

Package ‘SuperpixelImageSegmentation’

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Type Package

Title Superpixel Image Segmentation

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BugReports <https://github.com/mlampros/SuperpixelImageSegmentation/issues>

URL <https://github.com/mlampros/SuperpixelImageSegmentation>

Description Image Segmentation using Superpixels, Affinity Propagation and Kmeans Clustering. The R code is based primarily on the article ``Image Segmentation using SLIC Superpixels and Affinity Propagation Clustering, Bao Zhou, International Journal of Science and Research (IJSR), 2013" <<https://www.ijsr.net/archive/v4i4/SUB152869.pdf>>.

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Depends R(>= 3.2)

Imports Rcpp (>= 0.12.10), R6, OpenImageR, grDevices, lattice

LinkingTo Rcpp, RcppArmadillo (>= 0.9.1), ClusterR, OpenImageR

Suggests testthat, covr, knitr, rmarkdown

RoxygenNote 7.1.2

NeedsCompilation yes

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Image_Segmentation *Segmentation of images based on Superpixels, Affinity propagation and Kmeans clustering*

Description

Segmentation of images based on Superpixels, Affinity propagation and Kmeans clustering

Segmentation of images based on Superpixels, Affinity propagation and Kmeans clustering

Usage

```
# init <- Image_Segmentation$new()
```

Details

sim_wL, *sim_wA*, *sim_wB* are the weights of the three channels. They keep balance so as to be consistent with human perception.

The quantity *colorradius* adjusts the number of clusters, and if its value is low, the number of targets would increase, which leads to more detailed segmentation results.

If the *adjust_centroids_and_return_masks* parameter is set to FALSE then the output *kmeans_image_data* will be an RGB image, otherwise it will be a black-and-white image.

colour_type parameter: RGB (Red-Green-Blue), LAB (Lightness, A-colour-dimension, B-colour-dimension) or HSV (Hue, Saturation, Value) colour.

Higher resolution images give better results.

The *affinity propagation* algorithm is used here with default parameter values.

By setting the *sim_normalize* parameter to TRUE, the affinity propagation algorithm requires less iterations to complete. However, the *colorradius* parameter does not have an effect if the similarity matrix is normalized.

Regarding the *use_median* parameter in the Rcpp I use the following steps: *1st.* I compute the superpixels and extract the labels, *2nd.* each superpixel label consists of multiple pixels and for these superpixels I have to compute a dissimilarity matrix therefore each superpixel must correspond to a single value, *3rd.* to come to this single value for each superpixel the R user has the option to either use the 'mean' or the 'median of multiple image pixels (per superpixel)

kmeans initializers

optimal_init : this initializer adds rows of the data incrementally, while checking that they do not already exist in the centroid-matrix

quantile_init : initialization of centroids by using the cumulative distance between observations and by removing potential duplicates

kmeans++ : kmeans++ initialization. Reference : <http://theory.stanford.edu/~sergei/papers/kMeansPP-soda.pdf> AND <http://stackoverflow.com/questions/5466323/how-exactly-does-k-means-work>

random : random selection of data rows as initial centroids

Methods

```
Image_Segmentation$new()  
_____  
spixel_segmentation()  
_____  
spixel_masks_show()  
_____  
spixel_clusters_show()  
_____
```

Methods**Public methods:**

- `Image_Segmentation$new()`
- `Image_Segmentation$spixel_segmentation()`
- `Image_Segmentation$spixel_masks_show()`
- `Image_Segmentation$spixel_clusters_show()`
- `Image_Segmentation$clone()`

Method new():

Usage:

```
Image_Segmentation$new()
```

Method spixel_segmentation():

Usage:

```
Image_Segmentation$spixel_segmentation(  
  input_image,  
  method = "slic",  
  superpixel = 200,  
  kmeans_method = "",  
  AP_data = FALSE,  
  use_median = TRUE,  
  minib_kmeans_batch = 10,  
  minib_kmeans_init_fraction = 0.5,  
  kmeans_num_init = 3,  
  kmeans_max_iters = 100,  
  kmeans_initializer = "kmeans++",  
  colour_type = "RGB",  
  compactness_factor = 20,  
  adjust_centroids_and_return_masks = FALSE,  
  return_labels_2_dimensionsional = FALSE,  
  sim_normalize = FALSE,  
  sim_wL = 3,  
  sim_wA = 10,
```

```

    sim_wB = 10,
    sim_color_radius = 20,
    ap_maxits = 1000,
    ap_convits = 100,
    ap_dampfact = 0.9,
    ap_nonoise = 0,
    verbose = FALSE
)

```

Arguments:

`input_image` a 3-dimensional input image (the range of the pixel values should be preferably in the range 0 to 255)
`method` a character string specifying the superpixel method. It can be either "slic" or "slico"
`superpixel` a numeric value specifying the number of superpixels
`kmeans_method` a character string specifying the kmeans method. If not empty ("") then it can be either "kmeans" or "mini_batch_kmeans"
`AP_data` a boolean. If TRUE then the affinity propagation image data will be computed and returned
`use_median` a boolean. If TRUE then the median will be used rather than the mean value for the inner computations (see the details section for more information)
`minib_kmeans_batch` the size of the mini batches
`minib_kmeans_init_fraction` percentage of data to use for the initialization centroids (applies if initializer is *kmeans++* or *optimal_init*). Should be a float number between 0.0 and 1.0.
`kmeans_num_init` number of times the algorithm will be run with different centroid seeds
`kmeans_max_iters` the maximum number of clustering iterations
`kmeans_initializer` the method of initialization. One of, *optimal_init*, *quantile_init*, *kmeans++* and *random*. See details for more information
`colour_type` a character string specifying the colour type. It can be one of "RGB", "LAB" or "HSV"
`compactness_factor` a numeric value specifying the compactness parameter in case that *method* is "slic"
`adjust_centroids_and_return_masks` a boolean. If TRUE and the *kmeans_method* parameter is NOT empty ("") then the centroids will be adjusted and image-masks will be returned. This will allow me to plot the masks using the *spixel_masks_show* method.
`return_labels_2_dimensionsional` a boolean. If TRUE then a matrix of labels based on the output superpixels in combination with the Affinity Propagation clusters will be returned
`sim_normalize` a boolean. If TRUE then the constructed similarity matrix will be normalised to have unit p-norm (see the armadillo documentation for more details)
`sim_wL` a numeric value specifying the weight for the "L" channel of the image (see the details section for more information)
`sim_wA` a numeric value specifying the weight for the "A" channel of the image (see the details section for more information)
`sim_wB` a numeric value specifying the weight for the "B" channel of the image (see the details section for more information)
`sim_color_radius` a numeric value specifying the *colorradius* (see the details section for more information)

ap_maxits a numeric value specifying the maximum number of iterations for the Affinity Propagation Clustering (defaults to 1000)
 ap_convits a numeric value. If the estimated exemplars stay fixed for convits iterations, the affinity propagation algorithm terminates early (defaults to 100)
 ap_dampfact a float number specifying the update equation damping level in [0.5, 1). Higher values correspond to heavy damping, which may be needed if oscillations occur in the Affinity Propagation Clustering (defaults to 0.9)
 ap_nonoise a float number. The affinity propagation algorithm adds a small amount of noise to *data* to prevent degenerate cases; this disables that.
 verbose a boolean. If TRUE then information will be printed in the console (spixel_masks_show method)

Method spixel_masks_show():

Usage:

```
Image_Segmentation$spixel_masks_show(
  delay_display_seconds = 3,
  display_all = FALSE,
  margin_btwn_plots = 0.15,
  verbose = FALSE
)
```

Arguments:

delay_display_seconds a numeric value specifying the seconds to delay the display of the next image (It displays the images consecutively). This parameter applies only if the *display_all* is set to FALSE (spixel_masks_show method)
 display_all a boolean. If TRUE then all images will be displayed in a grid (spixel_masks_show method)
 margin_btwn_plots a float number specifying the margins between the plots if the *display_all* parameter is set to TRUE (spixel_masks_show method)
 verbose a boolean. If TRUE then information will be printed in the console (spixel_masks_show method)

Method spixel_clusters_show():

Usage:

```
Image_Segmentation$spixel_clusters_show(
  spix_labels,
  color_palette = grDevices::rainbow,
  parameter_list_png = NULL
)
```

Arguments:

spix_labels a matrix. I can retrieve the "spix_labels" parameter by setting the "return_labels_2_dimensionsional" parameter to TRUE in the "spixel_segmentation" method (spixel_clusters_show method)
 color_palette one of the color palettes. Use ?grDevices::topo.colors to see the available color palettes
 parameter_list_png either NULL or a list of parameters passed to the ?grDevices::png function, such as list(filename = 'img.png', width = 100, height = 100, units = "px", pointsize = 12, bg = "white", type = "quartz")

Method clone(): The objects of this class are cloneable with this method.

Usage:

```
Image_Segmentation$clone(deep = FALSE)
```

Arguments:

deep Whether to make a deep clone.

References

<https://www.ijsr.net/archive/v4i4/SUB152869.pdf> , "Image Segmentation using SLIC Superpixels and Affinity Propagation Clustering", Bao Zhou, 2013, International Journal of Science and Research (IJSR)

Examples

```
library(SuperpixelImageSegmentation)

path = system.file("images", "BSR_bsds500_image.jpg", package = "SuperpixelImageSegmentation")

im = OpenImageR::readImage(path)

init = Image_Segmentation$new()

num_spix = 10           # for illustration purposes
# num_spix = 600         # recommended number of superpixels

spx = init$spixel_segmentation(input_image = im,
                                superpixel = num_spix,
                                AP_data = TRUE,
                                use_median = TRUE,
                                return_labels_2_dimensional = TRUE,
                                sim_color_radius = 10)

#.....
# plot the superpixel labels
#.....

plt = init$spixel_clusters_show(spix_labels = spx$spix_labels,
                                  color_palette = grDevices::rainbow,
                                  parameter_list_png = NULL)

# plt

#.....
# create a binary image for a specified cluster label
#.....

pix_values = spx$spix_labels

target_cluster = 3          # determine clusters visually ('plt' variable)
```

```
pix_values[pix_values != target_cluster] = 0      # set all other values to 0 (background)
pix_values[pix_values == target_cluster] = 1      # set the target_cluster to 1 (binary image)

# OpenImageR::imageShow(pix_values)
```

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