Package 'patternize'

August 22, 2023

Title Quantification of Color Pattern Variation

Version 0.0.5

Maintainer Steven Van Belleghem <vanbelleghemsteven@hotmail.com>

Description Quantification of variation in organismal color patterns as obtained from image data. Patternize defines homology between pattern positions across images either through fixed landmarks or image registration. Pattern identification is performed by categorizing the distribution of colors using RGB thresholds or image segmentation.

BugReports https://github.com/StevenVB12/patternize/issues

URL https://github.com/StevenVB12/patternize

Depends R (>= 3.5.0)

Imports raster, sp, sf, abind, Morpho, dplyr, imager, magrittr, methods, purrr, vegan, RNiftyReg, geomorph, ClusterR

License GPL-3

Encoding UTF-8

LazyData true

RoxygenNote 7.2.3

NeedsCompilation no

Author Steven Van Belleghem [aut, cre]

Repository CRAN

Date/Publication 2023-08-22 14:40:05 UTC

R topics documented:

| alignLan | 3 |
|-------------------------|---|
| alignReg | 4 |
| colorChecker | 5 |
| colorChecker_customGray | 6 |
| colorChecker_half | 7 |
| createPhenotype | 8 |

| createTarget |
|------------------------|
| extdata |
| GMMImage |
| imageList |
| kImage 13 |
| kImageHSV |
| lanArray |
| landmarkArray |
| landmarkList |
| makeList |
| maskOutline |
| patArea |
| patGMM 21 |
| patK |
| patK_HSV |
| patLanHSV |
| patLanK |
| patLanK_HSV 27 |
| patLanRGB |
| patLanW |
| patPCA |
| patRDA |
| patRegHSV |
| patRegK |
| patRegK_HSV 43 |
| patRegRGB |
| patRegW 46 |
| patternize |
| plotHeat |
| plotRasterstackAsImage |
| rasterList_lanK |
| rasterList_lanRGB |
| rasterList_regK |
| rasterList_regRGB |
| redRes |
| sampleLandmarks |
| sampleRGB |
| setMask |
| sumRaster |

Index

alignLan

Description

Align images using landmarks

Usage

```
alignLan(
  imageList,
  landList,
  IDlist = NULL,
  adjustCoords = FALSE,
  resampleFactor = NULL,
  res = c(300, 300),
  transformRef = "meanshape",
  transformType = "tps",
 maskOutline = NULL,
  removebg = NULL,
  removebgColOffset = 0.1,
  inverse = FALSE,
  cartoonID = NULL,
  refImage = NULL,
 plotTransformed = FALSE,
  format = "imageJ"
)
```

| imageList | List of RasterStack objects. |
|----------------|---|
| landList | Landmark list as returned by makeList. |
| IDlist | List of sample IDs should be specified when masking outline and transformRef is 'meanshape'. |
| adjustCoords | Adjust landmark coordinates in case they are reversed compared to pixel coordinates (default = FALSE). |
| resampleFactor | Integer for downsampling used by redRes. |
| res | Resolution vector $c(x,y)$ for output rasters (default = $c(300,300)$). This should be reduced if the number of pixels in the image is lower than th raster. |
| transformRef | ID or landmark matrix of reference sample for shape to which color patterns will be transformed to. Can be 'meanshape' for transforming to mean shape of Procrustes analysis. |
| transformType | Transformation type as used by computeTransform (default ='tps'). |
| maskOutline | When outline is specified, everything outside of the outline will be masked for the color extraction (default = NULL). This can be a list of multiple outlines. |

| removebg | Integer or RGB vector indicating the range of RGB threshold to remove from image (e.g. 100 removes pixels with average RGB > 100; default = NULL). | |
|-----------------|--|--|
| removebgColOffs | et | |
| | Color offset for color background extraction (default = 0.10). | |
| inverse | If TRUE, areas withing the outline will be masked. If maskOutline is a list, this should also be a list. | |
| cartoonID | ID of the sample for which the cartoon was drawn and will be used for masking (should be set when transformRef = 'meanShape'). | |
| refImage | Image (RasterStack) used for target. Use raster::stack('filename'). | |
| plotTransformed | | |
| | Plot transformed image (default = FALSE). | |
| format | ImageJ (Fiji) or tps format (default = 'imageJ'). | |
| | | |

List of aligned RasterStack objects.

| alignReg | Aligns images using niftyreg utilities for automated image registra- tion |
|----------|--|
| | |

Description

Aligns images using niftyreg utilities for automated image registration..

Usage

```
alignReg(
 sampleList,
 target,
 resampleFactor = NULL,
 useBlockPercentage = 75,
 crop = c(0, 0, 0, 0),
 removebgR = NULL,
 maskOutline = NULL,
 plotTransformed = FALSE
)
```

| sampleList | List of RasterStack objects. | |
|--------------------|--|--|
| target | Image imported as RasterStack used as target for registration. | |
| resampleFactor | Integer for downsampling used by redRes (default = NULL). | |
| useBlockPercentage | | |
| | Block percentage as used in niftyreg (default = 75). | |

colorChecker

| crop | Vector c(xmin, xmax, ymin, ymax) that specifies the pixel coordinates to crop the original image. |
|-----------------|--|
| removebgR | Integer indicating the range RGB treshold to remove from image (e.g. 100 removes pixels with average RGB > 100; default = NULL) for registration analysis. This works only to remove a white background. |
| maskOutline | When outline is specified, everything outside of the outline will be masked for the color extraction (default = NULL). |
| plotTransformed | |
| | Whether to plot transformed images while processing (default = FALSE). |

Value

List of raster objects.

| colorChecker | Calibrate images using ColorChecker. |
|--------------|--------------------------------------|
|--------------|--------------------------------------|

Description

Calibrate images using ColorChecker.

Usage

```
colorChecker(
   IDlist,
   prepath = NULL,
   extension = NULL,
   colorCheckerType = "X-Rite",
   fixedCorners = FALSE,
   patchSize = 0.6,
   colorCheckerXY = NULL,
   fixedModel = NULL,
   resampleFactor = NULL,
   adjustCoords = FALSE
)
```

| IDlist | List of sample IDs. | |
|------------------|---|--|
| prepath | Prepath (default = NULL). | |
| extension | Extension (default = NULL). | |
| colorCheckerType | | |
| | Type of colorChecker. Options are 'X-Rite ' and 'ColorGauge Micro Analyzer' (default = 'X-Rite '). | |
| fixedCorners | Specify whether to set the coordinates of the colorChecker corners for every image (default = FALSE). | |

| patchSize | Proportion of ColorChecker patch that will be used for observed RGB values (default = 0.6). |
|----------------|---|
| colorCheckerXY | Landmark list of colorChecker corners as returned by ${\tt makeList}.$ The image will not be plotted. |
| fixedModel | Precalculated model to adjust colors. Should be a list a model for R, G and B (the colorChecker function gives as output such a list obtained from the last image in the analysis). |
| resampleFactor | Integer for downsampling used by redRes. |
| adjustCoords | Adjust landmark coordinates in case they are reversed compared to pixel coordinates (default = FALSE). |

Calibrated image(s) ('filename_calibrated.jpg')

colorChecker_customGray

Calibrate images using ColorChecker.

Description

Calibrate images using ColorChecker.

Usage

```
colorChecker_customGray(
   IDlist,
   prepath = NULL,
   extension = NULL,
   colorCheckerType = "X-Rite",
   fixedCorners = FALSE,
   patchSize = 0.6,
   colorCheckerXY = NULL,
   fixedModel = NULL,
   resampleFactor = NULL
)
```

| IDlist | List of sample IDs. |
|-----------------|---|
| prepath | Prepath (default = NULL). |
| extension | Extension (default = NULL). |
| colorCheckerTyp | e |
| | Type of colorChecker. Options are 'X-Rite ' and 'ColorGauge Micro Analyzer' |
| | (default = 'X-Rite '). |

| fixedCorners | Specify whether to set the coordinates of the colorChecker corners for every image (default = FALSE). |
|----------------|---|
| patchSize | Proportion of ColorChecker patch that will be used for observed RGB values (default = 0.6). |
| colorCheckerXY | Landmark list of colorChecker corners as returned by makeList. The image will not be plotted. |
| fixedModel | Precalculated model to adjust colors. Should be a list a model for R, G and B (the colorChecker function gives as output such a list obtained from the last image in the analysis). |
| resampleFactor | Integer for downsampling used by redRes. |

Calibrated image(s) ('filename_calibrated.jpg')

| colorChecker_half Calibrate images using (right) half of ColorChecker. Only works for X-Rite. | |
|---|--|
|---|--|

Description

Calibrate images using (right) half of ColorChecker. Only works for X-Rite.

Usage

```
colorChecker_half(
  IDlist,
  prepath = NULL,
  extension = NULL,
  colorCheckerType = "X-Rite",
  fixedCorners = FALSE,
  patchSize = 0.6,
  colorCheckerXY = NULL,
  fixedModel = NULL,
  resampleFactor = NULL,
  adjustCoords = FALSE
)
```

| IDlist | List of sample IDs. |
|-----------------|--|
| prepath | Prepath (default = NULL). |
| extension | Extension (default = NULL). |
| colorCheckerTyp | e |
| | Type of colorChecker. Options are 'X-Rite ' and 'ColorGauge Micro Analyzer (default = 'X-Rite'). |

| fixedCorners | Specify whether to set the coordinates of the colorChecker corners for every image (default = FALSE). |
|----------------|---|
| patchSize | Proportion of ColorChecker patch that will be used for observed RGB values (default = 0.6). |
| colorCheckerXY | Landmark list of colorChecker corners as returned by ${\tt makeList}.$ The image will not be plotted. |
| fixedModel | Precalculated model to adjust colors. Should be a list f a model for R, G and B (the colorChecker function gives as output such a list obtained from the last image in the analysis). |
| resampleFactor | Integer for downsampling used by redRes. |
| adjustCoords | Adjust landmark coordinates in case they are reversed compared to pixel coordinates (default = FALSE). |

Calibrated image(s) ('filename_calibrated.jpg')

createPhenotype

Plot color pattern prediction for specified PCA values

Description

Plot color pattern prediction for specified PCA values

Usage

```
createPhenotype(
 PCAdata,
 PCApredict,
  IDlist,
  rasterList,
  colpalette = NULL,
 plotCartoon = FALSE,
  refShape = NULL,
  outline = NULL,
  lines = NULL,
  landList = NULL,
  adjustCoords = FALSE,
  cartoonID = NULL,
  normalized = TRUE,
  crop = c(0, 0, 0, 0),
  flipRaster = NULL,
  flipOutline = NULL,
  imageList = NULL,
  cartoonOrder = "above",
  lineOrder = "above",
```

createPhenotype

```
cartoonCol = "gray",
cartoonFill = NULL,
legendTitle = "Proportion",
zlim = NULL
)
```

| PCApredict A vector with the PCA values for which to predict the phenotype. This vector only needs to include the values upto the last PCA axis to predict along, other values are set to zero. IDlist List of sample IDs. rasterList rasterList used for PCA. | |
|--|----|
| - | |
| rasterlist rasterList used for PCA | |
| | |
| <pre>colpalette Vector of colors for color palette (default = c("white","lightblue","blue","green "yellow","red"))</pre> | ", |
| plotCartoon Whether to plot a cartoon. This cartoon should be drawn on one of the sample used in the analysis. | es |
| refShape This can be 'target' in case the reference shape is a single sample (for registration analysis) or 'mean' if the images were transformed to a mean shape (only for meanshape when using landmark transformation) | |
| outline xy coordinates that define outline. | |
| lines list of files with xy coordinates of line objects to be added to cartoon. | |
| landList Landmark landmarkList. | |
| adjustCoords Adjust landmark coordinates. | |
| cartoonID ID of the sample for which the cartoon was drawn. | |
| normalized Set this to true in case the summed rasters are already devided by the samp number. | e |
| crop Vector c(xmin, xmax, ymin, ymax) that specifies the pixel coordinates to cro the original image used in landmark or registration analysis. | р |
| flipRaster Whether to flip raster along xy axis (in case there is an inconsistency betwee raster and outline coordinates). | n |
| flipOutline Whether to flip plot along x, y or xy axis. | |
| imageList List of images should be given if one wants to flip the outline or adjust landmar coordinates. | k |
| cartoonOrder Whether to plot the cartoon outline 'above' or 'under' the pattern raster (defau = 'above'). Set to 'under' for filled outlines. | lt |
| lineOrder Whether to plot the cartoon lines 'above' or 'under' the pattern raster (default 'above'). | = |
| cartoonCol Outline and line color for cartoon (deafault = 'gray'). | |
| cartoonFill Fill color for outline of cartoon (default = NULL). | |
| legendTitle Title of the raster legend (default = 'Proportion'). | |
| zlim zlim values for predicted pattern. | |

createTarget

Description

Create a target image (RasterStack) from a polygon.

Usage

```
createTarget(
   outline,
   image,
   res = 300,
   colorFill = "black",
   colorBG = "white",
   sigma = 10,
   plot = FALSE
)
```

Arguments

| outline | xy coordinates that define outline. |
|-----------|--|
| image | Image imported as RasterStack used in the analysis. This is used to extract the extent and dimensions for the raster layers. |
| res | Resolution for RasterStack (default = 300). |
| colorFill | Color for the fill of the polygon (default = 'black'). |
| colorBG | Color for the background (default = 'white'). |
| sigma | Size of sigma for Gaussian blurring (default = 10). |
| plot | Whether to plot the created target image (default = FALSE). |

Value

RasterStack

Examples

```
## Not run:
outline_BC0077 <- read.table(paste(system.file("extdata", package = 'patternize'),
'/BC0077_outline.txt', sep=''), header = FALSE)
data(imageList)
target <- createTarget(outline_BC0077, imageList[[1]], plot = TRUE)
## End(Not run)
```

extdata

Description

Raw image, landmark and cartoon data of Heliconius erato hydara wings.

Format

Raw JPG images, landmark and cartoon data.

BC0077.JPG jpeg image BC0071.JPG jpeg image BC0050.JPG jpeg image BC0049.JPG jpeg image BC0004.JPG jpeg image BC0077_landmarks_LFW.Txt xy landmark coordinates BC0071_landmarks_LFW.Txt xy landmark coordinates BC0050_landmarks_LFW.Txt xy landmark coordinates BC0049_landmarks_LFW.Txt xy landmark coordinates BC0004_landmarks_LFW.Txt xy landmark coordinates BC0077_outline.txt xy outline coordinates BC0077_vein1.txt xy vein coordinates BC0077_vein2.txt xy vein coordinates BC0077_vein3.txt xy vein coordinates BC0077_vein4.txt xy vein coordinates BC0077_vein5.txt xy vein coordinates BC0077_vein6.txt xy vein coordinates BC0077_vein7.txt xy vein coordinates BC0077_vein8.txt xy vein coordinates BC0077_vein9.txt xy vein coordinates BC0077_vein10.txt xy vein coordinates BC0077_vein11.txt xy vein coordinates

GMMImage

Description

GMM clustering of image imported as a RasterStack.

Usage

```
GMMImage(image, k = 5, maskToNA = NULL, kmeansOnAll = FALSE)
```

Arguments

| image | Image imported as a RasterStack for clustering. |
|-------------|---|
| k | Integer for number of k clusters (default = 3). |
| maskToNA | Replace the color value used for masking (i.e. 0 or 255) with NA. |
| kmeansOnAll | Whether to perform the kmeans clusters on the combined set of pixels of all images first (default = FALSE). |

Value

List including the clustered RasterSatck returned as an array and object of class "GMM".

|--|--|--|

Description

List of RasterStacks as returned by makeList.

Usage

imageList

Format

A list of 5 RasterStack objects of Heliconius erato hydara dorsal forewings.

Examples

```
## Not run:
data(imageList)
summary(imageList)
```

End(Not run)

kImage

kmeans clustering of image imported as a RasterStack. This function is used by patLanK and patRegK.

Description

 ${\sf kmeans}$ clustering of image imported as a RasterStack. This function is used by patLanK and patRegK.

Usage

```
kImage(image, k = 5, startCenter = NULL, maskToNA = NULL, kmeansOnAll = FALSE)
```

Arguments

| image | Image imported as a RasterStack for k-means clustering. |
|-------------|---|
| k | Integer for number of k-means clusters (default = 3). |
| startCenter | A matrix of cluster centres to start k-means clustering from (default = NULL). |
| maskToNA | Replace the color value used for masking (i.e. 0 or 255) with NA. |
| kmeansOnAll | Whether to perform the kmeans clusters on the combined set of pixels of all images first (default = FALSE). |

Value

List including the k-means clustered RasterSatck returned as an array and object of class "kmeans".

Examples

```
image <- raster::stack(system.file("extdata", "BC0077.jpg", package = "patternize"))
out <- kImage(image, 6)</pre>
```

| kImageHSV | kmeans clustering of image imported as a RasterStack. This function |
|-----------|---|
| | is used by patLanK and patRegK. |

Description

 ${\sf kmeans}$ clustering of image imported as a RasterStack. This function is used by patLanK and patRegK.

Usage

```
kImageHSV(
    image,
    k = 5,
    startCenter = NULL,
    maskToNA = NULL,
    kmeansOnAll = FALSE,
    ignoreHSVvalue = FALSE
)
```

Arguments

| image | HSV image imported as a RasterStack for k-means clustering. |
|----------------|---|
| k | Integer for number of k-means clusters (default = 3). |
| startCenter | A matrix of cluster centres to start k-means clustering from (default = NULL). |
| maskToNA | Replace the color value used for masking (i.e. 0 or 255) with NA. |
| kmeansOnAll | Whether to perform the kmeans clusters on the combined set of pixels of all images first (default = FALSE). |
| ignoreHSVvalue | Whether to ignore the HSV value (~darkness). |

Value

List including the k-means clustered RasterSatck returned as an array and object of class "kmeans".

Examples

```
image <- raster::stack(system.file("extdata", "BC0077.jpg", package = "patternize"))
out <- kImage(image, 6)</pre>
```

lanArray

Build landmark array for Morpho.

Description

Build landmark array for Morpho.

Usage

```
lanArray(sampleList, adjustCoords = FALSE, imageList = NULL, imageIDs = NULL)
```

14

landmarkArray

Arguments

| sampleList | List of landmark matrices as returned by makeList. |
|--------------|--|
| adjustCoords | Adjust landmark coordinates in case they are reversed compared to pixel coordinates (default = FALSE). |
| imageList | List of RasterStacks as returned by makeList should be given when adjustCoords = TRUE. |
| imageIDs | A list of IDs to match landmarks to images if landmarkList and imageList don't have the same length. |

Value

X x Y x n array, where X and Y define the coordinates of the landmark points and n is the sample size.

Examples

```
## Not run:
IDlist <- c('BC0077','BC0071','BC0050','BC0049','BC0004')
prepath <- system.file("extdata", package = 'patternize')
extension <- '_landmarks_LFW.txt'
landmarkList <- makeList(IDlist, 'landmark', prepath, extension)
landmarkArray <- lanArray(landmarkList)
## End(Not run)
```

landmarkArray *landmarkArray*

Description

Array of landmarks as returned by lanArray and used by link[Morpho]{procsym}.

Usage

landmarkArray

Format

An array of landmarks for 5 Heliconius erato hydara dorsal forewings.

Examples

```
## Not run:
data(landmarkArray)
summary(landmarkArray)
```

End(Not run)

landmarkList landmarkList

Description

List of landmarks as returned by makeList.

Usage

landmarkList

Format

A list of landmarks for 5 Heliconius erato hydara dorsal forewings.

Examples

```
## Not run:
data(landmarkList)
summary(landmarkList)
```

End(Not run)

```
makeList
```

Build list of landmarks or RasterStacks from images using filepath and file extension.

Description

Build list of landmarks or RasterStacks from images using filepath and file extension.

Usage

```
makeList(
   IDlist,
   type,
   prepath = NULL,
   extension = NULL,
   format = "imageJ",
   tpsFile = NULL,
   skipLandmark = NULL
)
```

16

maskOutline

Arguments

| IDlist | List of sample IDs. |
|--------------|---|
| type | 'landmark' or 'image' depending on what type of list to make. |
| prepath | Prepath (default = NULL). |
| extension | Extension (default = NULL). |
| format | ImageJ (Fiji) or tps format (default = 'imageJ'). |
| tpsFile | Provide filename of tps file ff format is 'tps'. |
| skipLandmark | Vector of rownumbers of landmarks to skip. |

Value

Landmark or RasterStack list.

Examples

IDlist <- c('BC0077','BC0071','BC0050','BC0049','BC0004')
prepath <- system.file("extdata", package = 'patternize')
extension <- '_landmarks_LFW.txt'
landmarkList <- makeList(IDlist, 'landmark', prepath, extension)
extension <- '.jpg'
imageList <- makeList(IDlist, 'image', prepath, extension)</pre>

maskOutline

Intersects a RasterStack with an outline. Everything outside of the outline will be removed from the raster.

Description

Intersects a RasterStack with an outline. Everything outside of the outline will be removed from the raster.

Usage

```
maskOutline(
  RasterStack,
  outline,
  refShape,
  landList = NULL,
  adjustCoords = FALSE,
  cartoonID = NULL,
  IDlist = NULL,
  crop = c(0, 0, 0, 0),
```

```
flipRaster = NULL,
flipOutline = NULL,
imageList = NULL,
maskColor = 0,
inverse = FALSE
)
```

Arguments

| RasterStack | RasterStack to be masked. |
|--------------|--|
| outline | xy coordinates that define outline. |
| refShape | This can be 'target' in case the reference shape is a single sample (for registration analysis) or 'mean' if the images were transformed to a mean shape (only for meanshape when using landmark transformation) |
| landList | Landmark list to be given when type = 'mean'. |
| adjustCoords | Adjust landmark coordinates in case they are reversed compared to pixel coordinates (default = FALSE). |
| cartoonID | ID of the sample for which the cartoon was drawn. Only has to be given when refShape is 'mean'. |
| IDlist | List of sample IDs should be specified when refShape is 'mean'. |
| crop | Vector c(xmin, xmax, ymin, ymax) that specifies the pixel coordinates to crop the original image used in landmark or registration analysis. |
| flipRaster | Whether to flip raster along xy axis (in case there is an inconsistency between raster and outline coordinates). |
| flipOutline | Whether to flip plot along x, y or xy axis. |
| imageList | List of image as obtained from makeList should be given if one wants to flip the outline or adjust landmark coordinates. |
| maskColor | Color the masked area gets. Set to 0 for black (default) or 255 for white. |
| inverse | If TRUE, areas withing the outline will be masked. |

Examples

```
## Not run:
data(imageList)
outline_BC0077 <- read.table(paste(system.file("extdata", package = 'patternize'),
'/BC0077_outline.txt', sep=''), header = FALSE)
masked <- maskOutline(imageList[[1]], outline_BC0077, refShape = 'target', flipOutline = 'y')
## End(Not run)
```

18

patArea

This function calculates the area in which the color pattern is expressed in each sample as the relative proportion using the provided outline of the considered trait or structure.

Description

This function calculates the area in which the color pattern is expressed in each sample as the relative proportion using the provided outline of the considered trait or structure.

Usage

```
patArea(
   rList,
   IDlist,
   refShape,
   type,
   outline = NULL,
   landList = NULL,
   adjustCoords = FALSE,
   cartoonID = NULL,
   crop = c(0, 0, 0, 0),
   flipRaster = NULL,
   flipOutline = NULL,
   imageList = NULL
)
```

| rList | List of RasterLayers as obtained from the main patternize functions. |
|--------------|---|
| IDlist | List of sample IDs. |
| refShape | This can be 'target' in case the reference shape is a single sample (for registra- tion analysis) or 'mean' if the images were transformed to a mean shape using landmark transformation. |
| type | Type of rasterlist; 'RGB' or 'k' (result from RGB or k-means analysis, respec- tively). |
| outline | xy coordinates that define outline. |
| landList | Landmark list as returned by makeList. |
| adjustCoords | Adjust landmark coordinates in case they are reversed compared to pixel coordinates (default = FALSE). |
| cartoonID | ID of the sample for which the cartoon was drawn. |
| crop | Vector c(xmin, xmax, ymin, ymax) that specifies the pixel coordinates to crop the original image used in landmark or registration analysis. |
| flipRaster | Whether to flip raster along xy axis (in case there is an inconsistency between raster and outline coordinates). |

patArea

| flipOutline | Whether to flip plot along x, y or xy axis. |
|-------------|---|
| imageList | List of images as obtained from makeList should be given if one wants to flip the outline or adjust landmark coordinates. |

Value

Table or list of tables with sample IDs and relative area of color pattern or kmeans cluster.

Examples

```
data(rasterList_lanRGB)
#data(rasterList_regRGB)
#data(rasterList_lanK)
#data(rasterList_regK)
data(imageList)
IDlist <- c('BC0077','BC0071','BC0050','BC0049','BC0004')</pre>
outline_BC0077 <- read.table(paste(system.file("extdata", package = 'patternize'),</pre>
'/BC0077_outline.txt', sep=''), header = FALSE)
prepath <- system.file("extdata", package = 'patternize')</pre>
extension <- '_landmarks_LFW.txt'</pre>
landmarkList <- makeList(IDlist, 'landmark', prepath, extension)</pre>
## Not run:
area_lanRGB <- patArea(rasterList_lanRGB, IDlist, refShape = 'mean', type = 'RGB',</pre>
outline = outline_BC0077, landList = landmarkList, adjustCoords = TRUE,
imageList = imageList, cartoonID = 'BC0077')
area_regRGB <- patArea(rasterList_regRGB, IDlist, refShape = 'target', type = 'RGB',
outline = outline_BC0077, crop = c(100,400,40,250), adjustCoords = TRUE,
imageList = imageList, cartoonID = 'BC0077', flipRaster = 'xy')
areaList_lanK <- patArea(rasterList_lanK, IDlist, refShape = 'mean', type = 'k',</pre>
outline = outline_BC0077, landList = landmarkList, adjustCoords = TRUE,
imageList = imageList, cartoonID = 'BC0077')
areaList_regK <- patArea(rasterList_regK, IDlist, refShape = 'target', type = 'k',
outline = outline_BC0077, crop = c(100,400,40,250), adjustCoords = TRUE,
imageList = imageList, cartoonID = 'BC0077', flipRaster = 'xy')
## End(Not run)
```

patGMM

Description

Extract colors using GMM clustering (for pre-aligned images).

Usage

```
patGMM(
  sampleList,
  k = 3,
  resampleFactor = NULL,
  maskOutline = NULL,
  plot = FALSE,
  focal = FALSE,
  sigma = 3,
  maskToNA = NULL,
  kmeansOnAll = FALSE
)
```

Arguments

| sampleList | List of RasterStack objects. |
|----------------|--|
| k | Integere for defining number of clusters (default = 3). |
| resampleFactor | Integer for downsampling used by redRes. |
| maskOutline | When outline is specified, everything outside of the outline will be masked for the color extraction (default = NULL). |
| plot | Whether to plot transformed color patterns while processing (default = FALSE). |
| focal | Whether to perform Gaussian blurring (default = FALSE). |
| sigma | Size of sigma for Gaussian blurring (default = 3). |
| maskToNA | Replace the color value used for masking (i.e. 0 or 255) with NA. |
| kmeansOnAll | Whether to perform the kmeans clusters on the combined set of pixels of all images first (default = FALSE). |

Value

List of summed raster for each k-means cluster objects.

patK

Description

Extract colors using k-means clustering (for pre-aligned images).

Usage

```
patK(
  sampleList,
  k = 3,
  fixedStartCenter = NULL,
  resampleFactor = NULL,
  maskOutline = NULL,
  plot = FALSE,
  focal = FALSE,
  sigma = 3,
  maskToNA = NULL,
  kmeansOnAll = FALSE
)
```

Arguments

| sampleList | List of RasterStack objects. |
|-----------------|--|
| k | Integere for defining number of k-means clusters (default = 3). |
| fixedStartCente | er |
| | Specify a dataframe with start centers for k-means clustering. |
| resampleFactor | Integer for downsampling used by redRes. |
| maskOutline | When outline is specified, everything outside of the outline will be masked for the color extraction (default = NULL). |
| plot | Whether to plot transformed color patterns while processing (default = FALSE). |
| focal | Whether to perform Gaussian blurring (default = FALSE). |
| sigma | Size of sigma for Gaussian blurring (default = 3). |
| maskToNA | Replace the color value used for masking (i.e. 0 or 255) with NA. |
| kmeansOnAll | Whether to perform the kmeans clusters on the combined set of pixels of all images first (default = FALSE). |

Value

List of summed raster for each k-means cluster objects.

patK_HSV

Description

Extract colors using k-means clustering (for pre-aligned images).

Usage

```
patK_HSV(
   sampleList,
   k = 3,
   fixedStartCenter = NULL,
   resampleFactor = NULL,
   maskOutline = NULL,
   plot = FALSE,
   focal = FALSE,
   sigma = 3,
   maskToNA = NULL,
   kmeansOnAll = FALSE,
   ignoreHSVvalue = FALSE
)
```

Arguments

| sampleList | List of RasterStack objects. |
|------------------|--|
| k | Integere for defining number of k-means clusters (default = 3). |
| fixedStartCenter | |
| | Specify a dataframe with start centers for k-means clustering. |
| resampleFactor | Integer for downsampling used by redRes. |
| maskOutline | When outline is specified, everything outside of the outline will be masked for the color extraction (default = NULL). |
| plot | Whether to plot transformed color patterns while processing (default = FALSE). |
| focal | Whether to perform Gaussian blurring (default = FALSE). |
| sigma | Size of sigma for Gaussian blurring (default = 3). |
| maskToNA | Replace the color value used for masking (i.e. 0 or 255) with NA. |
| kmeansOnAll | Whether to perform the kmeans clusters on the combined set of pixels of all images first (default = FALSE). |
| ignoreHSVvalue | Whether to ignore the HSV value (~darkness). |

Value

List of summed raster for each k-means cluster objects.

patLanHSV

Description

Aligns images usings transformations obtained from fixed landmarks and extracts colors using a predefined RGB values and cutoff value.

Usage

```
patLanHSV(
  sampleList,
  landList,
 HSV,
  resampleFactor = NULL,
  colOffset = 0.1,
  crop = FALSE,
  cropOffset = c(0, 0, 0, 0),
  res = 300,
  transformRef = "meanshape",
  transformType = "tps",
  adjustCoords = FALSE,
 plot = NULL,
  focal = FALSE,
  sigma = 3,
  iterations = 0,
  ignoreHSVvalue = FALSE,
  patternsToFile = NULL
)
```

| sampleList | List of RasterStack objects. |
|----------------|--|
| landList | Landmark list as returned by makeList. |
| HSV | HSV values for color pattern extraction specified as vector. |
| resampleFactor | Integer for downsampling used by redRes. |
| colOffset | Color offset for color pattern extraction (default = 0.10). |
| crop | Whether to use the landmarks range to crop the image. This can speed up the analysis (default = FALSE). |
| cropOffset | Vector c(xmin, xmax, ymin, ymax) that specifies the number of pixels you want the cropping to be offset from the landmarks (in case the landmarks do not sur- round the entire color pattern). The values specified should present the percent- age of the maximum landmark value along the x and y axis. |

patLanHSV

| res | Resolution for color pattern raster (default = 300). This should be reduced if the number of pixels in the image is lower than th raster. |
|----------------|--|
| transformRef | ID of reference sample for shape to which color patterns will be transformed to. Can be 'meanshape' for transforming to mean shape of Procrustes analysis. |
| transformType | Transformation type as used by computeTransform (default ='tps'). |
| adjustCoords | Adjust landmark coordinates in case they are reversed compared to pixel coordinates (default = FALSE). |
| plot | Whether to plot transformed color patterns while processing (default = NULL). Transformed color patterns can be plot on top of each other ('stack') or next to the original image for each sample ('compare'). |
| focal | Whether to perform Gaussian blurring (default = FALSE). |
| sigma | Size of sigma for Gaussian blurring (default = 3). |
| iterations | Number of iterations for recalculating average color. |
| ignoreHSVvalue | Whether to ignore the HSV value (~darkness). |
| patternsToFile | Name of directory to which the color pattern of each individual will be outputted (default = NULL). |

Value

List of raster objects.

Examples

```
## Not run:
IDlist <- c('BC0077','BC0071','BC0050','BC0049','BC0004')
prepath <- system.file("extdata", package = 'patternize')
extension <- '_landmarks_LFW.txt'
landmarkList <- makeList(IDlist, 'landmark', prepath, extension)
extension <- '.jpg'
imageList <- makeList(IDlist, 'image', prepath, extension)
HSV <- c(0.025,1,0.45)
rasterList_lanHSV <- patLanRGB(imageList, landmarkList, HSV,
coloffset = 0.15, crop = TRUE, res = 100, adjustCoords = TRUE, plot = 'stack')
## End(Not run)
```

patLanK

patLanK

Description

Aligns images usings transformations obtained from fixed landmarks and extracts colors using kmeans clustering.

Usage

```
patLanK(
  sampleList,
  landList,
  k = 3,
  fixedStartCenter = NULL,
  resampleFactor = NULL,
  crop = FALSE,
  cropOffset = c(0, 0, 0, 0),
  res = 300,
  transformRef = "meanshape",
  transformType = "tps",
  removebg = NULL,
  removebgColOffset = 0.1,
  adjustCoords = FALSE,
 plot = FALSE,
  focal = FALSE,
  sigma = 3
)
```

| sampleList | List of RasterStack objects. | |
|------------------|---|--|
| landList | Landmark list as returned by makeList. | |
| k | Integere for defining number of k-means clusters (default = 3). | |
| fixedStartCenter | | |
| | Specify a dataframe with start centers for k-means clustering. | |
| resampleFactor | Integer for downsampling used by redRes. | |
| crop | Whether to use the landmarks range to crop the image. This can significantly speed up the analysis (default = FALSE). | |
| cropOffset | Vector c(xmin, xmax, ymin, ymax) that specifies the number of pixels you want the cropping to be offset from the landmarks (in case the landmarks do not surround the entire color pattern). The values specified should present the percentage of the maximum landmark value along the x and y axis. | |

patLanK_HSV

| res | Resolution for color pattern raster (default = 300). This should be reduced if the number of pixels in the image is lower than th raster. | |
|-------------------|---|--|
| transformRef | ID of reference sample for shape to which color patterns will be transformed to. Can be 'meanshape' for transforming to mean shape of Procrustes analysis. | |
| transformType | Transformation type as used by computeTransform (default ='tps'). | |
| removebg | Integer or RGB vector indicating the range of RGB threshold to remove from image (e.g. 100 removes pixels with average RGB > 100; default = NULL). | |
| removebgColOffset | | |
| | Color offset for color background extraction (default = 0.10). | |
| adjustCoords | Adjust landmark coordinates in case they are reversed compared to pixel coordinates (default = FALSE). | |
| plot | Whether to plot transformed color patterns while processing (default = FALSE). | |
| focal | Whether to perform Gaussian blurring (default = FALSE). | |
| sigma | Size of sigma for Gaussian blurring (default = 3). | |

Value

List of summed raster for each k-means cluster objects.

Examples

```
## Not run:
IDlist <- c('BC0077','BC0071','BC0050','BC0049','BC0004')
prepath <- system.file("extdata", package = 'patternize')
extension <- '_landmarks_LFW.txt'
landmarkList <- makeList(IDlist, 'landmark', prepath, extension)
extension <- '.jpg'
imageList <- makeList(IDlist, 'image', prepath, extension)
# Note that this example only aligns two images with the target,
# remove [1:2] to run a full examples.
rasterList_lanK <- patLanK(imageList[1:2], landmarkList[1:2], k = 4, crop = TRUE,
res = 100, removebg = 100, adjustCoords = TRUE, plot = TRUE)
```

End(Not run)

patLanK_HSV

Aligns images usings transformations obtained from fixed landmarks and extracts colors using k-means clustering.

Description

Aligns images usings transformations obtained from fixed landmarks and extracts colors using kmeans clustering.

Usage

```
patLanK_HSV(
  sampleList,
  landList,
 k = 3,
 fixedStartCenter = NULL,
  resampleFactor = NULL,
  crop = FALSE,
  cropOffset = c(0, 0, 0, 0),
  res = 300,
  transformRef = "meanshape",
  transformType = "tps",
  removebgK = NULL,
  adjustCoords = FALSE,
 plot = FALSE,
  focal = FALSE,
  sigma = 3,
  ignoreHSVvalue = FALSE
)
```

Arguments

| sampleList | List of RasterStack objects. |
|-----------------|--|
| landList | Landmark list as returned by makeList. |
| k | Integere for defining number of k-means clusters (default = 3). |
| fixedStartCente | er |
| | Specify a dataframe with start centers for k-means clustering. |
| resampleFactor | Integer for downsampling used by redRes. |
| crop | Whether to use the landmarks range to crop the image. This can significantly speed up the analysis (default = FALSE). |
| cropOffset | Vector c(xmin, xmax, ymin, ymax) that specifies the number of pixels you want the cropping to be offset from the landmarks (in case the landmarks do not sur- round the entire color pattern). The values specified should present the percent- age of the maximum landmark value along the x and y axis. |
| res | Resolution for color pattern raster (default = 300). This should be reduced if the number of pixels in the image is lower than th raster. |
| transformRef | ID of reference sample for shape to which color patterns will be transformed to. Can be 'meanshape' for transforming to mean shape of Procrustes analysis. |
| transformType | Transformation type as used by computeTransform (default ='tps'). |
| removebgK | Integer indicating the range RGB treshold to remove from image (e.g. 100 removes pixels with average RGB > 100; default = NULL) for k-means analysis. This works only to remove a white background. |
| adjustCoords | Adjust landmark coordinates in case they are reversed compared to pixel coordinates (default = FALSE). |
| plot | Whether to plot transformed color patterns while processing (default = FALSE). |

28

patLanRGB

| focal | Whether to perform Gaussian blurring (default = FALSE). |
|----------------|---|
| sigma | Size of sigma for Gaussian blurring (default = 3). |
| ignoreHSVvalue | Whether to ignore the HSV value (~darkness). |

Value

List of summed raster for each k-means cluster objects.

Examples

```
## Not run:
IDlist <- c('BC0077','BC0071','BC0050','BC0049','BC0004')
prepath <- system.file("extdata", package = 'patternize')
extension <- '_landmarks_LFW.txt'
landmarkList <- makeList(IDlist, 'landmark', prepath, extension)
extension <- '.jpg'
imageList <- makeList(IDlist, 'image', prepath, extension)
# Note that this example only aligns two images with the target,
# remove [1:2] to run a full examples.
rasterList_lanK <- patLanK(imageList[1:2], landmarkList[1:2], k = 4, crop = TRUE,
res = 100, removebgK = 100, adjustCoords = TRUE, plot = TRUE)
```

End(Not run)

| patLanRGB | Aligns images usings transformations obtained from fixed landmarks |
|-----------|---|
| | and extracts colors using a predefined RGB values and cutoff value. |

Description

Aligns images usings transformations obtained from fixed landmarks and extracts colors using a predefined RGB values and cutoff value.

Usage

```
patLanRGB(
   sampleList,
   landList,
   RGB = NULL,
   sampleRGB = FALSE,
   sampleRGBtype = "point",
   resampleFactor = NULL,
   colOffset = 0.1,
   crop = FALSE,
   cropOffset = c(0, 0, 0, 0),
   res = 300,
```

```
transformRef = "meanshape",
transformType = "tps",
adjustCoords = FALSE,
plot = NULL,
focal = FALSE,
sigma = 3,
iterations = 0,
imageIDs = NULL,
patternsToFile = NULL
)
```

Arguments

| sampleList | List of RasterStack objects. |
|----------------|--|
| landList | Landmark list as returned by makeList. |
| RGB | RGB values for color pattern extraction specified as vector. |
| sampleRGB | Whether to set RGB for each image manually. |
| sampleRGBtype | Whether to pick a point or area (defined by left bottom and top right) for sampleRGB. |
| resampleFactor | Integer for downsampling used by redRes. |
| colOffset | Color offset for color pattern extraction (default = 0.10). |
| crop | Whether to use the landmarks range to crop the image. This can speed up the analysis (default = FALSE). |
| cropOffset | Vector c(xmin, xmax, ymin, ymax) that specifies the number of pixels you want the cropping to be offset from the landmarks (in case the landmarks do not sur- round the entire color pattern). The values specified should present the percent- age of the maximum landmark value along the x and y axis. |
| res | Resolution for color pattern raster (default = 300). This should be reduced if the number of pixels in the image is lower than th raster. |
| transformRef | ID of reference sample for shape to which color patterns will be transformed to. Can be 'meanshape' for transforming to mean shape of Procrustes analysis. |
| transformType | Transformation type as used by computeTransform (default ='tps'). |
| adjustCoords | Adjust landmark coordinates in case they are reversed compared to pixel coor- dinates (default = FALSE). |
| plot | Whether to plot transformed color patterns while processing (default = NULL). Transformed color patterns can be plot on top of each other ('stack') or next to the original image for each sample ('compare'). |
| focal | Whether to perform Gaussian blurring (default = FALSE). |
| sigma | Size of sigma for Gaussian blurring (default = 3). |
| iterations | Number of iterations for recalculating average color. |
| imageIDs | A list of IDs to match landmarks to images if landmarkList and imageList don't have the same length. |
| patternsToFile | Name of directory to which the color pattern of each individual will be outputted (default = NULL). |

30

patLanW

Value

List of raster objects.

Examples

```
## Not run:
IDlist <- c('BC0077','BC0071','BC0050','BC0049','BC0004')
prepath <- system.file("extdata", package = 'patternize')
extension <- '_landmarks_LFW.txt'
landmarkList <- makeList(IDlist, 'landmark', prepath, extension)
extension <- '.jpg'
imageList <- makeList(IDlist, 'image', prepath, extension)
RGB <- c(114,17,0)
rasterList_lanRGB <- patLanRGB(imageList, landmarkList, RGB,
colOffset = 0.15, crop = TRUE, res = 100, adjustCoords = TRUE, plot = 'stack')
## End(Not run)
```

| patLanW | Extracts color pattern from landmark transformed image using water- shed segmentation. This function works interactively by allowing to pick a starting pixel within each pattern element from which the wa- tershed will extract the pattern. This function works best for patterns with sharp boundaries. |
|---------|---|
| | |

Description

Extracts color pattern from landmark transformed image using watershed segmentation. This function works interactively by allowing to pick a starting pixel within each pattern element from which the watershed will extract the pattern. This function works best for patterns with sharp boundaries.

Usage

```
patLanW(
   sampleList,
   landList,
   IDlist = NULL,
   adjustCoords = FALSE,
   transformRef = "meanshape",
   resampleFactor = NULL,
   transformType = "tps",
   maskOutline = NULL,
   cartoonID = NULL,
```

```
correct = FALSE,
blur = TRUE,
sigma = 3,
bucketfill = TRUE,
cleanP = NULL,
splitC = NULL,
plotTransformed = FALSE,
plotCorrect = FALSE,
plotEdges = FALSE,
plotEdges = FALSE,
plotWS = FALSE,
plotBF = FALSE,
plotFinal = FALSE
)
```

Arguments

| sampleList | List of RasterStack objects. | |
|-----------------|---|--|
| landList | Landmark list as returned by makeList. | |
| IDlist | List of sample IDs should be specified when masking outline and transformRef is 'meanshape'. | |
| adjustCoords | Adjust landmark coordinates in case they are reversed compared to pixel coordinates (default = FALSE). | |
| transformRef | ID or landmark matrix of reference sample for shape to which color patterns will be transformed to. Can be 'meanshape' for transforming to mean shape of Procrustes analysis. | |
| resampleFactor | Integer for downsampling image used by redRes. | |
| transformType | Transformation type as used by computeTransform (default ='tps'). | |
| maskOutline | When outline is specified, everything outside of the outline will be masked for the color extraction (default = NULL). | |
| cartoonID | ID of the sample for which the cartoon was drawn and will be used for masking (should be set when transformRef = 'meanShape'). | |
| correct | Correct image illumination using a linear model (default = FALSE). | |
| blur | Blur image for priority map extraction (default = TRUE). | |
| sigma | Size of sigma for Gaussian blurring (default = 5). | |
| bucketfill | Use a bucket fill on the background to fill holes (default = TRUE). | |
| cleanP | Integer to remove spurious areas with width smaller than cleanP (default = NULL). | |
| splitC | Integer to split selected patterns into connected components and remove ones with areas smaller than splitC (default = NULL). | |
| plotTransformed | | |
| | Plot transformed image (default = FALSE). | |
| plotCorrect | Plot corrected image, corrected for illumination using a linear model (default = FALSE). | |

32

patPCA

| plotEdges | Plot image gradient (default = FALSE). |
|--------------|--|
| plotPriority | Plot priority map (default = FALSE). |
| plotWS | Plot watershed result (default = FALSE). |
| plotBF | Plot bucketfill (default = FALSE). |
| plotFinal | Plot extracted patterns (default = FALSE). |

Value

List of raster objects.

Examples

```
## Not run:
IDlist <- c('BC0077','BC0071','BC0050','BC0049','BC0004')
prepath <- system.file("extdata", package = 'patternize')
extension <- '_landmarks_LFW.txt'
landmarkList <- makeList(IDlist, 'landmark', prepath, extension)
extension <- '.jpg'
imageList <- makeList(IDlist, 'image', prepath, extension)
outline_BC0077 <- read.table(paste(system.file("extdata", package = 'patternize'),
'/BC0077_outline.txt', sep=''), header = FALSE)
rasterList_W <- patLanW(imageList, landmarkList, IDlist, transformRef = 'meanshape',
adjustCoords = TRUE, plotTransformed = FALSE, correct = TRUE, plotCorrect = FALSE, blur = FALSE,
sigma = 2, bucketfill = FALSE, cleanP = 0, splitC = 10, plotPriority = TRUE, plotWS = TRUE,
plotBF = TRUE, plotFinal = TRUE, maskOutline = outline_BC0077, cartoonID = 'BC0077')
```

End(Not run)

patPCA

This function transforms the individual color pattern rasters as obtained by the main patternize functions to a dataframe of 0 and 1 values that can be used for Principal Component Analysis (prcomp). This function also allows to plot the analysis including a visualization of the shape changes along the axis. Pixel values are predicted by multiplying the rotation matrix (eigenvectors) with a vector that has the same length as the number of rows in the rotation matrix and in which all values are set to zero except for the PC value for which we want to

predict the pixel values.

Description

This function transforms the individual color pattern rasters as obtained by the main patternize functions to a dataframe of 0 and 1 values that can be used for Principal Component Analysis (prcomp). This function also allows to plot the analysis including a visualization of the shape changes along the axis. Pixel values are predicted by multiplying the rotation matrix (eigenvectors) with a vector that has the same length as the number of rows in the rotation matrix and in which all values are set to zero except for the PC value for which we want to predict the pixel values.

Usage

```
patPCA(
  rList,
  popList,
  colList,
  symbolList = NULL,
  rListPredict = NULL,
  popListPredict = NULL,
  colListPredict = NULL,
  pcaListPredict = NULL,
  pcaPopListPredict = NULL,
  pcaColPredict = "red",
  symbolListPredict = NULL,
  plot = FALSE,
  plotType = "points",
  plotChanges = FALSE,
  PCx = 1,
  PCy = 2,
  plotCartoon = FALSE,
  refShape = NULL,
  outline = NULL,
  lines = NULL,
  landList = NULL,
  adjustCoords = FALSE,
  crop = c(0, 0, 0, 0),
  flipRaster = NULL,
  flipOutline = NULL,
  imageList = NULL,
  cartoonID = NULL,
  refImage = NULL,
  colpalette = NULL,
  normalized = NULL,
  cartoonOrder = "above",
  lineOrder = "above",
  cartoonCol = "gray",
  cartoonFill = NULL,
  plotLandmarks = FALSE,
  landCol = "black",
  zlim = c(-1, 1),
```

34

patPCA

```
legendTitle = "Predicted",
xlab = "",
ylab = "",
main = "",
...
```

| 18 | Junicitis | |
|----|-----------------|--|
| | rList | List of raster objects. |
| | popList | List of vectors including sampleIDs for each population. |
| | colList | List of colors for each population. |
| | symbolList | List with graphical plotting symbols (default = NULL). |
| | rListPredict | List of raster objects to predict into PCA space (default = NULL). |
| | popListPredict | List of vectors including sampleIDs for each set of predict samples (default = NULL). Note to that this also has to be a list if only one population is included. |
| | colListPredict | List of colors for each set of predict samples (default = NULL). |
| | pcaListPredict | Points to plot within PCA space. |
| | pcaPopListPredi | |
| | | List of population symbols for plotting additional PCA values. |
| | pcaColPredict | Color for additional PCA values. |
| | symbolListPredi | List with graphical plotting symbols for predict sets (default = NULL). |
| | nlat | |
| | plot | Whether to plot the PCA analysis (default = FALSE). |
| | plotType | Plot 'points' or sample 'labels' (default = 'points') |
| | plotChanges | Wether to include plots of the changes along the PC axis (default = FALSE). |
| | PCx | PC axis to be presented for x-axis (default PC1). |
| | РСу | PC axis to be presented for y-axis (default PC2). |
| | plotCartoon | Whether to plot a cartoon. This cartoon should be drawn on one of the samples used in the analysis. |
| | refShape | This can be 'target' in case the reference shape is a single sample (for registration analysis) or 'mean' if the images were transformed to a mean shape (only for meanshape when using landmark transformation) |
| | outline | xy coordinates that define outline. |
| | lines | list of files with xy coordinates of line objects to be added to cartoon. |
| | landList | Landmark landmarkList. |
| | adjustCoords | Adjust landmark coordinates. |
| | crop | Vector c(xmin, xmax, ymin, ymax) that specifies the pixel coordinates to crop the original image used in landmark or registration analysis. |
| | flipRaster | Whether to flip raster along xy axis (in case there is an inconsistency between raster and outline coordinates). |
| | flipOutline | Whether to flip plot along x, y or xy axis. |
| | | |

| imageList | List of image should be given if one wants to flip the outline or adjust landmark coordinates. |
|---------------|--|
| cartoonID | ID of the sample for which the cartoon was drawn. |
| refImage | Image (RasterStack) used for target. Use raster::stack('filename'). |
| colpalette | Vector of colors for color palette (default = c("white","lightblue","blue","green", "yellow","red")) |
| normalized | Set this to true in case the summed rasters are already devided by the sample number. |
| cartoonOrder | Whether to plot the cartoon outline 'above' or 'under' the pattern raster (default = 'above'). Set to 'under' for filled outlines. |
| lineOrder | Whether to plot the cartoon lines 'above' or 'under' the pattern raster (default = 'above'). |
| cartoonCol | Outline and line color for cartoon (deafault = 'gray'). |
| cartoonFill | Fill color for outline of cartoon (default = NULL). |
| plotLandmarks | Whether to plot the landmarks from the target image or mean shape landmarks (default = FALSE). |
| landCol | Color for plotting landmarks (default = 'black'). |
| zlim | z-axis limit (default = $c(0,1)$) |
| legendTitle | Title of the raster legend (default = 'Proportion') |
| xlab | Optional x-axis label. |
| ylab | Optional y-axis label. |
| main | Optional main title. |
| | additional arguments for PCA plot function. |

If plot = TRUE: List including a [1] dataframe of the binary raster values that can be used for principle component analysis, [2] a dataframe of sample IDs and specified population colors and [3] prcomp results. If plot = FALSE: prcomp result.

See Also

prcomp

Examples

```
data(rasterList_lanRGB)
```

```
pop1 <- c('BC0077','BC0071')
pop2 <- c('BC0050','BC0049','BC0004')
popList <- list(pop1, pop2)
colList <- c("red", "blue")
pcaOut <- patPCA(rasterList_lanRGB, popList, colList, plot = TRUE)</pre>
```
This function transforms the individual color pattern rasters as obtained by the main patternize functions to a dataframe of 0 and 1 values that can be used for constrained Redundancy Analysis (RDA) (rda). This function also allows to plot the analysis including a visualization of the shape changes along the axis.

Description

patRDA

This function transforms the individual color pattern rasters as obtained by the main patternize functions to a dataframe of 0 and 1 values that can be used for constrained Redundancy Analysis (RDA) (rda). This function also allows to plot the analysis including a visualization of the shape changes along the axis.

Usage

```
patRDA(
  rList,
  popList,
  colList,
  symbolList = NULL,
  rListPredict = NULL,
  popListPredict = NULL,
  colListPredict = NULL,
  symbolListPredict = NULL,
  plot = FALSE,
  plotType = "points",
  plotChanges = FALSE,
  PCx = 1,
  PCy = 2,
  plotCartoon = FALSE,
  refShape = NULL,
  outline = NULL,
  lines = NULL,
  landList = NULL,
  adjustCoords = FALSE,
  crop = c(0, 0, 0, 0),
  flipRaster = NULL,
  flipOutline = NULL,
  imageList = NULL,
  cartoonID = NULL,
  colpalette = NULL,
  normalized = NULL,
  cartoonOrder = "above",
  lineOrder = "above",
  cartoonCol = "gray",
  cartoonFill = NULL,
```

```
plotLandmarks = FALSE,
landCol = "black",
zlim = c(-1, 1),
legendTitle = "Predicted",
xlab = "",
ylab = "",
main = ""
```

Arguments

| rList | List of raster objects. |
|-----------------------------------|--|
| popList | List of vectors including sampleIDs for each population. |
| colList | List of colors for each population. |
| symbolList | List with graphical plotting symbols (default = NULL). |
| rListPredict | List of raster objects to predict into DFA space (default = NULL). |
| popListPredict | List of vectors including sampleIDs for each set of predict samples (default = NULL). Note to that this also has to be a list if only one population is included. |
| colListPredict symbolListPredi | List of colors for each set of predict samples (default = NULL). ct |
| | List with graphical plotting symbols for predict sets (default = NULL). |
| plot | Whether to plot the PCA analysis (default = FALSE). |
| plotType | Plot 'points' or sample 'labels' (default = 'points') |
| plotChanges | Wether to include plots of the changes along the PC axis (default = FALSE). |
| PCx | PC axis to be presented for x-axis (default PC1). |
| РСу | PC axis to be presented for y-axis (default PC2). |
| plotCartoon | Whether to plot a cartoon. This cartoon should be drawn on one of the samples used in the analysis. |
| refShape | This can be 'target' in case the reference shape is a single sample (for registration analysis) or 'mean' if the images were transformed to a mean shape (only for meanshape when using landmark transformation) |
| outline | xy coordinates that define outline. |
| lines | list of files with xy coordinates of line objects to be added to cartoon. |
| landList | Landmark landmarkList. |
| adjustCoords | Adjust landmark coordinates. |
| crop | Vector c(xmin, xmax, ymin, ymax) that specifies the pixel coordinates to crop the original image used in landmark or registration analysis. |
| flipRaster | Whether to flip raster along xy axis (in case there is an inconsistency between raster and outline coordinates). |
| flipOutline | Whether to flip plot along x, y or xy axis. |
| imageList | List of image should be given if one wants to flip the outline or adjust landmark coordinates. |

patRDA

| cartoonID | ID of the sample for which the cartoon was drawn. |
|---------------------|--|
| colpalette | Vector of colors for color palette (default = c("white","lightblue","blue","green", "yellow","red")) |
| normalized | Set this to true in case the summed rasters are already devided by the sample number. |
| cartoonOrder | Whether to plot the cartoon outline 'above' or 'under' the pattern raster (default = 'above'). Set to 'under' for filled outlines. |
| lineOrder | Whether to plot the cartoon lines 'above' or 'under' the pattern raster (default = 'above'). |
| cartoonCol | Outline and line color for cartoon (deafault = 'gray'). |
| cartoonFill | Fill color for outline of cartoon (default = NULL). |
| plotLandmarks | Whether to plot the landmarks from the target image or mean shape landmarks (default = FALSE). |
| landCol | Color for plotting landmarks (default = 'black'). |
| zlim | z-axis limit (default = $c(0,1)$) |
| | |
| legendTitle | Title of the raster legend (default = 'Proportion') |
| legendTitle xlab | Title of the raster legend (default = 'Proportion') Optional x-axis label. |
| - | |
| xlab | Optional x-axis label. |
| xlab | Optional x-axis label. |

Value

If plot = TRUE: List including a [1] dataframe of the binary raster values that can be used for discriminant function analysis, [2] a dataframe of sample IDs and specified population colors and [3] lda results. if rListPredict not empty: [4] class prediction of samples. If plot = FALSE: lda result only.

See Also

lda

Examples

```
data(rasterList_lanRGB)
```

```
pop1 <- c('BC0077','BC0071')
pop2 <- c('BC0050','BC0049','BC0004')
popList <- list(pop1, pop2)
colList <- c("red", "blue")
pcaOut <- patRDA(rasterList_lanRGB, popList, colList, plot = TRUE)</pre>
```

patRegHSV

Aligns images using niftyreg utilities for automated image registration and extracts colors using a predefined HSV values and cutoff value.

Description

Aligns images using niftyreg utilities for automated image registration and extracts colors using a predefined HSV values and cutoff value.

Usage

```
patRegHSV(
  sampleList,
  target,
 HSV,
  resampleFactor = NULL,
  useBlockPercentage = 75,
  colOffset = 0.1,
  crop = c(0, 0, 0, 0),
  removebgR = NULL,
 maskOutline = NULL,
  plot = FALSE,
  focal = FALSE,
  sigma = 3,
  iterations = 0,
  ignoreHSVvalue = FALSE,
  patternsToFile = NULL
)
```

| sampleList | List of RasterStack objects. | |
|--------------------|--|--|
| target | Image imported as RasterStack used as target for registration. | |
| HSV | Values for color pattern extraction specified as HSV vector. | |
| resampleFactor | Integer for downsampling used by redRes (default = NULL). | |
| useBlockPercentage | | |
| | Block percentage as used in niftyreg (default = 75). | |
| colOffset | Color offset for color pattern extraction (default = 0.10). | |
| crop | Vector c(xmin, xmax, ymin, ymax) that specifies the pixel coordinates to crop the original image. | |
| removebgR | Integer indicating the range RGB treshold to remove from image (e.g. 100 removes pixels with average RGB > 100; default = NULL) for registration analysis. This works only to remove a white background. | |

patRegK

| maskOutline | When outline is specified, everything outside of the outline will be masked for the color extraction (default = NULL). |
|----------------|--|
| plot | Whether to plot transformed color patterns while processing (default = FALSE). Transformed color patterns can be plot on top of each other ('stack') or next to the original image for each sample ('compare'). |
| focal | Whether to perform Gaussian blurring (default = FALSE). |
| sigma | Size of sigma for Gaussian blurring (default = 3). |
| iterations | Number of iterations for recalculating average color (default = 0). If set, the RGB value for pattern extraction will be iteratively recalculated to be the average of the extracted area. This may improve extraction of distinct color pattern, but fail for more gradually distributed (in color space) patterns. |
| ignoreHSVvalue | Whether to ignore the HSV value (~darkness). |
| patternsToFile | Name of directory to which the color pattern of each individual will be outputted (default = NULL). |

Value

List of raster objects.

Examples

```
## Not run:
IDlist <- c('BC0077','BC0071','BC0050','BC0049','BC0004')
prepath <- system.file("extdata", package = 'patternize')
extension <- '.jpg'
imageList <- makeList(IDlist, 'image', prepath, extension)
target <- imageList[[1]]
HSV <- c(0.025,1,0.45)
# Note that this example only aligns one image with the target,
# remove [2] to run a full examples.
rasterList_regHSV <- patRegRGB(imageList[2], target, HSV,
colOffset= 0.15, crop = c(100,400,40,250), removebgR = 100, plot = 'stack')
## End(Not run)
```

patRegK

Aligns images using niftyreg utilities for automated image registration and extracts colors using k-means clustering.

Description

Aligns images using niftyreg utilities for automated image registration and extracts colors using k-means clustering.

Usage

```
patRegK(
  sampleList,
  target,
 k = 3,
  fixedStartCenter = NULL,
  resampleFactor = NULL,
  useBlockPercentage = 75,
  crop = c(0, 0, 0, 0),
  removebgR = NULL,
  removebgK = NULL,
 maskOutline = NULL,
 maskColor = 0,
 plot = FALSE,
  focal = FALSE,
  sigma = 3
)
```

Arguments

| sampleList | List of RasterStack objects. |
|-----------------|--|
| target | Image imported as RasterStack used as target for registration. |
| k | Integere for defining number of k-means clusters (default = 3). |
| fixedStartCente | er |
| | Specify a dataframe with start centers for k-means clustering. |
| resampleFactor | Integer for downsampling used by redRes (default = NULL). |
| useBlockPercent | tage |
| | Block percentage as used in niftyreg (default = 75). |
| crop | Vector c(xmin, xmax, ymin, ymax) that specifies the pixel coordinates to crop the original image. |
| removebgR | Integer indicating the range RGB treshold to remove from image (e.g. 100 removes pixels with average RGB > 100; default = NULL) for registration analysis. This works only to remove a white background. |
| removebgK | Integer indicating the range RGB treshold to remove from image (e.g. 100 removes pixels with average RGB > 100; default = NULL) for k-means analysis. This works only to remove a white background. |
| maskOutline | When outline is specified, everything outside of the outline will be masked for the color extraction (default = NULL). |
| maskColor | Color the masked area gets. Set to 0 for black (default) or 255 for white. |
| plot | Whether to plot k-means clustered image while processing (default = FALSE). |
| focal | Whether to perform Gaussian blurring (default = FALSE). |
| sigma | Size of sigma for Gaussian blurring (default = 3). |

Value

List of rasters for each k-means cluster objects.

patRegK_HSV

Examples

```
IDlist <- c('BC0077','BC0071','BC0050','BC0049','BC0004')
prepath <- system.file("extdata", package = 'patternize')
extension <- '.jpg'
imageList <- makeList(IDlist, 'image', prepath, extension)
target <- imageList[[1]]
## Not run:
rasterList_regK <- patRegK(imageList[3], target, k = 5,
crop = c(100,400,40,250), removebgR = 100, plot = TRUE)
## End(Not run)</pre>
```

| patRegK_HSV | Aligns images using niftyreg utilities for automated image registra- |
|-------------|--|
| | tion and extracts colors using k-means clustering. |

Description

Aligns images using niftyreg utilities for automated image registration and extracts colors using k-means clustering.

Usage

```
patRegK_HSV(
  sampleList,
  target,
  k = 3,
  fixedStartCenter = NULL,
  resampleFactor = NULL,
  useBlockPercentage = 75,
  crop = c(0, 0, 0, 0),
  removebgR = NULL,
  removebgK = NULL,
  maskOutline = NULL,
 maskColor = 0,
  plot = FALSE,
  focal = FALSE,
  sigma = 3,
  ignoreHSVvalue = FALSE
)
```

Arguments

| sampleList | List of RasterStack objects. |
|-----------------|--|
| target | Image imported as RasterStack used as target for registration. |
| k | Integere for defining number of k-means clusters (default = 3). |
| fixedStartCente | er |
| | Specify a dataframe with start centers for k-means clustering. |
| resampleFactor | Integer for downsampling used by redRes (default = NULL). |
| useBlockPercent | age |
| | Block percentage as used in niftyreg (default = 75). |
| crop | Vector c(xmin, xmax, ymin, ymax) that specifies the pixel coordinates to crop the original image. |
| removebgR | Integer indicating the range RGB treshold to remove from image (e.g. 100 removes pixels with average RGB > 100; default = NULL) for registration analysis. This works only to remove a white background. |
| removebgK | Integer indicating the range RGB treshold to remove from image (e.g. 100 removes pixels with average RGB > 100; default = NULL) for k-means analysis. This works only to remove a white background. |
| maskOutline | When outline is specified, everything outside of the outline will be masked for the color extraction (default = NULL). |
| maskColor | Color the masked area gets. Set to 0 for black (default) or 255 for white. |
| plot | Whether to plot k-means clustered image while processing (default = FALSE). |
| focal | Whether to perform Gaussian blurring (default = FALSE). |
| sigma | Size of sigma for Gaussian blurring (default = 3). |
| ignoreHSVvalue | Whether to ignore the HSV value (~darkness). |

Value

List of rasters for each k-means cluster objects.

Examples

```
IDlist <- c('BC0077','BC0071','BC0050','BC0049','BC0004')
prepath <- system.file("extdata", package = 'patternize')
extension <- '.jpg'
imageList <- makeList(IDlist, 'image', prepath, extension)
target <- imageList[[1]]
## Not run:
rasterList_regK <- patRegK(imageList[3], target, k = 5,
crop = c(100,400,40,250), removebgR = 100, plot = TRUE)
## End(Not run)</pre>
```

patRegRGB

Aligns images using niftyreg utilities for automated image registration and extracts colors using a predefined RGB values and cutoff value.

Description

Aligns images using niftyreg utilities for automated image registration and extracts colors using a predefined RGB values and cutoff value.

Usage

```
patRegRGB(
  sampleList,
  target,
 RGB,
  resampleFactor = NULL,
  useBlockPercentage = 75,
  colOffset = 0.1,
  crop = c(0, 0, 0, 0),
  removebgR = NULL,
 maskOutline = NULL,
 plot = FALSE,
  focal = FALSE,
  sigma = 3,
  iterations = 0,
  patternsToFile = NULL
)
```

| sampleList | List of RasterStack objects. |
|----------------|--|
| target | Image imported as RasterStack used as target for registration. |
| RGB | Values for color pattern extraction specified as RGB vector. |
| resampleFactor | Integer for downsampling used by redRes (default = NULL). |
| useBlockPercen | tage |
| | Block percentage as used in niftyreg (default = 75). |
| colOffset | Color offset for color pattern extraction (default = 0.10). |
| crop | Vector c(xmin, xmax, ymin, ymax) that specifies the pixel coordinates to crop the original image. |
| removebgR | Integer indicating the range RGB treshold to remove from image (e.g. 100 removes pixels with average RGB > 100; default = NULL) for registration analysis. This works only to remove a white background. |
| maskOutline | When outline is specified, everything outside of the outline will be masked for the color extraction (default = NULL). |

| plot | Whether to plot transformed color patterns while processing (default = FALSE). Transformed color patterns can be plot on top of each other ('stack') or next to the original image for each sample ('compare'). |
|----------------|--|
| focal | Whether to perform Gaussian blurring (default = FALSE). |
| sigma | Size of sigma for Gaussian blurring (default = 3). |
| iterations | Number of iterations for recalculating average color (default = 0). If set, the RGB value for pattern extraction will be iteratively recalculated to be the average of the extracted area. This may improve extraction of distinct color pattern, but fail for more gradually distributed (in color space) patterns. |
| patternsToFile | Name of directory to which the color pattern of each individual will be outputted (default = NULL). |

Value

List of raster objects.

Examples

```
## Not run:
IDlist <- c('BC0077','BC0071','BC0050','BC0049','BC0004')
prepath <- system.file("extdata", package = 'patternize')
extension <- '.jpg'
imageList <- makeList(IDlist, 'image', prepath, extension)
target <- imageList[[1]]
RGB <- c(114,17,0)
# Note that this example only aligns one image with the target,
# remove [2] to run a full examples.
rasterList_regRGB <- patRegRGB(imageList[2], target, RGB,
colOffset= 0.15, crop = c(100,400,40,250), removebgR = 100, plot = 'stack')
```

End(Not run)

patRegW

Aligns images using niftyreg utilities for automated image registration and extracts color pattern using watershed segmentation. This function works interactively by allowing to pick a starting pixel within each pattern element from which the watershed will extract the pattern. This function works best for patterns with sharp boundaries.

patRegW

Description

Aligns images using **niftyreg** utilities for automated image registration and extracts color pattern using watershed segmentation. This function works interactively by allowing to pick a starting pixel within each pattern element from which the watershed will extract the pattern. This function works best for patterns with sharp boundaries.

Usage

```
patRegW(
  sampleList,
  target,
  resampleFactor = NULL,
  useBlockPercentage = 75,
  crop = c(0, 0, 0, 0),
  removebgR = NULL,
 maskOutline = NULL,
  cartoonID = NULL,
  correct = FALSE,
  blur = TRUE,
  sigma = 3,
  bucketfill = TRUE,
  cleanP = NULL,
  splitC = NULL,
  plotTransformed = FALSE,
  plotCorrect = FALSE,
  plotEdges = FALSE,
  plotPriority = FALSE,
  plotWS = FALSE,
  plotBF = FALSE,
  plotFinal = FALSE
)
```

| sampleList | List of RasterStack objects. |
|----------------|--|
| target | Image imported as RasterStack used as target for registration. |
| resampleFactor | Integer for downsampling image used by redRes. |
| useBlockPercen | tage |
| | Block percentage as used in niftyreg (default = 75). |
| crop | Vector c(xmin, xmax, ymin, ymax) that specifies the pixel coordinates to crop the original image. |
| removebgR | Integer indicating the range RGB treshold to remove from image (e.g. 100 removes pixels with average RGB > 100; default = NULL) for registration analysis. This works only to remove a white background. |
| maskOutline | When outline is specified, everything outside of the outline will be masked for the color extraction (default = NULL). |

| cartoonID | ID of the sample for which the cartoon was drawn and will be used for masking. | | |
|----------------|---|--|--|
| correct | Correct image illumination using a linear model (default = FALSE). | | |
| blur | Blur image for priority map extraction (default = TRUE). | | |
| sigma | Size of sigma for Gaussian blurring (default = 5). | | |
| bucketfill | Use a bucket fill on the background to fill holes (default = TRUE). | | |
| cleanP | Integer to remove spurious areas with width smaller than cleanP (default = NULL). | | |
| splitC | Integer to split selected patterns into connected components and remove ones with areas smaller than splitC (default = NULL). | | |
| plotTransforme | plotTransformed | | |
| | Plot transformed image (default = FALSE). | | |
| plotCorrect | Plot corrected image, corrected for illumination using a linear model (default = FALSE). | | |
| plotEdges | Plot image gradient (default = FALSE). | | |
| plotPriority | Plot priority map (default = FALSE). | | |
| plotWS | Plot watershed result (default = FALSE). | | |
| plotBF | Plot bucketfill (default = FALSE). | | |
| plotFinal | Plot extracted patterns (default = FALSE). | | |

Value

List of raster objects.

Examples

End(Not run)

patternize

patternize - An R package for quantifying color pattern variation.

Description

Quantifying variation in color patterns to study and compare the consistency of their expression necessitates the homologous alignment and color-based segmentation of images. Patternize is an R package that quantifies variation in color patterns as obtained from image data. Patternize defines homology between pattern positions across specimens either through fixed landmarks or image registration. Pattern identification is performed by categorizing the distribution of colors using either an RGB threshold or an unsupervised image segmentation. The quantification of the color patterns can be visualized as heat maps and compared between sets of samples.

patternize main functions

The package has six main functions depending on how you want the alignment of the iamges and the color extraction to be performed.

patLanRGB

Aligns images by transformations obtained from fixed landmarks and extracts colors using a predefined RGB values and cutoff value.

patLanK

Aligns images by transformations obtained from fixed landmarks and extracts colors using k-means clustering.

patLan₩

Aligns images by transformations obtained from fixed landmarks and extracts color patterns by watershed segmentation using imager utilities.

patRegRGB

Aligns images using niftyreg utilities for automated image registration and extracts colors using a predefined RGB values and cutoff value.

patRegK

Aligns images using niftyreg utilities for automated image registration and extracts colors using k-means clustering.

patRegW

Aligns images using **niftyreg** utilities for automated image registration and extracts color patterns by watershed segmentation using **imager** utilities.

patternize preprocessing functions

The input for the main patternize functions are RasterStack objects and when landmark transformation is used, landmark arrays.

makeList

This function returns a list of RasterStacks or a list of landmarks depending on the input provided.

sampleLandmarks Sample landmarks in an image.

lanArray

This function creates a landmark array as used by procSym in the package Morpho.

patternize postprocessing functions

sumRaster

This function sums the individual color pattern rasters as obtained by the main patternize functions.

plotHeat

Plots the color pattern heatmaps from sumRaster output.

patPCA

This function transforms the individual color pattern rasters as obtained by the main patternize functions to a dataframe of 0 and 1 values that can be used for Principal Component Analysis (prcomp). This function also allows to plot the analysis including a visualization of the shape changes along the axis.

patRDA

This function transforms the individual color pattern rasters as obtained by the main patternize functions to a dataframe of 0 and 1 values that can be used for constrained Redundancy Analysis (rda). This function also allows to plot the analysis including a visualization of the shape changes along the axis.

patArea

This function calculates the area in which the color pattern is expressed in each sample as the relative proportion using the provided outline of the considered trait or structure.

patternize miscellaneous functions

redRes

Reduces the resolution of the RasterStack objects to speed up analysis.

kImage

Performs k-means clustering of images.

sampleRGB

Interactive function to sample RGB value from pixel or area in an image.

createTarget

Creates an artificial target images using a provided outline that can be used for image registration (experimental).

maskOutline

Intersects a RasterStack with an outline. Everything outside of the outline will be removed from the raster.

colorChecker Calibrate images using ColorChecker.

Author(s)

Steven M. Van Belleghem

plotHeat

See Also

raster, stack, procSym, computeTransform, niftyreg imager

Jon Clayden, Marc Modat, Benoit Presles, Thanasis Anthopoulos and Pankaj Daga (2017). RNiftyReg: Image Registration Using the 'NiftyReg' Library. R package version 2.5.0. https://CRAN.Rproject.org/package=RNiftyReg

Stefan Schlager (2016). Morpho: Calculations and Visualisations Related to Geometric Morphometrics. R package version 2.4.1.1. https://github.com/zarquon42b/Morpho

Simon Barthelmé (2017). imager: Image processing library based on 'CImg'. R package version 0.40.2. https://CRAN.R-project.org/package=imager

plotHeat

Plots the color pattern heatmaps from sumRaster output.

Description

Plots the color pattern heatmaps from sumRaster output.

Usage

```
plotHeat(
  summedRaster,
  IDlist.
  colpalette = NULL,
 plotCartoon = FALSE,
  refShape = NULL,
 outline = NULL,
 lines = NULL,
  landList = NULL,
  adjustCoords = FALSE,
  cartoonID = NULL,
  normalized = FALSE,
  crop = c(0, 0, 0, 0),
  flipRaster = NULL,
  flipOutline = NULL,
  imageList = NULL,
  refImage = NULL,
  cartoonOrder = "above",
  lineOrder = "above",
  cartoonCol = "gray",
  cartoonFill = NULL,
  plotLandmarks = FALSE,
  landCol = "black",
```

```
zlim = c(0, 1),
legend = TRUE,
legendTitle = "Proportion",
legend.side = 4,
xlab = "",
ylab = "",
main = "",
plotType = "multi",
imageIDs = NULL,
format = "imageJ"
)
```

Arguments

| summedRaster | Summed raster or summedRasterList. |
|--------------|--|
| IDlist | List of sample IDs. |
| colpalette | Vector of colors for color palette (default = c("white","lightblue","blue","green", "yellow","red")) |
| plotCartoon | Whether to plot a cartoon. This cartoon should be drawn on one of the samples used in the analysis. |
| refShape | This can be 'target' in case the reference shape is a single sample (for registration analysis) or 'mean' if the images were transformed to a mean shape (only for meanshape when using landmark transformation) |
| outline | xy coordinates that define outline. |
| lines | list of files with xy coordinates of line objects to be added to cartoon. |
| landList | Landmark landmarkList. |
| adjustCoords | Adjust landmark coordinates. |
| cartoonID | ID of the sample for which the cartoon was drawn. |
| normalized | Set this to true in case the summed rasters are already devided by the sample number. |
| crop | Vector c(xmin, xmax, ymin, ymax) that specifies the pixel coordinates to crop the original image used in landmark or registration analysis. |
| flipRaster | Whether to flip raster along xy axis (in case there is an inconsistency between raster and outline coordinates). |
| flipOutline | Whether to flip plot along x, y or xy axis. |
| imageList | List of images should be given if one wants to flip the outline or adjust landmark coordinates. |
| refImage | Image (RasterStack) used for target. Use raster::stack('filename'). |
| cartoonOrder | Whether to plot the cartoon outline 'above' or 'under' the pattern raster (default = 'above'). Set to 'under' for filled outlines. |
| lineOrder | Whether to plot the cartoon lines 'above' or 'under' the pattern raster (default = 'above'). |
| cartoonCol | Outline and line color for cartoon (deafault = 'gray'). |

plotHeat

| cartoonFill | Fill color for outline of cartoon (default = NULL). |
|---------------|---|
| plotLandmarks | Whether to plot the landmarks from the target image or mean shape landmarks (default = FALSE). |
| landCol | Color for ploting landmarks (default = 'black'). |
| zlim | z-axis limit (default = $c(0,1)$) |
| legend | Whether to plot legend with heatmaps. |
| legendTitle | Title of the raster legend (default = 'Proportion') |
| legend.side | Side to plot legend (default = 4) |
| xlab | Optional x-axis label. |
| ylab | Optional y-axis label. |
| main | Optional main title. |
| plotType | Set as 'PCA' when visualizing shape changes along PCA axis in \ codepatPCA, as 'one' when visualizing single image or as 'multi' for multi plotting or when setting customized margins (default = 'multi'). |
| imageIDs | A list of IDs to match landmarks to images if landmarkList and imageList don't have the same length. |
| format | ImageJ (Fiji) or tps format (default = 'imageJ'). |

Examples

```
data(rasterList_lanRGB)
IDlist <- c('BC0077','BC0071','BC0050','BC0049','BC0004')
outline_BC0077 <- read.table(paste(system.file("extdata", package = 'patternize'),
'/BC0077_outline.txt', sep=''), header = FALSE)
lines_BC0077 <- list.files(path=paste(system.file("extdata", package = 'patternize')),
pattern='vein', full.names = TRUE)</pre>
```

```
summedRaster_regRGB <- sumRaster(rasterList_regRGB, IDlist, type = 'RGB')
data(imageList)</pre>
```

```
plotHeat(summedRaster_regRGB, IDlist, plotCartoon = TRUE, refShape = 'target',
outline = outline_BC0077, lines = lines_BC0077, crop = c(100,400,40,250),
flipRaster = 'xy', imageList = imageList, cartoonOrder = 'under', cartoonID = 'BC0077',
cartoonFill = 'black', main = 'registration_example')
```

```
## Not run:
data(rasterList_lanK)
IDlist <- c('BC0077','BC0071','BC0050','BC0049','BC0004')
summedRasterList <- sumRaster(rasterList_lanK, IDlist, type = 'k')
plotHeat(summedRasterList, IDlist)
```

```
summedRasterList_regK <- sumRaster(rasterList_regK, IDlist, type = 'k')
plotHeat(summedRasterList_regK, IDlist, plotCartoon = TRUE, refShape = 'target',
outline = outline_BC0077, lines = lines_BC0077, crop = c(100,400,40,250),
flipRaster = 'y', imageList = imageList, cartoonOrder = 'under',
cartoonFill = 'black', main = 'kmeans_example')</pre>
```

```
plotHeat(summedRasterList_regK[[1]], IDlist, plotCartoon = TRUE, refShape = 'target',
outline = outline_BC0077, lines = lines_BC0077, crop = c(100,400,40,250),
flipRaster = 'y', imageList = imageList, cartoonOrder = 'under',
cartoonFill = 'black', main = 'kmeans_example')
prepath <- system.file("extdata", package = 'patternize')</pre>
extension <- '_landmarks_LFW.txt'</pre>
landmarkList <- makeList(IDlist, 'landmark', prepath, extension)</pre>
summedRaster_lanRGB <- sumRaster(rasterList_lanRGB, IDlist, type = 'RGB')</pre>
plotHeat(summedRaster_lanRGB, IDlist, plotCartoon = TRUE, refShape = 'mean',
outline = outline_BC0077, lines = lines_BC0077, landList = landmarkList,
adjustCoords = TRUE, imageList = imageList, cartoonID = 'BC0077',
cartoonOrder = 'under', cartoonFill= 'black', main = 'Landmark_example')
summedRaster_lanK <- sumRaster(rasterList_lanK, IDlist, type = 'k')</pre>
plotHeat(summedRaster_lanK, IDlist, plotCartoon = TRUE, refShape = 'mean',
outline = outline_BC0077, lines = lines_BC0077, landList = landmarkList,
adjustCoords = TRUE, imageList = imageList, cartoonID = 'BC0077',
cartoonOrder = 'under', cartoonFill= 'black', main = 'Landmark_example')
plotHeat(summedRaster_lanK[[2]], IDlist, plotCartoon = TRUE, refShape = 'mean',
outline = outline_BC0077, lines = lines_BC0077, landList = landmarkList,
adjustCoords = TRUE, imageList = imageList, cartoonID = 'BC0077',
cartoonOrder = 'under', cartoonFill= 'black', main = 'Landmark_example')
```

End(Not run)

plotRasterstackAsImage

Plot rasterStack as image.

Description

Plot rasterStack as image.

Usage

plotRasterstackAsImage(rasterStack, flipY = FALSE)

Arguments

| rasterStack | A single rasterStack. |
|-------------|--|
| flipY | Whether to flip the raster along the Y-axis. |

rasterList_lanK rasterList_lanK

Description

List of RasterLayers as returned by patLanK.

Usage

rasterList_lanK

Format

A list of RasterLayers including the red color pattern extracted from 5 Heliconius erato hydara dorsal forewings using patLanK.

Examples

```
## Not run:
data(rasterList_lanK)
summary(rasterList_lanL)
```

End(Not run)

rasterList_lanRGB rasterList_lanRGB

Description

List of RasterLayers as returned by patLanRGB.

Usage

rasterList_lanRGB

Format

A list of RasterLayers including the red color pattern extracted from 5 Heliconius erato hydara dorsal forewings using patLanRGB.

Examples

```
## Not run:
data(rasterList_lanRGB)
summary(rasterList_lanRGB)
```

End(Not run)

rasterList_regK rasterList_regK

Description

List of RasterLayers as returned by patRegK.

Usage

rasterList_regK

Format

A list of RasterLayers including the red color pattern extracted from 5 Heliconius erato hydara dorsal forewings using patRegK.

Examples

```
## Not run:
data(rasterList_regK)
summary(rasterList_regK)
```

End(Not run)

rasterList_regRGB rasterList_regRGB

Description

List of RasterLayers as returned by patRegRGB.

Usage

rasterList_regRGB

Format

A list of RasterLayers including the red color pattern extracted from 5 Heliconius erato hydara dorsal forewings using patRegRGB.

Examples

```
## Not run:
data(rasterList_regRGB)
summary(rasterList_regRGB)
```

End(Not run)

redRes

Reduce the resolution of an image imported as a RasterStack by down-sampling.

Description

Reduce the resolution of an image imported as a RasterStack by downsampling.

Usage

```
redRes(image, resampleFactor)
```

Arguments

| image | RasterStack for downsampling. |
|----------------|-------------------------------|
| resampleFactor | Integer for downsampling. |

Value

Downsampled RasterStack

Examples

```
image <- raster::stack(system.file("extdata", "BC0077.jpg", package = "patternize"))
image_reduced <- redRes(image, 5)</pre>
```

sampleLandmarks Sample landmarks in an image.

Description

Sample landmarks in an image.

Usage

```
sampleLandmarks(sampleList, resampleFactor = NULL, crop = c(0, 0, 0, 0))
```

| sampleList | RasterStack or list of RasterStack objects as obtained by makeList. |
|----------------|---|
| resampleFactor | Integer for downsampling the image(s) used by redRes. |
| crop | Vector c(xmin, xmax, ymin, ymax) that specifies the pixel coordinates to crop the original image. |

Value

landmark matrix or landmark list

Examples

```
## Not run:
IDlist <- c('BC0077','BC0071')
prepath <- system.file("extdata", package = 'patternize')
extension <- '.jpg'
imageList <- makeList(IDlist, 'image', prepath, extension)
landmarkList <- sampleLandmarks(imageList)
## End(Not run)
```

| sampleRGB | Interactive function to sample RGB value from pixel or square area in |
|-----------|---|
| | an image. |

Description

Interactive function to sample RGB value from pixel or square area in an image.

Usage

```
sampleRGB(image, resampleFactor = NULL, crop = c(0, 0, 0, 0), type = "point")
```

Arguments

| image | Image imported as a RasterStack. |
|----------------|---|
| resampleFactor | Integer for downsampling used by redRes. |
| crop | Vector c(xmin, xmax, ymin, ymax) that specifies the pixel coordinates to crop the original image. |
| type | Set 'point' to extract RGB from a single point or 'area' to extract from a square area defined by setting two points (default = 'point'). |

Value

RGB vector

Examples

```
image <- raster::stack(system.file("extdata", "BC0077.jpg", package = "patternize"))
RGB <- sampleRGB(image, resampleFactor = 1)</pre>
```

setMask

Description

Interactive function to to draw an outline for masking.

Usage

```
setMask(summedRaster, IDlist, filename, ...)
```

Arguments

| summedRaster | Summed raster of extracted patterns. |
|--------------|---|
| IDlist | List of sample IDs. |
| filename | Name of file to which mask will be written. |
| | additional arguments for plotHeat function. |

Value

file

| sumRaster | This function sums the individual color pattern RasterLayes as ob- |
|-----------|--|
| | tained by the main patternize functions. |

Description

This function sums the individual color pattern RasterLayes as obtained by the main patternize functions.

Usage

```
sumRaster(rList, IDlist, type)
```

| rList | List of RasterLayers or list of RasterLayers for each k-means cluster. |
|--------|---|
| IDlist | List of sample IDs. |
| type | Type of rasterlist; 'RGB' or 'k' (result from RGB or k-means analysis, respectively). |

Examples

```
data(rasterList_lanRGB)
IDlist <- c('BC0077','BC0071','BC0050','BC0049','BC0004')
summedRaster <- sumRaster(rasterList_lanRGB, IDlist, type = 'RGB')
data(rasterList_lanK)
IDlist <- c('BC0077','BC0071','BC0050','BC0049','BC0004')
summedRasterList <- sumRaster(rasterList_lanK, IDlist, type = 'k')</pre>
```

Index

* datasets imageList, 12 landmarkArray, 15 landmarkList, 16 rasterList_lanK, 55 rasterList_lanRGB, 55 rasterList_regK, 56 rasterList_regRGB, 56 alignLan, 3 alignReg, 4 colorChecker, 5 colorChecker_customGray, 6 colorChecker_half, 7 computeTransform, 3, 25, 27, 28, 30, 32, 51 createPhenotype, 8 createTarget, 10 extdata, 11 GMM, 12 GMMImage, 12 imageList, 12 imager, 49, 51 kImage, 13 kImageHSV, 13 kmeans, 13 lanArray, 14 landmarkArray, 15 landmarkList, 16 1da, 39 makeList, 3, 6-8, 15, 16, 18-20, 24, 26, 28, 30, 32, 57 maskOutline, 17 Morpho, 14 niftyreg, 4, 40-47, 49, 51

patArea, 19 patGMM, 21 patK, 22 patK_HSV, 23 patLanHSV, 24 patLanK, 26 patLanK_HSV, 27 patLanRGB, 29 patLanW, 31 patPCA, 33, 53 patRDA, 37 patRegHSV, 40 patRegK, 41 patRegK_HSV, 43 patRegRGB, 45 patRegW, 46 patternize, 49 patternize-package (patternize), 49 plotHeat, 51 plotRasterstackAsImage, 54 prcomp, 33, 34, 36, 50 procSym, 50, 51

sampleLandmarks, 57
sampleRGB, 58
setMask, 59
stack, 51
sumRaster, 59