Package 'handwriter'

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```
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about_variable

About Varialbe

Description

about_variable() returns information about the model variable.

Usage

```
about_variable(variable, model)
```

Arguments

variable A variable in the fitted model output by fit_model()

model A fitted model created by fit_model()

Value

Text that explains the variable

Examples

```
about_variable(
  variable = "mu[1,2]",
  model = example_model
)
```

 ${\sf addToFeatures}$

add To Features

Description

addToFeatures

Usage

```
addToFeatures(FeatureSet, LetterList, vectorDims)
```

Arguments

FeatureSet The current list of features that have been calculated

LetterList List of all letters and their information

vectorDims Vectors with image Dims

Value

A list consisting of current features calculated in FeatureSet as well as measures of compactness, loop count, and loop dimensions

```
analyze_questioned_documents Analyze\ Questioned\ Documents
```

Description

analyze_questioned_documents() estimates the posterior probability of writership for the questioned documents using Markov Chain Monte Carlo (MCMC) draws from a hierarchical model created with fit_model().

Usage

```
analyze_questioned_documents(
  main_dir,
  questioned_docs,
  model,
  num_cores,
  writer_indices,
  doc_indices
)
```

Arguments

Value

A list of likelihoods, votes, and posterior probabilities of writership for each questioned document.

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Examples

```
## Not run:
main_dir <- "/path/to/main_dir"
questioned_docs <- "/path/to/questioned_images"
analysis <- analyze_questioned_documents(
    main_dir = main_dir,
    questioned_docs = questioned_docs,
    model = model,
    num_cores = 2,
    writer_indices = c(2, 5),
    doc_indices = c(7, 18)
)
analysis$posterior_probabilities
## End(Not run)</pre>
```

calculate_accuracy

Calculate Accuracy

Description

Fit a model with fit_model() and calculate posterior probabilities of writership with analyze_questioned_documents() of a set of test documents where the ground truth is known. Then use calculate_accuracy() to measure the accuracy of the fitted model on the test documents. Accuracy is calculated as the average posterior probability assigned to the true writer.

Usage

```
calculate_accuracy(analysis)
```

Arguments

analysis

Writership analysis output by analyze_questioned_documents

Value

The model's accuracy on the test set as a number

```
# calculate the accuracy for example analysis performed on test documents and a model with 1 chain
calculate_accuracy(example_analysis)
```

```
## Not run:
main_dir <- "/path/to/main_dir"
test_images_dir <- "/path/to/test_images"
analysis <- analyze_questioned_documents(
   main_dir = main_dir,</pre>
```

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```
questioned_docs = test_images_dir,
model = model,
num_cores = 2,
writer_indices = c(2, 5),
doc_indices = c(7, 18)
)
calculate_accuracy(analysis)
## End(Not run)
```

 ${\tt cleanBinaryImage}$

cleanBinaryImage

Description

Removes alpha channel from png image.

Usage

```
cleanBinaryImage(img)
```

Arguments

img

A matrix of 1s and 0s.

Value

png image with the alpha channel removed

csafe

Cursive written word: csafe

Description

Cursive written word: csafe

Usage

csafe

Format

Binary image matrix. 111 rows and 410 columns.

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Examples

```
csafe_document <- list()
csafe_document$image <- csafe
plotImage(csafe_document)
csafe_document$thin <- thinImage(csafe_document$image)
plotImageThinned(csafe_document)
csafe_processList <- processHandwriting(csafe_document$thin, dim(csafe_document$image))</pre>
```

drop_burnin

Drop Burn-In

Description

drop_burnin() removes the burn-in from the Markov Chain Monte Carlo (MCMC) draws.

Usage

```
drop_burnin(model, burn_in)
```

Arguments

model A list of MCMC draws from a model fit with fit_model().

burn_in An integer number of starting iterations to drop from each MCMC chain.

Value

A list of data frames of MCMC draws with burn-in dropped.

Examples

```
model <- drop_burnin(model = example_model, burn_in = 25)
plot_trace(variable = "mu[1,2]", model = example_model)</pre>
```

example_analysis

Example of writership analysis

Description

Example of writership analysis

Usage

```
example_analysis
```

Format

The results of analyze_questioned_documents() stored in a named list with 5 items:

graph_measurements A data frame of that shows the writer, document name, cluster assignment, slope, principle component rotation angle, and wrapped principle component rotation angle for each training graph in each questioned documents.

cluster_fill_counts A data frame of the cluster fill counts for each questioned document.

likelihoods A list of data frames where each data frame contains the likelihoods for a questioned document for each MCMC iteration.

votes A list of vote tallies for each questioned document.

posterior_probabilites A list of posterior probabilities of writership for each questioned document and each known writer in the closed set used to train the hierarchical model.

Examples

```
plot_cluster_fill_counts(formatted_data = example_analysis)
plot_posterior_probabilities(analysis = example_analysis)
```

example_cluster_template

Example cluster template

Description

An example cluster template created with make_clustering_template(). The cluster template was created from handwriting samples "w0016_s01_pLND_r01.png", "w0080_s01_pLND_r01.png", "w0124_s01_pLND_r01.png", "w0138_s01_pLND_r01.png", and "w0299_s01_pLND_r01.png" from the CSAFE Handwriting Database. The template has K=5 clusters.

Usage

```
example_cluster_template
```

Format

A list containing a single cluster template created by make_clustering_template(). The cluster template was created by sorting a random sample of 1000 graphs from 10 training documents into 10 clusters with a K-means algorithm. The cluster template is a named list with 16 items:

centers_seed An integer for the random number generator.

cluster A vector of cluster assignments for each graph used to create the cluster template.

centers The final cluster centers produced by the K-Means algorithm.

K The number of clusters to build (10) with the K-means algorithm.

n The number of training graphs to use (1000) in the K-means algorithm.

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docnames A vector that lists the training document from which each graph originated.

writers A vector that lists the writer of each graph.

iters The maximum number of iterations for the K-means algorithm (3).

changes A vector of the number of graphs that changed clusters on each iteration of the K-means algorithm.

outlierCutoff A vector of the outlier cutoff values calculated on each iteration of the K-means algorithm.

stop_reason The reason the K-means algorithm terminated.

wcd A matrix of the within cluster distances on each iteration of the K-means algorithm. More specifically, the distance between each graph and the center of the cluster to which it was assigned on each iteration.

wcss A vector of the within-cluster sum of squares on each iteration of the K-means algorithm.

Examples

```
# view cluster fill counts for template training documents
template_data <- format_template_data(example_cluster_template)
plot_cluster_fill_counts(template_data, facet = TRUE)</pre>
```

example_model

Example of a hierarchical model

Description

Example of a hierarchical model

Usage

example_model

Format

A hierarchical model created by fit_model with a single chain of 100 MCMC iterations. It is a named list of 4 objects:

graph_measurements A data frame of model training data that shows the writer, document name, cluster assignment, slope, principle component rotation angle, and wrapped principle component rotation angle for each training graph.

cluster_fill_counts A data frame of the cluster fill counts for each model training document.

rjags_data The model training information from graph_measurements and cluster_fill_counts formatted for RJAGS.

fitted_model A model fit using the rjags_data and the RJAGS and coda packages. It is an MCMC list that contains a single MCMC object.

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Examples

```
# convert to a data frame and view all variable names
df <- as.data.frame(coda::as.mcmc(example_model$fitted_model))</pre>
colnames(df)
# view a trace plot
plot_trace(variable = "mu[1,1]", model = example_model)
# drop the first 25 MCMC iterations for burn-in
model <- drop_burnin(model = example_model, burn_in = 25)</pre>
## Not run:
# analyze questioned documents
main_dir <- /path/to/main_dir</pre>
questioned_docs <- /path/to/questioned_documents_directory</pre>
analysis <- analyze_questioned_documents(</pre>
   main_dir = main_dir,
   questioned_docs = questioned_docs
   model = example_model
   num\_cores = 2
analysis$posterior_probabilities
## End(Not run)
```

extractGraphs

Extract Graphs

Description

```
'r lifecycle::badge("superseded")'
```

Usage

```
extractGraphs(source_folder = getwd(), save_folder = getwd())
```

Arguments

```
source_folder path to folder containing .png images
save_folder path to folder where graphs are saved to
```

Details

Development on 'extractGraphs()' is complete. We recommend using 'process_batch_dir()' instead.

Extracts graphs from .png images and saves each by their respective writer.

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Value

saves graphs in an rds file

Examples

```
## Not run:
sof = "path to folder containing .png images"
saf = "path to folder where graphs will be saved to"
extractGraphs(sof, saf)
## End(Not run)
```

fit_model

Fit Model

Description

fit_model() fits a Bayesian hierarchical model to the model training data in model_docs and draws samples from the model as Markov Chain Monte Carlo (MCMC) estimates.

Usage

```
fit_model(
   main_dir,
   model_docs,
   num_iters,
   num_chains = 1,
   num_cores,
   writer_indices,
   doc_indices,
   a = 2,
   b = 0.25,
   c = 2,
   d = 2,
   e = 0.5
)
```

Arguments

main_dir	A directory that contains a cluster template created by make_clustering_template()
model_docs	A directory containing model training documents
num_iters	An integer number of iterations of MCMC.
num_chains	An integer number of chains to use.
num_cores	An integer number of cores to use for parallel processing clustering assignments. The model fitting is not done in parallel.

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writer_indices	A vector of the start and stop character of the writer ID in the model training file names. E.g., if the file names are writer0195_doc1, writer0210_doc1, writer0033_doc1 then writer_indices is 'c(7,10)'.
doc_indices	A vector of the start and stop character of the "document name" in the model training file names. This is used to distinguish between two documents written by the same writer. E.g., if the file names are writer0195_doc1, writer0195_doc2, writer0033_doc1, writer0033_doc2 then doc_indices are 'c(12,15)'.
a	The shape parameter for the Gamma distribution in the hierarchical model
b	The rate parameter for the Gamma distribution in the hierarchical model
С	The first shape parameter for the Beta distribution in the hierarchical model
d	The second shape parameter for the Beta distribution in the hierarchical model
e	The scale parameter for the hyper prior for mu in the hierarchical model

Value

A list of training data used to fit the model and the fitted model

```
## Not run:
main_dir <- "/path/to/main_dir"
model_docs <- "path/to/model_training_docs"</pre>
questioned_docs <- "path/to/questioned_docs"</pre>
model <- fit_model(</pre>
 main_dir = main_dir,
  model_docs = model_docs,
  num\_iters = 100,
 num_chains = 1,
 num\_cores = 2,
  writer_indices = c(2, 5),
  doc_indices = c(7, 18)
)
model <- drop_burnin(model = model, burn_in = 25)</pre>
analysis <- analyze_questioned_documents(</pre>
  main_dir = main_dir,
  questioned_docs = questioned_docs,
  model = model,
  num\_cores = 2
)
analysis$posterior_probabilities
## End(Not run)
```

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```
format_template_data Format Template Data
```

Description

format_template_data() formats the template data for use with plot_cluster_fill_counts(). The output is a list that contains a data frame called cluster_fill_counts.

Usage

```
format_template_data(template)
```

Arguments

template

A single cluster template created by make_clustering_template()

Value

List that contains the cluster fill counts

Examples

```
template_data <- format_template_data(template = example_cluster_template)
plot_cluster_fill_counts(formatted_data = template_data, facet = TRUE)</pre>
```

```
get_clusters_batch
```

get_clusters_batch

Description

```
get_clusters_batch
```

Usage

```
get_clusters_batch(
  template,
  input_dir,
  output_dir,
  writer_indices = NULL,
  doc_indices = NULL,
  num_cores = 1,
  save_master_file = FALSE
)
```

Arguments

template A cluster template created with make_clustering_template input_dir A directory containing graphs created with process_batch_dir

output_dir Output directory for cluster assignments

writer_indices Optional. A Vector of start and end indices for the writer id in the graph file

names.

doc_indices Optional. Vector of start and end indices for the document id in the graph file

names.

num_cores Integer number of cores to use for parallel processing

save_master_file

TRUE or FALSE. If TRUE, a master file named 'all_clusters.rds' containing the cluster assignments for all documents in the input directory will be saved to the output directory. If FASLE, a master file will not be saved, but the individual files for each document in the input directory will still be saved to the output directory.

Value

A list of cluster assignments

Examples

```
## Not run:
template <- readRDS('path/to/template.rds')
get_clusters_batch(template=template, input_dir='path/to/dir', output_dir='path/to/dir',
writer_indices=c(2,5), doc_indices=c(7,18), num_cores=1)
get_clusters_batch(template=template, input_dir='path/to/dir', output_dir='path/to/dir',
writer_indices=c(1,4), doc_indices=c(5,10), num_cores=5)
## End(Not run)</pre>
```

```
get_cluster_fill_counts
```

Get Cluster Fill Counts

Description

get_cluster_fill_counts() creates a data frame that shows the number of graphs in each cluster for each input document.

Usage

```
get_cluster_fill_counts(df)
```

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Arguments

df

A data frame with columns docname and cluster. Each row corresponding to a graph and lists the document from which the graph was obtained, and the cluster to which that graph is assigned. Optionally, the data frame might also have writer and doc columns. If present, writer lists the writer ID of each document and doc is an identifier to distinguish between different documents from the same writer.

Value

A dataframe of cluster fill counts for each document in the input data frame.

Examples

```
docname <- c(rep('doc1',20), rep('doc2', 20), rep('doc3', 20))
writer <- c(rep(1, 20), rep(2, 20), rep(3, 20))
doc <- c(rep(1, 20), rep(2, 20), rep(3, 20))
cluster <- sample(3, 60, replace=TRUE)
df <- data.frame(docname, writer, doc, cluster)
get_cluster_fill_counts(df)</pre>
```

```
get_credible_intervals
```

Get Credible Intervals

Description

In a model created with fit_model() the pi parameters are the estimate of the true cluster fill count for a particular writer and cluster. The function get_credible_intervals() calculates the credible intervals of the pi parameters for each writer in the model.

Usage

```
get_credible_intervals(model, interval_min = 0.05, interval_max = 0.95)
```

Arguments

model A model output by fit_model()

interval_min The lower bound for the credible interval. The number must be between 0 and

1.

interval_min and must be less than 1.

Value

A list of data frames. Each data frame lists the credible intervals for a single writer.

Examples

```
get_credible_intervals(model=example_model)
get_credible_intervals(model=example_model, interval_min=0.05, interval_max=0.95)
```

Description

Get the posterior probabilities for questioned document analyzed with analyze_questioned_documents().

Usage

```
get_posterior_probabilities(analysis, questioned_doc)
```

Arguments

analysis

The output of analyze_questioned_documents(). If more than one questioned document was analyzed with this function, then the data frame analysis\$posterior_probabilities lists the posterior probabilities for all questioned documents. get_posterior_probabilities() creates a data frame of the posterior probabilities for a single questioned document and sorts the known writers from the most likely to least likely to have written the questioned document.

questioned_doc The filename of the questioned document

Value

A data frame of posterior probabilities for the questioned document

```
get_posterior_probabilities(
  analysis = example_analysis,
  questioned_doc = "w0030_s03_pWOZ_r01"
)
```

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graphToPrototype

Convert graph to a prototype

Description

A graph prototype consists of the starting and ending points of each path in the graph, as well as and evenly spaced points along each path. The prototype also stores the center point of the graph. All points are represented as xy-coordinates and the center point is at (0,0).

Usage

```
graphToPrototype(graph, numPathCuts = 8)
```

Arguments

graph A graph from a handwriting sample

numPathCuts Number of segments to cut the path(s) into

Value

List of pathEnds, pathQuarters, and pathCenters given as (x,y) coordinates with the graph centroid at (0,0). The returned list also contains path lengths. pathQuarters gives the (x,y) coordinates of the path at the cut points and despite the name, the path might not be cut into quarters.

london

Cursive written word: London

Description

Cursive written word: London

Usage

london

Format

Binary image matrix. 148 rows and 481 columns.

```
london_document <- list()
london_document$image <- london
plotImage(london_document)
london_document$thin <- thinImage(london_document$image)
plotImageThinned(london_document)
london_processList <- processHandwriting(london_document$thin, dim(london_document$image))</pre>
```

Description

make_clustering_template() applies a K-means clustering algorithm to the input handwriting samples pre-processed with process_batch_dir() and saved in the input folder main_dir > data > template_graphs. The K-means algorithm sorts the graphs in the input handwriting samples into groups, or *clusters*, of similar graphs.

Usage

```
make_clustering_template(
  main_dir,
  template_docs,
  writer_indices,
  centers_seed,
  K = 40,
  num_dist_cores = 1,
  max_iters = 25
)
```

Arguments

Main directory that will store template files main_dir A directory containing template training images template_docs writer_indices A vector of the starting and ending location of the writer ID in the file name. centers_seed Integer seed for the random number generator when selecting starting cluster centers. Κ Integer number of clusters num_dist_cores Integer number of cores to use for the distance calculations in the K-means algorithm. Each iteration of the K-means algorithm calculates the distance between each input graph and each cluster center. max_iters Maximum number of iterations to allow the K-means algorithm to run

Value

List containing the cluster template

```
## Not run:
main_dir <- "path/to/folder"
template_docs <- "path/to/template_training_docs"
template_list <- make_clustering_template(</pre>
```

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```
main_dir = main_dir,
  template_docs = template_docs,
  writer_indices = c(2, 5),
  K = 10,
  num_dist_cores = 2,
  max_iters = 25,
  centers_seed = 100,
)
## End(Not run)
```

message

Full page image of the handwritten London letter.

Description

Full page image of the handwritten London letter.

Usage

message

Format

Binary image matrix. 1262 rows and 1162 columns.

```
message_document <- list()
message_document$image <- message
plotImage(message_document)

## Not run:
message_document <- list()
message_document$image <- message
plotImage(message_document)
message_document$thin <- thinImage(message_document$image)
plotImageThinned(message_document)
message_processList <- processHandwriting(message_document$thin, dim(message_document$image))

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

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nature1	Full page image of the 4th sample (nature) of handwriting from the first writer.
nature1	

Description

Full page image of the 4th sample (nature) of handwriting from the first writer.

Usage

nature1

Format

Binary image matrix. 811 rows and 1590 columns.

Examples

```
nature1_document <- list()
nature1_document$image <- nature1
plotImage(nature1_document)

## Not run:
nature1_document <- list()
nature1_document$image <- nature1
plotImage(nature1_document)
nature1_document$thin <- thinImage(nature1_document$image)
plotImageThinned(nature1_document)
nature1_processList <- processHandwriting(nature1_document$thin, dim(nature1_document$image))

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

plotImage

Plot Image

Description

This function plots a basic black and white image.

Usage

```
plotImage(doc)
```

Arguments

doc

A document processed with processDocument() or a binary matrix (all entries are 0 or 1)

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Value

```
ggplot plot
```

Examples

```
csafe_document <- list()
csafe_document$image <- csafe
plotImage(csafe_document)

## Not run:
document <- processDocument('path/to/image.png')
plotImage(document)

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

plotImageThinned

Plot Thinned Image

Description

This function returns a plot with the full image plotted in light gray and the thinned skeleton printed in black on top.

Usage

```
plotImageThinned(doc)
```

Arguments

doc

A document processed with processHandwriting()

Value

gpplot plot of thinned image

```
csafe_document <- list()
csafe_document$image <- csafe
csafe_document$thin <- thinImage(csafe_document$image)
plotImageThinned(csafe_document)</pre>
```

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|--|

Description

This function returns a plot of a single graph extracted from a document. It uses the letterList parameter from the processHandwriting() or processDocument() function and accepts a single value as whichLetter. Dims requires the dimensions of the entire document, since this isn't contained in processHandwriting() or processDocument().

Usage

```
plotLetter(
   doc,
   whichLetter,
   showPaths = TRUE,
   showCentroid = TRUE,
   showSlope = TRUE,
   showNodes = TRUE
)
```

Arguments

doc A document processed with processHandwriting() or processDocument()

whichLetter Single value in 1:length(letterList) denoting which letter to plot.

showPaths Whether the calculated paths on the letter should be shown with numbers.

showCentroid Whether the centroid should be shown showSlope Whether the slope should be shown showNodes Whether the nodes should be shown

Value

Plot of single letter.

```
twoSent_document = list()
twoSent_document$image = twoSent
twoSent_document$thin = thinImage(twoSent_document$image)
twoSent_document$process = processHandwriting(twoSent_document$thin, dim(twoSent_document$image))
plotLetter(twoSent_document, 1)
plotLetter(twoSent_document, 4, showPaths = FALSE)
```

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Description

This function returns a plot of a single line extracted from a document. It uses the letterList parameter from the processHandwriting function and accepts a single value as whichLetter. Dims requires the dimensions of the entire document, since this isn't contained in processHandwriting.

Usage

```
plotLine(letterList, whichLine, dims)
```

Arguments

letterList Letter list from processHandwriting function

whichLine Single value denoting which line to plot - checked if too big inside function.

dims Dimensions of the original document

Value

ggplot plot of single line

Examples

```
twoSent_document = list()
twoSent_document$image = twoSent
twoSent_document$thin = thinImage(twoSent_document$image)
twoSent_processList = processHandwriting(twoSent_document$thin, dim(twoSent_document$image))
dims = dim(twoSent_document$image)
plotLine(twoSent_processList$letterList, 1, dims)
```

plotNodes Plot Nodes

Description

This function returns a plot with the full image plotted in light gray and the skeleton printed in black, with red triangles over the vertices. Also called from plotPath, which is a more useful function, in general.

Usage

```
plotNodes(doc, plot_break_pts = FALSE, nodeSize = 3, nodeColor = "red")
```

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Arguments

doc A document processed with processHandwriting()

plot_break_pts Logical value as to whether to plot nodes or break points. plot_break_pts=FALSE

plots nodes and plot_break_pts=TRUE plots break point.

nodeSize Size of triangles printed. 3 by default. Move down to 2 or 1 for small text

images.

nodeColor Which color the nodes should be

Value

Plot of full and thinned image with vertices overlaid.

Examples

```
csafe_document <- list()
csafe_document$image <- csafe
csafe_document$thin <- thinImage(csafe_document$image)
csafe_document$process <- processHandwriting(csafe_document$thin, dim(csafe_document$image))
plotNodes(csafe_document)
plotNodes(csafe_document, nodeSize=6, nodeColor="black")</pre>
```

Description

Plot the cluster centers of a cluster template created with make_clustering_template. This function uses a K-Means type algorithm to sort graphs from training documents into clusters. On each iteration of the algorithm, it calculates the mean graph of each cluster and finds the graph in each cluster that is closest to the mean graph. The graphs closest to the mean graphs are used as the cluster centers for the next iteration. Handwriter stores the cluster centers of a cluster template as graph prototypes. A graph prototype consists of the starting and ending points of each path in the graph, as well as and evenly spaced points along each path. The prototype also stores the center point of the graph. All points are represented as xy-coordinates and the center point is at (0,0).

Usage

```
plot_cluster_centers(template, plot_graphs = FALSE, size = 100)
```

Arguments

template A cluster template created with make_clustering_template

plot_graphs TRUE plots all graphs in each cluster in addition to the cluster centers. FALSE

only plots the cluster centers.

size The size of the output plot

plot_cluster_fill_counts 25

Value

A plot

Examples

```
# plot cluster centers from example template
plot_cluster_centers(example_cluster_template)
plot_cluster_centers(example_cluster_template, plot_graphs = TRUE)
```

```
{\tt plot\_cluster\_fill\_counts}
```

Plot Cluster Fill Counts

Description

Plot the cluster fill counts for each document in formatted_data.

Usage

```
plot_cluster_fill_counts(formatted_data, facet = TRUE)
```

Arguments

Value

ggplot plot of cluster fill counts

```
# Plot cluster fill counts for template training documents
template_data <- format_template_data(example_cluster_template)
plot_cluster_fill_counts(formatted_data = template_data, facet = TRUE)

# Plot cluster fill counts for model training documents
plot_cluster_fill_counts(formatted_data = example_model, facet = TRUE)

# Plot cluster fill counts for questioned documents
plot_cluster_fill_counts(formatted_data = example_analysis, facet = FALSE)</pre>
```

```
plot_cluster_fill_rates

Plot Cluster Fill Rates
```

Description

Plot the cluster fill rates for each document in formatted_data.

Usage

```
plot_cluster_fill_rates(formatted_data, facet = FALSE)
```

Arguments

Value

ggplot plot of cluster fill rates

Examples

```
# Plot cluster fill rates for template training documents
template_data <- format_template_data(example_cluster_template)
plot_cluster_fill_rates(formatted_data = template_data, facet = TRUE)

# Plot cluster fill rates for model training documents
plot_cluster_fill_rates(formatted_data = example_model, facet = TRUE)

# Plot cluster fill rates for questioned documents
plot_cluster_fill_rates(formatted_data = example_analysis, facet = FALSE)</pre>
```

```
plot_credible_intervals
```

Plot Credible Intervals

Description

Plot credible intervals for the model's pi parameters that estimate the true writer cluster fill counts.

plot_graphs 27

Usage

```
plot_credible_intervals(
  model,
  interval_min = 0.025,
  interval_max = 0.975,
  facet = FALSE
)
```

Arguments

model A model created by fit_model()

interval_min The lower bound of the credible interval. It must be greater than zero and less

than 1.

interval_max The upper bound of the credible interval. It must be greater than the interval

minimum and less than 1.

facet TRUE uses facet_wrap to create a subplot for each writer. FALSE plots the data

on a single plot.

Value

ggplot plot credible intervals

Examples

```
plot_credible_intervals(model = example_model)
plot_credible_intervals(model = example_model, facet = TRUE)
```

plot_graphs

Plot Graphs

Description

Use processDocument() to split handwritting into component shapes called *graphs*. plot_graphs() creates a plot that displays the graphs. ggplot2::facet_wrap() places each graph in its own facet, and ncol sets the number of columns of facets.

Usage

```
plot_graphs(doc, ncol = NULL)
```

Arguments

doc A PNG image of handwriting processed with processDocument().

ncol Optionally, set the number of columns in the output plot. The default is NULL

which allows ggplot2::facet_wrap() to automatically choose the number of

columns.

Value

A plot of all graphs in the document

Examples

```
image_path <- system.file("extdata", "phrase_example.png", package = "handwriter")
doc <- processDocument(image_path)
plot_graphs(doc)</pre>
```

```
plot_posterior_probabilities
```

Plot Posterior Probabilities

Description

Creates a tile plot of posterior probabilities of writership for each questioned document and each known writer analyzed with analyze_questioned_documents().

Usage

```
plot_posterior_probabilities(analysis)
```

Arguments

analysis

A named list of analysis results from analyze_questioned_documents().

Value

A tile plot of posterior probabilities of writership.

```
plot_posterior_probabilities(analysis = example_analysis)
```

plot_trace 29

plot_trace

Plot Trace

Description

Create a trace plot for all chains for a single variable of a fitted model created by fit_model(). If the model contains more than one chain, the chains will be combined by pasting them together.

Usage

```
plot_trace(variable, model)
```

Arguments

variable The name of a variable in the model model A model created by fit_model()

Value

A trace plot

Examples

```
plot_trace(model = example_model, variable = "pi[1,1]")
plot_trace(model = example_model, variable = "mu[2,3]")
```

processDocument

Process Document

Description

Load a handwriting sample from a PNG image. Then binarize, thin, and split the handwriting into graphs.

Usage

```
processDocument(path)
```

Arguments

path

File path for handwriting document. The document must be in PNG file format.

Value

The processed document as a list

30 processHandwriting

Examples

```
image_path <- system.file("extdata", "phrase_example.png", package = "handwriter")
doc <- processDocument(image_path)
plotImage(doc)
plotImageThinned(doc)
plotNodes(doc)</pre>
```

processHandwriting

Process Handwriting by Component

Description

The main driver of handwriting processing. Takes in an image of thinned handwriting created with thinImage() and splits the the handwriting into shapes called *graphs*. Instead of processing the entire document at once, the thinned writing is separated into connected components and each component is split into graphs.

Usage

```
processHandwriting(img, dims)
```

Arguments

img Thinned binary image created with thinImage().

dims Dimensions of thinned binary image.

Value

A list of the processed image

```
twoSent_document <- list()
twoSent_document$image <- twoSent
twoSent_document$thin <- thinImage(twoSent_document$image)
twoSent_processList <- processHandwriting(twoSent_document$thin, dim(twoSent_document$image))</pre>
```

process_batch_dir 31

process_batch_dir

Process Batch Directory

Description

Process a list of handwriting samples saved as PNG images in a directory: (1) Load the image and convert it to black and white with readPNGBinary() (2) Thin the handwriting to one pixel in width with thinImage() (3) Run processHandwriting() to split the handwriting into parts called *edges* and place *nodes* at the ends of edges. Then combine edges into component shapes called *graphs*. (4) Save the processed document in an RDS file. (5) Optional. Return a list of the processed documents.

Usage

```
process_batch_dir(input_dir, output_dir = ".", skip_docs_on_retry = TRUE)
```

Arguments

Logical whether to skip documents in input_dir that caused errors on a previous run. The errors and document names are stored in output_dir > problems.txt. If this is the first run, process_batch_list will attempt to process all documents in input_dir.

Value

No return value, called for side effects

Examples

```
## Not run:
process_batch_dir("path/to/input_dir", "path/to/output_dir")
## End(Not run)
```

process_batch_list

Process Batch List

Description

Process a list of handwriting samples saved as PNG images: (1) Load the image and convert it to black and white with readPNGBinary() (2) Thin the handwriting to one pixel in width with thinImage() (3) Run processHandwriting() to split the handwriting into parts called *edges* and place *nodes* at the ends of edges. Then combine edges into component shapes called *graphs*. (4) Save the processed document in an RDS file. (5) Optional. Return a list of the processed documents.

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Usage

```
process_batch_list(images, output_dir, skip_docs_on_retry = TRUE)
```

Arguments

Logical whether to skip documents in the images arguement that caused errors on a previous run. The errors and document names are stored in output_dir > problems.txt. If this is the first run, process_batch_list will attempt to process all documents in the images arguement.

Value

No return value, called for side effects

Examples

```
## Not run:
images <- c('path/to/image1.png', 'path/to/image2.png', 'path/to/image3.png')
process_batch_list(images, "path/to/output_dir", FALSE)
process_batch_list(images, "path/to/output_dir", TRUE)
## End(Not run)</pre>
```

readPNGBinary

Read PNG Binary

Description

This function reads in and binarizes a PNG image.

Usage

```
readPNGBinary(
  path,
  cutoffAdjust = 0,
  clean = TRUE,
  crop = TRUE,
  inversion = FALSE
)
```

read_and_process 33

Arguments

path File path for image.

cutoffAdjust Multiplicative adjustment to the K-means estimated binarization cutoff.

clean Whether to fill in white pixels with 7 or 8 neighbors. This will help a lot when

thinning – keeps from getting little white bubbles in text.

crop Logical value dictating whether or not to crop the white out around the image.

TRUE by default.

inversion Logical value dictating whether or not to flip each pixel of binarized image.

Flipping happens after binarization. FALSE by default.

Value

Returns image from path. 0 represents black, and 1 represents white by default.

Examples

```
image_path <- system.file("extdata", "phrase_example.png", package = "handwriter")
csafe_document <- list()
csafe_document$image = readPNGBinary(image_path)
plotImage(csafe_document)</pre>
```

read_and_process

Read and Process

Description

[Superseded]

Development on read_and_process() is complete. We recommend using processDocument(). read_and_process(image_name, "document") is equivalent to processDocument(image_name).

Usage

```
read_and_process(image_name, transform_output)
```

Arguments

 $image_name \qquad \quad The \ file \ path \ to \ an \ image$

transform_output

The type of transformation to perform on the output

Value

A list of the processed image components

34 rgba2rgb

Examples

```
# use handwriting example from handwriter package
image_path <- system.file("extdata", "phrase_example.png", package = "handwriter")
doc <- read_and_process(image_path, "document")</pre>
```

rgb2grayscale

rgba2grayscale

Description

Changes RGB image to grayscale

Usage

```
rgb2grayscale(img)
```

Arguments

img

A 3D array with slices R, G, and B

Value

img as a 3D array as grayscale

rgba2rgb

rgba2rgb

Description

Removes alpha channel from png image.

Usage

```
rgba2rgb(img)
```

Arguments

img

A 3-d array with slices R, G, B, and alpha.

Value

img as a 3D array with alpha channel removed

thinImage 35

thinImage

thinImage

Description

This function returns a vector of locations for black pixels in the thinned image. Thinning done using Zhang - Suen algorithm.

Usage

```
thinImage(img)
```

Arguments

img

A binary matrix of the text that is to be thinned.

Value

A thinned, one pixel wide, image.

twoSent

Two sentence printed example handwriting

Description

Two sentence printed example handwriting

Usage

twoSent

Format

Binary image matrix. 396 rows and 1947 columns

```
twoSent_document <- list()
twoSent_document$image <- twoSent
plotImage(twoSent_document)

## Not run:
twoSent_document <- list()
twoSent_document$image <- twoSent
plotImage(twoSent_document)
twoSent_document$thin <- thinImage(twoSent_document$image)
plotImageThinned(twoSent_document)</pre>
```

36 whichToFill

```
twoSent_processList <- processHandwriting(twoSent_document$thin, dim(twoSent_document$image))
## End(Not run)</pre>
```

whichToFill

whichToFill

Description

Finds pixels in the plot that shouldn't be white and makes them black. Quick and helpful cleaning for before the thinning algorithm runs.

Usage

```
whichToFill(img)
```

Arguments

img

A binary matrix.

Value

A cleaned up image.

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