TRUE,
  initCenter = TRUE,
  v1 = NULL,
  v2 = NULL,
  v3 = NULL,
 normal = NULL,
 mc.cores = 2
)
```

Arguments

k x 3 matrix or mesh3d Χ integer: number of iterations to match reflected configuration onto original one icpiter subsample integer: use only a subset for icp matching pcAlign if TRUE, the icp will be preceded by an alignment of the principal axis (only used if icpiter > 0), currently only works for 3D data. mirroraxis integer: which axis to mirror at initPC logical: if TRUE the data will be prealigned by its principal axes. initCenter logical: if TRUE and initPC=FALSE, x will be translated to its centroid before mirroring. v1 point on plane v2 if normal=NULL, the plane will be defined by three points v1, v2, v3 if normal=NULL, the plane will be defined by three points v1, v2, v3 v3 plane normal (overrides specification by v2 and v3) normal

use parallel processing to find best alignment to original shape. mc.cores

Details

reflect a mesh configuration at the plane spanned by its first 2 principal axis, then try to rigidily register the reflected configuration onto the original one using iterative closest point search to establish correspondences. Also, if a reflection plane is defined, pcAlign, initPC, initCenter and mirroraxis will be ignored and the object will be mirrored on the defined plane (and optionally aligned using an ICP approach).

Value

returns the reflected object

```
data(boneData)
boneMir <- mirror(boneLM[,,1],icpiter=50,mc.cores=2,mirroraxis=3)</pre>
### mirror on 3 midsaggital landmarks and then optimize it with an ICP
boneMirPlane <- mirror(boneLM[,,1],v1=boneLM[1,,1],v2=boneLM[2,,1],v3=boneLM[9,,1])</pre>
## 2D Example:
```

mirror2plane 87

mirror2plane

mirror points or mesh on an arbitrary plane

Description

mirror points or mesh on an arbitrary plane

Usage

```
mirror2plane(x, v1, normal = NULL, v2 = NULL, v3 = NULL)
## S3 method for class 'matrix'
mirror2plane(x, v1, normal = NULL, v2 = NULL, v3 = NULL)
## S3 method for class 'mesh3d'
mirror2plane(x, v1, normal = NULL, v2 = NULL, v3 = NULL)
```

Arguments

X	x 3D-vector or a k x 3 matrix with 3D vectors stored in rows. Or a triangular mesh of class mesh3d
v1	point on plane
normal	plane normal (overrides specification by v2 and v3)
v2	if pNorm=NULL, the plane will be defined by three points v1, v2, v3
v3	if pNorm=NULL, the plane will be defined by three points v1, v2, v3

Value

mirrored coordinates mesh

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Examples

```
# mirror mesh on plane spanned by 3 midsagital landmarks
data(boneData)
mirrmesh <- mirror2plane(skull_0144_ch_fe.mesh,v1=boneLM[1,,1],v2=boneLM[9,,1],v3=boneLM[10,,1])</pre>
```

name2factor

extract data from array names

Description

extract data from array names

Usage

```
name2factor(x, sep = "_-", which, collapse = sep, as.factor = TRUE)
name2num(x, sep = "_-", which, collapse = sep, dif = TRUE)
```

Arguments

Х	data, can be a three-dimensional array, a matrix, a named list or a vector containing names to split
sep	character by which to split the strings
which	integer or vector of integers, if more entries are selected, they will be concatenated by the string specified with the option 'collapse'.
collapse	character by which to collapse data if two strings are to be concatenated
as.factor	logical: if TRUE, a factor vector will be returned, strings otherwise.
dif	logical: calculate difference if two fields containing numbers are selected.

Details

extract data from array names and convert to factors or numbers

If an array is used as input, the data info is expected to be in the 3rd dimension, for a matrix, rownames are used.

Value

returns a vector containing factors or numbers

Author(s)

Stefan Schlager

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Examples

```
data <- matrix(rnorm(200),100,2)
id <- paste("id",1:100,sep="")
pop <- c(rep("pop1",50),rep("pop2",50))
sex <- c(rep("male",50),rep("female",50))
age <- floor(rnorm(100,mean=50,sd=10))
rownames(data) <- paste(id,pop,sex,age,sep="_")
infos <- data.frame(pop=name2factor(data,which=2))
infos$age <- name2num(data,which=4)
infos$pop.sex <- name2factor(data,which=2:3)</pre>
```

NNshapeReg

Estimate the shape by averaging the shape of the nearest neighbours.

Description

Estimate the shape of one set of landmarks by averaging the shape of the nearest neighbours obtained by a second set of landmarks. Weights are calculated either form Mahalanobis or Procrustes distances. This can be useful for data with missing landmarks.

Usage

```
NNshapeReg(
   x,
   y = NULL,
   n = 3,
   mahalanobis = FALSE,
   mc.cores = parallel::detectCores()
)
```

Arguments

an array or matrix (one row per specim) with data used for estimating weights.

y an array or matrix (one row per specim) with landmark data on which the weighted averaging is applied for prediction. If NULL, x will be used for both tasks.

n amount of nearest neighbours to consider

mahalanobis logical: use mahalanobis distance

mc.cores integer: amount of cores used for parallel processing.

Details

This function calculates weights from one set of shape data and then estimates the shape of another (or same) set of landmarks. CAUTION: landmark data has to be registered beforehand.

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Value

matrix or array of estimates.

See Also

```
proc.weight, fixLMtps
```

Examples

```
if (require(shapes)) {
proc <- procSym(gorf.dat)
#use the closest 3 specimen based on the first 4 landmarks
#to estimate the shape
estim <- NNshapeReg(proc$rotated[1:4,,],proc$rotated,n=3,mc.cores=1)
#compare estimation and true config
plot(proc$rotated[,,1],asp=1)
points(estim[,,1],col=2)
}</pre>
```

nose

landmarks and a triangular mesh representing a human nose

Description

triangular mesh representing a human nose and two matrices containing landmark data

Format

shortnose.mesh: A triangular mesh of class 'mesh3d'.

shortnose.lm: matrix containing example landmark data placed on shortnose.mesh.

longnose.lm: matrix containing example landmark data representing a caricaturesquely deformed human nose.

pcAlign

align two 3D-pointclouds/meshes by their principal axes

Description

align two 3D-pointclouds/meshes by their principal axes

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Usage

```
pcAlign(x, y, optim = TRUE, subsample = NULL, iterations = 10, mc.cores = 2)
## S3 method for class 'matrix'
pcAlign(x, y, optim = TRUE, subsample = NULL, iterations = 10, mc.cores = 2)
## S3 method for class 'mesh3d'
pcAlign(x, y, optim = TRUE, subsample = NULL, iterations = 10, mc.cores = 2)
```

Arguments

X	matrix or mesh3d
У	matrix or mesh3d, if missing, x will be centered by its centroid and aligned by its princial axis.
optim	logical if TRUE, the RMSE between reference and target will be minimized testing all possible axes alignments and (if iterations > 0) followed by a rigid ICP procedure.
subsample	integer: use subsampled points to decrease computation time of optimization.
iterations	integer: number of iterations for optimization (the higher the more accurate but also more time consuming).
mc.cores	use parallel processing to find best alignment to original shape.

Details

x and y will first be centered and aligned by their PC-axes. If optim=TRUE, all possible 8 ordinations of PC-axes will be tested and the one with the smallest RMSE between the transformed version of x and the closest points on y will be used. Then the rotated version of x is translated to the original center of mass of y.

Value

rotated and translated version of x to the center and principal axes of y.

```
data(boneData)
blm1 <- pcAlign(boneLM[,,1],boneLM[,,2])
## Not run:
require(rgl)
spheres3d(boneLM[,,1])#original position
spheres3d(blm1,col=2)#aligned configuration
spheres3d(boneLM[,,2],col=3)#target
## End(Not run)</pre>
```

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pcaplot3d

visualization of shape variation

Description

visualization of shape change

Usage

```
pcaplot3d(x, ...)
## S3 method for class 'symproc'
pcaplot3d(
  х,
 pcshow = c(1, 2, 3),
 mag = 3,
 color = 4,
 lwd = 1,
  sym = TRUE,
 legend = TRUE,
  type = c("spheres", "points"),
)
## S3 method for class 'nosymproc'
pcaplot3d(
 pcshow = c(1, 2, 3),
 mag = 3,
  color = 4,
  lwd = 1,
 legend = TRUE,
  type = c("spheres", "points"),
)
```

Arguments

X	a object derived from the function procSym calculated on 3D coordinates.
	Additional parameters which will be passed to the methods.
pcshow	a vector containing the PCscores to be visualized.
mag	a vector or an integer containing which standard deviation of which PC has to be visualized.
color	color of the 3d points/spheres.
lwd	width of the lines representing the shape change.

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logical: if TRUE the symmetric component of shape is displayed. Otherwise the sym

asymmetric one.

logical: if TRUE a legend explaining the color coding of the PCs is plotted. legend

character: for type="spheres", the landmarks will be rendered using rgl's type

spheres3d function and for type="points" by points3d respectivly.

Details

visualization of the shape changes explained by Principal components

Value

returns an invisible array containing the shapes associated with the Principal components selected.

See Also

procSym

Examples

```
## Not run:
data(boneData)
proc <- procSym(boneLM)</pre>
pcaplot3d(proc,pcshow=1:3,mag=-3)#only one PC available
## End(Not run)
```

PCdist

correlation between a reduced space and the original space

Description

Calculates the correlation between distances in a reduced space and the original space

Usage

```
PCdist(PCs, PCscores, x = 5, plot.type = "b")
```

Arguments

PCs	m x k matrix of Principal Components where m is the k is the number of PCs.
PCscores	${\bf n}$ x m matrix of Principal Component scores where ${\bf n}$ is the number of observations.
X	integer: increment for every x-th PC the subspace to fullspace correlation will be calculated.

"b"=barplot of correlation values, "s"=line between correlation values. plot.type

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Value

a vector of R-squared values between subspace and fullspace distances and a barplot depicting the correlations belonging to the subspace.

Author(s)

Stefan Schlager

Examples

```
if (require(shapes)) {
a <- procSym(gorf.dat)
PCdist(a$PCs, a$PCscores, x = 2)
}</pre>
```

permudist

performs permutation testing for group differences.

Description

This function compares the distance between two groupmeans to the distances obtained by random assignment of observations to this groups.

Usage

```
permudist(
  data,
  groups,
  rounds = 1000,
  which = NULL,
  p.adjust.method = "none",
  median = FALSE
)
```

Arguments

data array or matrix containing data groups factors determining grouping. rounds number of permutations

which integer (optional): in case the factor levels are > 2 this determins which factor-

levels to use

p.adjust.method

method to adjust p-values for multiple comparisons see p.adjust.methods for

options.

median logical: if TRUE, comparison will be median instead of mean.

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Value

```
dist distance matrix with distances between actual group means

p.adjust.method method used for p-value adjustion

p.value distance matrix containing pairwise p-values obtained by comparing the actual distance to randomly acquired distances
```

Examples

```
data(boneData)
proc <- procSym(boneLM)
groups <- name2factor(boneLM,which=3)
perm <- permudist(proc$PCscores[,1:10], groups=groups, rounds=10000)

## now we concentrate only on sex dimorphism between Europeans
groups <- name2factor(boneLM,which=3:4)
levels(groups)
perm1 <- permudist(proc$PCscores, groups=groups,which=3:4, rounds=10000)</pre>
```

permuvec

perfom permutation testing on angles and distances between subgroups of two major groups.

Description

perform permutation test on length and angle of the vectors connecting the subgroup means of two groups: e.g. compare if length and angle between sex related differences in two populations differ significantly.

Usage

```
permuvec(
  data,
  groups,
  subgroups = NULL,
  rounds = 9999,
  scale = TRUE,
  tol = 1e-10,
  mc.cores = parallel::detectCores()
)
```

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Arguments

data array or matrix containing data.

groups factors of firs two grouping variables.

subgroups factors of the subgrouping.

rounds number of requested permutation rounds

scale if TRUE: data will be scaled by pooled within group covarivance matrix. Other-

wise Euclidean distance will be used for calculating distances.

tol threshold for inverting covariance matrix.

mc.cores integer: determines how many cores to use for the computation. The default

is autodetect. But in case, it doesn't work as expected cores can be set manu-

ally.Parallel processing is disabled on Windows due to occasional errors.

Details

This function calculates means of all four subgroups and compares the residual vectors of the major grouping variables by angle and distance.

Value

angle angle between the vectors of the subgroups means

distances between subgroups

meanvec matrix containing the means of all four subgroups

permutangles vector containing angles (in radians) from random permutation

permudists vector containing distances from random permutation

p.angle p-value of angle between residual vectors

p.distp-value of length difference between residual vectorssubdistlength of residual vectors connecting the subgroups

means.

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placePatch

Project semi-landmarks from a predefined atlas onto all specimen in a sample

Description

Project semi-landmarks from a predefined atlas onto all specimen in a sample. Various mechanisms are implemented to avoid errorneous placement on the wrong surface layer (e.g. inside the bone).

Usage

```
placePatch(
   atlas,
   dat.array,
   path,
   prefix = NULL,
   fileext = ".ply",
   ray = TRUE,
   inflate = NULL,
   tol = inflate,
   relax.patch = TRUE,
   keep.fix = NULL,
   rhotol = NULL,
   silent = FALSE,
   mc.cores = 1
)
```

Arguments

atlas

object of class "atlas" created by createAtlas

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dat.array k x 3 x n array containing reference landmarks of the sample or a matrix in case of only one target specimen. character: specify the directory where the surface meshes of the sample are path stored. prefix character: prefix to the specimens names (stored in dimnames (dat.array)[[3]]) to match the corresponding file names. If dat.array has no dimnames (e.g. because it is a matrix - see example below), this can also be a character vector containing the filenames to which fileext will be appended. character: file extension of the surface meshes. fileext logical: projection will be along surface normals instead of simple closest point ray search. inflate (or deflate - if negative sign) the semilandmarks along the normals of the inflate deformed atlas to make sure that they stay on the outside (inside) of the target mesh. tol numeric: threshold to follow the ray back after inflation. See details below. If no surface is hit after to 1 mm, the simple closest point will be used. logical: request relaxation minimising bending energy toward the atlas. relax.patch keep.fix integer: rowindices of those landmarks that are not allowed to be relaxed in case relax.patch=TRUE. If not specified, all landmarks will be kept fix. This is preferably set during atlas creation with createAtlas: In case you specified corrCurves on the atlas, you should define explicitly which landmarks (also on the curves) are supposed to fix to prevent them from sliding. rhotol numeric: maximum amount of deviation a hit point's normal is allowed to deviate from the normal defined on the atlas. If relax.patch=TRUE, those points exceeding this value will be relaxed freely (i.e. not restricted to tangent plane). silent logical: suppress messages. mc.cores run in parallel (experimental stuff now even available on Windows). On windows this will only lead to a significant speed boost for many configurations, as all required packages (Morpho and Rvcg) need to be loaded by each newly spawned process.

Details

This function allows the (relatively) easy projection of surface points defined on an atlas onto all surface of a given sample by Thin-Plate Spline deformation and additional mechanisms to avoid distortions. The algorithm can be outlined as followed.

- 1. relax curves (if specified) against atlas.
- 2. deform atlas onto targets by TPS based on predefined landmarks (and curves).
- 3. project coordinates on deformed atlas onto target mesh
- 4. 'inflate' or 'deflate' configuration along their normals to make sure all coordinates are on the outside/inside
- 5. Project inflated points back onto surface along these normals.
- 6. Check if normals are roughly pointing into the same direction as those on the (deformed) atlas.
- 7. Relax all points against atlas.
- 8. the predefined coordinates will note change afterwards!

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Value

array containing the projected coordinates appended to the data.array specified in the input. In case dat.array is a matrix only a matrix is returned.

Author(s)

Stefan Schlager

References

Schlager S. 2013. Soft-tissue reconstruction of the human nose: population differences and sexual dimorphism. PhD thesis, Universitätsbibliothek Freiburg. URL: http://www.freidok.uni-freiburg.de/volltexte/9181/.

See Also

```
createAtlas, relaxLM, checkLM, slider3d, tps3d
```

```
## Not run:
data(nose)
require(rgl)
###create mesh for longnose
longnose.mesh <- tps3d(shortnose.mesh,shortnose.lm,longnose.lm,threads=1)</pre>
## create atlas
fix <- c(1:5,20:21)
atlas <- createAtlas(shortnose.mesh, landmarks =</pre>
           shortnose.lm[fix,], patch=shortnose.lm[-c(1:5,20:21),])
## view atlas
plotAtlas(atlas)
## create landmark array with only fix landmarks
data <- bindArr(shortnose.lm[fix,], longnose.lm[fix,], along=3)</pre>
dimnames(data)[[3]] <- c("shortnose", "longnose")</pre>
### write meshes to disk
mesh2ply(shortnose.mesh, filename="shortnose")
mesh2ply(longnose.mesh, filename="longnose")
patched <- placePatch(atlas, data, path="./", inflate=5)</pre>
## now browse through placed patches
checkLM(patched, path="./", atlas=atlas)
## same example with only one target specimen
data <- longnose.lm[fix, ]</pre>
patched <- placePatch(atlas, data, prefix="longnose", path="./", inflate=5)</pre>
wire3d(longnose.mesh,col=3)
```

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```
spheres3d(patched)
## End(Not run)
```

plot.slider3d

plot the result of slider3d

Description

plot the result of slider3d

Usage

```
## S3 method for class 'slider3d'
plot(
    x,
    cols = 2:4,
    pt.size = NULL,
    point = c("sphere", "point"),
    specimen = 1,
    add = TRUE,
    ...
)
```

Arguments

Х	result of slider3d call
cols	vector containing colors for each coordinate type cols[1]=landmarks, cols[2]=surface landmarks, cols[3]=outlines.
pt.size	size of plotted points/spheres. If point="s". pt.size defines the radius of the spheres. If point="p" it sets the variable size used in point3d.
point	how to render landmarks.
specimen	integer: select the specimen to plot
add	logical: if TRUE, a new rgl window is opened.
	additional, currently unused parameters

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plotAtlas visualize an atlas defined by createAtlas

Description

visualize an atlas defined by createAtlas

Usage

```
plotAtlas(
  atlas,
  pt.size = NULL,
  alpha = 1,
  render = c("w", "s"),
  point = c("s", "p"),
  meshcol = "white",
  add = TRUE,
  legend = TRUE,
  cols = 2:5
)
```

Arguments

atlas	object of class atlas created by createAtlas.
pt.size	size of plotted points/spheres. If point="s". pt.size defines the radius of the spheres. If point="p" it sets the variable size used in point3d.
alpha	value between 0 and 1. Sets transparency of mesh 1=opaque 0= fully transparent.
render	if render="w", a wireframe will be drawn, if render="s", the mesh will be shaded.
point	how to render landmarks. "s"=spheres, "p"=points.
meshcol	color to render the atlas mesh
add	logical: if TRUE, a new rgl window is opened.
legend	logical: request plot of legend specifying landmark coloring.
cols	vector containing colors for each coordinate type cols[1]=landmarks, cols[2]=patch, cols[3]=corrCurves, cols[4]=patchCurves.

Details

If legend=TRUE, a plot with a legend will open where coloring of the 3D-spheres is specified.

Value

returns invisible vector containing rgl.id of rendered objects.

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See Also

```
placePatch, createAtlas
```

Examples

plotNormals

plots the normals of a triangular surface mesh.

Description

visualises the vertex normals of a triangular surface mesh of class mesh3d. If no normals are contained, they are computed.

Usage

```
plotNormals(x, length = 1, lwd = 1, col = 1, ...)
```

Arguments

X	object of class "mesh3d"
length	either a single numeric value or a numeric vector defining per-normals lenght (default is 1)
lwd	width of the normals
col	color of the normals
	addtional parameters, currently not in use.

Author(s)

Stefan Schlager

```
## Not run:
require(rgl)
data(nose)
plotNormals(shortnose.mesh,col=4,length=0.01)
shade3d(shortnose.mesh,col=3)
```

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```
## End(Not run)
```

pls2B

Two-Block partial least square regression.

Description

Performs a Two-Block PLS on two sets of data and assesses the significance of each score by permutation testing

Usage

```
pls2B(
    x,
    y,
    tol = 1e-12,
    same.config = FALSE,
    rounds = 0,
    useCor = FALSE,
    cv = FALSE,
    cvlv = NULL,
    mc.cores = parallel::detectCores(),
    ...
)
```

Arguments

x	array containing superimposed landmark data second block. Matrices are also allowed but the option 'same.config' will not work.
у	array containing superimposed landmark data of the first block. Matrices are also allowed but the option 'same.config' will not work.
tol	threshold for discarding singular values.
same.config	logical: if TRUE each permutation includes new superimposition of permuted landmarks. This is necessary if both blocks originate from landmarks that are superimposed together.
rounds	rounds of permutation testing.
useCor	if TRUE, the correlation matrix instead of the covariance matrix is used.
cv	logical: if TRUE, a leave-one-out cross-validation is performed
cvlv	integer: number of latent variables to test
mc.cores	integer: determines how many cores to use for the
	arguments passed to ProcGPA computation. The default is autodetect. But in case, it doesn't work as expected cores can be set manually. Parallel processing

is disabled on Windows due to occasional errors.

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Details

The Two-Block PLS tries to find those linear combinations in each block maximising the covariance between blocks. The significance of each linear combination is assessed by comparing the singular value to those obtained from permuted blocks. If both blocks contain landmarks superimposed TOGETHER, the option same.config=TRUE requests superimposition of the permuted configurations (i.e. where the landmarks of block x are replaced by corresponding landmarks of other specimen.

Value

svd singular value decomposition (see svd) of the 'common' covariance block

Xscores PLS-scores of x Yscores PLS-scores of y

CoVar Dataframe containing singular values, explained covariation, correlation coef-

fictient between PLS-scores and p-values for singular values obtained from per-

mutation testing

xlm linear model: lm(Xscores ~ Yscores - 1)
ylm linear model: lm(Yscores ~ Xscores - 1)

predicted.x array containing matrices of cross-validated predictions for x(landmarks arrays

will be vectorized using vecx)

predicted.y array containing matrices of cross-validated predictions for y (landmarks arrays

will be vectorized using vecx)

rv RV-coefficient

p.value.RV p-value for RV-coefficient determined by permutation testing

Author(s)

Stefan Schlager

References

Rohlf FJ, Corti M. 2000. Use of two-block partial least-squares to study covariation in shape. Systematic Biology 49:740-753.

See Also

```
plsCoVar, getPLSfromScores, predictPLSfromScores, getPLSscores, predictPLSfromData,svd, plsCoVarCommonShape, getPLSCommonShape
```

```
if (require(shapes)) {
### very arbitrary test:
### check if first 4 landmarks covaries with the second 4
proc <- procSym(gorf.dat)
## we do only 50 rounds to minimize computation time</pre>
```

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```
## Not run: #same.config takes too long for CRAN check
pls1 <- pls2B(proc$rotated[1:4,,],proc$rotated[5:8,,],</pre>
                                          same.config=TRUE,rounds=50,mc.cores=2)
## End(Not run)
pls1 <- pls2B(proc$rotated[1:4,,],proc$rotated[5:8,,],</pre>
                                          same.config=FALSE,rounds=50,mc.cores=1)
 layout(matrix(1:4,2,2,byrow=TRUE))
 for(i in 1:4)
   plot(pls1$Xscores[,i]~pls1$Yscores[,i])
 ## predict first 4 landmarks from second 4 for first config
 layout(1)
predPLS <- predictPLSfromData(pls1,y=proc$rotated[5:8,,1])</pre>
## show differences between prediction and original
deformGrid2d(predPLS,proc$rotated[1:4,,1],pch=19)
 ##plot the complete first config
points(proc$rotated[,,1])
 ##show effects of first latent variable
plsEffects <- plsCoVar(pls1,i=1)</pre>
deformGrid2d(plsEffects\$x[,,1],plsEffects\$x[,,2])\#\#show\ on\ x
\label{lem:condition} deform Grid 2d (plsEffects \$y[,,1], plsEffects \$y[,,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,,1], plsEffects \$y[,,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,,1], plsEffects \$y[,,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,,1], plsEffects \$y[,,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,,1], plsEffects \$y[,,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,2], add = TRUE, pch = 19) \# show on y deform Grid 2d (plsEffects \$y[,2], add = TRUE, pch = 19) \# show of y deform Grid 2d (plsEffects \$y[,2], add = TRUE, pch 
 ##show effects of 2nd latent variable
plsEffects2 <- plsCoVar(pls1,i=2)
\label{lem:condition} deform \texttt{Grid2d(plsEffects2$x[,,1],plsEffects2$x[,,2])\#\#show\ on\ x}
\label{lem:condition} deform \texttt{Grid2d(plsEffects2\$y[,,1],plsEffects2\$y[,,2],add=TRUE,pch=19)\#\#show\ on\ y}
```

plsCoVar

Get the shape changes from pls2B associated with each latent variable

Description

Get the shape changes from pls2B associated with each latent variable

Usage

```
plsCoVar(pls, i, sdx = 3, sdy = 3)
```

Arguments

pls	output of pls2B
i	integer: which latent variable to show. E.g. i=3 will show the changes associated with the 3rd latent variable.
sdx	standard deviation on the xscores. sdx=3 will show the effecs of -3sd vs +3sd
sdy	standard deviation on the yscores. sdy=3 will show the effecs of -3sd vs +3sd

Value

Χ	matrix/array	with reconstructed x
X	mau ix/amay	with reconstructed x

y matrix/array with reconstructed y, with each prediction named accordingly: e.g.

 $neg_x_sd_3$ means the prediction of x at a score of -3*sd(Xscores)

.

See Also

 $\verb|pls2B|, getPLS from Scores|, predictPLS from Scores|, getPLS scores|, predictPLS from Data, svd, \\ \verb|plsCoVarCommonShape|$ |

plsCoVarCommonShape

Compute the shape changes along the common axis of deformations

Description

Compute the shape changes between two blocks of 2D or 3D shape coordinates along the common axis of deformations defined by each dimension of the latent space

Usage

```
plsCoVarCommonShape(pls, i, sdcommon = 1)
```

Arguments

pls object of class "pls2B"

i integer: dimension of latent space to show shape changes for

sdcommon standard deviations derived from scores scaled to a consensus scale

Value

returns an $k \times m \times 2$ array with the common shape changes associated with +-sdcommon SD of the i-th latent dimension

Note

this give the same results as plsCoVar, however, using common shape vectors as suggested by Mitteroecker and Bookstein (2007)

References

Mitteroecker P, Bookstein F. 2007. The conceptual and statistical relationship between modularity and morphological integration. Systematic Biology 56(5):818-836.

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See Also

 $\verb|pls2B|, getPLSfromScores|, predictPLSfromScores|, getPLSscores|, predictPLSfromData, svd, plsCoVar, getPLSCommonShape|$

Examples

```
data(boneData)
proc <- procSym(boneLM)
pls <- pls2B(proc$orpdata[1:4,,],proc$orpdata[5:10,,])
commShape <- getPLSCommonShape(pls)
## get common shape for first latent dimension at +-2 sd of the scores
pred <- plsCoVarCommonShape(pls,1,2)
## Not run:
deformGrid3d(pred[,,1],pred[,,2])
## End(Not run)</pre>
```

points2plane

projects a 3D coordinate orthogonally onto a plane

Description

projects a 3D coordinate orthogonally onto a plane

Usage

```
points2plane(x, v1, normal = NULL, v2 = NULL, v3 = NULL)
```

Arguments

X	3D-vector or a k x 3 matrix with 3D vectors stored in rows
v1	point on plane
normal	plane normal (overrides specification by v2 and v3)
v2	if pNorm=NULL, the plane will be defined by three points v1, v2, v3 $$
v3	if pNorm=NULL, the plane will be defined by three points v1, v2, v3

Value

projected point

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Examples

```
data(boneData)
##project rhinion onto plane spanned by Nasion and both Nariales
rpro <- points2plane(boneLM[10,,1],v1=boneLM[9,,1],v2=boneLM[3,,1],v3=boneLM[4,,1])</pre>
## Not run:
require(rgl)
#visualize
wire3d(skull_0144_ch_fe.mesh,col="white")
##get plane normal
normal <- crossProduct(boneLM[3,,1]-boneLM[9,,1],boneLM[4,,1]-boneLM[9,,1])</pre>
#' ## get plane offset
d <- norm(points2plane(c(0,0,0),v1=boneLM[9,,1],normal=normal),"2")</pre>
spheres3d(boneLM[,,1],radius=0.5)
spheres3d(boneLM[c(3,4,9),,1],radius=0.6,col=3)
##original position of Rhinion
spheres3d(boneLM[10,,1],radius=0.6,col=2)
##projected onto plane
spheres3d(rpro,radius=0.9,col=6)
lines3d(rbind(rpro,boneLM[10,,1]),lwd=3)
##plot plane
planes3d(normal[1],normal[2],normal[3],d=d,col=2,alpha=0.5)
##now we project all points onto that plane:
spheres3d(points2plane(boneLM[,,1],v1=boneLM[9,,1],v2=boneLM[3,,1],v3=boneLM[4,,1]),col=3)\\
## and finally project the vertices of the mesh onto the plane
meshpro <- points2plane(vert2points(skull_0144_ch_fe.mesh),v1=boneLM[9,,1],normal=normal)</pre>
points3d(meshpro,col=2)
## End(Not run)
```

prcompfast

fast Principal Component Analysis (PCA)

Description

```
fast Principal Component Analysis (PCA)
```

Usage

```
prcompfast(x, retx = TRUE, center = TRUE, scale. = FALSE, tol = NULL, ...)
```

Arguments

a numeric or complex matrix (or data frame) which provides the data for the Х principal components analysis.

a logical value indicating whether the rotated variables should be returned

retx

predict.bgPCA 109

center a logical value indicating whether the variables should be shifted to be zero

centered. Alternately, a vector of length

scale. a logical value indicating whether the variables should be scaled to have unit

variance before the analysis takes place. The default is FALSE for consistency with S, but in general scaling is advisable. Alternatively, a vector of length equal the number of columns of x can be supplied. The value is passed to scale. equal the number of columns of x can be supplied. The value is passed to scale.

tol a value indicating the magnitude below which components should be omitted.

(Components are omitted if their standard deviations are less than or equal to tol times the standard deviation of the first component.) With the default null setting, no components are omitted. Other settings for tol could be tol = 0 or tol = sqrt(.Machine\$double.eps), which would omit essentially constant

components.

... arguments passed to or from other methods.

Value

prcomp returns a list with class prcomp containing the followin components:

sdev the standard deviations of the principal components (i.e., the square roots of

the eigenvalues of the covariance/correlation matrix, though the calculation is

actually done with the singular values of the data matrix).

rotation: the matrix of variable loadings (i.e., a matrix whose columns contain the eigen-

vectors). The function princomp returns this in the element loadings.

x: if retx is true the value of the rotated data (the centred (and scaled if requested)

data multiplied by the rotation matrix) is returned. Hence, cov(x) is the diagonal matrix diag(sdev^2). For the formula method, napredict() is applied

to handle the treatment of values omitted by the na. action.

center, scale: the centering and scaling used, or FALSE

. pcafast <- prcompfast(iris[,1:4]) pcadefault <- prcompfast(iris[,1:4]) ## check if both results are idential (ignoring the sign) all.equal(lapply(pcafast,abs),lapply(pcadefault,abs))

Note

this function returns the same results as prcomp (apart from sign differences) but uses smarter matrix decompositions making it faster for $nrow(x) \gg ncol(x)$ and $nrow(x) \ll ncol(x)$.

predict.bgPCA Compute between-group-PC scores from new data

Description

Compute between-group-PC scores from new data

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Usage

```
## S3 method for class 'bgPCA'
predict(object, newdata, ...)
```

Arguments

object of class bgPCA returned from groupPCA

newdata matrix or 3D array containing data in the same format as originally used to

compute groupPCA

... currently not used.

Value

returns the between-group-PC scores for new data

Examples

```
data(boneData)

boneLMPart <- boneLM[,,-(1:2)]
procPart <- procSym(boneLMPart)
pop_sex <- name2factor(boneLMPart, which=3:4)
## compute group PCA without first 2 specimens
gpcaPart <- groupPCA(procPart$orpdata, groups=pop_sex, rounds=0, mc.cores=2,cv=FALSE)
## align new data to Procrustes analysis
newdata <- align2procSym(procPart,boneLM[,,1:2])
## get scores for new data
newscores <- predict(gpcaPart,newdata)</pre>
```

predict.CVA

Compute CV-scores from new data

Description

Compute CV-scores from new data

Usage

```
## S3 method for class 'CVA'
predict(object, newdata, ...)
```

Arguments

object of class CVA

newdata matrix or 3D array containing data in the same format as originally used to

compute CVA

... currently not used.

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Value

returns the CV-scores for new data

predictPLSfromData

predict 2 Block-PLS from new data

Description

predict 2 Block-PLS from new data

Usage

```
predictPLSfromData(pls, x, y, ncomp = NULL)
```

Arguments

pls	output of pls2B
x	data in the same format as in original pls2B (for landmarks this can be an array or a matrix and for other data a matrix of a vector)
У	data in the same format as in original pls2B (for landmarks this can be an array or a matrix and for other data a matrix of a vector)
ncomp	number of (latent) components to use for prediction.

Value

returns an array/matrix/vector of predictions - depending on input for computing pls

Note

either x or y must be missing

See Also

```
pls2B, getPLSscores,predictPLSfromScores
```

```
##see examples in pls2B
```

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predictPLSfromScores predict data from 2-Block PLS-scores

Description

predict data from 2-Block PLS-scores

Usage

```
predictPLSfromScores(pls, x, y)
```

Arguments

pls output of pls2B

x scores associated with dataset x in original pls2B scores associated with dataset y in original pls2B

Value

returns an array/matrix of landmarks or original values, depending on input for computing pls

Note

either x or y must be missing. If x-scores are provided, the yscores will be estimated and the predictions calculated.

See Also

pls2B, getPLSscores,predictPLSfromData

predictRelWarps

predict relative warps for data not included in the training data set

Description

predict relative warps for data not included in the training data set

Usage

```
predictRelWarps(x, newdata, noalign = FALSE)
```

Arguments

x output from relWarps

newdata k x m x n array holding new landmark data

noalign logical: if TRUE, data is assumed to be already aligned to training data and

alignment is skipped.

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Details

This function aligns the new data to the mean from x and transforms it into the relative warp space computed from the training data.

Value

```
returns a list containing
```

```
bescores relative warp scores (PC-scores if alpha = 0)
uniscores uniform scores, NULL if alpha = 0
```

Examples

```
data(boneData)
set.seed(42)
training <- sample(1:80,size=60)
rW1 <- relWarps(boneLM[,,training], alpha = -1)
## predict scores for the entire sample
predAll <- predictRelWarps(rW1,boneLM)

## now compare the scores predicted scores to the original ones
layout(matrix(1:4,2,2))
for (i in 1:2) {
   plot(rW1$bescores[,i],predAll$bescores[training,i],main=paste("RW",i))
   plot(rW1$uniscores[,i],predAll$uniscores[training,i],main=paste("UC",i))
}</pre>
```

predictShape.lm

Predict shapes based on linear models calculated from PCscores

Description

Predict shapes based on linear models calculated from PCscores.

Usage

```
predictShape.lm(fit, datamod, PC, mshape)
```

Arguments

fit	model of class 1m where the PCscores are fitted onto
datamod	a one-sided "model" formula, of the form $\sim x1 + x2 + \ldots + xk$, corresponding to the right hand term in the model used in fit. If omitted, the predicted shapes of all specimen are calculated based on the fitted values.
PC	Matrix/vector containing Principal components (rotation matrix) corresponding to PC-scores used in fit.
mshape	matrix of the meanshape's landmarks by which the data was centered before rotation in covariance eigenspace.

114 predictShape.lm

Details

This function predicts the landmarks based on models calculated from PCscores.

Value

```
predicted array or matrix containing predicted landmark coordinates
predictedPC matrix containing predicted PC-scores
```

Warning

Make sure that the levels of the variables used in datamod correspond exactly to those used in fit. Otherwise model matrix will be calculated erroneous.

See Also

```
model.matrix, lm, formula
```

```
data(boneData)
proc <- procSym(boneLM)</pre>
pop <- name2factor(boneLM,which=3)</pre>
##easy model with only one factor based on the first four PCs
fit <- lm(proc$PCscores[,1:4] ~ pop)</pre>
## get shape for Europeans only
datamod <- ~as.factor(levels(pop))[2]</pre>
Eu <- predictShape.lm(fit,datamod, proc$PCs[,1:4],proc$mshape)</pre>
## get shape for Europeans and Chinese
datamod <- ~as.factor(levels(pop))</pre>
pred <- predictShape.lm(fit,datamod, proc$PCs[,1:4],proc$mshape)</pre>
deformGrid3d(pred$predicted[,,1], pred$predicted[,,2], ngrid = 0)
## End(Not run)
## more complicated model
sex <- name2factor(boneLM, which=4)</pre>
fit <- lm(proc$PCscores[,1:4] ~ pop*sex)</pre>
## predict female for chinese and European
datamod <- ~(as.factor(levels(pop))*rep(as.factor(levels(sex))[1],2))</pre>
pred <- predictShape.lm(fit,datamod, proc$PCs[,1:4],proc$mshape)</pre>
## predict female and malefor chinese and European
popmod <- factor(c(rep("eu",2),rep("ch",2)))</pre>
sexmod <- rep(as.factor(levels(sex)),2)</pre>
datamod <- ~(popmod*sexmod)</pre>
pred <- predictShape.lm(fit,datamod, proc$PCs[,1:4],proc$mshape)</pre>
```

proc.weight 115

```
## add some (randomly generated) numeric covariate
somevalue <- rnorm(80,sd=10)
fit <- lm(proc$PCscores[,1:4] ~ pop+somevalue)
probs <- quantile(somevalue, probs=c(0.05, 0.95))
## make model for European at 5% and 95% quantile
popmod <- rep(factor(levels(pop))[2],2)
datamod <- ~(popmod+probs)
pred <- predictShape.lm(fit,datamod, proc$PCs[,1:4],proc$mshape)</pre>
```

proc.weight

calculate weights inverse to the distances from the specified observa-

Description

for calculation of a shape model by averaging the observations neighbouring the configuration in question, it is necessary to calculate weights by similarity.

Usage

```
proc.weight(
  data,
  number,
  ref,
  report = TRUE,
  reg = 0,
  log = FALSE,
  mahalanobis = FALSE,
  weightfun = NULL
)
```

Arguments

data array containing landmark configurations

number integer: how many of the neighbours are to be involved.

ref integer: position in the array that is used as reference.

report logical: require report about name of the reference.

reg numeric: regularise mahalanobis distance by adding reg to the diagonal of eigen-

values of the covariance matrix.

log logical: use the logarithm of the distances.

mahalanobis logical: use mahalanobis distance.

weightfun custom function that operates on a vector of distances (see examples) and gen-

erates weights accordingly.

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Details

distances of zero will get a weight of 1e12 (this is scaled to all weights summing to one), thus weights for observations further away are converging to zero.

Value

data dataframe containing id, procrustes/mahalanobis distance and weight according

to the reference

reference returns observations' names if available

rho.all dataframe containing distances to references of all observations

Examples

```
if (require(shapes)) {
proc <- procSym(gorf.dat)</pre>
##get weights for the the four specimen closest to the first observation.
weights <- proc.weight(proc$rotated,4,1)</pre>
##estimate the first specimen by weighted neighbour shapes.
estim <- proc$mshape*0;</pre>
for (i in 1:4)
{estim <-estim+proc$rotated[,,weights$data$nr[i]]*weights$data$weight[i]}</pre>
### visualise
plot(estim,asp=1)## show estimation
points(proc$rotated[,,1],col=3)##show original
## use a gaussian smoother to compute weights using a bandwidth of 0.05
gaussWeight <- function(r,sigma=0.05) {</pre>
   sigma <- 2*sigma^2
   return(exp(-r^2/ sigma))
weights <- proc.weight(proc$rotated,4,1,weightfun=gaussWeight)</pre>
}
```

procAOVsym

Procrustes ANOVA for structures with object symmetry

Description

Procrustes ANOVA for structures with object symmetry, currently only supporting the factors 'specimen', 'side' and the interaction term.

Usage

```
procAOVsym(symproc, indnames = NULL)
```

procAOVsym 117

Arguments

symproc object returned by procSym, where pairedLM is specified

indnames vector containing specimen identifiers. Only necessary, if data does not contain

dimnames containing identifiers

Details

performs a Procrustes ANOVA for configurations with object symmetry (as described in Klingenberg et al. 2002).

Value

returns a dataframe containing Sums of Squares for each factor.

Note

In future releases the implementation of support for bilateral symmetry and more factors is intended.

Author(s)

Stefan Schlager

References

Klingenberg CP, Barluenga M, Meyer A. 2002. Shape analysis of symmetric structures: quantifying variation among individuals and asymmetry. Evolution 56:1909-20.

See Also

procSym

```
data(boneData)
left <- c(4,6,8)
## determine corresponding Landmarks on the right side:
# important: keep same order
right <- c(3,5,7)
pairedLM <- cbind(left,right)
symproc <- procSym(boneLM, pairedLM=pairedLM)
procAOVsym(symproc)</pre>
```

ProcGPA

ProcGPA	Workhorse function for procSym, responsible for Procrustes registra-
	tion

Description

Workhorse function for procSym, responsible for Procrustes registration

Usage

```
ProcGPA(
  dat.array,
  tol = 1e-05,
  scale = TRUE,
  CSinit = FALSE,
  silent = TRUE,
  weights = NULL,
  centerweight = FALSE,
  reflection = TRUE,
  pcAlign = TRUE
)
```

Arguments

dat.array	Input $k \times m \times n$ real array, where k is the number of points, m is the number of dimensions, and n is the sample size.
tol	numeric: Threshold for convergence during iterative superimpositioning.
scale	logical: indicating if scaling is requested
CSinit	logical: if TRUE, all configurations are initially scaled to Unit Centroid Size.
silent	logical: suppress output of elapsed time.
weights	numeric vector: assign per landmark weights.
centerweight	logical: if TRUE, the landmark configuration is scaled according to weights during the rotation process, instead of being scaled to the Centroid size.
reflection	logical: allow reflections.
pcAlign	logical: if TRUE, the shapes are aligned by the principal axis of the first specimen, otherwise the orientation of the first specimen is used.

Value

returns a list with

rotated k x m x n array of the rotated configurations

mshape sample meanshape

Author(s)

Stefan Schlager

References

Goodall C. 1991. Procrustes methods in the statistical analysis of shape. Journal of the Royal Statistical Society. Series B. Statistical Methodology 53:285-239.

Dryden IL, Mardia KV. 1998. Statistical shape analysis. John Wiley and sons, Chichester.

See Also

```
procSym, rotonto
```

Examples

```
data(boneData)
proc <- ProcGPA(boneLM, CSinit=TRUE, silent=TRUE)
#now we landmarks 5 - 9 double the weight as the others
weights <- c(rep(1,4),rep(2,5),1)
proc.wt <- ProcGPA(boneLM, CSinit=TRUE, weights=weights, silent=TRUE)</pre>
```

procSym

Procrustes registration

Description

procSym performs Procrustes superimposition including sliding of semi-landmarks on curves/outlines in 2D and 3D.

Usage

```
procSym(
  dataarray,
  scale = TRUE,
  reflect = TRUE,
  CSinit = TRUE,
  orp = TRUE,
  proctol = 1e-05,
  tol = 1e-05,
  pairedLM = NULL,
  sizeshape = FALSE,
  use.lm = NULL,
  center.part = FALSE,
  weights = NULL,
  centerweight = FALSE,
```

```
pcAlign = TRUE,
distfun = c("angle", "riemann"),
SMvector = NULL,
outlines = NULL,
deselect = FALSE,
recursive = TRUE,
iterations = 0,
initproc = FALSE,
bending = TRUE,
stepsize = 1
)
```

Arguments

dataarray	Input $k \times m \times n$ real array, where k is the number of points, m is the number of dimensions, and n is the sample size.
scale	logical: indicating if scaling is requested to minimize the General Procrustes distance. To avoid all scaling, one has to set CSinit=FALSE, too.
reflect	logical: allow reflections.
CSinit	logical: if TRUE, all configurations are initially scaled to Unit Centroid Size.
orp	logical: if TRUE, an orthogonal projection at the meanshape into tangent space is performed.
proctol	numeric: Threshold for convergence in the alignment process
tol	numeric: Threshold for convergence in the sliding process
pairedLM	A X x 2 matrix containing the indices (rownumbers) of the paired LM. E.g. the left column contains the lefthand landmarks, while the right side contains the corresponding right hand landmarks.
sizeshape	Logical: if TRUE, a log transformed variable of Centroid Size will be added to the shapedata as first variable before performing the PCA.
use.lm	vector of integers to define a subset of landmarks to be used for Procrustes registration.
center.part	Logical: if TRUE, the data superimposed by the subset defined by use.lm will be centered according to the centroid of the complete configuration. Otherwise orp will be set to FALSE to avoid erroneous projection into tangent space.
weights	numeric vector: assign per landmark weights.
centerweight	logical: if TRUE, the landmark configuration is scaled according to weights during the rotation process, instead of being scaled to the Centroid size.
pcAlign	logical: if TRUE, the shapes are aligned by the principal axis of the first specimen
distfun	character: "riemann" requests a Riemannian distance for calculating distances to mean, while "angle" uses an approximation by calculating the angle between rotated shapes on the unit sphere.
SMvector	A vector containing the landmarks on the curve(s) that are allowed to slide

outlines	A vector (or if threre are several curves) a list of vectors (containing the rowindices) of the (Semi-)landmarks forming the curve(s) in the successive position on the curve - including the beginning and end points, that are not allowed to slide.
deselect	Logical: if TRUE, the SMvector is interpreted as those landmarks, that are not allowed to slide.
recursive	Logical: if TRUE, during the iterations of the sliding process, the outcome of the previous iteration will be used. Otherwise the original configuration will be used in all iterations.
iterations	integer: select manually how many iterations will be performed during the sliding process (usefull, when there is very slow convergence). 0 means iteration until convergence.
initproc	Logical: indicating if the first Relaxation step is performed against the mean of an initial Procrustes superimposition. Symmetric configurations will be relaxed against a perfectly symmetrical mean.
bending	if TRUE, bending energy will be minimized, Procrustes distance otherwise (not suggested with large shape differences)
stepsize	integer: dampening factor for the sliding. Useful to keep semi-landmarks from sliding too far off the surface. The displacement is calculated as stepsize * displacement.

Details

This function performs Procrustes registration, allowing a variety of options, including scaling, orthogonal projection into tangentspace and relaxation of semi-landmarks on curves (without reprojection onto the surface/actual outline). It also allows the superimpositioning to be performed using only a subset of the available landmark. For taking into account object symmetry, pairedLM needs to be set. This generates an object of class "symproc". Otherwise an object of class "nosymproc".

Value

size	a vector containing the Centroid Size of the configurations
rotated	k x m x n array of the rotated configurations
Sym	$k\ x\ m\ x$ n array of the Symmetrical component - only available for the "Symmetry"-Option (when pairedLM is defined)
Asym	k x m x n array of the Asymmetrical component. It contains the per-landmark asymmetric displacement for each specimen. Only available for the "Symmetry"-Option (when pairedLM is defined)
asymmean	k x m matrix of mean asymmetric deviation from symmetric mean
mshape	sample meanshape
symmean	meanshape of symmetrized configurations
tan	if orp=TRUE: Residuals in tangentspace else, Procrustes residuals - only available without the "Symmetrie"-Option
PCs	Principal Components - if sizeshape=TRUE, the first variable of the PCs is size information (as log transformed Centroid Size)

PCsym Principal Components of the Symmetrical Component
PCasym Principal Components of the Asymmetrical Component

PCscores PC scores

PCscore_sym PC scores of the Symmetrical Component
PCscore_asym PC scores of the Asymmetrical Component

eigenvalues eigenvalues of the Covariance matrix

eigensym eigenvalues of the "Symmetrical" Covariance matrix
eigenasym eigenvalues of the "Asymmetrical" Covariance matrix

Variance Table of the explained Variance by the PCs

SymVar Table of the explained "Symmetrical" Variance by the PCs

AsymVar Table of the explained "Asymmetrical" Variance by the PCs

orpdata k x m x n array of the rotated configurations projected into tangent space

rho vector of Riemannian distance from the mean

dataslide array containing slidden Landmarks in the original space - not yet processed by

a Procrustes analysis. Only available if a sliding process was requested

meanlogCS mean log-transformed centroid size

Note

For processing of surface landmarks or including the reprojection of slid landmarks back onto 3D-surface representations, the usage of slider3d is recommended.

Author(s)

Stefan Schlager

References

Dryden IL, and Mardia KV. 1998. Statistical shape analysis. Chichester.

Klingenberg CP, Barluenga M, and Meyer A. 2002. Shape analysis of symmetric structures: quantifying variation among individuals and asymmetry. Evolution 56(10):1909-1920.

Gunz, P., P. Mitteroecker, and F. L. Bookstein. 2005. Semilandmarks in Three Dimensions, in Modern Morphometrics in Physical Anthropology. Edited by D. E. Slice, pp. 73-98. New York: Kluwer Academic/Plenum Publishers.

See Also

slider3d

projRead 123

Examples

```
require(rgl)
data(boneData)
### do an analysis of symmetric landmarks
## visualize landmarks on surface
## Not run:
spheres3d(boneLM[,,1])
wire3d(skull_0144_ch_fe.mesh,col=3)
## get landmark numbers
text3d(boneLM[,,1],text=paste(1:10),adj = 1, cex=3)
## End(Not run)
## determine paired Landmarks left side:
left <- c(4,6,8)
## determine corresponding Landmarks on the right side:
# important: keep same order
right <- c(3,5,7)
pairedLM <- cbind(left,right)</pre>
symproc <- procSym(boneLM, pairedLM=pairedLM)</pre>
## Not run:
## visualize first 3 PCs of symmetric shape
pcaplot3d(symproc, sym=TRUE)
## visualize first 3 PCs of asymmetric shape
pcaplot3d(symproc, sym=FALSE)
## visualze distribution of symmetric PCscores population
pop <- name2factor(boneLM, which=3)</pre>
if (require(car)) {
spm(~symproc$PCscore_sym[,1:5], groups=pop)
## visualze distribution of asymmetric PCscores population
spm(~symproc$PCscore_asym[,1:5], groups=pop)
## End(Not run)
```

projRead

Project points onto the closest point on a mesh

Description

project points onto a given surface and return projected points and normals.

Usage

```
projRead(lm, mesh, readnormals = TRUE, smooth = FALSE, sign = TRUE, ...)
```

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Arguments

1m m x 3 matrix containing 3D coordinates.mesh character: specify path to mesh file.

readnormals logical: return normals of projected points.

smooth logical: rerturn smoothed normals.
sign logical: request signed distances.

... additional arguments currently not used.

Value

if readnormals = FALSE, a m x 3 matrix containing projected points is returned, otherwise a list, where

vb 3 x m matrix containing projected points

normals 3 x m matrix containing normals

quality vector containing distances

Author(s)

Stefan Schlager

References

Detection of inside/outside uses the algorithm proposed in:

Baerentzen, Jakob Andreas. & Aanaes, H., 2002. Generating Signed Distance Fields From Triangle Meshes. Informatics and Mathematical Modelling.

See Also

closemeshKD

```
data(nose)
## Not run:
repro <- projRead(shortnose.lm,shortnose.mesh)
## End(Not run)</pre>
```

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qqmat

Q-Q plot to assess normality of data

Description

qqmat plots Mahalanobisdistances of a given sample against those expected from a Gaussian distribution

Usage

```
qqmat(x, output = FALSE, square = FALSE)
```

Arguments

x sample data: matrix or vector

output logical: if TRUE results are returned

square plot in a square window - outliers might be cut off.

Value

if output=TRUE, the following values are returned

x distances from an expected Gaussian distribution

y observed distances - sorted
d observed distances - unsorted

Author(s)

Stefan Schlager

See Also

qqplot

```
require(MASS)
### create normally distributed data
data <- mvrnorm(100,mu=rep(0,5),Sigma = diag(5:1))
qqmat(data)

###create non normally distributed data
data1 <- rchisq(100,df=3)
qqmat(data1,square=FALSE)</pre>
```

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quad2trimesh

converts a mesh containing quadrangular faces into one only consisting of triangles

Description

converts a mesh containing quadrangular faces into one only consisting of triangles

Usage

```
quad2trimesh(mesh, updateNormals = TRUE)
```

Arguments

mesh object of class "mesh3d"

updateNormals logical: request recalculation of (angle weighted) vertex normals.

Value

triangular mesh with updated normals

Examples

```
Sigma <- diag(3:1) #create a 3D-covariance matrix
require(rgl)
quadmesh <- ellipse3d(Sigma)##create quadmesh
trimesh <- quad2trimesh(quadmesh)# convert to trimesh</pre>
```

r2morphoj

Export data to MorphoJ and Morphologika

Description

Export data to MorphoJ and Morphologika

Usage

```
r2morphoj(x, file, id.string = NULL)
r2morphologika(x, file = file, labels = NULL, labelname = NULL, ...)
```

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Arguments

x 3-dimensionla array containing landmark data. E.g. the input/output from procSym.

file character: name the output file

id.string a string with ids or factors to append

labels character vector specify labels to create for Morphologika

character: name the labels for Morphologika.

... unused at the moment

Details

Export data to MorphoJ and Morphologika

Examples

```
if (require(shapes)) {
  r2morphoj(gorf.dat,file="gorf.dat")

data <- bindArr(gorf.dat, gorm.dat, along=3)
  datalabels <- c(rep("female",dim(gorf.dat)[3]),
  rep("male",dim(gorm.dat)[3]))
  labelname <- "sex"
  r2morphologika(data, labels=datalabels, labelname= labelname, file="data.dat")
  ## cleanup
  unlink(c("gorf.dat","data.dat"))
}</pre>
```

ray2mesh

projects the vertices of a mesh along its normals onto the surface of another one.

Description

projects the vertices of a mesh onto the surface of another one by searching for the closest point along vertex normals on the target by for each vertex.

Usage

```
ray2mesh(mesh1, tarmesh, tol = 1e+12, inbound = FALSE, mindist = FALSE, ...)
```

Arguments

mesh1 mesh to project. Can be an object of class "mesh3d" or path to an external mesh

file (ply, obj, stl).

tarmesh mesh to project onto. Can be an object of class "mesh3d" or path to an external

mesh file (ply, obj, stl).

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tol numeric: maximum distance to search along ray, closest Euclidean distance will

be used, if tol is exceeded.

inbound inverse search direction along rays.

mindist search both ways (ray and -ray) and select closest point.

... additional arguments not used at the moment.

Value

returns projected mesh with additional list entries:

quality integer vector containing a value for each vertex of x: 1 indicates that a ray has

intersected 'tarmesh' within the given threshold, while 0 means not

distance numeric vector: distances to intersection

Author(s)

Stefan Schlager

See Also

```
ply2mesh, closemeshKD
```

read.csv.folder

batch import data from files

Description

imports all data files contained in a specified folder.

Usage

```
read.csv.folder(
  folder,
  x,
  y = 2:4,
  rownames = NULL,
  header = TRUE,
  dec = ".",
  sep = ";",
  pattern = "csv",
  addSpec = NULL,
  back = TRUE
)
```

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Arguments

folder character: path to folder

x either a vector specifing which rows are to be imported, or character vector

containing variable names to be sought for.

y a vector specifiing, which columns of the speradsheet ist to be imported.

rownames integer: specifies columns, where variable names are stored.

header logical: if spreadsheet contains header-row.

dec character: defines decimal sepearator.
sep character: defines column seperator.
pattern character: specify file format (e.g. csv).

addSpec character: add a custom specifier to the dimnames of the array.

back logical: where to place the specifier.

Value

arr array containing imported data

NAs vector containing position of observations with NAs

NA.list list: containing vectors containing information which LMs are missing in which

observation

Author(s)

Stefan Schlager

See Also

read.table

read.fcsv	read fiducials from slicer4	
-----------	-----------------------------	--

Description

read fiducials from slicer4

Usage

```
read.fcsv(x, na = NULL, lps2ras = FALSE)
```

Arguments

x filename

na value to be replaced by NA

lps2ras logical: if the coordinate system is LPS and lps2ras=TRUE, the data will be

rotated into the RAS space by inverting the first two dimensions using LPS2RAS.

read.mpp

Value

a k x 3 matrix with landmarks

read dta files	read.lmdta
----------------	------------

Description

reads .dta files created by the software Landmark http://graphics.idav.ucdavis.edu/research/EvoMorph

Usage

```
read.lmdta(file = "x", na = 9999)
```

Arguments

file a dta file

na specifies a value that indicates missing values

Value

arr array containing landmarks dimnames will be Information of ID and landmark

names specified in Landmark

info Information extracted from the header of the dta file

idnames character vector containing the names of the individuals as specified in the dta

file

read.mpp	Read saved pick-points from meshlab
----------	-------------------------------------

Description

imports pick points selected with meshlab

Usage

```
read.mpp(file, info = FALSE)
```

Arguments

file file to import

info logical: if TRUE, additional infos are returned

read.pts 131

Value

if info=FALSE:

a matrix containing picked-points coordinates (only those tagged as active).

if info=TRUE: a list containing

data matrix containing coordinates - including points tagged as inactive

info additional info contained in file.

Author(s)

Stefan Schlager

See Also

read.pts

read.pts

reads pts files

Description

reads Landmark data exported from the software Landmark from http://graphics.idav.ucdavis.edu/research/EvoMorph

Usage

```
read.pts(file = "x", na = 9999)
```

Arguments

file pts file

na specifies a value that indicates missing values

Value

matrix matrix containing landmark information rownames will be the names given to

the landmarks in Landmark

See Also

read.pts

```
data(nose)
write.pts(shortnose.lm, filename="shortnose")
data <- read.pts("shortnose.pts")</pre>
```

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read.slicerjson	read Landmarks from Slicer in Json format
i caa. Siicci joon	read Editaliarks from Succi in 350h format

Description

read Landmarks from Slicer in Json format

Usage

```
read.slicerjson(x, lps2ras = FALSE, na = NULL)
```

Arguments

x path to json file

lps2ras logical: if the coordinate system is LPS and lps2ras=TRUE, the data will be

rotated into the RAS space by inverting the first two dimensions using LPS2RAS.

na value to be replaced by NA

Value

returns matrix or list of matrices with imported landmark coordinates

readallTPS	Import landmarks and outlines from TPS files	
------------	--	--

Description

Imports outlines and landmarks from files generated by tpsdig2

Usage

```
readallTPS(file, scale = TRUE)
```

Arguments

file	A TPS-file generated	by tpsdig2
------	----------------------	------------

scale logical: if TRUE the data will be scaled according to the SCALE entry.

Value

ID	Specimen	IDs read	from	TPS file

LM list of landmarks contained in the TPS-file

outlines a list containing sublists for each specimen with all the outlines read from TPS

file

SCALE vector containing the scale factors for each landmark config.

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Note

currently only landmarks, ID and outlines are read from the TPS-file

Author(s)

Stefan Schlager

References

http://life.bio.sunysb.edu/ee/rohlf/software.html

See Also

```
read.lmdta, read.pts
```

readLandmarks.csv

import landmark data from csv files

Description

import landmark data from csv files

Usage

```
readLandmarks.csv(
  file,
  x,
  y = 2:4,
  rownames = NULL,
  header = TRUE,
  dec = ".",
  sep = ";"
)
```

Arguments

file	character: path to file containing landmark data.
X	either a vector specifiing which rows are to be imported, or character vector containing variable names to be sought for.
У	a vector specifiing, which columns of the speradsheet ist to be imported.
rownames	integer: specifies columns, where variable names are stored.
header	logical: if spreadsheet contains header-row.
dec	character: defines decimal sepearator.
sep	character: defines column seperator.

regdist regdist

Value

LM matrix containing imported data

NAs vector containing rows containing NAs

Author(s)

Stefan Schlager

See Also

```
read.table
```

regdist

correlation between shape space and tangent space

Description

performs a partial Procrustes superimposition of landmark data and calculates the correlation between tangent and shape space.

Usage

```
regdist(
  dataarray,
  plot = TRUE,
  main = "",
  rho = "angle",
  dist.mat.out = FALSE,
  ...
)
```

Arguments

dataarray Input k x m x n real array, where k is the number of points, m is the number of

dimensions, and n is the sample size.

plot Logical: whether to plot the distances between observations.

main character string: Title of the plot.

rho chose how to calculate distances in shape space. Options: "riemdist"=Riemannian

distance (function from the shapes package-takes along time to calculate), "angle"=calculates the angle between shape vectors, "sindist"=sinus of length of

residual vector between shape vectors.

dist.mat.out Logical: If TRUE, output will contain distance matrices.

... additional parameters passed to procSym

RegScore 135

Value

cor	correlation coefficient between distances in shape space and tangent space
procSS	Procrustes Sums of Squares (of full procrustes distance)
tanSS	Tangent Sums of Squares
rhoSS	Procrustes Sums of Squares (of angle)
euc.dist	distance matrix of euclidean distance in Tangent space
proc.dist	distance matrix of Procrustes distance in Shape space

1m linear model regressing tangent space distances onto Procrustes distances

Author(s)

Stefan Schlager

See Also

```
regdist
```

Examples

```
if (require(shapes)) {
regdist(gorf.dat)
}
```

RegScore

calulate regression scores for linear model

Description

calulate regression scores for linear model as specified in Drake & Klingenberg(2008)

Usage

```
RegScore(model, x = NULL)
```

Arguments

model linear model

x optional: matrix containing fitted data to be projected onto the regression lines.

If omitted the model's fitted values will be used.

Details

the data are orthogonally projected onto the regression lines associated with each factor.

Value

returns a n x m matrix containing the regression scores for each specimen.

Warning

if model contains factors with more than 2 levels, R calculates one regression line per 2 factors. Check the colnames of the returned matrix to select the appropriate one. See examples for details.

References

Drake, AG. & Klingenberg, CP. The pace of morphological change: historical transformation of skull shape in St Bernard dogs. Proceedings of the Royal Society B: Biological Sciences, The Royal Society, 2008, 275, 71-76.

Examples

```
model <- lm(as.matrix(iris[,1:3]) ~ iris[,4])</pre>
rs <- RegScore(model)</pre>
plot(rs,iris[,4])
##now use a random subsample for model fitting
rand <- sample(nrow(iris))</pre>
x <- iris[rand[1:100],4]
newmod <- lm(as.matrix(iris[rand[1:100],1:3]) \sim x)
##predict the rest of data and get their regression scores
rsPred <- RegScore(newmod,as.matrix(iris[rand[101:150],1:3]))</pre>
plot(rsPred, iris[rand[101:150], 4])
## Not run:
data(boneData)
proc <- procSym(boneLM)</pre>
pop.sex <- name2factor(boneLM,which=3:4) # generate a factor with 4 levels</pre>
lm.ps.size <- lm(proc$PCscores ~ pop.sex+proc$size)</pre>
rs <- RegScore(lm.ps.size)</pre>
colnames(rs) # in this case, the last column contains the regression
# scores associated with proc$size
## validate by using a subsample for fitting
rand <- sample(dim(boneLM)[3])</pre>
lm.ps.size0 <- lm(proc$PCscores[rand[1:50],] ~ proc$size[rand[1:50]])</pre>
rs0 <- RegScore(lm.ps.size0,proc$PCscores[rand[-c(1:50)],])
plot(rs0,proc$size[rand[-c(1:50)]])
## End(Not run)
```

relaxLM

relax one specific landmark configuration against a reference

Description

relax one specific landmark configuration against a reference (e.g. a sample mean)

Usage

```
relaxLM(lm, ...)
## S3 method for class 'matrix'
relaxLM(
  lm,
 reference,
 SMvector,
 outlines = NULL,
  surp = NULL,
  sur.name = NULL,
 mesh = NULL,
  tol = 1e-05,
 deselect = FALSE,
  inc.check = TRUE,
  iterations = 0,
  fixRepro = TRUE,
 missing = NULL,
 bending = TRUE,
  stepsize = ifelse(bending, 1, 0.5),
 use.lm = NULL,
 silent = FALSE,
)
## S3 method for class 'mesh3d'
relaxLM(
 lm,
 reference,
  tol = 1e-05,
 deselect = FALSE,
  inc.check = TRUE,
  iterations = 0,
  fixRepro = TRUE,
 missing = NULL,
 bending = FALSE,
  stepsize = ifelse(bending, 1, 0.5),
 use.lm = NULL,
 silent = FALSE,
)
```

Arguments

```
    lm k x 3 or k x 2 matrix containing landmark data to be slidden - or a triangular mesh of class "mesh3d". See details
    ... additional arguments - currently unused
```

reference	$k \times 3$ or $k \times 2$ matrix containing landmark of the reference, or a mesh with the same amount of vertices as there are landmarks in $1m$.
SMvector	A vector containing the row indices of (semi-) landmarks on the curve(s) that are allowed to slide
outlines	A vector (or if threre are several curves) a list of vectors (containing the rowindices) of the (Semi-)landmarks forming the curve(s) in the successive position on the curve - including the beginning and end points, that are not allowed to slide.
surp	integer vector containing the row indices of semi-landmarks positioned on surfaces.
sur.name	character: containing the filename of the corresponding surface. When specified, mesh has to be NULL. If sur.name=NULL and mesh=NULL, the tangent planes will be estimated from the point cloud.
mesh	triangular mesh of class "mesh3d" loaded into the R workspace, when specified, "sur.name" has to be NULL.
tol	numeric: Threshold for convergence in the sliding proces. Full Procrustes distance between actual result and previous iteration.
deselect	Logical: if TRUE, the SMvector is interpreted as those landmarks, that are not allowed to slide.
inc.check	Logical: if TRUE, the program stops when convergence criterion starts increasing and reports result from last iteration.
iterations	integer: maximum amounts the algorithm runs - even when 'tol' is not reached. When iterations=0, the algorithm runs until convergence.
fixRepro	logical: if TRUE, fix landmarks will also be projected onto the surface. If you have landmarks not on the surface, select fixRepro=FALSE
missing	vector of integers, specifying row indices of missing (semi-)landmarks. They will be relaxed freely in 3D and not projected onto the target (works only for 2D data).
bending	if TRUE, bending energy will be minimized, Procrustes distance otherwise (not suggested with large shape differences)
stepsize	integer: dampening factor for the amount of sliding. Useful to keep semilandmarks from sliding too far off the surface. The displacement is calculated as $\Upsilon = \Upsilon^0 + stepsize * UT$. Default is set to 1 for bending=TRUE and 0.5 for bending=FALSE.
use.lm	indices specifying a subset of (semi-)landmarks to be used in the rotation step only used if bending=FALSE.
silent	logical: if TRUE, console output is suppressed.

Details

if 1m is a surface mesh, all vertices will be treated as semilandmarks and a allowed to freely slide along the surface.

Value

returns kx3 matrix of slidden landmarks

Author(s)

Stefan Schlager

References

Gunz, P., P. Mitteroecker, and F. L. Bookstein. 2005. Semilandmarks in Three Dimensions, in Modern Morphometrics in Physical Anthropology. Edited by D. E. Slice, pp. 73-98. New York: Kluwer Academic/Plenum Publishers.

See Also

slider3d

```
require(rgl)
data(nose)
### relax shornose against longnose
# define fix landmarks
fix <- c(1:5,20:21)
# define surface patch by specifying row indices of matrices
# all except those defined as fix
surp <- c(1:dim(shortnose.lm)[1])[-fix]</pre>
relax <- relaxLM(shortnose.lm,</pre>
         longnose.lm, mesh=shortnose.mesh, iterations=1,
         SMvector=fix, deselect=TRUE, surp=surp)
## example minimizing Procrustes distance when displacement is not
## dampened by stepsize
relaxProcD <- relaxLM(shortnose.lm,</pre>
         longnose.lm, mesh=shortnose.mesh, iterations=1,
         SMvector=fix, deselect=TRUE, surp=c(1:623)[-fix],bending=FALSE,stepsize=1)
## Not run:
# visualize differences red=before and green=after sliding
deformGrid3d(shortnose.lm, relax, ngrid=0)
# visualize differences minimizing Procrusted distances red=before and green=after sliding
deformGrid3d(shortnose.lm, relaxProcD, ngrid=0)
## no smooth displacement, now let's check the distances:
rot2ref <- rotonto(relaxProcD,longnose.lm)</pre>
angle.calc(rot2ref$X,rot2ref$Y)
# 0.2492027 Procrustes distance between reference and slided shape
# (minimizing Procrustes distance)
rot2refBend <- rotonto(relax,longnose.lm)</pre>
angle.calc(rot2refBend$X,rot2refBend$Y)
# 0.2861322 Procrustes distance between reference and slided shape
```

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```
# (minimizing bending energy)

rot2refOrig <- rotonto(shortnose.lm,longnose.lm)
angle.calc(rot2refOrig$X,rot2refOrig$Y)
# 0.3014957 Procrustes distance between reference and original shape
##result: while minimizing Procrustes distance, displacement is not
##guaranteed to be smooth

# add surface
wire3d(shortnose.mesh, col="white")

## finally relax two meshes with corresponding vertices:
mediumnose.mesh <- tps3d(shortnose.mesh,shortnose.lm, (shortnose.lm+longnose.lm)/2,threads=1)
## we use Procrustes distance as criterion as bending energy is pretty slow because
## of too many coordinates (more than 3000 is very unreasonable).
relaxMesh <- relaxLM(shortnose.mesh,mediumnose.mesh,iterations=2,bending=FALSE,stepsize=0.05)
## End(Not run)</pre>
```

relWarps

calculate relative Warp analysis

Description

After Procrustes registration the data is scaled by the bending energy or its inverse to emphasize global/local differences when exploring a sample's shape.

Usage

```
relWarps(
  data,
  scale = TRUE,
  CSinit = TRUE,
  alpha = 1,
  tol = 1e-10,
  orp = TRUE,
  pcAlign = TRUE,
  computeBasis = TRUE,
  noalign = FALSE
)
```

Arguments

data Input k x m x n real array, where k is the number of points, m is the number of

dimensions, and n is the sample size.

scale Logical: indicating if scaling is requested

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CSinit Logical: if TRUE, all configurations are initially scaled to Unit Centroid Size.

alpha integer: power of the bending energy matrix. If alpha = 0 then standard Procrustes PCA is carried out. If alpha = 1 then large scale differences are empha-

sized, if alpha = -1 then small scale variations are emphasised.

tol tolerance for the eigenvalues of the bending energy matrix to be zero

orp logical: request orthogonal projection into tangent space.

pcAlign logical: if TRUE, the shapes are aligned by the principal axis of the first speci-

men

computeBasis logical: whether to compute the basis of the resulting vector space (takes a lot

of memory and time for configurations with > 1000 coordinates.

noalign logical: if TRUE, data is assumed to be already aligned and alignment and or-

thogonal projection are skipped.

Value

bescores relative warp scores (PC-scores if alpha = 0)

uniscores uniform scores, NULL if alpha = 0

Var non-affine variation explained by each relative warp

mshape sample's conensus shape rotated Procrustes superimposed data

bePCs vector basis of nonaffine shape variation- relative warps (plain PCs if alpha =

0)

uniPCs vector basis of affine shape variation - uniform component. NULL if alpha = 0

Author(s)

Stefan Schlager

References

Bookstein FL 1989. Principal Warps: Thin-plate splines and the decomposition of deformations. IEEE Transactions on pattern analysis and machine intelligence 11.

Bookstein FL, 1991. Morphometric tools for landmark data. Geometry and biology. Cambridge Univ. Press, Cambridge.

Rohlf FJ, Bookstein FL 2003. Computing the Uniform Component of Shape Variation. Systematic Biology 52:66-69.

```
data(boneData)
pop <- name2factor(boneLM,which=3)
rW <- relWarps(boneLM, alpha = -1)
## Not run:
if (require(car)) {
# plot first 5 relative warps scores grouped by population</pre>
```

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```
spm(rW$bescores[,1:5],group=pop)
# plot uniform component scores grouped by population
spm(rW$uniscores[,1:5],group=pop)
##plot non-affine variance associated with each relative warp
barplot(rW$Var[,2], xlab="relative Warps")
## visualize first relative warp +-3 sd of the scores
rw1 <- restoreShapes(as.matrix(c(-3,3)*sd(rW$bescores[,1])),rW$bePCs[,1,drop=FALSE],rW$mshape)
deformGrid3d(rw1[,,1],rw1[,,2],ngrid=5)
## 2D example:
if (require(shapes)) {
data <- bindArr(gorf.dat, gorm.dat, along=3)</pre>
sex <- factor(c(rep("fem", dim(gorf.dat)[3]), rep("male",dim(gorm.dat)[3])))</pre>
rW <- relWarps(data, alpha = -1)
if (require(car)) {
# plot first 3 relative warps scores grouped by population
spm(rW$bescores[,1:3],group=sex)
# plot uniform component scores grouped by population
spm(rW$uniscores[,1:2],group=sex)
}
##plot non-affine variance associated with each relative warp
barplot(rW$Var[,2], xlab="relative Warps")
## visualize first relative warp +-3 sd of the scores
 rw1 \leftarrow restoreShapes(as.matrix(c(-3,3)*sd(rW\$bescores[,1])), rW\$bePCs[,1,drop=FALSE], rW\$mshape) 
deformGrid2d(rw1[,,1],rw1[,,2],ngrid=10)
## End(Not run)
```

render

plot or save the results of meshDist

Description

plot or save the results of meshDist

Usage

```
render(x, ...)
## S3 method for class 'meshDist'
render(
    x,
    from = NULL,
    to = NULL,
    steps = NULL,
    ceiling = NULL,
    uprange = NULL,
```

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```
tol = NULL,
  tolcol = NULL,
  rampcolors = NULL,
 NAcol = NULL,
  displace = FALSE,
  shade = TRUE,
  sign = NULL,
  add = FALSE,
  scaleramp = NULL,
  titleplot = "Distance in mm",
)
## S3 method for class 'matrixDist'
render(
  х,
  from = NULL,
  to = NULL,
  steps = NULL,
  ceiling = NULL,
  uprange = NULL,
  tol = NULL,
  tolcol = NULL,
  type = c("s", "p"),
  radius = NULL,
  rampcolors = NULL,
 NAcol = NULL,
  displace = FALSE,
  sign = NULL,
  add = FALSE,
  scaleramp = FALSE,
  titleplot = "Distance in mm",
)
export(x, ...)
## S3 method for class 'meshDist'
export(
 Х,
  file = "default",
  imagedim = "100x800",
  titleplot = "Distance in mm",
)
```

Arguments

x object of class meshDist

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... for render.meshDist: additional arguments passed to shade3d. See material3d

for details.

from numeric: minimum distance to color; default is set to 0 mm

to numeric: maximum distance to color; default is set to the maximum distance

steps integer: determines how many intermediate colors the color ramp has.

ceiling logical: if TRUE, the next larger integer of "to" is used

uprange numeric between 0 and 1: restricts "to" to a quantile of "to", if to is NULL.

tol numeric: threshold to color distances within this threshold according to tolcol.

tolcol a custom color to color vertices below a threshold defined by tol. Default is

green.

rampcolors character vector: specify the colors which are used to create a colorramp.

NAcol character: specify color for values outside the range defined by from and to.

displace logical: if TRUE, displacement vectors between original and closest points are

drawn colored according to the distance.

shade logical: if FALSE, the rendering of the colored surface will be supressed.

sign logical: request signed distances to be visualised.

add logical: if TRUE, visualization will be added to the rgl window currently in

focus

scaleramp if TRUE the ramp colors get scaled symmetrically into positive and negative

direction.

titleplot character: axis description of heatmap.

type character: "s" shows coordinates as spheres, while "p" shows 3D dots.

radius determines size of spheres; if not specified, optimal radius size will be estimated

by centroid size of the configuration.

file character: filename for mesh and image files produced. E.g. "mydist" will

produce the files mydist.ply and mydist.png

imagedim character of pattern "100x200" where 100 determines the width and 200 the

height of the image.

Details

Visualise or save the results of meshDist to disk.

render.meshDist renders the colored mesh and displays the color ramp and returns an object of class "meshDist". export.meshDist exports the colored mesh as ply file and the color chart as png file.

Author(s)

Stefan Schlager

See Also

meshDist, shade3d

resampleCurve 145

resampleCurve	Resample a curve equidistantly	

Description

Resample a curve equidistantly (optionally with smoothing)

Usage

```
resampleCurve(x, n, smooth = FALSE, smoothn = n, open = TRUE)
```

Arguments

x matrix containing coordinates

n number of resulting points on the resampled curve

smooth logical: if TRUE, the resulting curve will be smoothed by using bezier curves.

smoothn integer: define the refinement of the bezier curve. The higher this value, the

closer the final curve will be to the original.

open logical: define whether it is a closed curve or not.

Value

returns a matrix containing the resampled curve

Examples

```
data(nose)
x <- shortnose.lm[c(304:323),]
xsample <- resampleCurve(x,n=50)</pre>
```

restoreFromPCA

restore original data from PCA

Description

restore original data from PCA by reverting rotation and centering

Usage

```
restoreFromPCA(scores, rotation, center)
```

Arguments

scores matrix containing the PC-scores rotation matrix containing the PCs center vector containing the center

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Examples

```
myirispca <- prcomp(iris[,1:4])
myirisRecovered <- restoreFromPCA(myirispca$x,myirispca$rotation,myirispca$center)
all.equal(myirisRecovered,as.matrix(iris[,1:4]))</pre>
```

restoreShapes

restore shapes from PC-Scores or similar projections

Description

restore shapes from PC-Scores or similar projections

Usage

```
restoreShapes(
   scores,
   PC,
   mshape,
   sizeshape = FALSE,
   origsize = FALSE,
   meanlogCS
)
```

Arguments

scores	vector of PC-scores, or matrix with rows containing PC-scores
PC	Principal components (eigenvectors of the covariance matrix) associated with 'scores'.
mshape	matrix containing the meanshape's landmarks (used to center the data by the PCA)
sizeshape	logical: if TRUE, it is assumed that the data is the output of procSym run with sizeshape=TRUE.
origsize	logical: if sizeshape = TRUE, this will apply the scaling to the original size from the corresponding entry from the PC basis matrix.
meanlogCS	numeric: provide the average log Centroid Size of the original sample (see examples below). Only needed if sizeshape = TRUE and origsize = TRUE

Details

Rotates and translates PC-scores (or similar) derived from shape data back into configuration space.

Value

returns matrix or array containing landmarks

retroDeform3d 147

Author(s)

Stefan Schlager

See Also

```
prcomp, procSym
getPCscores
```

Examples

retroDeform3d

symmetrize a bilateral landmark configuration

Description

symmetrize a bilateral landmark configuration by removing bending and stretching

Usage

```
retroDeform3d(mat, pairedLM, hmult = 5, alpha = 0.01)
```

Arguments

mat	matrix with bilateral landmarks
pairedLM	2-column integer matrix with the 1st columns containing row indices of left side landmarks and 2nd column the right hand landmarks
hmult	factor controlling the bandwidth for calculating local weights (which will be hmult * average distance between landmarks and their closest neighbour).
alpha	factor controlling spacing along x-axis

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Value

deformed matrix containing deformed landmarks
orig matrix containing original landmarks

References

Ghosh, D.; Amenta, N. & Kazhdan, M. Closed-form Blending of Local Symmetries. Computer Graphics Forum, Wiley-Blackwell, 2010, 29, 1681-1688

retroDeformMesh

symmetrize a triangular mesh

Description

symmetrize a triangular mesh

Usage

```
retroDeformMesh(
  mesh,
  mat,
  pairedLM,
  hmult = 5,
  alpha = 0.01,
  rot = TRUE,
  lambda = 1e-08,
  threads = 0
)
```

Arguments

mesh triangular mesh of class mesh3d mat matrix with bilateral landmarks

pairedLM 2-column integer matrix with the 1st columns containing row indices of left side

landmarks and 2nd column the right hand landmarks

hmult damping factor for calculating local weights which is calculated as humult times

the average squared distance between a landmark and its closest neighbor (on

each side).

alpha factor controlling spacing along x-axis

rot logical: if TRUE the deformed landmarks are rotated back onto the original ones

lambda control parameter passed to tps3d

threads integer: number of threads to use for TPS deform

rotaxis3d 149

Details

this function performs retroDeform3d and deforms the mesh accordingly using the function tps3d.

Value

mesh symmetrized mesh

landmarks a list containing the deformed and original landmarks

rotaxis3d

Rotate an object (matrix or mesh) around an arbitrary axis in 3D

Description

Rotate an object around an arbitrary axis in 3D

Usage

```
rotaxis3d(x, pt1, pt2 = c(0, 0, 0), theta)

## S3 method for class 'matrix'
rotaxis3d(x, pt1, pt2 = c(0, 0, 0), theta)

## S3 method for class 'mesh3d'
rotaxis3d(x, pt1, pt2 = c(0, 0, 0), theta)
```

Arguments

X	k x 3 matrix containing 3D-coordinates or a triangular mesh of class "mesh3d".
pt1	numeric vector of length 3, defining first point on the rotation axis.
pt2	numeric vector of length 3, defining second point on the rotation axis.
theta	angle to rotate in radians. With pt1 being the viewpoint, the rotation is counterclockwise.

Details

Rotate an object (matrix or triangular mesh) around an 3D-axis defined by two points.

Value

returns rotated object (including updated normals for mesh3d objects)

Author(s)

Stefan Schlager

150 rotaxisMat

References

http://en.wikipedia.org/wiki/Rotation_matrix

See Also

```
rotonto, rotmesh.onto
```

Examples

```
require(rg1)
data(nose)
shrot.rot <- rotaxis3d(shortnose.mesh,pt1=c(1,1,1),theta=pi)
## Not run:
shade3d(shortnose.mesh,col=3,specular=1)
shade3d(shrot.rot,col=2)
###print rotation axis
#' lines3d(rbind(rep(-0.1,3),rep(0.1,3)))
## End(Not run)</pre>
```

rotaxisMat

calculate a rotation matrix around an arbitrary axis through the origin in 3D

Description

calculate a rotation matrix around an arbitrary axis in 3D

Usage

```
rotaxisMat(u, theta, homogeneous = FALSE)
```

Arguments

u a vector around which to rotate

theta angle in radians to rotate

homogeneous logical: if TRUE a 4x4 rotation matrix is returned

Value

returns 3x3 rotation matrix

References

http://en.wikipedia.org/wiki/Rotation_matrix

rotmesh.onto 151

See Also

rotaxis3d

rotmesh.onto

rotate, scale and translate a mesh based on landmark information.

Description

rotates and reflects a mesh onto by calculating the transformation from two sets of referenced landmarks.

Usage

```
rotmesh.onto(
  mesh,
  refmat,
  tarmat,
  adnormals = FALSE,
  scale = FALSE,
  reflection = FALSE,
  ...
)
```

Arguments

mesh object of class mesh3d.

refmat k x m matrix with landmarks on the mesh

tarmat k x m matrix as target configuration

adnormals logical - if TRUE, vertex normals will be recomputed after rotation. If mesh has

normals and adnormals=FALSE, the existing normals are rotated by the same

rotation matrix as the mesh's vertices.

scale logical: if TRUE the mesh will be scaled according to the size of the target.

reflection logical: allow reflection.

... additional parameters passed on to rotonto.

Value

mesh rotated mesh yrot rotated refmat

trafo 4x4 transformation matrix

Author(s)

Stefan Schlager

152 rotonmat

See Also

file2mesh,tps3d ,rotonto,mesh2ply

Examples

rotonmat

rotate matrix of landmarks

Description

rotate matrix of landmarks by using a rotation determined by two matrices.

Usage

```
rotonmat(
   X,
   refmat,
   tarmat,
   scale = TRUE,
   reflection = FALSE,
   weights = NULL,
   centerweight = FALSE,
   getTrafo = FALSE
)
```

Arguments

X Matrix to be rotated

refmat reference matrix used to estimate rotation parameters tarmat target matrix used to estimate rotation parameters

rotonmat 153

scale logical: requests scaling to minimize sums of squared distances

reflection logical: if TRUE, reflections are allowed.

weights vector of length k, containing weights for each landmark.

centerweight logical: if weights are defined and centerweigths=TRUE, the matrix will be

centered according to these weights instead of the barycenter.

getTrafo logical: if TRUE, a 4x4 transformation matrix will also be returned.

Details

A matrix is rotated by rotation parameters determined by two different matrices. This is usefull, if the rotation parameters are to be estimated by a subset of landmark coordinates.

Value

if getTrafo=FALSE the transformed X will be returned, else alist containing:

Xrot the transformed matrix X trafo a 4x4 transformation matrix

Author(s)

Stefan Schlager

See Also

rotonto, rotmesh. onto

```
data(nose)
shortnose.rot <-
rotonmat(shortnose.lm, shortnose.lm[1:9,],longnose.lm[1:9,])
##view result
## Not run:
deformGrid3d(shortnose.rot, shortnose.lm, ngrid=0)
## End(Not run)</pre>
```

154 rotonto

rotonto

rotates, translates and scales one matrix onto an other using Procrustes fitting

Description

rotates, translates and scales one matrix onto an other using Procrustes fitting

Usage

```
rotonto(
    x,
    y,
    scale = FALSE,
    signref = TRUE,
    reflection = TRUE,
    weights = NULL,
    centerweight = FALSE,
    ...
)

rotreverse(mat, rot)

## S3 method for class 'matrix'
rotreverse(mat, rot)

## S3 method for class 'mesh3d'
rotreverse(mat, rot)
```

Arguments x

rot

	, ,
У	k x m matrix which will be rotated (reference matrix)
scale	logical: scale matrix to minimize sums of squares
signref	logical: report if reflections were involved in the rotation
reflection	allow reflections.
weights	vector of length k, containing weights for each landmark.
centerweight	logical or vector of weights: if weights are defined and centerweights=TRUE, the matrix will be centered according to these weights instead of the barycenter. If centerweight is a vector of length nrow(x), the barycenter will be weighted accordingly.
	currently not used
mat	matrix on which the reverse transformations have to be applied

an object resulting from the former application of rotonto

k x m matrix to be rotated onto (targetmatrix)

rotonto 155

Details

rotate a matrix onto an other without loosing information about the location of the targetmatrix and reverse this transformations using rotreverse

Value

yrot rotated and translated matrix

Y centred and rotated reference matrix

X centred target matrix

trans vector between original position of target and centered reference (during rotation

process)

transy vector between original position of reference and centered reference (during

rotation process)

gamm rotation matrix

bet scaling factor applied

reflect if reflect = 1, reflections are involved in the superimposition. Else, reflect = 0

Note

all lines containing NA, or NaN are ignored in computing the transformation.

Author(s)

Stefan Schlager

References

Lissitz, R. W., Schoenemann, P. H., & Lingoes, J. C. (1976). A solution to the weighted Procrustes problem in which the transformation is in agreement with the loss function. Psychometrika, 41,547-550.

See Also

rotmesh.onto

```
if (require(shapes)) {
  lims <- c(min(gorf.dat[,,1:2]),max(gorf.dat[,,1:2]))
  rot <- rotonto(gorf.dat[,,1],gorf.dat[,,2]) ### rotate the second onto the first config
  plot(rot$yrot,pch=19,xlim=lims,ylim=lims) ## view result
  points(gorf.dat [,,2],pch=19,col=2) ## view original config
  rev1 <- rotreverse(rot$yrot,rot)
  points(rev1,cex=2) ### show inversion by larger circles around original configuration
}</pre>
```

156 scalemesh

scalemesh

scale a mesh of class "mesh3d"

Description

scales (the vertices of a mesh by a scalar

Usage

```
scalemesh(mesh, size, center = c("bbox", "mean", "none"))
```

Arguments

mesh object of class "mesh3d" size numeric: scale factor

center character: method to position center of mesh after scaling: values are "bbox",

and "mean". See Details for more info.

Details

The mesh's center is determined either as mean of the bounding box (center="bbox") or mean of vertex coordinates (center="mean") and then scaled according to the scaling factor. If center="none", vertex coordinates will simply be multiplied by "size".

Value

returns a scaled mesh

Author(s)

Stefan Schlager

See Also

```
rotmesh.onto
```

```
data(nose)
#inflate mesh by factor 4
largenose <- scalemesh(shortnose.mesh,4)</pre>
```

slider2d 157

slider2d	slides Semilandmarks along curves 2D by minimising bending energy of a thin-plate spline deformation.

Description

slides Semilandmarks along curves 2D. The positions are sought by minimising bending energy (of a thin-plate spline deformation) or Procrustes distance

Usage

```
slider2d(
  dataframe,
  SMvector,
  outlines,
  tol = 1e-05,
  deselect = FALSE,
  recursive = TRUE,
  iterations = 0,
  initproc = FALSE,
  pairedLM = NULL,
  bending = TRUE,
  stepsize = 1,
  silent = FALSE
)
```

Arguments

dataframe	Input k x 2 x n real array, where k is the number of points and n is the sample size. Ideally the
SMvector	A vector containing the row indices of (semi-) landmarks on the curve(s) and surfaces that are allowed to slide
outlines	A vector (or if threre are several curves) a list of vectors (containing the rowindices) of the (Semi-)landmarks forming the curve(s) in the successive position on the curve - including the beginning and end points, that are not allowed to slide.
tol	numeric: Threshold for convergence in the sliding process
deselect	Logical: if TRUE, the SMvector is interpreted as those landmarks, that are not allowed to slide.
recursive	Logical: if TRUE, during the iterations of the sliding process, the outcome of the previous iteration will be used. Otherwise the original configuration will be used in all iterations.
iterations	integer: select manually the max. number of iterations that will be performed during the sliding process (usefull, when there is very slow convergence). 0 means iteration until convergence.
initproc	requests initial Procrustes fit before sliding.

pairedLM A X x 2 numeric matrix with the indices of the rows containing paired Landmarks. E.g. the left column contains the lefthand landmarks, while the right side contains the corresponding right hand landmarks. - This will ideally create

symmetric mean to get rid of assymetry.

bending if TRUE, bending energy will be minimized, Procrustes distance otherwise.

stepsize integer: dampening factor for the amount of sliding. Useful to keep semi-

landmarks from sliding too far off the surface. The displacement is calculated as $\Upsilon = \Upsilon^0 + stepsize * UT$. Default is set to 1 for bending=TRUE and 0.5 for

bending=FALSE.

silent logical: if TRUE, console output is suppressed.

Value

returns an array containing slided coorndinates in the original space - not yet processed by a Procrustes analysis.

Warning

Depending on the amount of landmarks this can use an extensive amount of your PC's resources, especially when running in parallel. As the computation time and RAM usage of matrix algebra involved is quadratic to the amount of landmarks used, doubling the amount of semi-landmarks will quadruple computation time and system resource usage. You can easily stall you computer with this function with inappropriate data.

Author(s)

Stefan Schlager

See Also

relaxLM, slider3d

slider3d	slides Semilandmarks along curves and surfaces in 3D by minimising
	bending energy of a thin-plate spline deformation.

Description

slides Semilandmarks along curves and surfaces in 3D. The positions on the surface are sought which minimise bending energy (of a thin-plate spline deformation)

Usage

```
slider3d(
  dat.array,
  SMvector,
 outlines = NULL,
  surp = NULL,
  sur.path = NULL,
  sur.name = NULL,
 meshlist = NULL,
  ignore = NULL,
  sur.type = "ply",
  tol = 1e-05,
  deselect = FALSE,
  inc.check = TRUE,
  recursive = TRUE,
  iterations = 0,
  initproc = TRUE,
  fullGPA = FALSE,
  pairedLM = 0,
  bending = TRUE,
  stepsize = ifelse(bending, 1, 0.5),
 mc.cores = parallel::detectCores(),
  fixRepro = TRUE,
 missingList = NULL,
 use.lm = NULL,
  smoothnormals = FALSE,
  silent = FALSE
)
```

Arguments

dat.array	Input k x m x n real array, where k is the number of points, m is the number of dimensions, and n is the sample size. Ideally the dimnames[[3]] vector contains the names of the surface model (without file extension) - e.g. if the model is named "surface.ply", the name of the corresponding matrix of the array would be "surface"
SMvector	A vector containing the row indices of (semi-) landmarks on the curve(s) and surfaces that are allowed to slide
outlines	A vector (or if threre are several curves) a list of vectors (containing the rowindices) of the (Semi-)landmarks forming the curve(s) in the successive position on the curve - including the beginning and end points, that are not allowed to slide.
surp	integer vector containing the row indices of semi-landmarks positioned on surfaces.
sur.path	Path to the surface models (e.g. ply, obj, stl files)
sur.name	character vector: containing the filenames of the corresponding surfaces - e.g. if the dat.array[,,i] belongs to surface_i.ply, sur.name[i] would be surface_i.ply.

Only necessary if dat.array does not contain surface names.

meshlist	list containing triangular meshes of class 'mesh3d', for example imported with mesh2ply or file2mesh in the same order as the specimen in the array (see examples below).
ignore	vector containing indices of landmarks that are to be ignored. Indices of outlines/surfaces etc will be updated automatically.
sur.type	character:if all surfaces are of the same file format and the names stored in dat.array, the file format will be specified here.
tol	numeric: Threshold for convergence in the sliding process
deselect	Logical: if TRUE, the SMvector is interpreted as those landmarks, that are not allowed to slide.
inc.check	Logical: if TRUE, the program stops when convergence criterion starts increasing and reports result from last iteration.
recursive	Logical: if TRUE, during the iterations of the sliding process, the outcome of the previous iteration will be used. Otherwise the original configuration will be used in all iterations.
iterations	integer: select manually the max. number of iterations that will be performed during the sliding process (usefull, when there is very slow convergence). 0 means iteration until convergence.
initproc	requests initial Procrustes fit before sliding.
fullGPA	Logical: if FALSE, only a partial procrustes fit will be performed.
pairedLM	A X x 2 numeric matrix with the indices of the rows containing paired Landmarks. E.g. the left column contains the lefthand landmarks, while the right side contains the corresponding right hand landmarks This will ideally create symmetric mean to get rid of assymetry.
bending	if TRUE, bending energy will be minimized, Procrustes distance otherwise.
stepsize	integer: dampening factor for the amount of sliding. Useful to keep semilandmarks from sliding too far off the surface. The displacement is calculated as $\Upsilon = \Upsilon^0 + stepsize * UT$. Default is set to 1 for bending=TRUE and 0.5 for bending=FALSE.
mc.cores	integer: determines how many cores to use for the computation. The default is autodetect. But in case, it doesn't work as expected cores can be set manually.
fixRepro	logical: if TRUE, fix landmarks will also be projected onto the surface. If you have landmarks not on the surface, select fixRepro=FALSE
missingList	a list of length samplesize containing integer vectors of row indices specifying missing landmars for each specimen. For specimens without missing landmarks enter $numeric(\emptyset)$.
use.lm	indices specifying a subset of (semi-)landmarks to be used in the rotation step-only used if bending=FALSE.
smoothnormals	logical: if TRUE, tangent planes will be computed from locally smoothed normals
silent	logical: if TRUE, console output is suppressed.

Value

dataslide array containing slidden Landmarks in the original space - not yet processed by

a Procrustes analysis

vn.array array containing landmark normals

Warning

Depending on the size of the suface meshes and especially the amount of landmarks this can use an extensive amount of your PC's resources, especially when running in parallel. As the computation time and RAM usage of matrix algebra involved is quadratic to the amount of landmarks used, doubling the amount of semi-landmarks will quadruple computation time and system resource usage. You can easily stall you computer with this function with inappropriate data.

Note

if sur.path = NULL and meshlist = NULL, surface landmarks are relaxed based on a surface normals approximated by the pointcloud, this can lead to bad results for sparse sets of semilandmarks. Obviously, no projection onto the surfaces will be occur and landmarks will likely be off the original surface.

Author(s)

Stefan Schlager

References

Klingenberg CP, Barluenga M, and Meyer A. 2002. Shape analysis of symmetric structures: quantifying variation among individuals and asymmetry. Evolution 56(10):1909-1920.

Gunz, P., P. Mitteroecker, and F. L. Bookstein. 2005. Semilandmarks in Three Dimensions, in Modern Morphometrics in Physical Anthropology. Edited by D. E. Slice, pp. 73-98. New York: Kluwer Academic/Plenum Publishers.

Schlager S. 2012. Sliding semi-landmarks on symmetric structures in three dimensions. American Journal of Physical Anthropology, 147(S52):261. URL: http://dx.doi.org/10.1002/ajpa.21502.

Schlager S. 2013. Soft-tissue reconstruction of the human nose: population differences and sexual dimorphism. PhD thesis, Universitätsbibliothek Freiburg. URL: http://www.freidok.uni-freiburg.de/volltexte/9181/.

See Also

```
relaxLM, createMissingList
```

```
## Not run:
data(nose)
###create mesh for longnose
longnose.mesh <- tps3d(shortnose.mesh,shortnose.lm,longnose.lm,threads=1)
### write meshes to disk</pre>
```

```
mesh2ply(shortnose.mesh, filename="shortnose")
mesh2ply(longnose.mesh, filename="longnose")
## create landmark array
data <- bindArr(shortnose.lm, longnose.lm, along=3)</pre>
dimnames(data)[[3]] <- c("shortnose", "longnose")</pre>
# define fix landmarks
fix <- c(1:5,20:21)
# define surface patch by specifying row indices of matrices
# all except those defined as fix
surp <- c(1:nrow(shortnose.lm))[-fix]</pre>
slide <- slider3d(data, SMvector=fix, deselect=TRUE, surp=surp,</pre>
                  sur.path=".",iterations=1,mc.cores=1)
                  # sur.path="." is the current working directory
# now one example with meshes in workspace
meshlist <- list(shortnose.mesh,longnose.mesh)</pre>
slide <- slider3d(data, SMvector=fix, deselect=TRUE, surp=surp,</pre>
                  iterations=1, meshlist=meshlist,
                  mc.cores=1,fixRepro=FALSE)
require(rgl)
## visualize sliding
deformGrid3d(slide$dataslide[,,1],shortnose.lm,ngrid = 0)
## these are fix
spheres3d(slide$dataslide[fix,,1],col=4,radius=0.7)
###finally an example with missing landmarks:
## we assume that coordinates 185:189, 205:209 and 225:229 are in the second config are missing
missingList <- createMissingList(2)</pre>
missingList[[2]] <- c(185:189,205:209,225:229)
slideMissing <- slider3d(data, SMvector=fix, deselect=TRUE, surp=surp,</pre>
                  iterations=1, meshlist=meshlist,
                  mc.cores=1,fixRepro=FALSE,missingList=missingList)
## example with two curves
## Example with surface semilandmarks and two curves
fix <- c(1:5,20:21)
outline1 <- c(304:323)
outline2 <- c(604:623)
outlines <- list(outline1,outline2)</pre>
surp <- c(1:623)[-c(fix,outline1,outline2)]</pre>
slideWithCurves <- slider3d(data, SMvector=fix, deselect=TRUE, surp=surp,</pre>
                             meshlist=meshlist,iterations=1,mc.cores=1,outlines=outlines)
deformGrid3d(slideWithCurves$dataslide[,,1],shortnose.lm,ngrid = 0)
plot(slideWithCurves)
## finally an example with sliding without meshes by estimating the surface from the
## semi-landmarks
```

solutionSpace 163

Description

returns the solution space (basis and translation vector) for an equation system

Usage

```
solutionSpace(A, b)
```

Arguments

A numeric matrix
b numeric vector

Details

For a linear equation system, Ax = b, the solution space then is

$$x = A^*b + (I - A^*A)y$$

where A^* is the Moore-Penrose pseudoinverse of A. The QR decomposition of $I - A^*A$ determines the dimension of and basis of the solution space.

Value

basis matrix containing the basis of the solution space

translate translation vector

```
A <- matrix(rnorm(21),3,7)
b <- c(1,2,3)
subspace <- solutionSpace(A,b)
dims <- ncol(subspace$basis) # we now have a 4D solution space
## now pick any vector from this space. E.g
y <- 1:dims
solution <- subspace$basis%*%y+subspace$translate # this is one solution for the equation above
A%*%solution ## pretty close
```

164 sortCurve

		\sim		
SO	rt	(11	irve	1

sort curvepoints by using the subsequent neighbours

Description

sort curvepoints by using the subsequent neighbours

Usage

```
sortCurve(x, k = 5, start = NULL)
```

Arguments

x k x m matrix containing the 2D or 3D coordinates

k number of nearest neighbours to look at. Set high for very irregularly clustered

curves.

start integer: which row of x to use as a starting point. If NULL, it is assumed that the

curve is open and the point where the angle between the two nearest neighbours

is closest will be chosen.

Value

xsorted matrix with coordinates sorted along a curve

index vector containing the sorting indices

```
## generate a curve from a polynome
x \leftarrow c(32,64,96,118,126,144,152.5,158)
y < -c(99.5,104.8,108.5,100,86,64,35.3,15)
fit <- lm(y\sim poly(x,2,raw=TRUE))
xx <- seq(30,160, length=50)
layout(matrix(1:3,3,1))
curve <- cbind(xx,predict(fit, data.frame(x=xx)))</pre>
## permute order
set.seed(42)
plot(curve); lines(curve)
curveunsort <- curve[sample(1:50),]</pre>
## now the curve is scrambled
plot(curveunsort);lines(curveunsort,col=2)
curvesort <- sortCurve(curveunsort)</pre>
## after sorting lines are nice again
plot(curvesort$xsorted);lines(curvesort$xsorted,col=3)
```

symmetrize 165

symmetrize

create a perfectly symmetric version of landmarks

Description

create a perfectly symmetric version of landmarks

Usage

```
symmetrize(x, pairedLM)
```

Arguments

k x m matrix or k x m x n array, with rows containing landmark coordinates

pairedLM A X x 2 matrix containing the indices (rownumbers) of the paired LM. E.g. the

left column contains the lefthand landmarks, while the right side contains the

corresponding right hand landmarks.

Details

the landmarks are reflected and relabled according to pairedLM and then rotated and translated onto x. Both configurations are then averaged to obtain a perfectly symmetric one.

Value

a symmetrized version of x

References

Klingenberg CP, Barluenga M, and Meyer A. 2002. Shape analysis of symmetric structures: quantifying variation among individuals and asymmetry. Evolution 56(10):1909-1920.

```
data(boneData)
left <- c(4,6,8)
right <- c(3,5,7)
pairedLM <- cbind(left,right)
symx <- symmetrize(boneLM[,,2],pairedLM)
## Not run:
deformGrid3d(symx,boneLM[,,2])
## End(Not run)</pre>
```

166 tps3d

	tps3d	thin plate spline mapping (2D and 3D) for coordinates and triangular meshes
--	-------	---

Description

maps landmarks or a triangular mesh via thin plate spline based on a reference and a target configuration in 2D and 3D

Usage

```
tps3d(x, refmat, tarmat, lambda = 1e-08, threads = 0, ...) tps2d(x, refmat, tarmat, lambda = 1e-08, threads = 0, ...)
```

Arguments

X	matrix - e.g. the matrix information of vertices of a given surface or a triangular mesh of class "mesh3d"
refmat	reference matrix - e.g. landmark configuration on a surface
tarmat	target matrix - e.g. landmark configuration on a target surface
lambda	numeric: regularisation parameter of the TPS.
threads	threads to be used for parallel execution in tps deformation.
	additional arguments, currently not used.

Value

returns the deformed input

Note

tps2d is simply an alias for tps3d that can handle both cases.

Author(s)

Stefan Schlager

References

Bookstein FL. 1989. Principal Warps: Thin-plate splines and the decomposition of deformations. IEEE Transactions on pattern analysis and machine intelligence 11(6).

See Also

```
computeTransform, applyTransform
```

typprob 167

Examples

```
data(nose)
## define some landmarks
refind <- c(1:3,4,19:20)
## use a subset of shortnose.lm as anchor points for a TPS-deformation
reflm <- shortnose.lm[refind,]</pre>
tarlm <- reflm
##replace the landmark at the tip of the nose with that of longnose.lm
tarlm[4,] <- longnose.lm[4,]</pre>
## deform a set of semilandmarks by applying a TPS-deformation
## based on 5 reference points
deformed <- tps3d(shortnose.lm, reflm, tarlm,threads=1)</pre>
## Not run:
##visualize results by applying a deformation grid
deformGrid3d(shortnose.lm,deformed,ngrid = 5)
data(nose)##load data
##warp a mesh onto another landmark configuration:
longnose.mesh <- tps3d(shortnose.mesh,shortnose.lm,longnose.lm,threads=1)</pre>
require(rgl)
shade3d(longnose.mesh,col=skin1)
## End(Not run)
data(boneData)
## deform mesh belonging to the first specimen
## onto the landmark configuration of the 10th specimen
## Not run:
warpskull <- tps3d(skull_0144_ch_fe.mesh,boneLM[,,1],</pre>
                     boneLM[,,10], threads=1)
## render deformed mesh and landmarks
shade3d(warpskull, col=2, specular=1)
spheres3d(boneLM[,,1])
## render original mesh
shade3d(skull_0144_ch_fe.mesh, col=3, specular=1)
spheres3d(boneLM[,,10])
## End(Not run)
```

typprob

calculate typicality probabilities

Description

calculate typicality probabilities

168 typprob

Usage

```
typprob(
 х,
 data,
  small = FALSE,
 method = c("chisquare", "wilson"),
 center = NULL,
  cova = NULL,
 robust = c("classical", "mve", "mcd"),
)
typprobClass(
 Х,
 data,
 groups,
  small = FALSE,
 method = c("chisquare", "wilson"),
 outlier = 0.01,
 sep = FALSE,
 cv = TRUE,
  robust = c("classical", "mve", "mcd"),
)
```

Arguments

x	vector or matrix of data the probability is to be calculated.
data	Reference data set. If missing x will be used.
small	adjustion of Mahalanobis D^2 for small sample sizes as suggested by Wilson (1981), only takes effect when method="wilson".
method	select method for probability estimation. Available options are "chisquare" (or any abbreviation) or "wilson". "chisquare" exploits simply the chisquare distribution of the mahalanobisdistance, while "wilson" uses the methods suggested by Wilson(1981). Results will be similar in general.
center	vector: specify custom vector to calculate distance to. If not defined, group mean will be used.
cova	covariance matrix to calculate mahalanobis-distance: specify custom covariance matrix to calculate distance.
robust	character: determines covariance estimation methods, allowing for robust estimations using MASS::cov.rob. Default is the standard product-moment covariance matrix.
• • •	additional parameters passed to MASS:: $cov.rob$ for robust covariance and mean estimations.
groups	vector containing grouping information.
outlier	probability threshold below which a specimen will not be assigned to any group-

typprob 169

sep logical: if TRUE, probability will be calculated from the pooled within group

covariance matrix.

cv logical: if data is missing and cv=TRUE, the resulting classification will be vali-

dated by leaving-one-out crossvalidation.

Details

get the probability for an observation belonging to a given multivariate nromal distribution

Value

typprob: returns a vector of probabilities.

typprobClass:

probs matrix of probabilities for each group

groupaffin vector of groups each specimen has been assigned to. Outliers are classified

"none"

probsCV cross-validated matrix of probabilities for each group

groupaffinCV cross-validated vector of groups each specimen has been assigned to. Outliers

are classified "none"

self logical: if TRUE, the data has been classified by self-inference.

Author(s)

Stefan Schlager

References

Albrecht G. 1992. Assessing the affinities of fossils using canonical variates and generalized distances Human Evolution 7:49-69.

Wilson S. 1981. On comparing fossil specimens with population samples Journal of Human Evolution 10:207 - 214.

```
if (require(shapes)) {
  data <- procSym(gorf.dat)$PCscores[,1:3]
  probas <- typprob(data,data,small=TRUE)### get probability for each specimen

### now we check how this behaves compared to the mahalanobis distance
maha <- mahalanobis(data,colMeans(data),cov(data))
  plot(probas,maha,xlab="Probability",ylab="Mahalanobis D^2")

data2 <- procSym(abind(gorf.dat,gorm.dat))$PCscores[,1:3]
  fac <- as.factor(c(rep("female",dim(gorf.dat)[3]),rep("male",dim(gorm.dat)[3])))
  typClass <- typprobClass(data2,groups=fac,method="w",small=TRUE,cv=TRUE)

## only 59 specimen is rather small.
  typClass2 <- typprobClass(data2,groups=fac,method="c",cv=TRUE)## use default settings</pre>
```

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```
### check results for first method:
typClass

### check results for second method:
typClass2
}
```

unrefVertex

some little helpers for vertex operations on triangular meshes

Description

some little helpers for vertex operations on triangular meshes

Usage

```
unrefVertex(mesh)
rmVertex(mesh, index, keep = FALSE)
vert2points(mesh)
rmUnrefVertex(mesh, silent = FALSE)
```

Arguments

mesh triangular mesh of class mesh3d.

index vector containing indices of vertices to be removed.

keep logical: if TRUE, the vertices specified by index are kept and the rest is removed.

silent logical: suppress output about info on removed vertices.

Details

extract vertex coordinates from meshes, find and/or remove (unreferenced) vertices from triangular meshes

unrefVertex finds unreferenced vertices in triangular meshes of class mesh3d or tmesh3d.

rmVertex removes specified vertices from triangular meshes.

vert2points extacts vertex coordinates from triangular meshes.

rmUnrefVertex removes unreferenced vertices from triangular meshes.

updateIndices 171

Value

```
unrefVertex: vector with indices of unreferenced vertices.

rmVertex: returns mesh with specified vertices removed and faces and normals updated.

vert2points: k x 3 matrix containing vertex coordinates.

rmUnrefVertex: mesh with unreferenced vertices removed.
```

Author(s)

Stefan Schlager

See Also

```
ply2mesh, file2mesh
```

Examples

```
require(rg1)
data(nose)
testmesh <- rmVertex(shortnose.mesh,1:50) ## remove first 50 vertices
## Not run:
shade3d(testmesh,col=3)### view result

## End(Not run)
testmesh$vb <- cbind(testmesh$vb,shortnose.mesh$vb[,1:50]) ## add some unreferenced vertices
## Not run:
points3d(vert2points(testmesh),col=2)## see the vertices in the holes?

## End(Not run)
cleanmesh <- rmUnrefVertex(testmesh)## remove those lonely vertices!
## Not run:
pop3d()
points3d(vert2points(cleanmesh),col=2) ### now the holes are empty!!

## End(Not run)</pre>
```

updateIndices

update a vector of indices after removal of some referenced items

Description

update a vector of indices after removal of some referenced items

Usage

```
updateIndices(x, ignore, indexrange)
```

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Arguments

x vector containing indices (e.g. to matrix rows)

ignore integer vector: remove those items from the original structure indexrange maximum range of the index in the referenced item structure

Examples

```
refItem <- matrix(1:10,5,2)
index <- c(1,3,5) # this indexes some rows of the matrix we are interested in
## now we want to ignore row 2 and 5 and want to update the index so it will still fit
indexNew <- updateIndices(index,c(2,5),indexrange=5)

## Here a more useful example:
data(boneData)
left <- c(4,6,8)

    ## determine corresponding Landmarks on the right side:
    # important: keep same order
    right <- c(3,5,7)
    pairedLM <- cbind(left,right)

## now we want to remove some landmarks and need to updated the pairedLM indices
ignore <- c(5,6)
mynewboneLM <- boneLM[-ignore,,]
pairedLMnew <- apply(pairedLM,2,updateIndices,ignore=ignore,indexrange=dim(boneLM)[1])</pre>
```

updateNormals

Compute face or vertex normals of a triangular mesh

Description

Compute face or vertex normals of a triangular mesh of class "mesh3d"

Usage

```
updateNormals(x, angle = TRUE)
facenormals(x)
```

Arguments

x triangular mesh of class "mesh3d"

angle logical: if TRUE, angle weighted normals are used.

Value

updateNormals returns mesh with updated vertex normals.

facenormals returns an object of class "mesh3d" with

vb faces' barycenters normals faces' normals vecx 173

Note

only supports triangular meshes

Author(s)

Stefan Schlager

References

Baerentzen, Jakob Andreas. & Aanaes, H., 2002. Generating Signed Distance Fields From Triangle Meshes. Informatics and Mathematical Modelling, .

See Also

ply2mesh

Examples

```
require(rgl)
require(Morpho)
data(nose)
### calculate vertex normals
shortnose.mesh$normals <- NULL ##remove normals</pre>
shade3d(shortnose.mesh,col=3)##render
## End(Not run)
shortnose.mesh <- updateNormals(shortnose.mesh)</pre>
## Not run:
clear3d()
shade3d(shortnose.mesh,col=3)##smoothly rendered now
## End(Not run)
## calculate facenormals
facemesh <- facenormals(shortnose.mesh)</pre>
## Not run:
plotNormals(facemesh,long=0.01)
points3d(vert2points(facemesh),col=2)
wire3d(shortnose.mesh)
## End(Not run)
```

vecx

convert an 3D array into a matrix and back

Description

converts a 3D-array (e.g. containing landmark coordinates) into a matrix, one row per specimen or reverse this.

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Usage

```
vecx(x, byrow = FALSE, revert = FALSE, lmdim)
```

Arguments

x array or matrix

byrow logical: if TRUE, the resulting vector for each specimen will be x1, y1, z1, x2, y2, z2, ...,

and $x1, x2, \dots, y1, y2, \dots, z1, z2, \dots$ otherwise (default). The same is for re-

verting the process: if the matrix contains the coordinates as rows like: x1,y1,z1,x2,y2,z2,...

set byrow=TRUE

revert revert the process and convert a matrix with vectorized landmarks back into an

array.

1mdim number of columns for reverting

Value

returns a matrix with one row per specimen

Author(s)

Stefan Schlager

Examples

```
if (require(shapes)) {
data <- vecx(gorf.dat)
#revert the procedure
gdat.restored <- vecx(data,revert=TRUE,lmdim=2)
range(gdat.restored-gorf.dat)
}</pre>
```

virtualMeshScan remove all parts of a triangular mesh, not visible from a set of view-points

Description

remove all parts of a triangular mesh, not visible from a set of viewpoints

Usage

```
virtualMeshScan(x, viewpoints, offset = 0.001, cores = 1)
```

Arguments

X	triangular mesh	of class	mesh3d	

viewpoints vector or k x 3 matrix containing a set of viewpoints

offset value to generate an offset at the meshes surface (see notes)
cores integer: number of cores to use (not working on windows)

Value

returns a list containing subsets of the original mesh

visible the parts visible from at least one of the viewpoints

invisible the parts not visible from the viewpoints

Note

The function tries to filter out all vertices where the line connecting each vertex with the viewpoints intersects with the mesh itself. As, technically speaking this always occurs at a distance of value=0, a mesh with a tiny offset is generated to avoid these false hits.

Examples

```
SCP1 <- file2mesh(system.file("extdata","SCP1.ply",package="Morpho"))
viewpoints <- read.fcsv(system.file("extdata","SCP1_Endo.fcsv",package="Morpho"))
## Create a quick endocast
quickEndo <- virtualMeshScan(SCP1,viewpoints)
## Not run:
rgl::shade3d(quickEndo$visible,col="orange")
rgl::shade3d(SCP1,col="white",alpha=0.5)
## End(Not run)</pre>
```

warpmovie3d

Creates a sequence of images showing predefined steps of warping two meshes or landmark configurations (2D and 3D) into each other

Description

Creates a sequence of images showing predefined steps of warping two meshes or landmark configurations (2D and 3D) into each other

Usage

```
warpmovie3d(
    x,
    y,
    n,
    col = "green",
```

```
palindrome = FALSE,
  folder = NULL,
 movie = "warpmovie",
)
## S3 method for class 'matrix'
warpmovie3d(
 Х,
 у,
 n,
  col = "green",
  palindrome = FALSE,
  folder = NULL,
 movie = "warpmovie",
  add = FALSE,
  close = TRUE,
  countbegin = 0,
  ask = TRUE,
  radius = NULL,
 links = NULL,
 lwd = 1,
)
warpmovie2d(
  х,
 у,
  col = "green",
  palindrome = FALSE,
  folder = NULL,
 movie = "warpmovie",
 links = NULL,
  lwd = 1,
  imagedim = "800x800",
 par = list(xaxt = "n", yaxt = "n", bty = "n"),
)
## S3 method for class 'mesh3d'
warpmovie3d(
 Х,
 у,
 n,
  col = NULL,
  palindrome = FALSE,
  folder = NULL,
```

```
movie = "warpmovie",
add = FALSE,
close = TRUE,
countbegin = 0,
ask = TRUE,
radius = NULL,
xland = NULL,
yland = NULL,
lmcol = "black",
...
)
```

Arguments

x mesh to start with (object of class mesh3d)

y resulting mesh (object of class mesh3d), having the same amount of vertices and

faces than the starting mesh

n integer: amount of intermediate steps.

col color of the mesh

palindrome logical: if TRUE, the procedure will go forth and back.

folder character: output folder for created images (optional)

movie character: name of the output files

... additional arguments passed to shade3d (3D) or points (2D).

add logical: if TRUE, the movie will be added to the focussed rgl-windows.

close logical: if TRUE, the rgl window will be closed when finished. width and 200

the height of the image.

countbegin integer: number to start image sequence.

ask logical: if TRUE, the viewpoint can be selected manually.

radius numeric: define size of spheres (overides atuomatic size estimation).

links vector or list of vectors containing wireframe information to connect landmarks

(optional).

lwd numeric: controls width of lines defined by "links".

imagedim character of pattern "100x200" where 100 determines the width and 200 the

height of the image.

par list of graphial parameters: details can be found here: par.

xland optional argument: add landmarks on mesh x yland optional argument: add landmarks on mesh y

lmcol optional argument: color of landmarks xland and yland

Details

given two landmark configurations or two meshes with the same amount of vertices and faces (e.g a mesh and its warped counterpart), the starting configuration/mesh will be subsequently transformed into the final configuration/mesh by splitting the differences into a predefined set of steps.

A series of png files will be saved to disk. These can be joined to animated gifs by external programs such as imagemagick or used to create animations in PDFs in a latex environment (e.g. latex package: animate).

Author(s)

Stefan Schlager

See Also

```
ply2mesh, file2mesh, mesh2ply, tps3d
```

```
###3D example
 data(nose)##load data
if (interactive()){
##warp a mesh onto another landmark configuration:
longnose.mesh <- tps3d(shortnose.mesh,shortnose.lm,longnose.lm,threads=1)</pre>
warpmovie3d(shortnose.mesh,longnose.mesh,n=15)## create 15 images.
### ad some landmarks
warpmovie3d(shortnose.mesh,longnose.mesh,n=15,xland=shortnose.lm,
            yland=longnose.lm)## create 15 images.
### restrict to landmarks
warpmovie3d(shortnose.lm,longnose.lm,n=15,movie="matrixmovie")## create 15 images.
### the images are now stored in your current working directory and can
### be concatenated to a gif using an external program such as
### imagemagick.
}
### 2D example
if (require(shapes)) {
bb <- procSym(gorf.dat)</pre>
### morph superimposed first specimen onto sample mean
warpmovie2d(bb$rotated[,,1],bb$mshape,n=20,links=c(1,5,4:2,8:6,1),imagedim="600x400")
## remove files
unlink("warpmovie00*")
}
```

write.fcsv 179

write.TCSV write nauciais in sucer4 forma	write.fcsv	write fiducials in slicer4 forma
---	------------	----------------------------------

Description

write fiducials in slicer4 format

Usage

```
write.fcsv(x, filename = dataname, description = NULL, slicer4.11 = FALSE)
```

Arguments

x matrix with row containing 2D or 3D coordinates

filename will be substituted with ".fcsv"

description optional: character vector containing a description for each landmark

slicer4.11 logical: Slicer changed their fiducial format in version >= 4.11. Set TRUE if

you use the latest Slicer version

Examples

```
require(Rvcg)
data(dummyhead)
write.fcsv(dummyhead.lm)
## remove file
unlink("dummyhead.lm.fcsv")
```

write.pts

exports a matrix containing landmarks into .pts format

Description

exports a matrix containing landmarks into .pts format that can be read by IDAV Landmark.

Usage

```
write.pts(x, filename = dataname, rownames = NULL, NA.string = 9999)
```

Arguments

	1			1 1 1	c
Y	k x m	matrix	confaming	landmark	configuration
^	1. / 111	munia	comuning	iuiiuiiuik	comingulation

filename character: Path/name of the requested output - extension will be added atuomat-

ically. If not specified, the file will be named as the exported object.

rownames provide an optional character vector with rownames

NA.string specify the string to use for encoding missing values

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Details

you can import the information into the program landmarks available at http://graphics.idav.ucdavis.edu/research/EvoMorph

Author(s)

Stefan Schlager

See Also

```
read.pts
```

Examples

```
data(nose)
write.pts(shortnose.lm, filename="shortnose")
unlink("shortnose.pts")
```

write.slicerjson

Export landmarks (or any 3D coordinates) to the new slicer json format

Description

Export landmarks (or any 3D coordinates) to the new slicer json format

Usage

```
write.slicerjson(
    x,
    filename = dataname,
    type = c("Fiducial", "Curve", "ClosedCurve"),
    coordinateSystem = c("LPS", "RAS"),
    labels = dataname
)
```

Arguments

labels character or character vector containing landmark labels.

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