Package 'colordistance'

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Title Distance Metrics for Image Color Similarity
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Description Loads and displays images, selectively masks specified background colors, bins pixels by color using either data-dependent or automatically generated color bins, quantitatively measures color similarity among images using one of several distance metrics for comparing pixel color clusters, and clusters images by object color similarity. Uses CIELAB, RGB, or HSV color spaces. Originally written for use with organism coloration (reef fish color diversity, butterfly mimicry, etc), but easily applicable for any image set.
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chiso	qDistance Chi-square distance between vectors	

Description

Computes the chi-squared distance between each element of a pair of vectors which must be of the same length. Good for comparing color histograms if you don't want to weight by color similarity. Probably hugely redundant; alas.

Usage

```
chisqDistance(a, b)
```

Arguments

- a Numeric vector.
- b Numeric vector; must be the same length as a.

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Value

Chi-squared distance, $(a - b)^2/(a + b)$, between vectors a and b. If one or both elements are NA/NaN, contribution is counted as a 0.

Examples

```
colordistance:::chisqDistance(rnorm(10), rnorm(10))
```

colorDistance

Sum of Euclidean distances between color clusters

Description

Calculates the Euclidean distance between each pair of points in two dataframes as returned by extractClusters or getImageHist and returns the sum of the distances.

Usage

```
colorDistance(T1, T2)
```

Arguments

T1 Dataframe (especially a dataframe as returned by extractClusters() or getImageHist(),

but first three columns must be coordinates).

T2 Another dataframe like T1.

Value

Sum of Euclidean distances between each pair of points (rows) in the provided dataframes.

```
cluster.list <- colordistance::getHistList(system.file("extdata",
  "Heliconius/Heliconius_B", package="colordistance"), lower=rep(0.8, 3),
  upper=rep(1, 3))
colordistance:::colorDistance(cluster.list[[1]], cluster.list[[2]])</pre>
```

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Average 3D color histograms by subdirectory

Description

Calculates color histograms for images in immediate subdirectories of a folder, and averages histograms for images in the same subdirectory.

Usage

```
combineClusters(folder, method = "mean", ...)
```

Arguments

folder Path to the folder containing subdirectories of images. Must be a character vec-

tor.

method Method for combining color histograms. Default is "mean", but other generic

functions ("median", "sum", etc) will work. String is evaluated using "eval"

so any appropriate R function is accepted.

... Additional arguments passed to getHistList, including number of bins, HSV

flag, etc.

Examples

```
combined_clusters <- colordistance::combineClusters(system.file("extdata",
"Heliconius", package="colordistance"), method="median", bins=2,
lower=rep(0.8, 3), upper=rep(1, 3))</pre>
```

combineList

Combine a list of cluster features into a single cluster set

Description

Combine a list of cluster features as returned by getHistList according to the specified method.

Usage

```
combineList(hist_list, method = "mean")
```

Arguments

hist_list A list of cluster dataframes as returned by getHistList.

method Method for combining color histograms. Default is "mean", but other generic

functions ("median", "sum", etc) will work. String is evaluated using "eval"

so any appropriate R function is accepted.

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Note

While the function can also accept clusters generated using kmeans (getKMeansList followed by extractClusters), this is not recommended, as kmeans does not provide explicit analogous pairs of clusters, and clusters are combined by row number (all row 1 clusters are treated as analogous, etc). Color histograms are appropriate because the bins are defined the same way for each image.

Examples

```
hist_list <- getHistList(system.file("extdata", "Heliconius/Heliconius_A",
package="colordistance"), lower=rep(0.8, 3), upper=rep(1, 3))
median_clusters <- combineList(hist_list, method="median")</pre>
```

convertColorSpace

Convert between color spaces

Description

Wrapper for convertColor that builds in random sampling, error messages, and removes default illuminant (D65) to enforce manual specification of a reference white.

Usage

```
convertColorSpace(color.coordinate.matrix, from = "sRGB", to = "Lab",
   sample.size = 1e+05, from.ref.white, to.ref.white)
```

Arguments

color.coordinate.matrix

A color coordinate matrix with rows as colors and channels as columns. If a color histogram (e.g. as returned by getImageHist) is passed, the 'Pct' column is ignored.

from, to Input and output color spaces, passed to convertColor. See details.

sample.size Number of pixels to be randomly sampled from filtered pixel array for con-

version. If not numeric or larger than number of colors provided (i.e. cluster matrix), all colors are converted. See details.

matrix), an colors are converted. See details

from.ref.white, to.ref.white

Reference whites passed to convertColor. Unlike convertColor, no default is provided. See details for explanation of different reference whites.

Details

```
Color spaces are all passed to convertColor, and can be any of: "XYZ", "sRGB", "Apple RGB", "CIE RGB", "Lab", or "Luv".
```

Lab and Luv color spaces are approximately perceptually uniform, meaning they usually do the best job of reflecting intuitive color distances without the non-linearity problems of more familiar RGB

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spaces. However, because they describe object colors, they require a reference 'white light' color (dimly and brightly lit photographs of the same object will have very different RGB palettes, but similar Lab palettes if appropriate white references are used). The idea here is that the apparent colors in an image depend not just on the "absolute" color of an object, but also on the available light in the scene. There are seven CIE standardized illuminants available in colordistance (A, B, C, E, and D50, D55, and D65), but the most common are:

- "A": Standard incandescent lightbulb
- "D65": Average daylight
- "D50": Direct sunlight

Color conversions will be highly dependent on the reference white used, which is why no default is provided. Users should look into standard illuminants to choose an appropriate reference for a dataset.

The conversion from RGB to a standardized color space (XYZ, Lab, or Luv) is approximate, non-linear, and relatively time-consuming. Converting a large number of pixels can be computationally expensive, so convertColorSpace will randomly sample a specified number of rows to reduce the time. The default sample size, 100,000 rows, takes about 5 seconds convert from sRGB to Lab space on an early 2015 Macbook with 8 GB of RAM. Time scales about linearly with number of rows converted.

Value

A 3- or 4-column matrix depending on whether color.coordinate.matrix included a 'Pct' column (as from getImageHist), with one column per channel.

```
# Convert a single RGB triplet and then back convert it
rgb_color <- c(0, 1, 0)
lab_color <- colordistance::convertColorSpace(rgb_color,</pre>
from="sRGB", to="Lab", to.ref.white="D65")
rgb_again <- colordistance::convertColorSpace(lab_color,</pre>
from="Lab", to="sRGB", from.ref.white="D65")
# Convert pixels from loadImage() function
img <- colordistance::loadImage(system.file("extdata",</pre>
"Heliconius/Heliconius_B/Heliconius_07.jpeg", package="colordistance"))
lab_pixels <- colordistance::convertColorSpace(img$filtered.rgb.2d,</pre>
from="sRGB", to="XYZ", sample.size=5000)
# Convert clusters
img <- colordistance::loadImage(system.file("extdata",</pre>
"Heliconius/Heliconius_B/Heliconius_07.jpeg", package="colordistance"))
img_hist <- colordistance::getImageHist(img, bins=2, plotting=FALSE)</pre>
lab_clusters <- colordistance::convertColorSpace(img_hist, to.ref.white="D55")
```

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EMDistance

Earth mover's distance between two sets of color clusters

Description

Calculates the Earth mover's distance (briefly, the amount of work required to move the data from one distribution to resemble the other distribution, or the amount of "dirt" you have to shovel weighted by how far you have to shovel it). Accounts for both color disparity and size disparity. Recommended unless binAvg is off for histogram generation.

Usage

```
EMDistance(T1, T2)
```

Arguments

T1 Dataframe (especially a dataframe as returned by link{extractClusters} or

getImageHist, but first three columns must be coordinates).

T2 Another dataframe like T1.

Value

Earth mover's distance between the two dataframes (metric of overall bin similarity for a pair of 3-dimensional histograms).

Examples

```
## Not run:
cluster.list <- colordistance::getHistList(system.file("extdata",
   "Heliconius/Heliconius_B", package="colordistance"), lower=rep(0.8, 3),
   upper=rep(1, 3))
colordistance:::EMDistance(cluster.list[[1]], cluster.list[[2]])
## End(Not run)</pre>
```

exportTree

Export a distance matrix as a tree object

Description

Converts a symmetrical distance matrix to a tree and saves it in newick format. Uses hclust to form clusters.

Usage

```
exportTree(getColorDistanceMatrixObject, file, return.tree = FALSE)
```

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Arguments

getColorDistanceMatrixObject

A distance matrix, especially as returned by getColorDistanceMatrix, but any

numeric symmetrical matrix will work.

file Character vector of desired filename for saving tree. Should end in ".newick".

return. tree Logical. Should the tree object be returned to the working environment in addi-

tion to being saved as a file?

Value

Newick tree saved in specified location and as.phylo tree object if return.tree=TRUE.

Examples

```
clusterList <- colordistance::getHistList(dir(system.file("extdata",
  "Heliconius/", package="colordistance"), full.names=TRUE), lower=rep(0.8, 3),
upper=rep(1, 3))
CDM <- colordistance::getColorDistanceMatrix(clusterList, method="emd",
plotting=FALSE)

# Tree is both saved in current working directory and stored in
# heliconius_tree variable
heliconius_tree <- colordistance::exportTree(CDM,
  "./HeliconiusColorTree.newick", return.tree=TRUE)</pre>
```

extractClusters

Extract cluster values and sizes from kmeans fit objects

Description

Extract a list of dataframes with the same format as those returned by getHistList, where each dataframe has 3 color attributes (R, G, B or H, S, V) and a size attribute (Pct) for every cluster.

Usage

```
extractClusters(getKMeansListObject, ordering = TRUE,
    normalize = FALSE)
```

Arguments

getKMeansListObject

A list of kmeans fit objects (especially as returned by getKMeansList).

ordering Logical. Should clusters by reordered by color similarity? If TRUE, the Hungar-

ian algorithm via solve_LSAP is applied to find the minimum sum of Euclidean distances between color pairs for every pair of cluster objects and colors are

reordered accordingly.

normalize Logical. Should each cluster be normalized to show R:G:B or H:S:V ratios

rather than absolute values? Can be helpful for inconsistent lighting, but reduces

variation. See normalizeRGB.

Value

A list of dataframes (same length as input list), each with 4 columns: R, G, B (or H, S, V) and Pct (cluster size), with one row per cluster.

Note

Names are inherited from the list passed to the function.

Examples

```
clusterList <- colordistance::getKMeansList(system.file("extdata",
"Heliconius/Heliconius_A", package="colordistance"), bins=3)
colordistance::extractClusters(clusterList)</pre>
```

getColorDistanceMatrix

Distance matrix for a list of color cluster sets

Description

Calculates a distance matrix for a list of color cluster sets as returned by extractClusters or getHistList based on the specified distance metric.

Usage

```
getColorDistanceMatrix(cluster.list, method = "emd",
  ordering = "default", size.weight = 0.5, color.weight = 0.5,
  plotting = TRUE, ...)
```

Arguments

cluster.list A list of identically sized dataframes with 4 columns each (R, G, B, Pct or H, S,

V, Pct) as output by extractClusters or getHistList.

method One of four possible comparison methods for calculating the color distances:

"emd" (uses EMDistance, recommended), "chisq" (uses chisqDistance), "color.dist"

(uses colorDistance; not appropriate if binAvg=F), or "weighted.pairs"

(weightedPairsDistance).

ordering Logical if not left as "default". Should the color clusters in the list be reordered

to minimize the distances between the pairs? If left as default, ordering depends on distance method: "emd" and "chisq" do not order clusters ("emd" orders on a case-by-case in the EMDistance function itself and reordering by size similarity would make chi-squared meaningless); "color.dist" and "weighted.pairs" use ordering. To override defaults, set to either T (for ordering) or F (for no

ordering).

size.weight Same as in weightedPairsDistance.

color.weight Same as in weightedPairsDistance.

plotting Logical. Should a heatmap of the distance matrix be displayed once the function

finishes running?

... Additional arguments passed on to heatmapColorDistance.

Details

Each cell represents the distance between a pair of color cluster sets as measured using either chi-squared distance (cluster size only), earth mover's distance (size and color), weighted pairs (size and color with user-specified weights for each), or color distance (Euclidean distance between clusters as 3-dimensional - RGB or HSV - color coordinates).

Earth mover's distance is recommended unless binAvg is set to false during cluster list generation (in which case all paired bins will have the same colors across datasets), in which case chi-squared is recommended. Weighted pairs or color distance may be appropriate depending on the question, but generally give poorer results.

Value

A distance matrix of image distance scores (the scales vary depending on the distance metric chosen, but for all four methods, higher scores = more different).

```
## Not run:
cluster.list <- colordistance::getHistList(c(system.file("extdata",</pre>
"Heliconius/Heliconius_A", package="colordistance"), system.file("extdata",
"Heliconius/Heliconius_B", package="colordistance")), lower=rep(0.8, 3),
upper=rep(1, 3))
# Default values - recommended!
colordistance::getColorDistanceMatrix(cluster.list, main="EMD")
# Without plotting
colordistance::getColorDistanceMatrix(cluster.list, plotting=FALSE)
# Use chi-squared instead
colordistance::getColorDistanceMatrix(cluster.list, method="chisq", main="Chi-squared")
# Override ordering (throws a warning if you're trying to do this with
# chisq!)
colordistance::getColorDistanceMatrix(cluster.list, method="chisq",
ordering=TRUE, main="Chi-squared w/ ordering")
# Specify high size weight/low color weight for weighted pairs
colordistance::getColorDistanceMatrix(cluster.list, method="weighted.pairs",
color.weight=0.1, size.weight=0.9, main="Weighted pairs")
# Color distance only
colordistance::getColorDistanceMatrix(cluster.list, method="color.dist",
ordering=TRUE, main="Color distance only")
```

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

getHistColors

Vector of hex colors for histogram bin coloration

Description

Gets a vector of colors for plotting histograms from getImageHist in helpful ways.

Usage

```
getHistColors(bins, hsv = FALSE)
```

Arguments

bins Number of bins for each channel OR a vector of length 3 with bins for each

channel. Bins = 3 will result in 3^3 = 27 bins; bins = c(2, 2, 3) will result in 2 *

2 * 3 = 12 bins (2 red, 2 green, 3 blue), etc.

hsv Logical. Should HSV be used instead of RGB?

Value

A vector of hex codes for bin colors.

Examples

```
colordistance:::getHistColors(bins = 3)
colordistance:::getHistColors(bins = c(8, 3, 3), hsv = TRUE)
```

getHistList

Generate a list of cluster sets for multiple images

Description

Applies getImageHist to every image in a provided set of image paths and/or directories containing images.

Usage

```
getHistList(images, bins = 3, bin.avg = TRUE, lower = c(0, 0.55, 0), upper = c(0.24, 1, 0.24), norm.pix = FALSE, plotting = FALSE, pausing = TRUE, hsv = FALSE, title = "path", img.type = FALSE, bounds = c(0, 1))
```

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Arguments

images	Character vector of directories, image paths, or both.
bins	Number of bins for each channel OR a vector of length 3 with bins for each channel. Bins=3 will result in $3^3 = 27$ bins; bins=c(2, 2, 3) will result in $2^2^3=12$ bins (2 red, 2 green, 3 blue), etc.
bin.avg	Logical. Should the returned color clusters be the average of the pixels in that bin (bin.avg=TRUE) or the center of the bin (FALSE)? If a bin is empty, the center of the bin is returned as the cluster color regardless.
lower	RGB or HSV triplet specifying the lower bounds for background pixels. Default upper and lower bounds are set to values that work well for a bright green background (RGB [0, 1, 0]).
upper	RGB or HSV triplet specifying the upper bounds for background pixels. Default upper and lower bounds are set to values that work well for a bright green background (RGB [0, 1, 0]). Determining these bounds may take some trial and error, but the following bounds may work for certain common background colors:
	• Black: lower=c(0, 0, 0); upper=c(0.1, 0.1, 0.1)
	• White: lower=c(0.8, 0.8, 0.8); upper=c(1, 1, 1)
	• Green: lower=c(0, 0.55, 0); upper=c(0.24, 1, 0.24)
	• Blue: lower=c(0, 0, 0.55); upper=c(0.24, 0.24, 1)
	If no background filtering is needed, set bounds to some non-numeric value (NULL, FALSE, "off", etc); any non-numeric value is interpreted as NULL.
norm.pix	Logical. Should RGB or HSV cluster values be normalized using normalizeRGB?
plotting	Logical. Should the histogram generated for each image be displayed?
pausing	Logical. If plotting=T, should the function pause between graphing and wait for user to hit [enter] before continuing? Useful for data/histogram inspection.
hsv	Logical. Should HSV be used instead of RGB?
title	String for what the title the plots if plotting is on; defaults to the image name.
img.type	Logical. Should the file extension for the images be retained when naming the output list elements? If FALSE, just the image name is used (so "Heliconius_01.png" becomes "Heliconius_01").
bounds	Upper and lower limits for the channels; R reads in images with intensities on a 0-1 scale, but 0-255 is common.

Value

A list of getImageHist dataframes, 1 per image, named by image name.

Note

For every image, the pixels are binned according to the specified bin breaks. By providing the bounds for the bins rather than letting an algorithm select centers (as in getKMeansList), clusters of nearly redundant colors are avoided.

So you don't end up with, say, 3 nearly-identical yellow clusters which are treated as unrelated just because there's a lot of yellow in your image; you just get a very large yellow cluster and empty non-yellow bins.

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Examples

```
## Not run:
# Takes >10 seconds if you run all examples
clusterList <- colordistance::getHistList(system.file("extdata",
   "Heliconius/Heliconius_B", package="colordistance"), upper = rep(1, 3),
lower = rep(0.8, 3))

clusterList <- colordistance::getHistList(c(system.file("extdata",
   "Heliconius/Heliconius_B", package="colordistance"), system.file("extdata",
   "Heliconius/Heliconius_A", package="colordistance")), pausing = FALSE,
   upper = rep(1, 3), lower = rep(0.8, 3))

clusterList <- colordistance::getHistList(system.file("extdata",
   "Heliconius/Heliconius_B", package = "colordistance"), plotting = TRUE,
   upper = rep(1, 3), lower = rep(0.8, 3))

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

getImageHist

Generate a 3D histogram based on color distribution in an image

Description

Computes a histogram in either RGB or HSV colorspace by sorting pixels into a specified number of bins.

Usage

```
getImageHist(image, bins = 3, bin.avg = TRUE, defaultClusters = NULL, lower = c(0, 0.55, 0), upper = c(0.24, 1, 0.24), as.vec = FALSE, norm.pix = FALSE, plotting = TRUE, hsv = FALSE, title = "path", bounds = c(0, 1), ...)
```

Arguments

image Path to a valid image (PNG or JPG) or a loadImage object.

Number of bins for each channel OR a vector of length 3 with bins for each

channel. Bins=3 will result in $3^3 = 27$ bins; bins=c(2, 2, 3) will result in

2*2*3=12 bins (2 red, 2 green, 3 blue), etc.

bin.avg Logical. Should the returned color clusters be the average of the pixels in that

bin (bin.avg=TRUE) or the center of the bin (FALSE)? If a bin is empty, the center

of the bin is returned as the cluster color regardless.

defaultClusters

Optional dataframe of default color clusters to be returned when a bin is empty. If NULL, the geometric centers of the bins are used.

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lower RGB or HSV triplet specifying the lower bounds for background pixels. Default upper and lower bounds are set to values that work well for a bright green

background (RGB [0, 1, 0]).

RGB or HSV triplet specifying the upper bounds for background pixels. Default upper and lower bounds are set to values that work well for a bright green background (RGB [0, 1, 0]). Determining these bounds may take some trial

and error, but the following bounds may work for certain common background

colors:

• Black: lower=c(0, 0, 0); upper=c(0.1, 0.1, 0.1)

• White: lower=c(0.8, 0.8, 0.8); upper=c(1, 1, 1)

• Green: lower=c(0, 0.55, 0); upper=c(0.24, 1, 0.24)

• Blue: lower=c(0, 0, 0.55); upper=c(0.24, 0.24, 1)

If no background filtering is needed, set bounds to some non-numeric value (NULL, FALSE, "off", etc); any non-numeric value is interpreted as NULL.

Logical. Should the bin sizes just be returned as a vector? Much faster if only as.vec

using chisqDistance for comparison metric.

norm.pix Logical. Should RGB or HSV cluster values be normalized using normalizeRGB?

plotting Logical. Should a histogram of the bin colors and sizes be plotted?

hsv Logical. Should HSV be used instead of RGB?

title String for what to title the plots if plotting is on; defaults to the image name.

Upper and lower limits for the channels; R reads in images with intensities on a bounds

0-1 scale, but 0-255 is common.

Optional arguments passed to the barplot function.

Details

If you choose 2 bins for each color channel, then each of R, G, and B will be divided into 2 bins each, for a total of $2^3 = 8$ bins.

Once all pixels have been binned, the function will return either the size of each bin, either in number of pixels or fraction of total pixels, and the color of each bin, either as the geometric center of the bin or as the average color of all pixels assigned to it.

For example, if you input an image of a red square and used 8 bins, all red pixels (RGB triplet of [1, 0, 0]) would be assigned to the bin with R bounds (0.5, 1], G bounds [0, 0.5) and B bounds [0, 0.5). The average color of the bin would be [0.75, 0.25, 0.25], but the average color of the pixels assigned to that bin would be [1, 0, 0]. The latter option is obviously more informative, but takes longer (about 1.5-2x longer depending on the images).

Value

A vector or dataframe (depending on whether as.vec=T) of bin sizes and color values.

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Examples

```
# generate HSV histogram for a single image
colordistance::getImageHist(system.file("extdata",
   "Heliconius/Heliconius_B/Heliconius_07.jpeg", package="colordistance"),
upper=rep(1, 3), lower=rep(0.8, 3), bins=c(8, 3, 3), hsv=TRUE, plotting=TRUE)
# generate RGB histogram
colordistance::getImageHist(system.file("extdata",
   "Heliconius/Heliconius_B/Heliconius_07.jpeg", package="colordistance"),
upper=rep(1, 3), lower=rep(0.8, 3), bins=2)
```

getImagePaths

Fetch paths to all valid images in a given directory

Description

Find all valid image paths (PNG and JPG) in a directory (does not search subdirectories). Will recover any image ending in .PNG, .JPG, or .JPEG, case-insensitive.

Usage

```
getImagePaths(path)
```

Arguments

path

Path to directory in which to search for images. Absolute or relative filepaths are fine.

Value

A vector of absolute filepaths to JPG and PNG images in the given directory.

Note

In the event that no compatible images are found in the directory, it returns a message to that effect instead of an empty vector.

```
im.dir <- colordistance::getImagePaths(system.file("extdata",
"Heliconius/Heliconius_A", package="colordistance"))
## Not run:
im.dir <- colordistance::getImagePaths("some/nonexistent/directory")
## End(Not run)
im.dir <- colordistance::getImagePaths(getwd())</pre>
```

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Fit pixels to clusters using KMeans clustering

Description

Uses KMeans clustering to determine color clusters that minimize the sum of distances between pixels and their assigned clusters. Useful for parsing common color motifs in an object.

Usage

```
getKMeanColors(path, n = 10, sample.size = 20000, plotting = TRUE, lower = c(0, 0.55, 0), upper = c(0.24, 1, 0.24), iter.max = 50, nstart = 5, return.clust = TRUE, color.space = "rgb", from = "sRGB", ref.white)
```

Arguments

path	Path to an image (JPG or PNG).
n	Number of KMeans clusters to fit. Unlike getImageHist, this represents the actual final number of bins, rather than the number of breaks in each channel.
sample.size	Number of pixels to be randomly sampled from filtered pixel array for performing fit. If set to FALSE, all pixels are fit, but this can be time-consuming, especially for large images.
plotting	Logical. Should the results of the KMeans fit (original image + histogram of colors and bin sizes) be plotted?
lower	RGB triplet specifying the lower bounds for background pixels. Default upper and lower bounds are set to values that work well for a bright green background (RGB [0, 1, 0]).
upper	RGB triplet specifying the upper bounds for background pixels. Default upper and lower bounds are set to values that work well for a bright green background (RGB [0, 1, 0]). Determining these bounds may take some trial and error, but the following bounds may work for certain common background colors:
	 Black: lower=c(0, 0, 0); upper=c(0.1, 0.1, 0.1) White: lower=c(0.8, 0.8, 0.8); upper=c(1, 1, 1) Green: lower=c(0, 0.55, 0); upper=c(0.24, 1, 0.24) Blue: lower=c(0, 0, 0.55); upper=c(0.24, 0.24, 1)
	If no background filtering is needed, set bounds to some non-numeric value (NULL, FALSE, "off", etc); any non-numeric value is interpreted as NULL.
iter.max	Inherited from kmeans. The maximum number of iterations allowed.
nstart	Inherited from kmeans. How many random sets should be chosen?
return.clust	Logical. Should clusters be returned? If FALSE, results are plotted but not returned.
color.space	The color space ("rgb", "hsv", or "lab") in which to cluster pixels.

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from Display color space of image if clustering in CIE Lab space, probably either

"sRGB" or "Apple RGB", depending on your computer.

ref.white The reference white passed to convertColorSpace; must be specified if using

CIE Lab space. See convertColorSpace.

Value

A kmeans fit object.

Examples

```
colordistance::getKMeanColors(system.file("extdata",
  "Heliconius/Heliconius_B/Heliconius_07.jpeg", package="colordistance"), n=3,
  return.clust=FALSE, lower=rep(0.8, 3), upper=rep(1, 3))
```

getKMeansList

Get KMeans clusters for every image in a set

Description

Performs getKMeanColors on every image in a set of images and returns a list of kmeans fit objects, where each dataframe contains the RGB coordinates of the clusters and the percentage of pixels in the image assigned to that cluster.

Usage

```
getKMeansList(images, bins = 10, sample.size = 20000,
  plotting = FALSE, lower = c(0, 0.55, 0), upper = c(0.24, 1, 0.24),
  iter.max = 50, nstart = 5, img.type = FALSE, color.space = "rgb",
  from = "sRGB", ref.white)
```

Arguments

images	A character vector of directories, image paths, or a combination of both. Takes either absolute or relative filepaths.
bins	Number of KMeans clusters to fit. Unlike getImageHist, this represents the actual final number of bins, rather than the number of breaks in each channel.
sample.size	Number of pixels to be randomly sampled from filtered pixel array for performing fit. If set to FALSE, all pixels are fit, but this can be time-consuming, especially for large images.
plotting	Logical. Should the results of the KMeans fit (original image + histogram of colors and bin sizes) be plotted for each image?
lower	RGB triplet specifying the lower bounds for background pixels. Default upper

and lower bounds are set to values that work well for a bright green background

(RGB [0, 1, 0]).

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u	a	p	e	r

RGB triplet specifying the upper bounds for background pixels. Default upper and lower bounds are set to values that work well for a bright green background (RGB [0, 1, 0]). Determining these bounds may take some trial and error, but the following bounds may work for certain common background colors:

Black: lower=c(0, 0, 0); upper=c(0.1, 0.1, 0.1)
White: lower=c(0.8, 0.8, 0.8); upper=c(1, 1, 1)
Green: lower=c(0, 0.55, 0); upper=c(0.24, 1, 0.24)

• Blue: lower=c(0, 0, 0.55); upper=c(0.24, 0.24, 1)

If no background filtering is needed, set bounds to some non-numeric value (NULL, FALSE, "off", etc); any non-numeric value is interpreted as NULL.

iter.max Inherited from kmeans. The maximum number of iterations allowed.

nstart Inherited from kmeans. How many random sets should be chosen?

img.type Logical. Should the image extension (.PNG or .JPG) be retained in the list

names?

color.space The color space ("rgb", "hsv", or "lab") in which to cluster pixels.

from Original color space of images if clustering in CIE Lab space, probably either

"sRGB" or "Apple RGB", depending on your computer.

ref.white The reference white passed to convertColorSpace; must be specified if using

CIE Lab space. See convertColorSpace.

Value

A list of kmeans fit objects, where the list element names are the original image names.

Examples

```
## Not run:
# Takes a few seconds to run
kmeans_list <- colordistance::getKMeansList(dir(system.file("extdata",
"Heliconius/", package="colordistance"), full.names=TRUE), bins=3,
lower=rep(0.8, 3), upper=rep(1, 3), plotting=TRUE)
## End(Not run)</pre>
```

getLabHist

Generate a 3D histogram based on CIE Lab color coordinates in an image

Description

Computes a histogram in CIE Lab color space by sorting pixels into specified bins.

getLabHist 19

Usage

```
getLabHist(image, bins = 3, sample.size = 10000, ref.white, from = "sRGB", bin.avg = TRUE, as.vec = FALSE, plotting = TRUE, lower = c(0, 0.55, 0), upper = c(0.24, 1, 0.24), title = "path", a.bounds = c(-128, 127), b.bounds = c(-128, 127), ...)
```

Arguments

image Path to a valid image (PNG or JPG) or a loadImage object.

bins Number of bins for each channel OR a vector of length 3 with bins for each

channel. Bins = 3 will result in 3^3 = 27 bins; bins = c(2, 2, 3) will result in 2 *

2 * 3 = 12 bins (2 L, 2 a, 3 b), etc.

sample.size Numeric. How many pixels should be randomly sampled from the non-background

part of the image and converted into CIE Lab coordinates? If non-numeric, all

pixels will be converted, but this can be very slow (see details).

ref.white Reference white passed to convertColorSpace. Unlike convertColor, no de-

fault is provided. See details for explanation of different reference whites.

from Original color space of image, probably either "sRGB" or "Apple RGB", de-

pending on your computer.

bin.avg Logical. Should the returned color clusters be the average of the pixels in that

bin (bin.avg=TRUE) or the center of the bin (FALSE)? If a bin is empty, the center

of the bin is returned as the cluster color regardless.

as.vec Logical. Should the bin sizes just be returned as a vector? Much faster if only

using chisqDistance for comparison metric.

plotting Logical. Should a histogram of the bin colors and sizes be plotted?

lower, upper RGB or HSV triplets specifying the lower and upper bounds for ba

RGB or HSV triplets specifying the lower and upper bounds for background pixels. Default upper and lower bounds are set to values that work well for a bright green background (RGB [0, 1, 0]). Determining these bounds may take some trial and error, but the following bounds may work for certain common

background colors:

• Black: lower=c(0, 0, 0); upper=c(0.1, 0.1, 0.1)

• White: lower=c(0.8, 0.8, 0.8); upper=c(1, 1, 1)

• Green: lower=c(0, 0.55, 0); upper=c(0.24, 1, 0.24)

• Blue: lower=c(0, 0, 0.55); upper=c(0.24, 0.24, 1)

If no background filtering is needed, set bounds to some non-numeric value (NULL, FALSE, "off", etc); any non-numeric value is interpreted as NULL.

title String for what the title the plot if plotting is on; defaults to the image name.

a.bounds, b.bounds

Numeric ranges for the a (green-red) and b (blue-yellow) channels of Lab color space. Technically, a and b have infinite range, but in practice nearly all values fall between -128 and 127 (the default). Many images will have an even narrower range than this, depending on the lighting conditions and conversion; setting narrower ranges will result in finer-scale binning, without generating empty bins at the edges of the channels.

. . Additional arguments passed to barplot.

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Details

getLabHist uses convertColorSpace to convert pixels into CIE Lab coordinates, which requires a references white. There are seven CIE standardized illuminants available in colordistance (A, B, C, E, and D50, D55, and D65), but the most common are:

• "A": Standard incandescent lightbulb

• "D65": Average daylight

• "D50": Direct sunlight

Color conversions will be highly dependent on the reference white used, which is why no default is provided. Users should look into standard illuminants to choose an appropriate reference for a dataset.

The conversion from RGB to a standardized color space (XYZ, Lab, or Luv) is approximate, non-linear, and relatively time-consuming. Converting a large number of pixels can be computationally expensive, so convertColorSpace will randomly sample a specified number of rows to reduce the time. The default sample size, 10,000 rows, takes about 1 second to convert from sRGB to Lab space on an early 2015 Macbook with 8 GB of RAM. Time scales about linearly with number of rows converted.

Unlike RGB or HSV color spaces, the three channels of CIE Lab color space do not all range between 0 and 1; instead, L (luminance) is always between 0 and 100, and the a (green-red) and b (blue-yellow) channels generally vary between -128 and 127, but usually occupy a narrower range depending on the reference white. To achieve the best results, ranges for a and b should be restricted to avoid generating empty bins.

Value

A vector or dataframe (depending on whether as .vec = TRUE) of bin sizes and color coordinates.

Examples

```
path <- system.file("extdata", "Heliconius/Heliconius_B/Heliconius_07.jpeg", package="colordistance") getLabHist(path, ref.white = "D65", bins = c(2, 3, 3), lower = rep(0.8, 3), upper = rep(1, 3), sample.size = 1000, ylim = c(0, 1))
```

getLabHistList

Generate a list of cluster sets in CIE Lab color space

Description

Applies getLabHist to every image in a provided set of image paths and/or directories containing images.

getLabHistList 21

Usage

```
getLabHistList(images, bins = 3, sample.size = 10000, ref.white,
  from = "sRGB", bin.avg = TRUE, as.vec = FALSE, plotting = FALSE,
  pausing = TRUE, lower = c(0, 0.55, 0), upper = c(0.24, 1, 0.24),
  title = "path", a.bounds = c(-128, 127), b.bounds = c(-128, 127),
  ...)
```

Arguments

images Character vector of directories, image paths, or both.

Number of bins for each channel OR a vector of length 3 with bins for each channel. Bins = 3 will result in $3^3 = 27$ bins; bins = c(2, 2, 3) will result in 2 *

2 * 3 = 12 bins (2 L, 2 a, 3 b), etc.

sample.size Numeric. How many pixels should be randomly sampled from the non-background

part of the image and converted into CIE Lab coordinates? If non-numeric, all

pixels will be converted, but this can be very slow (see details).

ref.white Reference white passed to convertColorSpace. Unlike convertColor, no de-

fault is provided. See details for explanation of different reference whites.

from Original color space of image, probably either "sRGB" or "Apple RGB", de-

pending on your computer.

bin.avg Logical. Should the returned color clusters be the average of the pixels in that

bin (bin.avg=TRUE) or the center of the bin (FALSE)? If a bin is empty, the center

of the bin is returned as the cluster color regardless.

as.vec Logical. Should the bin sizes just be returned as a vector? Much faster if only

using chisqDistance for comparison metric.

plotting Logical. Should a histogram of the bin colors and sizes be plotted?

pausing Logical. If plotting=T, should the function pause between graphing and wait

for user to hit [enter] before continuing? Useful for data/histogram inspection.

lower, upper RGB or HSV triplets specifying the lower and upper bounds for background pixels. Default upper and lower bounds are set to values that work well for a

bright green background (RGB [0, 1, 0]). Determining these bounds may take some trial and error, but the following bounds may work for certain common

background colors:

• Black: lower=c(0, 0, 0); upper=c(0.1, 0.1, 0.1)

• White: lower=c(0.8, 0.8, 0.8); upper=c(1, 1, 1)

• Green: lower=c(0, 0.55, 0); upper=c(0.24, 1, 0.24)

• Blue: lower=c(0, 0, 0.55); upper=c(0.24, 0.24, 1)

If no background filtering is needed, set bounds to some non-numeric value (NULL, FALSE, "off", etc); any non-numeric value is interpreted as NULL.

title String for what the title the plot if plotting is on; defaults to the image name.

a.bounds, b.bounds

Numeric ranges for the a (green-red) and b (blue-yellow) channels of Lab color space. Technically, a and b have infinite range, but in practice nearly all values

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fall between -128 and 127 (the default). Many images will have an even narrower range than this, depending on the lighting conditions and conversion; setting narrower ranges will result in finer-scale binning, without generating empty bins at the edges of the channels.

... Additional arguments passed to barplot.

Details

getLabHist uses convertColorSpace to convert pixels into CIE Lab coordinates, which requires a references white. There are seven CIE standardized illuminants available in colordistance (A, B, C, E, and D50, D55, and D65), but the most common are:

• "A": Standard incandescent lightbulb

• "D65": Average daylight

• "D50": Direct sunlight

Color conversions will be highly dependent on the reference white used, which is why no default is provided. Users should look into standard illuminants to choose an appropriate reference for a dataset.

The conversion from RGB to a standardized color space (XYZ, Lab, or Luv) is approximate, nonlinear, and relatively time-consuming. Converting a large number of pixels can be computationally expensive, so convertColorSpace will randomly sample a specified number of rows to reduce the time. The default sample size, 10,000 rows, takes about 1 second to convert from sRGB to Lab space on an early 2015 Macbook with 8 GB of RAM. Time scales about linearly with number of rows converted.

Unlike RGB or HSV color spaces, the three channels of CIE Lab color space do not all range between 0 and 1; instead, L (luminance) is always between 0 and 100, and the a (green-red) and b (blue-yellow) channels generally vary between -128 and 127, but usually occupy a narrower range depending on the reference white. The exception is reference white A (standard incandescent lighting), which tends to have lower values when converting with convertColor.

Value

A list of getLabHist dataframes, 1 per image, named by image name.

```
images <- system.file("extdata", "Heliconius/Heliconius_B",
package="colordistance")

colordistance::getLabHistList(images, bins = 2, sample.size = 1000, ref.white
= "D65", plotting = TRUE, pausing = FALSE, lower = rep(0.8, 3), upper =
rep(1, 3), a.bounds = c(-100, 100), b.bounds = c(-127, 100), ylim = c(0, 1))</pre>
```

heatmapColorDistance Plot a heatmap of a distance matrix

Description

Plots a heatmap of a symmetrical distance matrix in order to visualize similarity/dissimilarity in scores. Values are clustered by similarity using hclust.

Usage

```
heatmapColorDistance(clusterList_or_matrixObject, main = NULL,
  col = "default", margins = c(6, 8), ...)
```

Arguments

clusterList_or_matrixObject

Either a list of identically sized dataframes with 4 columns each (3 color channels + Pct) as output by extractClusters or getHistList, or a symmetrical

distance matrix as output by getColorDistanceMatrix.

main Title for heatmap plot.

col Color scale for heatmap from low to high. Default is colorRampPalette(c("royalblue4", "ghostwhi

where pink is more dissimilar and blue is more similar.

margins Margins for column and row labels.

... Additional arguments passed on to heatmap. 2.

Value

Heatmap representation of distance matrix.

```
## Not run:
# Takes a few seconds to run
cluster.list <- colordistance::getHistList(dir(system.file("extdata",
   "Heliconius/", package="colordistance"), full.names=TRUE), lower=rep(0.8, 3),
upper=rep(1, 3))

CDM <- colordistance::getColorDistanceMatrix(cluster.list, plotting=FALSE)

colordistance::heatmapColorDistance(CDM, main="Heliconius color similarity")
colordistance::heatmapColorDistance(cluster.list,
   col=colorRampPalette(c("red", "cyan", "blue"))(n=299))

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

imageClusterPipeline Generate and plot a color distance matrix from a set of images

Description

Takes images, computes color clusters for each image, and calculates distance matrix/dendrogram from those clusters.

Usage

```
imageClusterPipeline(images, cluster.method = "hist",
 distance.method = "emd", lower = c(0, 140/255, 0),
 upper = c(60/255, 1, 60/255), hist.bins = 3, kmeans.bins = 27,
 bin.avg = TRUE, norm.pix = FALSE, plot.bins = FALSE,
 pausing = TRUE, color.space = "rgb", ref.white, from = "sRGB",
 bounds = c(0, 1), sample.size = 20000, iter.max = 50, nstart = 5,
 img.type = FALSE, ordering = "default", size.weight = 0.5,
 color.weight = 0.5, plot.heatmap = TRUE,
 return.distance.matrix = TRUE, save.tree = FALSE,
 save.distance.matrix = FALSE, a.bounds = c(-127, 128),
 b.bounds = c(-127, 128))
```

Arguments

images

Character vector of directories, image paths, or both.

cluster.method Which method for getting color clusters from each image should be used? Must be either "hist" (predetermined bins generated by dividing each channel with equidistant bounds; calls getHistList) or "kmeans" (determine clusters using kmeans fitting on pixels; calls getKMeansList).

distance.method

One of four possible comparison methods for calculating the color distances: "emd" (uses EMDistance, recommended), "chisq" (uses chisqDistance), "color.dist" (uses colorDistance; not appropriate if bin.avg=F), or "weighted.pairs" (weightedPairsDistance).

lower

RGB or HSV triplet specifying the lower bounds for background pixels. Default upper and lower bounds are set to values that work well for a bright green background (RGB [0, 1, 0]).

upper

RGB or HSV triplet specifying the upper bounds for background pixels. Default upper and lower bounds are set to values that work well for a bright green background (RGB [0, 1, 0]). Determining these bounds may take some trial and error, but the following bounds may work for certain common background colors:

- Black: lower=c(0, 0, 0); upper=c(0.1, 0.1, 0.1)
- White: lower=c(0.8, 0.8, 0.8); upper=c(1, 1, 1)
- Green: lower=c(0, 0.55, 0); upper=c(0.24, 1, 0.24)

• Blue: lower=c(0, 0, 0.55); upper=c(0.24, 0.24, 1)

If no background filtering is needed, set bounds to some non-numeric value (NULL, FALSE, "off", etc); any non-numeric value is interpreted as NULL.

Only applicable if cluster.method="hist". Number of bins for each channel OR a vector of length 3 with bins for each channel. Bins=3 will result in 3^3 = 27 bins; bins=c(2, 2, 3) will result in 2*2*3=12 bins (2 red, 2 green, 3 blue), etc.

Passed to getHistList.

kmeans.bins Only applicable if cluster.method="kmeans". Number of KMeans clusters to

 $fit. \ Unlike \ {\tt getImageHist}, this \ represents \ the \ actual \ final \ number \ of \ bins, \ rather$

than the number of breaks in each channel.

bin.avg Logical. Should the color clusters used for the distance matrix be the average of

the pixels in that bin (bin.avg=TRUE) or the center of the bin (FALSE)? If a bin is empty, the center of the bin is returned as the cluster color regardless. Only applicable if cluster.method="hist", since kmeans clusters are at the center

of their assigned pixel clouds by definition.

norm.pix Logical. Should RGB or HSV cluster values be normalized using normalizeRGB?

plot.bins Logical. Should the bins for each image be plotted as they are calculated?

pausing Logical. If plot.bins=TRUE, pause and wait for user keystroke before plotting

bins for next image?

color.space The color space ("rgb", "hsv", or "lab") in which to plot pixels.

ref.white The reference white passed to convertColorSpace; must be specified if using

color.space = "lab".

from Display color space of image if clustering in CIE Lab space, probably either

"sRGB" or "Apple RGB", depending on your computer.

bounds Upper and lower limits for the channels; R reads in images with intensities on a

0-1 scale, but 0-255 is common.

sample.size Only applicable if cluster.method="kmeans". Number of pixels to be ran-

domly sampled from filtered pixel array for performing fit. If set to FALSE, all pixels are fit, but this can be time-consuming, especially for large images.

Passed to getKMeansList.

iter.max Only applicable if cluster.method="kmeans". Inherited from kmeans. The

maximum number of iterations allowed during kmeans fitting. Passed to getKMeansList.

nstart Only applicable if cluster.method="kmeans". Inherited from kmeans. How

many random sets should be chosen? Passed to getKMeansList.

img. type Logical. Should file extensions be retained with labels?

ordering Logical if not left as "default". Should the color clusters in the list be reordered

to minimize the distances between the pairs? If left as default, ordering depends on distance method: "emd" and "chisq" do not order clusters ("emd" orders on a case-by-case in the EMDistance function itself and reordering by size similarity would make chi-squared meaningless); "color.dist" and "weighted.pairs" use ordering. To override defaults, set to either T (for ordering) or F (for no

ordering).

size.weight Weight of size similarity in determining overall score and ordering (if ordering=T).

color.weight Weight of color similarity in determining overall score and ordering (if ordering=T).

Color and size weights do not necessarily have to sum to 1.

plot.heatmap Logical. Should a heatmap of the distance matrix be plotted?

return.distance.matrix

Logical. Should the distance matrix be returned to the R environment or just

plotted?

save.tree Either logical or a filepath for saving the tree; default if set to TRUE is to save in

current working directory as "ColorTree.newick".

save.distance.matrix

Either logical or filepath for saving distance matrix; default if set to TRUE is to save in current working directory as "ColorDistanceMatrix.csv"

a.bounds, b.bounds

Passed to getLabHistList.Numeric ranges for the a (green-red) and b (blue-yellow) channels of Lab color space. Technically, a and b have infinite range, but in practice nearly all values fall between -128 and 127 (the default). Many images will have an even narrower range than this, depending on the lighting conditions and conversion; setting narrower ranges will result in finer-scale binning, without generating empty bins at the edges of the channels.

Value

Color distance matrix, heatmap, and saved distance matrix and tree files if saving is TRUE.

Note

This is the fastest way to get a distance matrix for color similarity starting from a folder of images. Essentially, it just calls in a series of other package functions in order: input images -> getImagePaths -> getHistList or getKMeansList followed by extractClusters -> getColorDistanceMatrix -> plotting -> return/save distance matrix. Sort of railroads you, but good for testing different combinations of clustering methods and distance metrics.

```
## Not run:
colordistance::imageClusterPipeline(dir(system.file("extdata", "Heliconius/",
package="colordistance"), full.names=TRUE), color.space="hsv", lower=rep(0.8,
3), upper=rep(1, 3), cluster.method="hist", distance.method="emd",
hist.bins=3, plot.bins=TRUE, save.tree="example_tree.newick",
save.distance.matrix="example_DM.csv")

## End(Not run)
```

27 loadImage

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Import image and generate filtered 2D pixel array(s)

Description

Imports a single image and returns a list with the original image as a 3D array, a 2D matrix with background pixels removed, and the absolute path to the original image.

Usage

```
loadImage(path, lower = c(0, 0.55, 0), upper = c(0.24, 1, 0.24),
 hsv = TRUE, CIELab = FALSE, sample.size = 1e+05,
 ref.white = NULL)
```

Arguments

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path	Path to image (a ctring)
patii	i ani to illiage (a suma.

lower RGB or HSV triplet specifying the lower bounds for background pixels. De-

fault upper and lower bounds are set to values that work well for a bright green

background (RGB [0, 1, 0]).

upper RGB or HSV triplet specifying the upper bounds for background pixels. De-

fault upper and lower bounds are set to values that work well for a bright green background (RGB [0, 1, 0]). Determining these bounds may take some trial and error, but the following bounds may work for certain common background

colors:

• Black: lower=c(0, 0, 0); upper=c(0.1, 0.1, 0.1)

• White: lower=c(0.8, 0.8, 0.8); upper=c(1, 1, 1)

• Green: lower=c(0, 0.55, 0); upper=c(0.24, 1, 0.24)

• Blue: lower=c(0, 0, 0.55); upper=c(0.24, 0.24, 1)

If no background filtering is needed, set bounds to some non-numeric value (NULL, FALSE, "off", etc); any non-numeric value is interpreted as NULL.

Logical. Should HSV pixel array also be calculated? Setting to FALSE will shave

some time off the analysis, but not much (a few microseconds per image). Logical. Should CIEL*a*b color space pixels be calculated from RGB? Re-

quires specification of a reference white (see details).

Number of pixels to be randomly sampled from filtered pixel array for conversample.size

sion. If not numeric, all pixels are converted, but this can be time-consuming,

especially for large images. See details for speed.

ref.white String; white reference for converting from RGB to CIEL*a*b color space. Ac-

cepts any of the standard white references for convertColor (see details).

hsv

CIELab

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Details

The upper and lower limits for background pixel elimination set the inclusive bounds for which pixels should be ignored for the 2D arrays; while all background pixels are ideally a single color, images photographed against "uniform" backgrounds often contain some variation, and even segmentation done with photo editing software will produce some variance as a result of image compression.

The upper and lower bounds represent cutoffs: any pixel for which the first channel falls between the first upper and lower bounds, the second channel falls between the second upper and lower bounds, and the third channel falls between the third upper and lower bounds, will be ignored. For example, if you have a green pixel with RGB channel values [0.1, 0.9, 0.2], and your upper and lower bounds were (0.2, 1, 0.2) and (0, 0.6, 0) respectively, the pixel would be ignored because 0 <= 0.1 <= 0.2, 0.6 <= 0.9 <= 1, and 0 <= 0.2 <= 0.2. But a pixel with the RGB channel values [0.3, 0.9, 0.2] would not be considered background because 0.3 >= 0.2.

CIEL*a*b color space requires a reference 'white light' color (dimly and brightly lit photographs of the same object will have very different RGB palettes, but similar Lab palettes if appropriate white references are used). The idea here is that the apparent colors in an image depend not just on the "absolute" color of an object (whatever that means), but also on the available light in the scene. There are seven CIE standardized illuminants available in colordistance (A, B, C, E, and D50, D55, and D60), but the most common are:

- "A": Standard incandescent lightbulb
- "D65": Average daylight
- "D50": Direct sunlight

Color conversions will be highly dependent on the reference white used, which is why no default is provided. Users should look into standard illuminants to choose an appropriate reference for a dataset.

Value

A list with original image (\$original.rgb, 3D array), 2D matrix with background pixels removed (\$filtered.rgb.2d and \$filtered.hsv.2d), and path to the original image (\$path).

Note

The 3D array is useful for displaying the original image, while the 2D arrays (RGB and HSV) are treated as rows of data for clustering in the rest of the package.

```
loadedImg <- colordistance::loadImage(system.file("extdata",
  "Heliconius/Heliconius_A/Heliconius_01.jpeg", package="colordistance"),
  upper=rep(1, 3), lower=rep(0.8, 3))

loadedImgNoHSV <- colordistance::loadImage(system.file("extdata",
  "Heliconius/Heliconius_A/Heliconius_01.jpeg", package="colordistance"),
  upper=rep(1, 3), lower=rep(0.8, 3), hsv=FALSE)</pre>
```

normalizeRGB 29

normalizeRGB

Normalize pixel RGB ratios

Description

Converts clusters from raw channel intensity to their fraction of the intensity for that cluster

Usage

```
normalizeRGB(extractClustersObject)
```

Arguments

```
extractClustersObject
```

A list of color clusters such as those returned by extractClusters or getHistList. List must contain identically sized dataframes with color coordinates (R, G, B or H, S, V) as the first three columns.

Value

A list of the same size and structure as the input list, but with the cluster normalized as described.

Note

This is a useful option if your images have a lot of variation in lighting, but obviously comes at the cost of reducing variation (if darker and lighter colors are meaningful sources of variation in the dataset).

For example, a bright yellow (R=1, G=1, B=0) and a darker yellow (R=0.8, G=0.8, B=0) both have 50% red, 50% green, and 0% blue, so their normalized values would be equivalent.

A similar but less harsh alternative would be to use HSV rather than RGB for pixel binning and color similarity clustering by setting hsv=T in clustering functions and specifying a low number of 'value' bins (e.g. bins=c(8, 8, 2)).

```
cluster.list <- colordistance::getKMeansList(c(system.file("extdata",
"Heliconius/Heliconius_A", package="colordistance"), lower=rep(0.8, 3),
upper=rep(1, 3)))
cluster.list <- colordistance::extractClusters(cluster.list)
colordistance:::normalizeRGB(cluster.list)</pre>
```

30 orderClusters

orderClusters

Order color clusters to minimize overall color distance between pairs

Description

Reorders clusters to minimize color distance using the Hungarian algorithm as implemented by solve_LSAP.

Usage

```
orderClusters(extractClustersObject)
```

Arguments

```
extractClustersObject
```

A list of color clusters such as those returned by extractClusters or getHistList. List must contain identically sized dataframes with color coordinates (R, G, B or H, S, V) as the first three columns.

Details

Briefly: Euclidean distances between every possible pair of clusters across two dataframes are calculated, and pairs of clusters are chosen in order to minimize the total sum of color distances between the cluster pairs (i.e. A1-B1, A2-B2, etc).

For example, if dataframe A has a black cluster, a white cluster, and a blue cluster, in that order, and dataframe B has a white cluster, a blue cluster, and a grey cluster, in that order, the final pairs might be A1-B3 (black and grey), A2-B2 (blue and blue), and A3-B1 (white and white).

Rows are reordered so that paired rows are in the same row index (in the example, dataframe B would be reshuffled to go grey, blue, white instead of white, grey, blue).

Value

A list with identical data to the input list, but with rows in each dataframe reordered to minimize color distances per cluster pair.

```
cluster.list <- colordistance::getKMeansList(c(system.file("extdata",
"Heliconius/Heliconius_A", package="colordistance"), lower=rep(0.8, 3),
upper=rep(1, 3)))
cluster.list <- colordistance::extractClusters(cluster.list)
colordistance:::orderClusters(cluster.list)</pre>
```

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pause

Pause and wait for user input

Description

Tiny little function wrapper, mostly used for looping or when several plots are output by a single function. Waits for user keystroke to move on to next image or exit.

Usage

```
pause()
```

Examples

```
for (i in c(1:5)) {
   print(i)
   if (i < 5) {
      colordistance:::pause()
   }
}</pre>
```

plotClusters

Plot clusters in 3D color space

Description

Interactive, 3D plot_ly plots of cluster sizes and colors for each image in a list of cluster dataframes in order to visualize cluster output.

Usage

```
plotClusters(cluster.list, color.space = "rgb", p = "all",
   pausing = TRUE, ref.white, to = "sRGB")
```

Arguments

cluster.list	A list of identically sized dataframes with 4 columns each (R, G, B, Pct or H, S, V, Pct) as output by extractClusters or getHistList.
color.space	The color space ("rgb", "hsv", or "lab") in which to plot pixels.
p	Numeric vector of indices for which elements to plot; otherwise each set of clusters is plotted in succession.
pausing	Logical. Should the function pause and wait for user keystroke before plotting the next plot?
ref.white	The reference white passed to convertColorSpace; must be specified if using color.space = "lab".
to	Display color space of image if clustering in CIE Lab space, probably either "sRGB" or "Apple RGB", depending on your computer.

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Value

A 3D plot_ly plot of cluster sizes in the specified colorspace for each cluster dataframe provided.

Examples

```
## Not run:
# Takes >10 seconds
cluster.list <- colordistance::getHistList(dir(system.file("extdata",
    "Heliconius/", package="colordistance"), full.names=TRUE), plotting=FALSE,
lower=rep(0.8, 3), upper=rep(1, 3))

colordistance::plotClusters(cluster.list, p=c(1:3, 7:8), pausing=FALSE)

clusterListHSV <- colordistance::getHistList(dir(system.file("extdata",
    "Heliconius/", package="colordistance"), full.names=TRUE), hsv=TRUE,
plotting=FALSE, lower=rep(0.8, 3), upper=rep(1, 3))

colordistance::plotClusters(clusterListHSV, p=c(1:3, 7:8), hsv=TRUE,
    pausing=FALSE)

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

plotClustersMulti

Plot several different cluster sets together

Description

Plots cluster sets from several different dataframes on a single plot for easy comparison.

Usage

```
plotClustersMulti(cluster.list, color.space = "rgb", p = "all",
   title = "", ref.white, to = "sRGB")
```

Arguments

cluster.list	A list of identically sized dataframes with 4 columns each as output by extractClusters, getLabHistList, or getHistList.
color.space	The color space ("rgb", "hsv", or "lab") in which to plot pixels.
p	Numeric vector of indices for which elements to plot; otherwise all of the cluster sets provided will be plotted together.
title	Optional title for the plot.
ref.white	The reference white passed to convertColorSpace; must be specified if using color.space = "lab".
to	Display color space of image if clustering in CIE Lab space, probably either "sRGB" or "Apple RGB", depending on your computer.

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Value

A single plot_ly plot of every cluster in a list of cluster sets. Each cluster is colored by cluster color, proportional to cluster size, and labeled according to the image from which it originated.

Note

Each cluster plotted is colored according to its actual color, and labeled according to the image from which it originated.

Examples

```
## Not run:
# Takes >10 seconds
cluster.list <- colordistance::getHistList(dir(system.file("extdata",
    "Heliconius/", package="colordistance"), full.names=TRUE), plotting=FALSE,
lower=rep(0.8, 3), upper=rep(1, 3))

colordistance::plotClustersMulti(cluster.list, p=c(1:4), title="Orange and
black Heliconius")

colordistance::plotClustersMulti(cluster.list, p=c(5:8), title="Black, yellow,
and red Heliconius")

clusterListHSV <- colordistance::getHistList(dir(system.file("extdata",
    "Heliconius/", package="colordistance"), full.names=TRUE), hsv=TRUE,
plotting=FALSE, lower=rep(0.8, 3), upper=rep(1, 3))

colordistance::plotClustersMulti(clusterListHSV, p=c(1:3, 7:8), hsv=TRUE)

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

plotHist

Color histogram of binned image

Description

Plots a color histogram from a dataframe as returned by getImageHist, getHistList, or extractClusters. Bars are colored according to the color of the bin.

Usage

```
plotHist(histogram, pausing = TRUE, color.space = "rgb", ref.white,
  from = "sRGB", main = "default", ...)
```

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Arguments

histogram	A single dataframe or a list of dataframes as returned by getLabHist, getLabHistList, or extractClusters. First three columns must be color coordinates and fourth column must be cluster size.
pausing	Logical. Pause and wait for keystroke before plotting the next histogram?
color.space	The color space ("rgb", "hsv", or "lab") in which to plot cluster histogram.
ref.white	The reference white passed to convertColorSpace; must be specified if using CIE Lab space. See convertColorSpace.
from	Display color space of image if clustering in CIE Lab space, probably either "sRGB" or "Apple RGB", depending on your computer.
main	Title for plot. If "default", the name of the cluster histogram is used.
	Optional arguments passed to the barplot function.

Examples

```
color_df <- as.data.frame(matrix(rep(seq(0, 1, length.out=3), 3), nrow=3,
ncol=3))

color_df$Pct <- c(0.2, 0.5, 0.3)

colordistance::plotHist(color_df, main="Example plot")</pre>
```

plotImage

Display an image in a plot window

Description

Plots an image as an image.

Usage

```
plotImage(img)
```

Arguments

img

Either a path to an image or a loadImage object.

Details

Redundant, but a nice sanity check. Used in a few other functions in colordistance package. Takes either a path to an image (RGB or PNG) or an image object as read in by loadImage.

Value

A plot of the provided image in the current plot window.

plotPixels 35

Examples

```
colordistance::plotImage(system.file("extdata",
  "Heliconius/Heliconius_A/Heliconius_01.jpeg", package="colordistance"))
colordistance::plotImage(loadImage(system.file("extdata",
  "Heliconius/Heliconius_A/Heliconius_01.jpeg", package="colordistance"),
lower=rep(0.8, 3), upper=rep(1, 3)))
```

plotPixels

Plot pixels in color space

Description

Plots non-background pixels according to their color coordinates, and colors them according to their RGB or HSV values. Dimensions are either RGB or HSV depending on flags.

Usage

```
plotPixels(img, n = 10000, lower = c(0, 0.55, 0), upper = c(0.25, 1, 0.25), color.space = "rgb", ref.white = NULL, pch = 20, main = "default", from = "sRGB", xlim = "default", ylim = "default", zlim = "default", ...)
```

Arguments

upper

img Either a path to an image or a loadImage object.

n Number of randomly selected pixels to plot; recommend <20000 for speed. If n exceeds the number of non-background pixels in the image, all pixels are

plotted. If n is not numeric, all pixels are plotted.

lower RGB or HSV triplet specifying the lower bounds for background pixels. Default upper and lower bounds are set to values that work well for a bright green

background (RGB [0, 1, 0]).

RGB or HSV triplet specifying the upper bounds for background pixels. Default upper and lower bounds are set to values that work well for a bright green background (RGB [0, 1, 0]). Determining these bounds may take some trial and error, but the following bounds may work for certain common background

colors:

• Black: lower=c(0, 0, 0); upper=c(0.1, 0.1, 0.1)

• White: lower=c(0.8, 0.8, 0.8); upper=c(1, 1, 1)

• Green: lower=c(0, 0.55, 0); upper=c(0.24, 1, 0.24)

• Blue: lower=c(0, 0, 0.55); upper=c(0.24, 0.24, 1)

If no background filtering is needed, set bounds to some non-numeric value (NULL, FALSE, "off", etc); any non-numeric value is interpreted as NULL.

color.space

The color space ("rgb", "hsv", or "lab") to use for plotting.

ref.white

The reference white passed to convertColor; must be specified if img does not already contain CIE Lab pixels. See convertColorSpace.

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pch	Passed to scatterplot3d.	
main	Plot title. If left as "default", image name is used.	
from	Original color space of image if plotting in CIE Lab space, probably either "sRGB" or "Apple RGB", depending on your computer.	
xlim, ylim, zlim		
	Ranges for the X, Y, and Z axes. If "default", the widest ranges for each axis according to the specified color space (0-1 for RGB and HSV, 0-100 for L of Lab, -128-127 for a and b of Lab) are used.	
	Optional parameters passed to scatterplot3d.	

Value

3D plot of pixels in either RGB or HSV color space, colored according to their color in the image. Uses scatterplot3d function.

Note

If n is not numeric, then all pixels are plotted, but this is not recommended. Unless the image has a low pixel count, it takes much longer, and plotting this many points in the plot window can obscure important details.

There are seven CIE standardized illuminants available in colordistance (A, B, C, E, and D50, D55, and D65), but the most common are:

```
• "A": Standard incandescent lightbulb
```

"D65": Average daylight "D50": Direct sunlight

Examples

```
colordistance::plotPixels(system.file("extdata",
"Heliconius/Heliconius_B/Heliconius_07.jpeg", package="colordistance"),
n=20000, upper=rep(1, 3), lower=rep(0.8, 3), color.space = "rgb", angle = -45)
```

scatter3dclusters

Plot 3D clusters in a 2D plot

Description

Uses scatterplot3d to plot clusters in color space.

Usage

```
scatter3dclusters(clusters, color.space, ref.white = "D65",
   xlim = "default", ylim = "default", zlim = "default",
   main = "Color clusters", scaling = 10, opacity = 0.9,
   plus = 0.01, ...)
```

weightedPairsDistance 37

Arguments

clusters	A single dataframe or a list of dataframes as returned by getLabHist, getLabHistList, or extractClusters. First three columns must be color coordinates and fourth column must be cluster size.	
color.space	The color space ("rgb", "hsv", or "lab") in which to plot. If not specified, the function uses column names to guess the color space.	
ref.white	Standard reference white for converting lab coordinates to RGB coordinates for coloring clusters. One of either "A", "B", "C", "E", "D50", "D55", or "D65".	
xlim, ylim, zlim		
	X, Y, and Z-axis limits. If not specified, the defaults are 0-1 for all channels in RGB and HSV space, or 0-100 for L and -100-100 for a and b channels of CIE Lab space.	
main	Title for the plot.	
scaling	Scaling factor for size of clusters.	
opacity	Transparency value for plotting; must be between 0 and 1.	
plus	Amount to add to percent column for plotting; can help to make very small (or 0) clusters visible.	
	Additional parameters passed to scatterplot3d.	

See Also

plotClusters, plotClustersMulti

Examples

weightedPairsDistance Distance between color clusters with user-specified color/size weights

Description

Distance metric with optional user input for specifying how much the bin size similarity and color similarity should be weighted when pairing clusters from different color cluster sets.

Usage

```
weightedPairsDistance(T1, T2, ordering = FALSE, size.weight = 0.5,
  color.weight = 0.5)
```

Arguments

T1	Dataframe (especially a dataframe as returned by extractClusters or getImageHist, but first three columns must be coordinates).
T2	Another dataframe like T1.
ordering	Logical. Should clusters by paired in order to minimize overall distance scores or evaluated in the order given?
size.weight	Weight of size similarity in determining overall score and ordering (if ordering=T).
color.weight	Weight of color similarity in determining overall score and ordering (if ordering=T). Color and size weights do not necessarily have to sum to 1.

Value

Similarity score based on size and color similarity of each pair of points in provided dataframes.

Note

Use with caution, since weights can easily swing distance scores more dramatically than might be expected. For example, if size.weight = 1 and color.weight = 0, two clusters of identical color but different sizes would not be compared.

```
cluster.list <- colordistance::getKMeansList(system.file("extdata",
  "Heliconius/Heliconius_B", package="colordistance"), lower=rep(0.8, 3),
  upper=rep(1, 3))
  cluster.list <- colordistance::extractClusters(cluster.list, ordering=TRUE)
  colordistance:::weightedPairsDistance(cluster.list[[1]], cluster.list[[2]],
  size.weight=0.8, color.weight=0.2)</pre>
```

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