

Package ‘fsbrain’

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Type Package

Title Managing and Visualizing Brain Surface Data

Version 0.4.0

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Description Provides high-level access to 'FreeSurfer' <<http://freesurfer.net/>> neuroimaging data on the level of subjects and groups. Load morphometry data, surfaces and brain parcellations based on atlases. Mask data using labels, load data for specific atlas regions only, and visualize data and results directly in 'R'.

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Encoding UTF-8

LazyData true

URL <https://github.com/dfsp-spirit/fsbrain>

BugReports <https://github.com/dfsp-spirit/fsbrain/issues>

Imports reshape, freesurferformats (>= 0.1.12), pkgfilecache (>= 0.1.1), rgl, squash, fields, data.table

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R topics documented:

alphablend	5
annot.outline	6
apply.label.to.morphdata	7

apply.labeldata.to.morphdata	8
apply.transform	9
arrange.brainview.images	9
arrange.brainview.images.grid	10
brainviews	12
clip.data	13
cm.cbry	13
cm.div	14
cm.heat	14
cm.qual	15
cm.seq	15
collayer.bg	16
collayer.bg.atlas	16
collayer.bg.meancurv	18
collayer.bg.sulc	19
collayer.from.annot	20
collayer.from.annotdata	20
collayer.from.mask.data	21
collayer.from.morphlike.data	22
collayers.merge	23
coloredmesh.from.annot	23
coloredmesh.from.label	24
coloredmesh.from.mask	26
coloredmesh.from.morph.native	27
coloredmesh.from.morph.standard	28
coloredmesh.from.morphdata	30
coloredmesh.from.preloaded.data	31
coloredmesh.plot.colorbar.separate	31
coloredmeshes.from.color	33
colorlist.brain.clusters	34
combine.colorbar.with.brainview.animation	35
combine.colorbar.with.brainview.image	36
cube3D.tris	37
cubes3D.tris	38
delete_all_optional_data	39
demographics.to.fsgd.file	39
desaturate	40
download_fsaverage	41
download_fsaverage3	42
download_optional_data	42
download_optional_paper_data	43
export.coloredmesh.ply	43
face.edges	44
find.freesurferhome	45
find.subjectsdir.of	45
fs.coloredmesh	46
fs.home	47
fsaverage.path	47

fsbrain.set.default.figsize	48
fup	48
gen.test.volume	49
get.atlas.region.names	50
get.rglstyle	51
get.view.angle.names	51
getIn	52
get_optional_data_filepath	53
group.agg.atlas.native	53
group.agg.atlas.standard	55
group.annot	57
group.concat.measures.native	58
group.concat.measures.standard	59
group.label	60
group.label.from.annot	61
group.morph.agg.native	62
group.morph.agg.standard	64
group.morph.agg.standard.vertex	66
group.morph.native	67
group.morph.standard	68
group.morph.standard.sf	70
group.multimorph.agg.native	71
group.multimorph.agg.standard	72
group.surface	74
hasIn	75
hemilist.derive.hemi	76
hemilist.get.combined.data	76
hemilist.unwrap	77
hemilist.wrap	77
is.fs.coloredmesh	78
is.fs.coloredvoxels	78
is.hemilist	79
label.border	79
label.colFn	80
label.colFn.inv	81
label.from.annotdata	81
label.to.annot	82
labeldata.from.mask	83
limit_fun	84
limit_fun_na	85
list_optional_data	85
mask.from.labeldata.for.hemi	86
mesh.vertex.neighbors	87
mkco.cluster	88
mkco.div	89
mkco.heat	89
mkco.seq	90
principal.curvatures	90

print.fs.coloredmesh	91
print.fs.coloredvoxels	91
qc.for.group	92
qc.from.regionwise.df	93
qc.from.segstats.tables	94
qc.vis.failcount.by.region	94
ras2vox_tkr	95
read.colorcsv	96
read.md.demographics	96
read.md.subjects	97
read.md.subjects.from.fsgd	98
regions.to.ignore	99
report.on.demographics	99
rglactions	100
rglo	101
rglot	101
rglvoxels	102
shape.descriptor.names	103
shape.descriptors	103
shift.hemis.apart	104
spread.values.over.annot	105
spread.values.over.hemi	106
spread.values.over.subject	107
subject.annot	109
subject.atlas.agg	110
subject.filepath.morph.native	111
subject.filepath.morph.standard	112
subject.label	113
subject.label.from.annot	114
subject.lobes	116
subject.mask	117
subject.morph.native	118
subject.morph.standard	119
subject.num.verts	121
subject.surface	122
subject.volume	123
vdata.split.by.hemi	124
vis.color.on.subject	125
vis.coloredmeshes	126
vis.coloredmeshes.rotating	127
vis.colortable.legend	129
vis.data.on.fsaverage	129
vis.data.on.group.native	131
vis.data.on.group.standard	133
vis.data.on.subject	134
vis.export.from.coloredmeshes	136
vis.fs.surface	138
vis.group.annot	139

vis.group.coloredmeshes	140
vis.group.morph.native	141
vis.group.morph.standard	142
vis.labeldata.on.subject	144
vis.mask.on.subject	146
vis.path.along.verts	148
vis.region.values.on.subject	149
vis.subject.annot	151
vis.subject.label	153
vis.subject.morph.native	155
vis.subject.morph.standard	157
vis.symmetric.data.on.subject	159
vislayout.from.coloredmeshes	162
vol.boundary.box	163
vol.boundary.box.apply	164
vol.hull	165
vol.imagestack	165
vol.intensity.to.color	166
vol.mask.from.segmentation	167
vol.merge	168
vol.overlay.colors.from.activation	169
vol.overlay.colors.from.colortable	169
vol.planes	170
vol.slice	171
vol.vox.from.crs	172
volvis.contour	172
volvis.lightbox	173
volvis.voxels	174
vox2ras_tkr	176
write.group.morph.standard.sf	176
write.region.aggregated	177
write.region.values	178
write.region.values.fsaverage	179

Index**182**

alphablend	<i>Perform alpha blending for pairs of RGBA colors.</i>
------------	---

Description

Implements the **over** alpha blending operation.

Usage

```
alphablend(front_color, back_color, silent = TRUE)
```

Arguments

front_color	rgba color strings, the upper color layer or foreground
back_color	rgba color strings, the lower color layer or background
silent	logical, whether to suppress messages

Value

rgba color strings, the alpha-blended colors

References

see the **Alpha blending** section on https://en.wikipedia.org/wiki/Alpha_compositing

See Also

Other color functions: [desaturate\(\)](#)

annot.outline	<i>Compute outline vertex colors from annotation.</i>
---------------	---

Description

For each region in an atlas, compute the outer border and color the respective vertices in the region-specific color from the annot's colortable.

Usage

```
annot.outline(
  annotdata,
  surface_mesh,
  background = "white",
  silent = TRUE,
  expand_inwards = 0L,
  outline_color = NULL,
  limit_to_regions = NULL
)
```

Arguments

annotdata	an annotation, as returned by functions like subject.annot .
surface_mesh	brain surface mesh, as returned by functions like subject.surface or read.fs.surface .
background	color, the background color to assign to the non-border parts of the regions. Defaults to 'white'.
silent	logical, whether to suppress status messages.
expand_inwards	integer, additional thickness of the borders. Increases computation time, defaults to 0L.

`outline_color` NULL or a color string (like 'black' or '#000000'), the color to use for the borders. If left at the default value 'NULL', the colors from the annotation color lookup table will be used.

`limit_to_regions` vector of character strings or NULL, a list of regions for which to draw the outline (see [get.atlas.region.names](#)). If NULL, all regions will be used. If (and only if) this parameter is used, the 'outline_color' parameter can be a vector of color strings, one color per region.

Value

vector of colors, one color for each mesh vertex

Note

Sorry for the computational time, the mesh datastructure is not ideal for neighborhood search.

`apply.label.to.morphdata`

Load a label from file and apply it to morphometry data.

Description

This function will set all values in morphdata which are *not* part of the label loaded from the file to NA (or whatever is specified by 'masked_data_value'). This is typically used to ignore values which are not part of the cortex (or any other label) during your analysis.

Usage

```
apply.label.to.morphdata(
  morphdata,
  subjects_dir,
  subject_id,
  hemi,
  label,
  masked_data_value = NA
)
```

Arguments

`morphdata` numerical vector, the morphometry data for one hemisphere

`subjects_dir` string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.

`subject_id` string. The subject identifier

`hemi` string, one of 'lh', 'rh' or 'both'. The hemisphere name. Used to construct the names of the annotation and morphometry data files to be loaded.

label	string, 'fs.label' instance, or label vertex data. If a string, interpreted as the file name of the label file, without the hemi part (if any), optionally including the '.label' suffix. E.g., 'cortex.label' or 'cortex' for '?h.cortex.label'.
masked_data_value	numerical, the value to set for all morphometry data values of vertices which are <i>*not*</i> part of the label. Defaults to NA.

Value

numerical vector, the masked data.

See Also

Other label functions: [apply.labeldata.to.morphdata\(\)](#), [subject.lobes\(\)](#), [subject.mask\(\)](#), [vis.labeldata.on.subject\(\)](#), [vis.subject.label\(\)](#)

Other morphometry data functions: [apply.labeldata.to.morphdata\(\)](#), [group.morph.native\(\)](#), [group.morph.standard\(\)](#), [subject.morph.native\(\)](#), [subject.morph.standard\(\)](#)

`apply.labeldata.to.morphdata`

Apply a label to morphometry data.

Description

This function will set all values in `morphdata` which are **not** part of the `labeldata` to NA (or whatever is specified by `'masked_data_value'`). This is typically used to ignore values which are not part of the cortex (or any other label) during your analysis.

Usage

```
apply.labeldata.to.morphdata(morphdata, labeldata, masked_data_value = NA)
```

Arguments

<code>morphdata</code>	numerical vector, the morphometry data for one hemisphere
<code>labeldata</code>	integer vector or 'fs.label' instance. A label as returned by subject.label .
<code>masked_data_value</code>	numerical, the value to set for all morphometry data values of vertices which are <i>*not*</i> part of the label. Defaults to NA.

Value

numerical vector, the masked data.

See Also

Other label functions: [apply.label.to.morphdata\(\)](#), [subject.lobes\(\)](#), [subject.mask\(\)](#), [vis.labeldata.on.subject.vis.subject.label\(\)](#)

Other morphometry data functions: [apply.label.to.morphdata\(\)](#), [group.morph.native\(\)](#), [group.morph.standard\(\)](#), [subject.morph.native\(\)](#), [subject.morph.standard\(\)](#)

apply.transform	<i>Apply matmult transformation to input.</i>
-----------------	---

Description

Apply affine transformation, like a **vox2ras_tkr** transformation, to input. This is just matrix multiplication for different input objects.

Usage

```
apply.transform(object, matrix_fun)
```

Arguments

object	numerical vector/matrix or Triangles3D instance, the coordinates or object to transform.
matrix_fun	a 4x4 affine matrix or a function returning such a matrix. If 'NULL', the input is returned as-is. In many cases you may want to use a matrix computed from the header of a volume file, e.g., the 'vox2ras' matrix of the respective volume. See the 'mghheader.*' functions in the <i>*freesurferformats*</i> package to obtain these matrices.

Value

the input after application of the affine matrix (matrix multiplication)

arrange.brainview.images	<i>Combine several brainview images into a new figure.</i>
--------------------------	--

Description

Create a new image from several image tiles, the exact layout depends on the number of given images.

Usage

```

arrange.brainview.images(
  brainview_images,
  output_img,
  colorbar_img = NULL,
  silent = TRUE,
  grid_like = TRUE,
  border_geometry = "5x5",
  background_color = "white"
)

```

Arguments

brainview_images	vector of character strings, paths to the brainview images, usually in PNG format
output_img	path to output image that including the file extension
colorbar_img	path to the main image containing the separate colorbar, usually an image in PNG format
silent	logical, whether to suppress messages
grid_like	logical, whether to arrange the images in a grid-like fashion. If FALSE, they will all be merged horizontally.
border_geometry	string, a geometry string passed to <code>magick::image_border</code> to define the borders to add to each image tile. The default value adds 5 pixels, both horizontally and vertically.
background_color	string, a valid ImageMagick color string such as "white" or "#000080". The color to use when extending images (e.g., when creating the border).

Value

named list with entries: 'brainview_images': vector of character strings, the paths to the input images. 'output_img_path': character string, path to the output image. 'merged_img': the magick image instance.

```
arrange.brainview.images.grid
```

Combine several brainview images as a grid into a new figure.

Description

Create a new image from several image tiles, the exact layout is a grid with n per row.

Usage

```

arrange.brainview.images.grid(
    brainview_images,
    output_img,
    colorbar_img = NULL,
    silent = TRUE,
    num_per_row = 10L,
    border_geometry = "5x5",
    background_color = "white",
    captions = NULL
)

```

Arguments

brainview_images	vector of character strings, paths to the brainview images, usually in PNG format
output_img	path to output image that including the file extension
colorbar_img	path to the main image containing the separate colorbar, usually an image in PNG format
silent	logical, whether to suppress messages
num_per_row	positive integer, the number of image tiles per row.
border_geometry	string, a geometry string passed to <code>magick::image_border</code> to define the borders to add to each image tile. The default value adds 5 pixels, both horizontally and vertically.
background_color	string, a valid ImageMagick color string such as "white" or "#000080". The color to use when extending images (e.g., when creating the border).
captions	vector of character strings or NULL, the (optional) text annotations for the images. Useful to print the subject identifier onto the individual tiles. Length must match number of image tiles in 'brainview_images'.

Value

named list with entries: 'brainview_images': vector of character strings, the paths to the input images. 'output_img_path': character string, path to the output image. 'merged_img': the magick image instance.

Note

The tiles are written row-wise, in the order in which they occur in the parameter 'brainview_images'.

 brainviews

Show one or more views of the given meshes in rgl windows.

Description

Show one or more views of the given meshes in rgl windows.

Usage

```
brainviews(
  views,
  coloredmeshes,
  rgloptions = rglo(),
  rglactions = list(),
  style = "default",
  draw_colorbar = FALSE
)
```

Arguments

views	list of strings. Valid entries include: 'si': single interactive view. 'sd_<angle>': single view from angle <angle>. The <angle> part must be one of the strings returned by get.view.angle.names . Example: 'sd_caudal'. 'sr': single rotating view. 't4': tiled view showing the brain from 4 angles. 't9': tiled view showing the brain from 9 angles.
coloredmeshes	list of coloredmesh or renderable. A coloredmesh is a named list as returned by the coloredmesh.from.* functions. It has the entries 'mesh' of type tmesh3d, a 'col', which is a color specification for such a mesh.
rgloptions	option list passed to par3d . Example: rgloptions = list("windowRect"=c(50,50,1000,1000))
rglactions	named list. A list in which the names are from a set of pre-defined actions. The values can be used to specify parameters for the action.
style	character string, a rendering style, e.g., 'default', 'shiny' or 'semitransparent'.
draw_colorbar	logical, whether to draw a colorbar. WARNING: The colorbar is drawn to a subplot, and this only works if there is enough space for it. You will have to increase the plot size using the 'rgloptions' parameter for the colorbar to show up. Defaults to FALSE. See coloredmesh.plot.colorbar.separate for an alternative.

Value

list of coloredmeshes. The coloredmeshes used for the visualization.

See Also

[get.view.angle.names](#)

clip.data	<i>Clip data at quantiles to remove outliers.</i>
-----------	---

Description

Set all data values outside the given quantile range to the border values. This is useful to properly visualize morphometry data that includes outliers. These outliers negatively affect the colormap, as all the non-outlier values become hard to distinguish. This function can be used to filter the data before plotting it.

Usage

```
clip.data(data, lower = 0.05, upper = 0.95)
```

Arguments

data,	numeric vector. The input data. Can also be a hemi list.
lower,	numeric. The probability for the lower quantile, defaults to '0.05'.
upper,	numeric. The probability for the upper quantile, defaults to '0.95'.

Value

numeric vector. The output data.

Examples

```
full_data = rnorm(50, 3, 1);  
clipped = clip.data(full_data);
```

cm.cbry	<i>Get cyan blue red yellow colormap function.</i>
---------	--

Description

Get cyan blue red yellow colormap function.

Usage

```
cm.cbry()
```

Note

Returns a diverging palette with negative values in blue/cyan and positive ones in red/yellow, suitable for visualizing data that is centered around zero. Often used for clusters in neuroscience.

cm.div *Return the standard fsbrain diverging colormap.*

Description

Return the standard fsbrain diverging colormap.

Usage

```
cm.div(report = FALSE)
```

Arguments

report logical, whether to print a message with a name of the chosen colormap, in format `package::function#palette`.

Note

Returns some diverging palette, suitable for visualizing data that is centered around zero.

cm.heat *Return the standard fsbrain heat colormap.*

Description

Return the standard fsbrain heat colormap.

Usage

```
cm.heat(report = FALSE)
```

Arguments

report logical, whether to print a message with a name of the chosen colormap, in format `package::function#palette`.

Note

The heat palette is a sequential, single-hue palette.

cm.qual	<i>Return the standard fsbrain qualitative colormap.</i>
---------	--

Description

Return the standard fsbrain qualitative colormap.

Usage

```
cm.qual(report = FALSE)
```

Arguments

report	logical, whether to print a message with a name of the chosen colormap, in format <code>package::function#palette</code> .
--------	--

Note

Returns some qualitative palette, suitable for visualizing categorical data.

cm.seq	<i>Return the standard fsbrain sequential colormap.</i>
--------	---

Description

Return the standard fsbrain sequential colormap.

Usage

```
cm.seq(report = FALSE)
```

Arguments

report	logical, whether to print a message with a name of the chosen colormap, in format <code>package::function#palette</code> .
--------	--

Note

This returns a sequential, multi-hue palette.

collayer.bg *Compute binarized mean curvature surface color layer.*

Description

Compute a binarized mean curvature surface color layer, this is intended as a background color layer. You can merge it with your data layer using [collayers.merge](#).

Usage

```
collayer.bg(subjects_dir, subject_id, bg, hemi = "both")
```

Arguments

subjects_dir	character string, the FreeSurfer SUBJECTS_DIR.
subject_id	character string, the subject identifier.
bg	character string, a background name. One of 'curv', 'curv_light', 'sulc', 'sulc_light', or 'aparc'. If this is already a colorlayer in a hemilist, it will be returned as-is.
hemi	character string, one of 'lh', 'rh', or 'both'. The latter will merge the data for both hemis into a single vector.

Value

a color layer, i.e., vector of color strings in a hemilist

See Also

You can plot the return value using [vis.color.on.subject](#).

Other surface color layer: [collayer.bg.atlas\(\)](#), [collayer.bg.meancurv\(\)](#), [collayer.bg.sulc\(\)](#), [collayer.from.annotdata\(\)](#), [collayer.from.annot\(\)](#), [collayer.from.mask.data\(\)](#), [collayer.from.morphlike.d](#), [collayers.merge\(\)](#)

collayer.bg.atlas *Compute atlas or annotation surface color layer.*

Description

Compute atlas or annotation surface color layer.

Usage

```
collayer.bg.atlas(
  subjects_dir,
  subject_id,
  hemi = "both",
  atlas = "aparc",
  grayscale = FALSE,
  outline = FALSE,
  outline_surface = "white"
)
```

Arguments

subjects_dir	character string, the FreeSurfer SUBJECTS_DIR.
subject_id	character string, the subject identifier.
hemi	character string, one of 'lh', 'rh', or 'both'. The latter will merge the data for both hemis into a single vector.
atlas	character string, the atlas name. E.g., "aparc", "aparc.2009s", or "aparc.DKTatlas". Used to construct the name of the annotation file to be loaded.
grayscale	logical, whether to convert the atlas colors to grayscale
outline	logical, whether to draw an outline only instead of filling the regions. Defaults to 'FALSE'. Instead of passing 'TRUE', one can also pass a list of extra parameters to pass to <code>annot.outline</code> , e.g., <code>outline=list('outline_color'='#000000')</code> .
outline_surface	character string, the surface to load. Only relevant when 'outline' is used. (In that case the surface mesh is needed to compute the vertices forming the region borders.)

Value

a color layer, i.e., vector of color strings in a hemilist

Note

Using 'outline' mode is quite slow, and increasing the border thickness makes it even slower.

See Also

You can plot the return value using `vis.color.on.subject`.

Other surface color layer: `collayer.bg.meancurv()`, `collayer.bg.sulc()`, `collayer.bg()`, `collayer.from.annotdata()`, `collayer.from.annot()`, `collayer.from.mask.data()`, `collayer.from.morphlike.d`, `collayers.merge()`

collayer.bg.meancurv *Compute binarized mean curvature surface color layer.*

Description

Compute a binarized mean curvature surface color layer, this is intended as a background color layer. You can merge it with your data layer using [collayers.merge](#).

Usage

```
collayer.bg.meancurv(  
  subjects_dir,  
  subject_id,  
  hemi = "both",  
  cortex_only = FALSE,  
  bin_colors = c("#898989", "#5e5e5e"),  
  bin_thresholds = c(0)  
)
```

Arguments

subjects_dir	character string, the FreeSurfer SUBJECTS_DIR.
subject_id	character string, the subject identifier.
hemi	character string, one of 'lh', 'rh', or 'both'. The latter will merge the data for both hemis into a single vector.
cortex_only	logical, whether to restrict pattern computation to the cortex.
bin_colors	vector of two character strings, the two colors to use.
bin_thresholds	vector of 1 or 2 double values, the curvature threshold values used to separate gyri from sulci.

Value

a color layer, i.e., vector of color strings in a hemilist

See Also

You can plot the return value using [vis.color.on.subject](#).

Other surface color layer: [collayer.bg.atlas\(\)](#), [collayer.bg.sulc\(\)](#), [collayer.bg\(\)](#), [collayer.from.annotdata\(\)](#), [collayer.from.annot\(\)](#), [collayer.from.mask.data\(\)](#), [collayer.from.morphlike.data\(\)](#), [collayers.merge\(\)](#)

collayer.bg.sulc *Compute binarized sulcal depth surface color layer.*

Description

Compute a binarized sulcal depth surface color layer, this is intended as a background color layer. You can merge it with your data layer using [collayers.merge](#).

Usage

```
collayer.bg.sulc(  
  subjects_dir,  
  subject_id,  
  hemi = "both",  
  cortex_only = FALSE,  
  bin_colors = c("#898989", "#5e5e5e"),  
  bin_thresholds = c(0)  
)
```

Arguments

`subjects_dir` character string, the FreeSurfer SUBJECTS_DIR.
`subject_id` character string, the subject identifier.
`hemi` character string, one of 'lh', 'rh', or 'both'. The latter will merge the data for both hemis into a single vector.
`cortex_only` logical, whether to restrict pattern computation to the cortex.
`bin_colors` vector of two character strings, the two colors to use.
`bin_thresholds` vector of 1 or 2 double values, the curvature threshold values used to separate gyri from sulci.

Value

a color layer, i.e., vector of color strings in a hemilist

See Also

You can plot the return value using [vis.color.on.subject](#).

Other surface color layer: [collayer.bg.atlas\(\)](#), [collayer.bg.meancurv\(\)](#), [collayer.bg\(\)](#), [collayer.from.annotdata\(\)](#), [collayer.from.annot\(\)](#), [collayer.from.mask.data\(\)](#), [collayer.from.morphlike.d](#), [collayers.merge\(\)](#)

collayer.from.annot *Compute surface color layer from annotation or atlas data.*

Description

Compute surface color layer from annotation or atlas data.

Usage

```
collayer.from.annot(subjects_dir, subject_id, hemi, atlas)
```

Arguments

subjects_dir	character string, the FreeSurfer SUBJECTS_DIR.
subject_id	character string, the subject identifier.
hemi	character string, one of 'lh', 'rh', or 'both'.
atlas	character string, the atlas name. E.g., "aparc", "aparc.2009s", or "aparc.DKTatlas". Used to construct the name of the annotation file to be loaded.

Value

named hemi list, each entry is a vector of color strings, one color per surface vertex. The coloring represents the atlas data.

See Also

You can plot the return value using [vis.color.on.subject](#).

Other surface color layer: [collayer.bg.atlas\(\)](#), [collayer.bg.meancurv\(\)](#), [collayer.bg.sulc\(\)](#), [collayer.bg\(\)](#), [collayer.from.annotdata\(\)](#), [collayer.from.mask.data\(\)](#), [collayer.from.morphlike.data\(\)](#), [collayers.merge\(\)](#)

collayer.from.annotdata
Compute surface color layer from annotation or atlas data.

Description

Compute surface color layer from annotation or atlas data.

Usage

```
collayer.from.annotdata(lh_annotdata = NULL, rh_annotdata = NULL)
```

Arguments

lh_annotdata loaded annotation data for left hemi, as returned by `subject.annot`
rh_annotdata loaded annotation data for right hemi

Value

named hemi list, each entry is a vector of color strings, one color per surface vertex. The coloring represents the atlas data.

See Also

You can plot the return value using `vis.color.on.subject`.

Other surface color layer: `collayer.bg.atlas()`, `collayer.bg.meancurv()`, `collayer.bg.sulc()`, `collayer.bg()`, `collayer.from.annot()`, `collayer.from.mask.data()`, `collayer.from.morphlike.data()`, `collayers.merge()`

`collayer.from.mask.data`

Compute surface color layer from morph-like data.

Description

Compute surface color layer from morph-like data.

Usage

```
collayer.from.mask.data(
  lh_data = NULL,
  rh_data = NULL,
  makecmap_options = list(colFn = label.colFn)
)
```

Arguments

lh_data integer vector, can be NULL
rh_data numerical vector, can be NULL
makecmap_options named list of parameters to pass to `makecmap`. Must not include the unnamed first parameter, which is derived from 'measure'.

Value

named hemi list, each entry is a vector of color strings, one color per surface vertex. The coloring represents the label data.

See Also

You can plot the return value using [vis.color.on.subject](#).

Other surface color layer: [collayer.bg.atlas\(\)](#), [collayer.bg.meancurv\(\)](#), [collayer.bg.sulc\(\)](#), [collayer.bg\(\)](#), [collayer.from.annotdata\(\)](#), [collayer.from.annot\(\)](#), [collayer.from.morphlike.data\(\)](#), [collayers.merge\(\)](#)

collayer.from.morphlike.data

Compute surface color layer from morph-like data.

Description

Compute surface color layer from morph-like data.

Usage

```
collayer.from.morphlike.data(
  lh_morph_data = NULL,
  rh_morph_data = NULL,
  makecmap_options = list(colFn = cm.seq()),
  return_metadata = FALSE
)
```

Arguments

`lh_morph_data` numerical vector, can be NULL

`rh_morph_data` numerical vector, can be NULL

`makecmap_options` named list of parameters to pass to [makecmap](#). Must not include the unnamed first parameter, which is derived from 'measure'.

`return_metadata` logical, whether to return additional metadata as entry 'metadata' in the returned list

Value

named hemi list, each entry is a vector of color strings, one color per surface vertex. The coloring represents the morph data.

See Also

You can plot the return value using [vis.color.on.subject](#).

Other surface color layer: [collayer.bg.atlas\(\)](#), [collayer.bg.meancurv\(\)](#), [collayer.bg.sulc\(\)](#), [collayer.bg\(\)](#), [collayer.from.annotdata\(\)](#), [collayer.from.annot\(\)](#), [collayer.from.mask.data\(\)](#), [collayers.merge\(\)](#)

collayers.merge *Merge two or more color layers based on their transparency values.*

Description

Merge several color layers into one based on their transparency and alpha blending. In the final result, the lower layers are visible through the transparent or 'NA' parts (if any) of the upper layers.

Usage

```
collayers.merge(collayers, opaque_background = "#FFFFFF")
```

Arguments

`collayers` named list, the values must be vectors, matrices or arrays of color strings (as produced by [rgb](#)). The names are freeform and do not really matter. All values must have the same length.

`opaque_background` a single color string or 'NULL'. If a color string, this color will be used as a final opaque background layer to ensure that the returned colors are all opaque. Pass 'NULL' to skip this, which may result in a return value that contains non-opaque color values.

Value

a color layer, i.e., vector of color strings in a hemilist

See Also

Other surface color layer: [collayer.bg.atlas\(\)](#), [collayer.bg.meancurv\(\)](#), [collayer.bg.sulc\(\)](#), [collayer.bg\(\)](#), [collayer.from.annotdata\(\)](#), [collayer.from.annot\(\)](#), [collayer.from.mask.data\(\)](#), [collayer.from.morphlike.data\(\)](#)

`coloredmesh.from.annot` *Create a coloredmesh from an annotation of an atlas.*

Description

Create a coloredmesh from an annotation of an atlas.

Usage

```
coloredmesh.from.annot(
    subjects_dir,
    subject_id,
    atlas,
    hemi,
    surface = "white",
    outline = FALSE
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	string. The subject identifier.
atlas	string or a loaded annotation. If a string, interpreted as the atlas name that should be loaded to get the annotation. E.g., "aparc", "aparc.2009s", or "aparc.DKAtlas". Used to construct the name of the annotation file to be loaded.
hemi	string, one of 'lh' or 'rh'. The hemisphere name. Used to construct the names of the label data files to be loaded.
surface	character string or 'fs.surface' instance. The display surface. E.g., "white", "pial", or "inflated". Defaults to "white".
outline	logical, whether to draw an outline only instead of filling the regions. Defaults to FALSE. Only makes sense if you did not pass an outline already. The current implementation for outline computation is rather slow, so setting this to TRUE will considerably increase computation time.

Value

coloredmesh. A named list with entries: "mesh" the `tmesh3d` mesh object. "col": the mesh colors. "render", logical, whether to render the mesh. "hemi": the hemisphere, one of 'lh' or 'rh'.

See Also

Other coloredmesh functions: [coloredmesh.from.label\(\)](#), [coloredmesh.from.mask\(\)](#), [coloredmesh.from.morph.nat](#), [coloredmesh.from.morph.standard\(\)](#), [coloredmesh.from.morphdata\(\)](#), [coloredmeshes.from.color\(\)](#)

coloredmesh.from.label

Create a coloredmesh from a label.

Description

Create a coloredmesh from a label.

Usage

```
coloredmesh.from.label(
    subjects_dir,
    subject_id,
    label,
    hemi,
    surface = "white",
    colormap = NULL,
    makecmap_options = list(colFn = squash::rainbow2)
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	string. The subject identifier.
label	string or vector of integers. If a string, the name of the label file, without the hemi part (if any), but including the '.label' suffix. E.g., 'cortex.label' for '?h.cortex.label'. Alternatively, the already loaded label data as a vector of integers.
hemi	string, one of 'lh' or 'rh'. The hemisphere name. Used to construct the names of the label data files to be loaded.
surface	character string or 'fs.surface' instance. The display surface. E.g., "white", "pial", or "inflated". Defaults to "white".
colormap	a colormap function. See the squash package for some colormaps. Defaults to jet .
makecmap_options	named list of parameters to pass to makecmap . Must not include the unnamed first parameter, which is derived from 'measure'.

Value

coloredmesh. A named list with entries: "mesh" the [tmesh3d](#) mesh object. "col": the mesh colors. "render", logical, whether to render the mesh. "hemi": the hemisphere, one of 'lh' or 'rh'.

See Also

Other coloredmesh functions: [coloredmesh.from.annot\(\)](#), [coloredmesh.from.mask\(\)](#), [coloredmesh.from.morph.nat](#), [coloredmesh.from.morph.standard\(\)](#), [coloredmesh.from.morphdata\(\)](#), [coloredmeshes.from.color\(\)](#)

coloredmesh.from.mask *Create a coloredmesh from a mask.*

Description

Create a coloredmesh from a mask.

Usage

```
coloredmesh.from.mask(
  subjects_dir,
  subject_id,
  mask,
  hemi,
  surface = "white",
  colormap = NULL,
  surface_data = NULL,
  makecmap_options = list(colFn = squash::rainbow2)
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	string. The subject identifier.
mask	logical vector, contains one logical value per vertex.
hemi	string, one of 'lh' or 'rh'. The hemisphere name. Used to construct the names of the label data files to be loaded.
surface	character string or 'fs.surface' instance. The display surface. E.g., "white", "pial", or "inflated". Defaults to "white".
colormap	a colormap function. See the squash package for some colormaps. Defaults to jet .
surface_data	optional surface mesh object, as returned by subject.surface . If given, used instead of loading the surface data from disk (which users of this function may already have done). Defaults to NULL.
makecmap_options	named list of parameters to pass to makecmap . Must not include the unnamed first parameter, which is derived from 'measure'.

Value

coloredmesh. A named list with entries: "mesh" the [tmesh3d](#) mesh object. "col": the mesh colors. "render", logical, whether to render the mesh. "hemi": the hemisphere, one of 'lh' or 'rh'.

See Also

Other mask functions: [mask.from.labeldata.for.hemi\(\)](#), [vis.mask.on.subject\(\)](#)

Other coloredmesh functions: [coloredmesh.from.annot\(\)](#), [coloredmesh.from.label\(\)](#), [coloredmesh.from.morph.native\(\)](#), [coloredmesh.from.morph.standard\(\)](#), [coloredmesh.from.morphdata\(\)](#), [coloredmeshes.from.color\(\)](#)

coloredmesh.from.morph.native

Create a coloredmesh from native space morphometry data.

Description

Create a coloredmesh from native space morphometry data.

Usage

```
coloredmesh.from.morph.native(
    subjects_dir,
    subject_id,
    measure,
    hemi,
    surface = "white",
    colormap = NULL,
    clip = NULL,
    cortex_only = FALSE,
    makecmap_options = mkco.seq()
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	string. The subject identifier.
measure	string. The morphometry data to use. E.g., 'area' or 'thickness'. Pass NULL to render the surface in white, without any data. One can also pass the pre-loaded morphometry data as a numerical vector, the length of which must match the number of surface vertices.
hemi	string, one of 'lh' or 'rh'. The hemisphere name. Used to construct the names of the label data files to be loaded.
surface	character string or 'fs.surface' instance. The display surface. E.g., "white", "pial", or "inflated". Defaults to "white".
colormap	a colormap function. See the squash package for some colormaps. Defaults to jet .
clip	numeric vector of length 2 or NULL. If given, the 2 values are interpreted as lower and upper percentiles, and the morph data is clipped at the given lower and upper percentile (see clip.data). Defaults to NULL (no data clipping).

cortex_only	logical, whether to mask the medial wall, i.e., whether the morphometry data for all vertices which are <i>*not*</i> part of the cortex (as defined by the label file 'label/?h.cortex.label') should be replaced with NA values. In other words, setting this to TRUE will ignore the values of the medial wall between the two hemispheres. If set to true, the mentioned label file needs to exist for the subject. Defaults to FALSE.
makecmap_options	named list of parameters to pass to <code>makecmap</code> . Must not include the unnamed first parameter, which is derived from 'measure'.

Value

coloredmesh. A named list with entries: "mesh" the `tmesh3d` mesh object. "col": the mesh colors. "render", logical, whether to render the mesh. "hemi": the hemisphere, one of 'lh' or 'rh'.

See Also

Other coloredmesh functions: `coloredmesh.from.annot()`, `coloredmesh.from.label()`, `coloredmesh.from.mask()`, `coloredmesh.from.morph.standard()`, `coloredmesh.from.morphdata()`, `coloredmeshes.from.color()`

coloredmesh.from.morph.standard

Create a coloredmesh from standard space morphometry data.

Description

Create a coloredmesh from standard space morphometry data.

Usage

```
coloredmesh.from.morph.standard(
  subjects_dir,
  subject_id,
  measure,
  hemi,
  fwhm,
  surface = "white",
  template_subject = "fsaverage",
  template_subjects_dir = NULL,
  colormap = NULL,
  clip = NULL,
  cortex_only = FALSE,
  makecmap_options = mkco.seq()
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	string. The subject identifier.
measure	string. The morphometry data to use. E.g., 'area' or 'thickness'. Pass NULL to render the surface in white, without any data. One can also pass the pre-loaded morphometry data as a numerical vector, the length of which must match the number of surface vertices.
hemi	string, one of 'lh' or 'rh'. The hemisphere name. Used to construct the names of the label data files to be loaded.
fwhm	string, smoothing setting. The smoothing part of the filename, typically something like '0', '5', '10', ..., or '25'.
surface	character string or 'fs.surface' instance. The display surface. E.g., "white", "pial", or "inflated". Defaults to "white".
template_subject	The template subject used. This will be used as part of the filename, and its surfaces are loaded for data visualization. Defaults to 'fsaverage'.
template_subjects_dir	The template subjects dir. If 'NULL', the value of the parameter 'subjects_dir' is used. Defaults to NULL. If you have FreeSurfer installed and configured, and are using the standard fsaverage subject, try passing the result of calling 'file.path(Sys.getenv('FREESURFER_HOME'), 'subjects')'.
colormap	a colormap function. See the squash package for some colormaps. Defaults to jet .
clip	numeric vector of length 2 or NULL. If given, the 2 values are interpreted as lower and upper percentiles, and the morph data is clipped at the given lower and upper percentile (see clip.data). Defaults to NULL (no data clipping).
cortex_only	logical, whether to mask the medial wall, i.e., whether the morphometry data for all vertices which are *not* part of the cortex (as defined by the label file 'label/?h.cortex.label') should be replaced with NA values. In other words, setting this to TRUE will ignore the values of the medial wall between the two hemispheres. If set to true, the mentioned label file needs to exist for the subject. Defaults to FALSE.
makecmap_options	named list of parameters to pass to makecmap . Must not include the unnamed first parameter, which is derived from 'measure'.

Value

coloredmesh. A named list with entries: "mesh" the [tmesh3d](#) mesh object. "col": the mesh colors. "render", logical, whether to render the mesh. "hemi": the hemisphere, one of 'lh' or 'rh'.

See Also

Other coloredmesh functions: [coloredmesh.from.annot\(\)](#), [coloredmesh.from.label\(\)](#), [coloredmesh.from.mask\(\)](#), [coloredmesh.from.morph.native\(\)](#), [coloredmesh.from.morphdata\(\)](#), [coloredmeshes.from.color\(\)](#)

coloredmesh.from.morphdata

Create a coloredmesh from arbitrary data.

Description

Create a coloredmesh from arbitrary data.

Usage

```
coloredmesh.from.morphdata(
    subjects_dir,
    vis_subject_id,
    morph_data,
    hemi,
    surface = "white",
    colormap = NULL,
    makecmap_options = mkco.seq()
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
vis_subject_id	string. The subject identifier from which to obtain the surface for data visualization. Example: 'fsaverage'.
morph_data	string. The morphometry data to use. E.g., 'area' or 'thickness'.
hemi	string, one of 'lh' or 'rh'. The hemisphere name. Used to construct the names of the label data files to be loaded.
surface	character string or 'fs.surface' instance. The display surface. E.g., "white", "pial", or "inflated". Defaults to "white".
colormap	a colormap function. See the squash package for some colormaps. Defaults to jet .
makecmap_options	named list of parameters to pass to makecmap . Must not include the unnamed first parameter, which is derived from 'measure'.

Value

coloredmesh. A named list with entries: "mesh" the [tmesh3d](#) mesh object. "col": the mesh colors. "render", logical, whether to render the mesh. "hemi": the hemisphere, one of 'lh' or 'rh'.

See Also

Other coloredmesh functions: [coloredmesh.from.annot\(\)](#), [coloredmesh.from.label\(\)](#), [coloredmesh.from.mask\(\)](#), [coloredmesh.from.morph.native\(\)](#), [coloredmesh.from.morph.standard\(\)](#), [coloredmeshes.from.color\(\)](#)

 coloredmesh.from.preloaded.data

Generate coloredmesh from loaded data.

Description

Generate coloredmesh from loaded data.

Usage

```
coloredmesh.from.preloaded.data(
  fs_surface,
  morph_data = NULL,
  col = NULL,
  hemi = "lh",
  makecmap_options = mkco.seq()
)
```

Arguments

fs_surface	an fs.surface instance or a character string, which will be interpreted as the path to a file and loaded with <code>freesurferformats::read.fs.surface</code> .
morph_data	numerical vector, per-vertex data (typically morphometry) for the mesh. If given, takes precedence over 'col' parameter.
col	vector of colors, typically hex color strings like '#FF00FF'. The per-vertex-colors for the mesh. Alternative to morph_data.
hemi	character string, one of 'lh' or 'rh'. Metadata, the hemisphere. May be used by visualization functions to decide whether to draw the mesh in certain views.
makecmap_options	named list of parameters to pass to <code>makecmap</code> . Must not include the unnamed first parameter, which is derived from 'measure'.

Value

as fs.coloredmesh instance

 coloredmesh.plot.colorbar.separate

Draw colorbar for coloredmeshes in separate 2D plot.

Description

Draw a colorbar for the coloredmeshes to a separate 2D plot. Due to the suboptimal handling of colorbar drawing in the three-dimensional multi-panel views, it is often desirable to plot the colorbar in a separate window, export it from there and then manually add it to the final plot version in some image manipulation software like Inkscape. If you need more control over the colormap than offered by this function (e.g., setting the color value for NA values or making a symmetric colormap to ensure that the zero point for divergent colormaps is a neutral color), you should write custom code, and the return value from this function will come in handy to do that.

Usage

```
coloredmesh.plot.colorbar.separate(
  coloredmeshes,
  show = FALSE,
  image.plot_extra_options = list(horizontal = FALSE, legend.cex = 1.8, legend.width =
    2, legend.mar = 12, axis.args = list(cex.axis = 5)),
  png_options = list(filename = "fsbrain_cbar.png", width = 1400, height = 1400, bg =
    "#FFFFFF00"),
  silent = FALSE,
  trim_png = TRUE
)
```

Arguments

- | | |
|--------------------------|--|
| coloredmeshes | list of coloredmeshes. A coloredmesh is a named list as returned by the ‘coloredmesh.from’ functions. It has the entries ‘mesh’ of type tmesh3d, a ‘col’, which is a color specification for such a mesh. The ‘vis*’ functions (like vis.subject.morph.native) all return a list of coloredmeshes. |
| show | logical, Whether to open the resulting plot. Defaults to ‘TRUE’. |
| image.plot_extra_options | named list of extra options to pass to image.plot . This can be used to add a legend to the colorbar, rotate the colorbar, or whatever. The options "legend_only", "zlim", and "col" are computed and set for you by this function, so there is no need to pass these. Your list will be merged with the internal options, so you could overwrite named arguments if needed. |
| png_options | Options to pass to png , see the docs of that function for details. Allow you to save the plot as a png bitmap image. Example: png_options = list("filename"="fsbrain_cbar.png") Defaults to NULL, which will not save anything. |
| silent | logical, whether to suppress messages. Defaults to ‘FALSE’. |
| trim_png | logical, whether to trim the output PNG image using image magick, i.e., remove everything but the foreground. Ignored unless an output PNG image is actually written (see ‘png_options’) and the ‘magick’ package is installed. |

Value

named list, entries: ‘output_img_path’: character string, the path to the output file, or NULL.

Note

If you increase the output resolution of the colorbar (using 'png_options'), you will have to increase the font sizes as well (using 'image.plot_extra_options'), otherwise the axis and legend labels will be hard to read.

See Also

Other colorbar functions: `combine.colorbar.with.brainview.animation()`, `combine.colorbar.with.brainview.ima`

Examples

```
## Not run:
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
coloredmeshes = vis.subject.morph.native(subjects_dir, 'subject1',
  'thickness', 'lh', views=c('t4'));
coloredmesh.plot.colorbar.separate(coloredmeshes);

# Or plot a colorbar with a label:
coloredmesh.plot.colorbar.separate(coloredmeshes,
  image.plot_extra_options = list("legend.lab"="Thickness [mm]",
  horizontal=TRUE, legend.cex=1.5, legend.line=-3));

## End(Not run)
```

coloredmeshes.from.color

Create coloredmeshes for both hemis using pre-defined colors.

Description

Create coloredmeshes for both hemis using pre-defined colors.

Usage

```
coloredmeshes.from.color(
  subjects_dir,
  subject_id,
  color_data,
  hemi,
  surface = "white",
  metadata = list()
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	string. The subject identifier.
color_data	a hemilist containing vectors of hex color strings
hemi	string, one of 'lh' or 'rh'. The hemisphere name. Used to construct the names of the label data files to be loaded.
surface	character string or 'fs.surface' instance. The display surface. E.g., "white", "pial", or "inflated". Defaults to "white".
metadata	a named list, can contain whatever you want. Typical entries are: 'src_data' a hemilist containing the source data from which the 'color_data' was created, optional. If available, it is encoded into the coloredmesh and can be used later to plot a colorbar. 'makecmap_options': the options used to created the colormap from the data.

Value

named list of coloredmeshes. Each entry is a named list with entries: "mesh" the `tmesh3d` mesh object. "col": the mesh colors. "render", logical, whether to render the mesh. "hemi": the hemisphere, one of 'lh' or 'rh'.

See Also

Other coloredmesh functions: `coloredmesh.from.annot()`, `coloredmesh.from.label()`, `coloredmesh.from.mask()`, `coloredmesh.from.morph.native()`, `coloredmesh.from.morph.standard()`, `coloredmesh.from.morphdata()`

colorlist.brain.clusters

Return diverging color list

Description

Return diverging color list

Usage

```
colorlist.brain.clusters(num_colors)
```

Arguments

num_colors	integer, the number of colors you want
------------	--

Value

vector of colors

```
combine.colorbar.with.brainview.animation
```

Combine a colorbar and a brain animation in gif format into a new animation.

Description

Combine a colorbar and a brain animation in gif format into a new animation.

Usage

```
combine.colorbar.with.brainview.animation(  
    brain_animation,  
    colorbar_img,  
    output_animation,  
    offset = "+0+0",  
    extend_brainview_img_height_by = 0L,  
    silent = FALSE,  
    allow_colorbar_shrink = TRUE  
)
```

Arguments

brain_animation	path to the brain animation in gif format
colorbar_img	path to the main image containing the separate colorbar, usually an image in PNG format
output_animation	path to output image in gif format, must include the '.gif' file extension
offset	offset string passed to <code>magick::image_composite</code> . Allows you to shift the location of the colorbar in the final image.
extend_brainview_img_height_by	integer value in pixels, the size of the lower border to add to the <code>brainview_img</code> . Use this if the lower part of the colorbar is off the image canvas.
silent	logical, whether to silence all messages
allow_colorbar_shrink	logical, whether to shrink the colorbar to the width of the animation in case it is considerably wider (more than 20 percent). Defaults to TRUE.

See Also

Other colorbar functions: [coloredmesh.plot.colorbar.separate\(\)](#), [combine.colorbar.with.brainview.image\(\)](#)

```
combine.colorbar.with.brainview.image
```

Combine a colorbar and a brainview image into a new figure.

Description

Combine a colorbar and a brainview image into a new figure.

Usage

```
combine.colorbar.with.brainview.image(
    brainview_img = "fsbrain_arranged.png",
    colorbar_img = "fsbrain_cbar.png",
    output_img = "fsbrain_merged.png",
    offset = "+0+0",
    extend_brainview_img_height_by = NULL,
    silent = FALSE,
    allow_colorbar_shrink = TRUE,
    horizontal = FALSE
)
```

Arguments

<code>brainview_img</code>	path to the main image containing the view of the brain, usually an image in PNG format.
<code>colorbar_img</code>	path to the main image containing the separate colorbar, usually an image in PNG format.
<code>output_img</code>	path to output image, including the file extension.
<code>offset</code>	offset string passed to <code>magick::image_composite</code> . Allows you to shift the location of the colorbar in the final image.
<code>extend_brainview_img_height_by</code>	integer value in pixels, the size of the lower border to add to the <code>brainview_img</code> . Increase this if the lower part of the colorbar is off the image canvas.
<code>silent</code>	logical, whether to silence all messages
<code>allow_colorbar_shrink</code>	logical, whether to shrink the colorbar to the width of the animation in case it is considerably wider (more than 20 percent). Defaults to TRUE.
<code>horizontal</code>	logical, whether the colorbar is horizontal. If so, it will be added below the <code>'brainview_img'</code> . If it is vertical, it will be added to the right of the <code>'brainview_img'</code> .

Value

named list with entries `'output_img_path'`: character string, path to saved image. `'merged_img'`: magick image instance, the merged image

See Also

Other colorbar functions: [coloredmesh.plot.colorbar.separate\(\)](#), [combine.colorbar.with.brainview.animation\(\)](#)

cube3D.tris

Return triangles for a 3D cube or cuboid.

Description

Each row of the returned matrix encodes a point (the x, y, and z coordinates), and 3 consecutive rows encode a triangle. Obviously, a point will occur several times (as part of several triangles). The result can be passed to [triangles3d](#) to render a 3D box. The defaults for the parameters will create a cube with edge length 1 centered at (0, 0, 0).

Usage

```
cube3D.tris(
    xmin = -0.5,
    xmax = 0.5,
    ymin = -0.5,
    ymax = 0.5,
    zmin = -0.5,
    zmax = 0.5,
    center = NULL,
    edge_length = 1
)
```

Arguments

xmin	numeric, minimal x coordinate
xmax	numeric, maximal x coordinate
ymin	numeric, minimal y coordinate
ymax	numeric, maximal y coordinate
zmin	numeric, minimal z coordinate
zmax	numeric, maximal z coordinate
center	numeric vector of length 3 or NULL, coordinates where to center a cube with the edge length defined in parameter 'edge_length'. If this is not 'NULL', the parameters 'xmin', 'xmax', ... will be ignored, and their values will be computed for a cube based on the 'center' and 'edge_length'. Note that you can only create cubes using 'center' and 'edge_length', while the min/max methods allows the construction of cuboids.
edge_length	numeric, the edge length of the cube. Only used if parameter 'center' is used, ignored otherwise.

Value

numerical matrix with 36 rows and 3 columns, the 3D coordinates. Each row encodes a point (the x, y, and z coordinates), and 3 consecutive rows encode a triangle.

Examples

```
# Create a cube with edge length 2, centered at (3,4,5):
cube_coords = cube3D.tris(center=c(3,4,5), edge_length=2.0);
# Create the same cube using the min/max method:
cube_coords = cube3D.tris(xmin=2, xmax=4, ymin=3, ymax=5, zmin=4, zmax=6);
# Create a cuboid:
cuboid_coords = cube3D.tris(xmin=2, xmax=4, ymin=3, ymax=9, zmin=4, zmax=5);
# To render the cuboid:
#rgl::triangles3d(cuboid_coords, col="red");
```

cubes3D.tris

Vectorized version of cube3D.tris

Description

Vectorized version of cube3D.tris

Usage

```
cubes3D.tris(centers, edge_length = 1)
```

Arguments

centers	numerical matrix with 3 columns. Each column represents the x, y, z coordinates of a center at which to create a cube.
edge_length	numerical vector or scalar, the edge length. Must have length 1 (same edge length for all cubes), or the length must be identical to the number of rows in parameter 'centers'.

Value

matrix of triangle coordinates, see [cube3D.tris](#).

Examples

```
# Plot a 3D cloud of 20000 voxels:
centers = matrix(rnorm(20000*3)*100, ncol=3);
rgl::triangles3d(cubes3D.tris(centers));
```

```
delete_all_optional_data
```

Delete all data in the package cache.

Description

Delete all data in the package cache.

Usage

```
delete_all_optional_data()
```

Value

integer. The return value of the unlink() call: 0 for success, 1 for failure. See the unlink() documentation for details.

```
demographics.to.fsgd.file
```

Write FreeSurfer Group Descriptor (FSGD) file from demographics dataframe.

Description

Write FreeSurfer Group Descriptor (FSGD) file from demographics dataframe.

Usage

```
demographics.to.fsgd.file(
  filepath,
  demographics_df,
  group_column_name = "group",
  subject_id_column_name = "id",
  var_columns = NULL,
  ftitle = "OSGM",
  fsgd_flag_lines = c("DeMeanFlag 1", "ReScaleFlag 1")
)
```

Arguments

filepath character string, the path to the output file in FSGD format

demographics_df

data.frame, as returned by read.md.demographics or created manually. Note that the data.frame must not contain any character columns, they should be converted to factors.

group_column_name	character string, the column name of the group column in the 'demographics_df'
subject_id_column_name	character string, the column name of the subject identifier column in the 'demographics_df'
var_columns	vector of character strings, the column names to include as variables in the FSGD file. If NULL (the default), all columns will be included (with the exception of the group column and the subject id column).
ftitle	character string, freeform title for the FSGD file
fsgd_flag_lines	vector of character strings, extra flag lines to write to the file. The default setting will activate de-meaning and rescaling.

Value

vector of character strings, the lines written to the 'filepath', invisible.

See Also

Other metadata functions: [read.md.demographics\(\)](#), [read.md.subjects\(\)](#), [report.on.demographics\(\)](#)

desaturate *Perform simple desaturation or grayscale conversion of RGBA colors.*

Description

Perform simple desaturation or grayscale conversion of RGBA colors.

Usage

```
desaturate(color, gamma_correct = FALSE)
```

Arguments

color	rgba color strings
gamma_correct	logical, whether to apply non-linear gamma correction. First performs gamma expansion, then applies the gray-scale channel weights, then gamma compression.

Value

rgba color strings, the grayscale colors. The information from one of the three rgb channels would be enough. The alpha value is not touched.

Note

Assumes sRGB color space.

References

see https://en.wikipedia.org/wiki/Grayscale#Converting_color_to_grayscale

See Also

Other color functions: [alphablend\(\)](#)

download_fsaverage *Download the FreeSurfer v6 fsaverage subject.*

Description

Download some relevant files from the FreeSurfer v6 fsaverage subject. The files are subject to the FreeSurfer software license, see parameter 'accept_freesurfer_license' for details. This data is not required for the package to work. If you are working on a machine that has FreeSurfer installed, you already have this data anyways and do not need to download it. If not, it is very convenient to have it if you are using the fsaverage template subject to analyze your standard space data, as it is required for visualization of such data.

Usage

```
download_fsaverage(accept_freesurfer_license = FALSE)
```

Arguments

`accept_freesurfer_license`
logical, whether you accept the FreeSurfer license for fsaverage, available at <https://surfer.nmr.mgh.harvard.edu/fswiki/FreeSurferSoftwareLicense>. Defaults to FALSE.

Value

Named list. The list has entries: "available": vector of strings. The names of the files that are available in the local file cache. You can access them using `get_optional_data_file()`. "missing": vector of strings. The names of the files that this function was unable to retrieve.

download_fsaverage3 *Download the FreeSurfer v6 low-resolution fsaverage3 subject.*

Description

Download some relevant files from the FreeSurfer v6 fsaverage3 subject. The files are subject to the FreeSurfer software license, see parameter 'accept_freesurfer_license' for details. This data is not required for the package to work. If you are working on a machine that has FreeSurfer installed, you already have this data anyways and do not need to download it. Also downloads data for subject1 that has been mapped to fsaverage.

Usage

```
download_fsaverage3(accept_freesurfer_license = FALSE)
```

Arguments

accept_freesurfer_license
 logical, whether you accept the FreeSurfer license for fsaverage, available at <https://surfer.nmr.mgh.harvard.edu/fswiki/FreeSurferSoftwareLicense>. Defaults to FALSE.

Value

Named list. The list has entries: "available": vector of strings. The names of the files that are available in the local file cache. You can access them using `get_optional_data_file()`. "missing": vector of strings. The names of the files that this function was unable to retrieve.

Note

The subject fsaverage3 is a downsampled (low mesh resolution) version of the standard fsaverage. If you never heard about fsaverage3, you do not need it. You will have to manually re-sample your data in FreeSurfer if you want to use it with fsaverage3.

download_optional_data
 Download optional data for this package if required.

Description

Ensure that the optionl data is available locally in the package cache. Will try to download the data only if it is not available. This data is not required for the package to work, but it is used in the examples, in the unit tests and also in the example code from the vignette. Downloading it is highly recommended.

Usage

```
download_optional_data()
```

Value

Named list. The list has entries: "available": vector of strings. The names of the files that are available in the local file cache. You can access them using `get_optional_data_file()`. "missing": vector of strings. The names of the files that this function was unable to retrieve.

```
download_optional_paper_data
```

Download extra data to reproduce the figures from the fsbrain paper.

Description

Download extra data to reproduce the figures from the fsbrain paper.

Usage

```
download_optional_paper_data()
```

Value

Named list. The list has entries: "available": vector of strings. The names of the files that are available in the local file cache. You can access them using `get_optional_data_file()`. "missing": vector of strings. The names of the files that this function was unable to retrieve.

Note

Called for side effect of data download.

```
export.coloredmesh.ply
```

Export a coloredmeshes with vertexcolors in PLY format.

Description

Exports coloredmeshes with vertex coloring to standard mesh files in Stanford Triangle (PLY) format. This is very hand for rendering in external standard 3D modeling software like Blender.

Usage

```
export.coloredmesh.ply(filepath, coloredmesh)
```

Arguments

filepath	The export filepath, including file name and extension.
coloredmesh	an 'fs.coloredmesh' instance, as returned (silently) by all surface visualization functions, like vis.subject.morph.native .

Examples

```
## Not run:
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
coloredmeshes = vis.subject.morph.native(subjects_dir, 'subject1', 'thickness');
export.coloredmesh.ply('~/.subject1_thickness_1h.ply', coloredmeshed$1h);

## End(Not run)
```

face.edges	<i>Enumerate all edges of the given faces or mesh.</i>
------------	--

Description

Compute edges of a tri-mesh. Can compute all edges, or only a subset, given by the face indices in the mesh.

Usage

```
face.edges(surface_mesh, face_indices = "all")
```

Arguments

surface_mesh	surface mesh, as loaded by subject.surface or read.fs.surface .
face_indices	integer vector, the face indices. Can also be the character string 'all' to use all faces.

Value

integer matrix of size (n, 2) where n is the number of edges. The indices (source and target vertex) in each row are **not** sorted, and the edges are **not** unique. I.e., each undirected edge 'u, v' (with the exception of edges on the mesh border) will occur twice in the result: once as 'u, v' and once as 'v, u'.

See Also

Other surface mesh functions: [label.border\(\)](#), [mesh.vertex.included.faces\(\)](#), [mesh.vertex.neighbors\(\)](#), [subject.surface\(\)](#), [vis.path.along.verts\(\)](#)

find.freesurferhome *Find the FREESURFER_HOME directory on disk.*

Description

Try to find directory containing the FreeSurfer installation, based on environment variables and *educated guessing*.

Usage

```
find.freesurferhome(mustWork = FALSE)
```

Arguments

mustWork logical. Whether the function should with an error stop if the directory cannot be found. If this is TRUE, the return value will be only the 'found_at' entry of the list (i.e., only the path of the FreeSurfer installation dir).

Value

named list with the following entries: "found": logical, whether it was found. "found_at": Only set if found=TRUE, the path to the FreeSurfer installation directory (including the directory itself). See 'mustWork' for important information.

See Also

[fs.home](#)

find.subjectsdir.of *Find the subject directory containing the fsaverage subject (or others) on disk.*

Description

Try to find directory containing the fsaverage subject (or any other subject) by checking in the following places and returning the first path where it is found: first, the directory given by the environment variable SUBJECTS_DIR, then in the subir 'subjects' of the directory given by the environment variable FREESURFER_HOME, and finally the base dir of the package cache. See the function [download_fsaverage](#) if you want to download fsaverage to your package cache and ensure it always gets found, no matter whether the environment variables are set or not.

Usage

```
find.subjectsdir.of(subject_id = "fsaverage", mustWork = FALSE)
```

Arguments

subject_id	string, the subject id of the subject. Defaults to 'fsaverage'.
mustWork	logical. Whether the function should with an error stop if the directory cannot be found. If this is TRUE, the return value will be only the 'found_at' entry of the list (i.e., only the path of the subjects dir).

Value

named list with the following entries: "found": logical, whether it was found. "found_at": Only set if found=TRUE, the path to the fsaverage directory (NOT including the fsaverage dir itself). "found_all_locations": list of all locations in which it was found. See 'mustWork' for important information.

See Also

[fsaverage.path](#)

fs.coloredmesh	<i>fs.coloredmesh constructor</i>
----------------	-----------------------------------

Description

fs.coloredmesh constructor

Usage

```
fs.coloredmesh(
  mesh,
  col,
  hemi,
  render = TRUE,
  metadata = NULL,
  add_normals = FALSE
)
```

Arguments

mesh	a 'mesh3d' instance as returned by tmesh3d or an 'fs.surface' brain surface mesh as returned by functions like subject.surface .
col	vector of vertex colors for the mesh, one color per vertex. Expanded if exactly one color.
hemi	character string, one of 'lh' or 'rh'. This may be used by visualization functions to decide whether or not to show this mesh in a certain view.
render	logical, whether to render this mesh during visualization
metadata	optional, named list containing metadata
add_normals	logical, whether to compute normals and save them in the mesh.

Value

an 'fs.coloredmesh' instance. The only fields one should use in client code are 'mesh', 'hemi' and 'col', all others are considered internal and may change without notice.

fs.home	<i>Return FreeSurfer path.</i>
---------	--------------------------------

Description

Return FreeSurfer path.

Usage

fs.home()

Value

the FreeSurfer path, typically what the environment variable 'FREESURFER_HOME' points to.

Note

This function will stop (i.e., raise an error) if the directory cannot be found.

fsaverage.path	<i>Return path to fsaverage dir.</i>
----------------	--------------------------------------

Description

Return path to fsaverage dir.

Usage

fsaverage.path()

Value

the path to the fsaverage directory (NOT including the 'fsaverage' dir itself).

Note

This function will stop (i.e., raise an error) if the directory cannot be found.

```
fsbrain.set.default.figsizes
```

Set default figure size for fsbrain visualization functions.

Description

Set default figure size for fsbrain visualization functions.

Usage

```
fsbrain.set.default.figsizes(width, height, xstart = 50L, ystart = 50L)
```

Arguments

width	integer, default figure width in pixels
height	integer, default figure height in pixels
xstart	integer, default horizontal position of plot window on screen, left border is 0. The max value (right border) depends on your screen resolution.
ystart	integer, default vertical position of plot window on screen, upper border is 0. The max value (lower border) depends on your screen resolution.

Note

This function overwrites options("fsbrain.rgloptions"). Output size is limited by your screen resolution. To set your preferred figure size for future R sessions, you could call this function in your '~/.Rprofile' file.

```
fup
```

Transform first character of a string to uppercase.

Description

Transform first character of a string to uppercase. This is useful when labeling plots. Important: this function does not know about different encodings, languages or anything, it just calls [toupper](#) for the first character.

Usage

```
fup(word)
```

Arguments

word,	string. Any string.
-------	---------------------

Value

string, the input string with the first character transformed to uppercase.

Examples

```
word_up = fup("word");
```

gen.test.volume	<i>Generate test 3D volume of integers. The volume has an outer background area (intensity value 'bg') and an inner foreground areas (intensity value 200L).</i>
-----------------	--

Description

Generate test 3D volume of integers. The volume has an outer background area (intensity value 'bg') and an inner foreground areas (intensity value 200L).

Usage

```
gen.test.volume(vdim = c(256L, 256L, 256L), bg = NA)
```

Arguments

vdim	integer vector of length 3, the dimensions
bg	value to use for outer background voxels. Typically '0L' or 'NA'.

Value

a 3d array of integers

Note

This function exists for software testing purposes only, you should not use it in client code.

`get.atlas.region.names`*Determine atlas region names from a subject.*

Description

Determine atlas region names from a subject. WARNING: Not all subjects have all regions of an atlas. You should use an average subject like fsaverage to get all regions.

Usage

```
get.atlas.region.names(  
  atlas,  
  template_subjects_dir = NULL,  
  template_subject = "fsaverage",  
  hemi = "lh"  
)
```

Arguments

`atlas`, string. The atlas name. E.g., "aparc", "aparc.2009s", or "aparc.DKAtlas". Used to construct the name of the annotation file to be loaded.

`template_subjects_dir`, string. The directory containing the dir of the template_subject. E.g., the path to FREESURFER_HOME/subjects. If NULL, env vars will be searched for candidates, and the function will fail if they are not set correctly. Defaults to NULL.

`template_subject`, string. The subject identifier. Defaults to 'fsaverage'.

`hemi`, string, one of 'lh' or 'rh'. The hemisphere name. Used to construct the names of the annotation and morphometry data files to be loaded. Defaults to 'lh'. Should not matter much, unless you do not have the file for one of the hemis for some reason.

Value

vector of strings, the region names.

See Also

Other atlas functions: [group.agg.atlas.native\(\)](#), [group.agg.atlas.standard\(\)](#), [group.annot\(\)](#), [group.label.from.annot\(\)](#), [label.from.annotdata\(\)](#), [label.to.annot\(\)](#), [regions.to.ignore\(\)](#), [spread.values.over.annot\(\)](#), [spread.values.over.hemi\(\)](#), [spread.values.over.subject\(\)](#), [subject.annot\(\)](#), [subject.atlas.agg\(\)](#), [subject.label.from.annot\(\)](#), [subject.lobes\(\)](#)

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
atlas_regions = get.atlas.region.names('aparc',
  template_subjects_dir=subjects_dir, template_subject='subject1');
```

get.rglstyle *Get the default visualization style parameters as a named list.*

Description

Run [material3d](#) without arguments to see valid style keywords to create new styles.

Usage

```
get.rglstyle(style)
```

Arguments

style string. A style name. Available styles are one of: "default", "shiny", "semitransparent", "edges".

Value

a style, resolved to a parameter list compatible with [material3d](#).

See Also

[shade3d](#) can use the returned style

get.view.angle.names *Get list of valid view angle names.*

Description

The returned strings are used as constants to identify a view of type 'sd_<angle>'. They can be used to construct entries for the parameter 'views' of functions like [vis.subject.morph.native](#), or directly as parameter 'view_angles' for functions like [vislayout.from.coloredmeshes](#).

Usage

```
get.view.angle.names(angle_set = "all", add_sd_prefix = TRUE)
```

Arguments

- `angle_set` string, which view subset to return. Available subsets are: 'all' (or alias 't9'): for all 9 angles. 't4': for the t4 views. 'medial': the 2 medial views, one for each hemi. 'lateral': the 2 lateral views, one for each hemi. 'lh': medial and lateral for the left hemisphere. 'rh': medial and lateral for the right hemisphere.
- `add_sd_prefix` logical, whether the prefix 'sd_' should be added to the string. This will construct full view names. If set to false, only the substring after the prefix 'sd_' will be returned. This is used internally only and should not be needed in general.

Value

vector of character strings, all valid view angle strings.

<code>getIn</code>	<i>Retrieve values from nested named lists</i>
--------------------	--

Description

Retrieve values from nested named lists

Usage

```
getIn(named_list, listkeys, default = NULL)
```

Arguments

- `named_list` a named list
- `listkeys` vector of character strings, the nested names of the lists
- `default` the default value to return in case the requested value is 'NULL'.

Value

the value at the path through the lists, or 'NULL' (or the 'default') if no such path exists.

Examples

```
data = list("regions"=list("frontal"=list("thickness"=2.3, "area"=2345)));
getIn(data, c("regions", "frontal", "thickness"));      # 2.3
getIn(data, c("regions", "frontal", "nosuchentry"));    # NULL
getIn(data, c("regions", "nosuchregion", "thickness")); # NULL
getIn(data, c("regions", "nosuchregion", "thickness"), default=14); # 14
```

`get_optional_data_filepath`

Access a single file from the package cache by its file name.

Description

Access a single file from the package cache by its file name.

Usage

```
get_optional_data_filepath(filename, mustWork = TRUE)
```

Arguments

`filename`, string. The filename of the file in the package cache.
`mustWork`, logical. Whether an error should be created if the file does not exist. If `mustWork=FALSE` and the file does not exist, the empty string is returned.

Value

string. The full path to the file in the package cache or the empty string if there is no such file available. Use this in your application code to open the file.

`group.agg.atlas.native`

Aggregate native space morphometry data over brain atlas regions and subjects for a group of subjects.

Description

Aggregate native space morphometry data over brain atlas regions, e.g., compute the mean thickness value over all regions in an atlas for all subjects.

Usage

```
group.agg.atlas.native(  
  subjects_dir,  
  subjects_list,  
  measure,  
  hemi,  
  atlas,  
  agg_fun = mean,  
  cache_file = NULL  
)
```

Arguments

subjects_dir,	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subjects_list,	string vector. A vector of subject identifiers that match the directory names within subjects_dir.
measure,	string. Name of the vertex-wise measure of morphometry data file. E.g., "area" or "thickness". Used to construct the name of the morphometry file to be loaded.
hemi,	string, one of 'lh', 'rh', or 'both'. The hemisphere name. Used to construct the names of the annotation and morphometry data files to be loaded. If set to 'both', combined data for 'lh' and 'rh' will be used.
atlas,	string. The atlas name. E.g., "aparc", "aparc.2009s", or "aparc.DKAtlas". Used to construct the name of the annotation file to be loaded.
agg_fun,	function. An R function that aggregates data, typically max, mean, min or something similar. Note: this is NOT a string, put the function name without quotes. Defaults to mean.
cache_file,	string or NULL. If given, it is interpreted as path of a file, and the data will be cached in the file cache_file in RData format. If the file does not exist yet, the function will run and cache the data in the file. If the file exists, the function will load the data from the file instead of running. The filename should end in '.RData', but that is not enforced or checked in any way. WARNING: If cached data is returned, all parameters passed to this function (with the exception of 'cache_file') are ignored! Whether the cached data is for another subjects_list or hemi is NOT checked! You have to ensure this yourself, by using different filenames. Defaults to NULL.

Value

dataframe with aggregated values for all regions and subjects, with n columns and m rows, where n is the number of subjects and m is the number of regions.

See Also

Other aggregation functions: [group.agg.atlas.standard\(\)](#), [group.morph.agg.standard.vertex\(\)](#), [subject.atlas.agg\(\)](#)

Other atlas functions: [get.atlas.region.names\(\)](#), [group.agg.atlas.standard\(\)](#), [group.annot\(\)](#), [group.label.from.annot\(\)](#), [label.from.annotdata\(\)](#), [label.to.annot\(\)](#), [regions.to.ignore\(\)](#), [spread.values.over.annot\(\)](#), [spread.values.over.hemi\(\)](#), [spread.values.over.subject\(\)](#), [subject.annot\(\)](#), [subject.atlas.agg\(\)](#), [subject.label.from.annot\(\)](#), [subject.lobes\(\)](#)

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
agg = group.agg.atlas.native(subjects_dir, c('subject1', 'subject2'),
  'thickness', 'lh', 'aparc');
```

```
# Visualize the mean values. Could use any subject, typically
# one would use fsaverage. Here we use subject1:
agg$subject = NULL; # remove non-numeric column.
vis.region.values.on.subject(subjects_dir, 'subject1', 'aparc',
  lh_region_value_list=colMeans(agg), rh_region_value_list=NULL);
```

```
group.agg.atlas.standard
```

Aggregate standard space morphometry data over brain atlas regions and subjects for a group of subjects.

Description

Aggregate standard space morphometry data over brain atlas regions, e.g., compute the mean thickness value over all regions in an atlas for all subjects.

Usage

```
group.agg.atlas.standard(
  subjects_dir,
  subjects_list,
  measure,
  hemi,
  atlas,
  fwhm,
  agg_fun = mean,
  template_subject = "fsaverage",
  cache_file = NULL
)
```

Arguments

subjects_dir,	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subjects_list,	string vector. A vector of subject identifiers that match the directory names within subjects_dir.
measure,	string. Name of the vertex-wise measure of morphometry data file. E.g., "area" or "thickness". Used to construct the name of the morphometry file to be loaded.
hemi,	string, one of 'lh', 'rh', or 'both'. The hemisphere name. Used to construct the names of the annotation and morphometry data files to be loaded. If set to 'both', combined data for 'lh' and 'rh' will be used.
atlas,	string. The atlas name. E.g., "aparc", "aparc.2009s", or "aparc.DKAtlas". Used to construct the name of the annotation file to be loaded.

fwhm,	string. The smoothing setting which was applied when mapping data to the template subject. Usually one of '0', '5', '10', '15', '20', or '25'.
agg_fun,	function. An R function that aggregates data, typically max, mean, min or something similar. Note: this is NOT a string, put the function name without quotes. Defaults to mean.
template_subject,	string. The template subject name. Defaults to 'fsaverage'. Must have its data in subjects_dir.
cache_file,	string or NULL. If given, it is interpreted as path of a file, and the data will be cached in the file cache_file in RData format. If the file does not exist yet, the function will run and cache the data in the file. If the file exists, the function will load the data from the file instead of running. The filename should end in '.RData', but that is not enforced or checked in any way. WARNING: If cached data is returned, all parameters passed to this function (with the exception of 'cache_file') are ignored! Whether the cached data is for another subjects_list or hemi is NOT checked! You have to ensure this yourself, by using different filenames. Defaults to NULL.

Value

dataframe with aggregated values for all regions and subjects, with n columns and m rows, where n is the number of subjects and m is the number of regions.

See Also

Other aggregation functions: [group.agg.atlas.native\(\)](#), [group.morph.agg.standard.vertex\(\)](#), [subject.atlas.agg\(\)](#)

Other atlas functions: [get.atlas.region.names\(\)](#), [group.agg.atlas.native\(\)](#), [group.annot\(\)](#), [group.label.from.annot\(\)](#), [label.from.annotdata\(\)](#), [label.to.annot\(\)](#), [regions.to.ignore\(\)](#), [spread.values.over.annot\(\)](#), [spread.values.over.hemi\(\)](#), [spread.values.over.subject\(\)](#), [subject.annot\(\)](#), [subject.atlas.agg\(\)](#), [subject.label.from.annot\(\)](#), [subject.lobes\(\)](#)

Examples

```
## Not run:
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
agg = group.agg.atlas.standard(subjects_dir, c('subject1', 'subject2'),
  'thickness', 'lh', 'aparc', fwhm='10');
# Visualize the mean values. Could use any subject, typically
# one would use fsaverage. Here we use subject1:
agg$subject = NULL; # remove non-numeric column.
vis.region.values.on.subject(subjects_dir, 'subject1', 'aparc',
  lh_region_value_list=colMeans(agg), rh_region_value_list=NULL);

## End(Not run)
```

group.annot	<i>Load annotations for a group of subjects.</i>
-------------	--

Description

Load a brain surface annotation, i.e., a cortical parcellation based on an atlas, for a group of subjects.

Usage

```
group.annot(subjects_dir, subjects_list, hemi, atlas)
```

Arguments

subjects_dir,	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subjects_list,	vector of strings. The subject identifiers.
hemi,	string, one of 'lh' or 'rh'. The hemisphere name. Used to construct the names of the annotation and morphometry data files to be loaded.
atlas,	string. The atlas name. E.g., "aparc", "aparc.2009s", or "aparc.DKTatlas". Used to construct the name of the annotation file to be loaded.

Value

list of annotations, as returned by `freesurferformats::read.fs.annot()`. If hemi is 'both', the annotations are the results of merging over the hemispheres for each subject.

See Also

Other atlas functions: `get.atlas.region.names()`, `group.agg.atlas.native()`, `group.agg.atlas.standard()`, `group.label.from.annot()`, `label.from.annotdata()`, `label.to.annot()`, `regions.to.ignore()`, `spread.values.over.annot()`, `spread.values.over.hemi()`, `spread.values.over.subject()`, `subject.annot()`, `subject.atlas.agg()`, `subject.label.from.annot()`, `subject.lobes()`

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
subjects_list = c("subject1", "subject2");
annotations = group.annot(subjects_dir, subjects_list, "lh", "aparc");
```

`group.concat.measures.native`*Concatenate native space data for a group of subjects.*

Description

A measure is something like 'thickness' or 'area'. This function concatenates the native space data for all subjects into a single long vector for each measure. A dataframe is then created, in which each column is one such vector. This can be used to compute the correlation between measures on vertex level, for example.

Usage

```
group.concat.measures.native(  
  subjects_dir,  
  subjects_list,  
  measures,  
  hemi,  
  cortex_only = FALSE  
)
```

Arguments

<code>subjects_dir</code> ,	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
<code>subjects_list</code> ,	string vector. A vector of subject identifiers that match the directory names within <code>subjects_dir</code> .
<code>measures</code> ,	vector of strings. Names of the vertex-wise morphometry measures. E.g., <code>c("area", "thickness")</code> . Used to construct the names of the morphometry file to be loaded. The data of each measure will be one column in the resulting dataframe.
<code>hemi</code> ,	string, one of 'lh', 'rh' or 'both'. The hemisphere name. Used to construct the names of the annotation and morphometry data files to be loaded.
<code>cortex_only</code>	logical, whether to set non-cortex data to NA

Value

dataframe with concatenated vertex values. Each column contains the values for one measure, concatenated for all subjects. WARNING: This dataframe can get large if you have many subjects.

See Also

Other concatenation functions: [group.concat.measures.standard\(\)](#)

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
subjects_list = c('subject1', 'subject2');
cm = group.concat.measures.native(subjects_dir, subjects_list,
  c("thickness", "area"), "lh");
```

```
group.concat.measures.standard
```

Concatenate standard space data for a group of subjects.

Description

A measure is something like 'thickness' or 'area'. This function concatenates the standard space data for all subjects into a single long vector for each measure. A dataframe is then created, in which each column is one such vector. This can be used to compute the correlation between measures on vertex level, for example.

Usage

```
group.concat.measures.standard(
  subjects_dir,
  subjects_list,
  measures,
  hemi,
  fwhm_per_measure,
  cortex_only = FALSE
)
```

Arguments

subjects_dir,	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subjects_list,	string vector. A vector of subject identifiers that match the directory names within subjects_dir.
measures,	vector of strings. Names of the vertex-wise morphometry measures. E.g., c("area", "thickness"). Used to construct the names of the morphometry file to be loaded. The data of each measure will be one column in the resulting dataframe.
hemi,	string, one of 'lh', 'rh' or 'both'. The hemisphere name. Used to construct the names of the annotation and morphometry data files to be loaded.
fwhm_per_measure,	vector of strings. The fwhm settings to use, for each measure. If this is a string instead of a vector of strings, the same fwhm will be used for all measures.
cortex_only	logical, whether to set non-cortex data to NA

Value

dataframe with concatenated vertex values. Each column contains the values for one measure, concatenated for all subjects. The column names are a concatenation of the measure, "_fwhm", and the fwhm for that measure. **WARNING:** This dataframe can get large if you have many subjects.

See Also

Other concatenation functions: [group.concat.measures.native\(\)](#)

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
subjects_list = c('subject1', 'subject2');
cm = group.concat.measures.standard(subjects_dir, subjects_list,
  c("thickness", "area"), "lh", "10");
```

group.label

Retrieve label data for a group of subjects.

Description

Load a label (like 'label/lh.cortex.label') for a group of subjects from disk. Uses knowledge about the FreeSurfer directory structure to load the correct file.

Usage

```
group.label(
  subjects_dir,
  subjects_list,
  label,
  hemi,
  return_one_based_indices = TRUE
)
```

Arguments

subjects_dir, string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.

subjects_list, vector of strings. The subject identifiers.

label, string. Name of the label file, without the hemi part (if any), but including the '.label' suffix. E.g., 'cortex.label' for '?h.cortex.label'

hemi, string, one of 'lh' or 'rh'. The hemisphere name. Used to construct the names of the label data files to be loaded.

return_one_based_indices, logical. Whether the indices should be 1-based. Indices are stored zero-based in the file, but R uses 1-based indices. Defaults to TRUE, which means that 1 will be added to all indices read from the file before returning them.

Value

named list of integer vectors with label data: Each name is a subject identifier from subjects_list, and the values are lists of the vertex indices in the respective label. See 'return_one_based_indices' for important information.

See Also

Other label data functions: `labeldata.from.mask()`, `mask.from.labeldata.for.hemi()`, `subject.label()`

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
subjects_list = c("subject1", "subject2");
labels = group.label(subjects_dir, subjects_list, 'cortex.label', "lh");
```

group.label.from.annot

Extract a region from an atlas annotation as a label for a group of subjects.

Description

The returned label can be used to mask morphometry data, e.g., to set the values of a certain region to NaN or to extract only values from a certain region.

Usage

```
group.label.from.annot(
  subjects_dir,
  subjects_list,
  hemi,
  atlas,
  region,
  return_one_based_indices = TRUE,
  invert = FALSE,
  error_on_invalid_region = TRUE
)
```

Arguments

subjects_dir,	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subjects_list,	vector of string. The subject identifiers.
hemi,	string, one of 'lh' or 'rh'. The hemisphere name. Used to construct the names of the label data files to be loaded.
atlas,	string. The atlas name. E.g., "aparc", "aparc.2009s", or "aparc.DKTatlas". Used to construct the name of the annotation file to be loaded.
region,	string. A valid region name for the annotation, i.e., one of the regions of the atlas.
return_one_based_indices,	logical. Whether the indices should be 1-based. Indices are stored zero-based in label files, but R uses 1-based indices. Defaults to TRUE.
invert,	logical. If TRUE, return the indices of all vertices which are NOT part of the region. Defaults to FALSE.
error_on_invalid_region,	logical. Whether to throw an error if the given region does not appear in the region list of the annotation. If set to FALSE, this will be ignored and an empty vertex list will be returned. Defaults to TRUE.

Value

named list of integer vectors with label data: for each subject, the list of vertex indices in the label.

See Also

Other atlas functions: [get.atlas.region.names\(\)](#), [group.agg.atlas.native\(\)](#), [group.agg.atlas.standard\(\)](#), [group.annot\(\)](#), [label.from.annotdata\(\)](#), [label.to.annot\(\)](#), [regions.to.ignore\(\)](#), [spread.values.over.annot\(\)](#), [spread.values.over.hemi\(\)](#), [spread.values.over.subject\(\)](#), [subject.annot\(\)](#), [subject.atlas.agg\(\)](#), [subject.label.from.annot\(\)](#), [subject.lobes\(\)](#)

group.morph.agg.native

Aggregate native space morphometry data over one hemisphere for a group of subjects.

Description

Compute the mean (or other aggregates) over all vertices of a subject from native space morphometry data (like 'surf/lh.area'). Uses knowledge about the FreeSurfer directory structure to load the correct file.

Usage

```
group.morph.agg.native(
  subjects_dir,
  subjects_list,
  measure,
  hemi,
  agg_fun = mean,
  cast = TRUE,
  format = "curv",
  cortex_only = FALSE,
  agg_fun_extra_params = NULL
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subjects_list	string vector. A vector of subject identifiers that match the directory names within subjects_dir.
measure,	string. Name of the vertex-wise measure of morphometry data file. E.g., "area" or "thickness". Used to construct the name of the morphometry file to be loaded.
hemi,	string, one of 'lh', 'rh' or 'both'. The hemisphere name. Used to construct the names of the annotation and morphometry data files to be loaded.
agg_fun	function. An R function that aggregates data, typically <code>max</code> , <code>mean</code> , <code>min</code> or something similar. Note: this is NOT a string, put the function name without quotes. Defaults to <code>mean</code> .
cast	Whether a separate 'hemi' column should exist.
format	string. One of 'mgh', 'mgz', 'curv'. Defaults to 'mgh'.
cortex_only	logical, whether to mask the medial wall, i.e., whether the morphometry data for all vertices which are <i>not</i> part of the cortex (as defined by the label file 'label/?h.cortex.label') should be replaced with NA values. In other words, setting this to TRUE will ignore the values of the medial wall between the two hemispheres. If set to true, the mentioned label file needs to exist for the subjects. Also note that the aggregation function will need to be able to cope with NA values if you set this to TRUE. You can use 'agg_fun_extra_params' if needed to achieve that, depending on the function. For example, if you use the <code>mean</code> function, you could set <code>agg_fun_extra_params=list("na.rm"=TRUE)</code> to get the mean of the vertices which are not masked. Defaults to FALSE.
agg_fun_extra_params	named list, extra parameters to pass to the aggregation function 'agg_fun' besides the loaded morphometry data. This is useful if you have masked the data and need to ignore NA values in the <code>agg_fun</code> .

Value

dataframe with aggregated values for all subjects, with 3 columns and n rows, where n is the number of subjects. The 3 columns are 'subject_id', 'hemi', and '<measure>' (e.g., "thickness"), the latter

contains the aggregated data.

See Also

Other global aggregation functions: [group.morph.agg.standard\(\)](#), [group.multimorph.agg.native\(\)](#), [group.multimorph.agg.standard\(\)](#)

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
subjects_list = c("subject1", "subject2");
fulldata = group.morph.agg.native(subjects_dir, subjects_list, "thickness", "lh");
head(fulldata);
```

group.morph.agg.standard

Aggregate standard space (fsaverage) morphometry data over one hemisphere for a group of subjects.

Description

Compute the mean (or other aggregates) over all vertices of a subject from standard space morphometry data (like 'surf/lh.area.fwhm10.fsaverage.mgh'). Uses knowledge about the FreeSurfer directory structure to load the correct file.

Usage

```
group.morph.agg.standard(
  subjects_dir,
  subjects_list,
  measure,
  hemi,
  fwhm,
  agg_fun = mean,
  template_subject = "fsaverage",
  format = "mgh",
  cast = TRUE,
  cortex_only = FALSE,
  agg_fun_extra_params = NULL
)
```


Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subjects_list	string vector. A vector of subject identifiers that match the directory names within subjects_dir.
measure,	string. Name of the vertex-wise measure of morphometry data file. E.g., "area" or "thickness". Used to construct the name of the morphometry file to be loaded.
hemi,	string, one of 'lh', 'rh' or 'both'. The hemisphere name. Used to construct the names of the annotation and morphometry data files to be loaded.
fwhm	string. Smoothing as string, e.g. '10' or '25'.
agg_fun	function. An R function that aggregates data, typically <code>max</code> , mean, min or something similar. Note: this is NOT a string, put the function name without quotes. Defaults to mean.
template_subject	string. Template subject name, defaults to 'fsaverage'.
format	string. One of 'mgh', 'mgz', 'curv'. Defaults to 'mgh'.
cast	Whether a separate 'hemi' column should exist.
cortex_only	logical, whether to mask the medial wall, i.e., whether the morphometry data for all vertices which are <i>*not*</i> part of the cortex (as defined by the label file 'label/?h.cortex.label') should be replaced with NA values. In other words, setting this to TRUE will ignore the values of the medial wall between the two hemispheres. If set to true, the mentioned label file needs to exist for the subjects. Also not that the aggregation function will need to be able to cope with NA values if you set this to TRUE. You can use 'agg_fun_extra_params' if needed to achieve that, depending on the function. For example, if you use the <code>mean</code> function, you could set <code>agg_fun_extra_params=list("na.rm"=TRUE)</code> to get the mean of the vertices which are not masked. Defaults to FALSE.
agg_fun_extra_params	named list, extra parameters to pass to the aggregation function 'agg_fun' besides the loaded morphometry data. This is useful if you have masked the data and need to ignore NA values in the <code>agg_fun</code> .

Value

dataframe with aggregated values for all subjects, with 2 columns and n rows, where n is the number of subjects. The 2 columns are 'subject_id' and '<hemi>.<measure>' (e.g., "lh.thickness"), the latter contains the aggregated data.

See Also

Other global aggregation functions: `group.morph.agg.native()`, `group.multimorph.agg.native()`, `group.multimorph.agg.standard()`

```
group.morph.agg.standard.vertex
```

Aggregate standard space morphometry data over subjects.

Description

Aggregate vertex-wise values over subjects, leading to one aggregated measure per vertex.

Usage

```
group.morph.agg.standard.vertex(
  subjects_dir,
  subjects_list,
  measure,
  hemi,
  fwhm,
  agg_fun = mean,
  template_subject = "fsaverage",
  format = "mgh",
  cortex_only = FALSE,
  agg_fun_extra_params = NULL,
  split_by_hemi = FALSE
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subjects_list	string vector. A vector of subject identifiers that match the directory names within subjects_dir.
measure	string. Name of the vertex-wise measure of morphometry data file. E.g., "area" or "thickness". Used to construct the name of the morphometry file to be loaded.
hemi	string, one of 'lh', 'rh' or 'both'. The hemisphere name. Used to construct the names of the annotation and morphometry data files to be loaded.
fwhm	string. Smoothing as string, e.g. '10' or '25'.
agg_fun	function. An R function that aggregates data, typically <code>max</code> , <code>mean</code> , <code>min</code> or something similar. Note: this is NOT a string, put the function name without quotes. Defaults to <code>mean</code> .
template_subject	string. Template subject name, defaults to 'fsaverage'.
format	string. One of 'mgh', 'mgz', 'curv'. Defaults to 'mgh'.
cortex_only	logical, whether to mask the medial wall, i.e., whether the morphometry data for all vertices which are <i>not</i> part of the cortex (as defined by the label file 'label/?h.cortex.label') should be replaced with NA values. In other words, setting

this to TRUE will ignore the values of the medial wall between the two hemispheres. If set to true, the mentioned label file needs to exist for the subjects. Also not that the aggregation function will need to be able to cope with NA values if you set this to TRUE. You can use 'agg_fun_extra_params' if needed to achieve that, depending on the function. For example, if you use the `mean` function, you could set `agg_fun_extra_params=list("na.rm"=TRUE)` to get the mean of the vertices which are not masked. Defaults to FALSE.

`agg_fun_extra_params`

named list, extra parameters to pass to the aggregation function 'agg_fun' besides the loaded morphometry data. This is useful if you have masked the data and need to ignore NA values in the `agg_fun`.

`split_by_hemi` logical, whether to return a hemilist

See Also

Other aggregation functions: [group.agg.atlas.native\(\)](#), [group.agg.atlas.standard\(\)](#), [subject.atlas.agg\(\)](#)

`group.morph.native` *Retrieve native space morphometry data for a group of subjects.*

Description

Load native space morphometry data (like 'surf/lh.area') for a group of subjects from disk. Uses knowledge about the FreeSurfer directory structure to load the correct file.

Usage

```
group.morph.native(
  subjects_dir,
  subjects_list,
  measure,
  hemi,
  format = "curv",
  cortex_only = FALSE
)
```

Arguments

`subjects_dir`, string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.

`subjects_list`, vector of strings. The subject identifiers.

`measure`, string. Name of the vertex-wise measure of morphometry data file. E.g., "area" or "thickness". Used to construct the name of the morphometry file to be loaded.

`hemi`, string, one of 'lh', 'rh' or 'both'. The hemisphere name. Used to construct the names of the annotation and morphometry data files to be loaded.

format, string. One of 'mgh', 'mgz', 'curv'. Defaults to 'curv'.

cortex_only logical, whether to mask the medial wall, i.e., whether the morphometry data for all vertices which are **not** part of the cortex (as defined by the label file 'label/?h.cortex.label') should be replaced with NA values. In other words, setting this to TRUE will ignore the values of the medial wall between the two hemispheres. If set to true, the mentioned label file needs to exist for the subjects. Defaults to FALSE.

Value

named list with native space morph data, the names are the subject identifiers from the subjects_list, and the values are morphometry data vectors (of different length, as each subject has a different vertex count in native space).

See Also

Other morphometry data functions: [apply.label.to.morphdata\(\)](#), [apply.labeldata.to.morphdata\(\)](#), [group.morph.standard\(\)](#), [subject.morph.native\(\)](#), [subject.morph.standard\(\)](#)

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
subjects_list = c("subject1", "subject2");
data = group.morph.native(subjects_dir, subjects_list, "thickness", "lh");
```

group.morph.standard *Retrieve standard space morphometry data for a group of subjects.*

Description

Load standard space morphometry data (like 'surf/lh.area') for a group of subjects from disk. Uses knowledge about the FreeSurfer directory structure to load the correct file.

Usage

```
group.morph.standard(
  subjects_dir,
  subjects_list,
  measure,
  hemi = "both",
  fwhm = "10",
  template_subject = "fsaverage",
  format = "mgh",
  cortex_only = FALSE,
```

```

    df = FALSE
  )

```

Arguments

`subjects_dir`, string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.

`subjects_list`, vector of strings. The subject identifiers.

`measure`, string. Name of the vertex-wise measure of morphometry data file. E.g., "area" or "thickness". Used to construct the name of the morphometry file to be loaded.

`hemi`, string, one of 'lh', 'rh' or 'both'. The hemisphere name. Used to construct the names of the annotation and morphometry data files to be loaded.

`fwhm`, string. Smoothing as string, e.g. '10' or '25'.

`template_subject`, string. Template subject name, defaults to 'fsaverage'.

`format`, string. One of 'mgh', 'mgz', 'curv'. Defaults to 'mgh'.

`cortex_only` logical, whether to mask the medial wall, i.e., whether the morphometry data for all vertices which are *not* part of the cortex (as defined by the label file 'label/?h.cortex.label') should be replaced with NA values. In other words, setting this to TRUE will ignore the values of the medial wall between the two hemispheres. If set to true, the mentioned label file needs to exist for the template subject. Defaults to FALSE.

`df` logical, whether to return a dataframe instead of the named list. The dataframe will have one subject per column, and *n* rows, where *n* is the number of vertices of the template subject surface.

Value

named list with standard space morph data, the names are the subject identifiers from the `subjects_list`, and the values are morphometry data vectors (all with identical length, the data is mapped to a template subject).

See Also

Other morphometry data functions: [apply.label.to.morphdata\(\)](#), [apply.labeldata.to.morphdata\(\)](#), [group.morph.native\(\)](#), [subject.morph.native\(\)](#), [subject.morph.standard\(\)](#)

Examples

```

## Not run:
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
subjects_list = c("subject1", "subject2");
fulldata = group.morph.standard(subjects_dir, subjects_list, "thickness", "lh", fwhm='10');
mean(fulldata$subject1);

cortexdata = group.morph.standard(subjects_dir, subjects_list, "thickness",

```

```
"lh", fwhm='10', cortex_only=FALSE);  
mean(cortexdata$subject1, na.rm=TRUE);  
  
## End(Not run)
```

group.morph.standard.sf

Read combined data for a group from a single file.

Description

Read morphometry data for a group from a matrix in a single MGH or MGZ file.

Usage

```
group.morph.standard.sf(filepath, df = TRUE)
```

Arguments

filepath	character string, path to a file in MGH or MGZ format
df	logical, whether to return a data.frame, like <code>group.morph.standard</code> . If FALSE, the raw 4d matrix is returned.

Value

dataframe or 4d matrix, the morph data. See parameter 'df' for details.

Note

The file has typically been generated by running `mris_preproc` and/or `mri_surf2surf` on the command line, or written from R using [write.group.morph.standard.sf](#). The file contains no information on the subject identifiers, you need to know the subjects and their order in the file. Same goes for the hemisphere.

See Also

If you have created the input data file in FreeSurfer based on an FSGD file, you can read the subject identifiers from that FSGD file using [read.md.subjects.from.fsgd](#).

```
group.multimorph.agg.native
```

Aggregate native space morphometry data for multiple measures over hemispheres for a group of subjects.

Description

Compute the mean (or other aggregates) over all vertices of a subject from native space morphometry data (like 'surf/lh.area'). You can specify several measures and hemispheres. Uses knowledge about the FreeSurfer directory structure to load the correct files.

Usage

```
group.multimorph.agg.native(
  subjects_dir,
  subjects_list,
  measures,
  hemis,
  agg_fun = mean,
  format = "curv",
  cast = TRUE,
  cortex_only = FALSE,
  agg_fun_extra_params = NULL
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subjects_list	string vector. A vector of subject identifiers that match the directory names within subjects_dir.
measures	vector of strings. Names of the vertex-wise morphometry measures. E.g., c("area", "thickness"). Used to construct the names of the morphometry file to be loaded.
hemis	string, one of 'lh', 'rh' or 'both'. The hemisphere name. Used to construct the names of the annotation and morphometry data files to be loaded.
agg_fun	function. An R function that aggregates data, typically <code>max</code> , <code>mean</code> , <code>min</code> or something similar. Note: this is NOT a string, put the function name without quotes. Defaults to <code>mean</code> .
format	string. One of 'mgh', 'mgz', 'curv'. Defaults to 'mgh'.
cast	Whether a separate 'hemi' column should exist.
cortex_only	logical, whether to mask the medial wall, i.e., whether the morphometry data for all vertices which are <i>not</i> part of the cortex (as defined by the label file 'label/?h.cortex.label') should be replaced with NA values. In other words, setting

this to TRUE will ignore the values of the medial wall between the two hemispheres. If set to true, the mentioned label file needs to exist for the subjects. Also not that the aggregation function will need to be able to cope with NA values if you set this to TRUE. You can use 'agg_fun_extra_params' if needed to achieve that, depending on the function. For example, if you use the `mean` function, you could set `agg_fun_extra_params=list("na.rm"=TRUE)` to get the mean of the vertices which are not masked. Defaults to FALSE.

`agg_fun_extra_params`

named list, extra parameters to pass to the aggregation function 'agg_fun' besides the loaded morphometry data. This is useful if you have masked the data and need to ignore NA values in the `agg_fun`.

Value

dataframe with aggregated values over all measures and hemis for all subjects, with `m` columns and `n` rows, where `n` is the number of subjects. The `m` columns are 'subject_id' and '<hemi>.<measure>' (e.g., "lh.thickness") for all combinations of hemi and measure, the latter contains the aggregated data.

See Also

Other global aggregation functions: [group.morph.agg.native\(\)](#), [group.morph.agg.standard\(\)](#), [group.multimorph.agg.standard\(\)](#)

Examples

```
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
subjects_list = c("subject1", "subject2")
data = group.multimorph.agg.native(subjects_dir, subjects_list, c("thickness", "area"),
  c("lh", "rh"), cast=FALSE, cortex_only=TRUE, agg_fun=mean,
  agg_fun_extra_params=list("na.rm"=TRUE));
head(data);
```

group.multimorph.agg.standard

Aggregate standard space (fsaverage) morphometry data for multiple measures over hemispheres for a group of subjects.

Description

Compute the mean (or other aggregates) over all vertices of a subject from standard space morphometry data (like 'surf/lh.area.fwhm10.fsaverage.mgh'). You can specify several measures and hemispheres. Uses knowledge about the FreeSurfer directory structure to load the correct files.

Usage

```
group.multimorph.agg.standard(
  subjects_dir,
  subjects_list,
  measures,
  hemis,
  fwhm,
  agg_fun = mean,
  template_subject = "fsaverage",
  format = "mgh",
  cast = TRUE,
  cortex_only = FALSE,
  agg_fun_extra_params = NULL
)
```

Arguments

`subjects_dir`, string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.

`subjects_list`, string vector. A vector of subject identifiers that match the directory names within `subjects_dir`.

`measures`, vector of strings. Names of the vertex-wise morphometry measures. E.g., `c("area", "thickness")`. Used to construct the names of the morphometry file to be loaded.

`hemis`, string, one of 'lh', 'rh' or 'both'. The hemisphere name. Used to construct the names of the annotation and morphometry data files to be loaded.

`fwhm`, string. Smoothing as string, e.g. '10' or '25'.

`agg_fun`, function. An R function that aggregates data, typically `max`, `mean`, `min` or something similar. Note: this is NOT a string, put the function name without quotes. Defaults to `mean`.

`template_subject`, string. Template subject name, defaults to 'fsaverage'.

`format`, string. One of 'mgh', 'mgz', 'curv'. Defaults to 'mgh'.

`cast`, Whether a separate 'hemi' column should exist.

`cortex_only` logical, whether to mask the medial wall, i.e., whether the morphometry data for all vertices which are *not* part of the cortex (as defined by the label file 'label/?h.cortex.label') should be replaced with NA values. In other words, setting this to TRUE will ignore the values of the medial wall between the two hemispheres. If set to true, the mentioned label file needs to exist for the subjects. Also note that the aggregation function will need to be able to cope with NA values if you set this to TRUE. You can use 'agg_fun_extra_params' if needed to achieve that, depending on the function. For example, if you use the `mean` function, you could set `agg_fun_extra_params=list("na.rm"=TRUE)` to get the mean of the vertices which are not masked. Defaults to FALSE.

agg_fun_extra_params

named list, extra parameters to pass to the aggregation function 'agg_fun' besides the loaded morphometry data. This is useful if you have masked the data and need to ignore NA values in the agg_fun.

Value

dataframe with aggregated values over all measures and hemis for all subjects, with m columns and n rows, where n is the number of subjects. The m columns are 'subject_id' and '<hemi>.<measure>' (e.g., "lh.thickness") for all combinations of hemi and measure, the latter contains the aggregated data.

See Also

Other global aggregation functions: [group.morph.agg.native\(\)](#), [group.morph.agg.standard\(\)](#), [group.multimorph.agg.native\(\)](#)

group.surface	<i>Retrieve surface mesh data for a group of subjects.</i>
---------------	--

Description

Retrieve surface mesh data for a group of subjects.

Usage

```
group.surface(
  subjects_dir,
  subjects_list,
  surface,
  hemi = "both",
  force_hemilist = TRUE
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subjects_list	vector of strings. The subject identifiers.
surface	character string, the surface to load. Something like 'white' or 'pial'.
hemi	string, one of 'lh', 'rh' or 'both'. The hemisphere name. Used to construct the names of the mesh files to be loaded.
force_hemilist	logical, whether to force the individual values inside the named return value list to be hemilists (even if the 'hemi' parameter is not set to 'both'). If this is FALSE, the inner values will contain the respective (lh or rh) surface only.

Value

named list of surfaces: Each name is a subject identifier from `subjects_list`, and the values are hemilists of `'fs.surface'` instances.

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
subjects_list = c("subject1", "subject2");
surfaces = group.surface(subjects_dir, subjects_list, 'white', "both");
```

hasIn

Check for values in nested named lists

Description

Check for values in nested named lists

Usage

```
hasIn(named_list, listkeys)
```

Arguments

<code>named_list</code>	a named list
<code>listkeys</code>	vector of character strings, the nested names of the lists

Value

whether a non-NULL value exists at the path

Examples

```
data = list("regions"=list("frontal"=list("thickness"=2.3, "area"=2345)));
hasIn(data, c("regions", "nosuchregion")); # FALSE
```

`hemilist.derive.hemi` *Derive 'hemi' string from the data in a hemilist*

Description

Derive 'hemi' string from the data in a hemilist

Usage

```
hemilist.derive.hemi(hemilist)
```

Arguments

`hemilist` `hemilist`, an existing hemilist

Value

character string, one of 'lh', 'rh' or 'both'

`hemilist.get.combined.data`
Get combined data of hemi list

Description

Get combined data of hemi list

Usage

```
hemilist.get.combined.data(hemi_list)
```

Arguments

`hemi_list` named list, can have entries 'lh' and/or 'rh'

Value

the data combined with `c`, or NULL if both entries are NULL.

hemilist.unwrap	<i>Unwrap hemi data from a named hemi list.</i>
-----------------	---

Description

Unwrap hemi data from a named hemi list.

Usage

```
hemilist.unwrap(hemi_list, hemi = NULL, allow_null_list = FALSE)
```

Arguments

hemilist	named list, can have entries 'lh' and/or 'rh'
hemi	character string, the hemi data name to retrieve from the list. Can be NULL if the list only has a single entry.
allow_null_list	logical, whether to silently return NULL instead of raising an error if 'hemilist' is NULL

Value

the data

hemilist.wrap	<i>Wrap data into a named hemi list.</i>
---------------	--

Description

Wrap data into a named hemi list.

Usage

```
hemilist.wrap(data, hemi, hemilist = NULL)
```

Arguments

data	something to wrap, typically some data for a hemisphere, e.g., a vector of morphometry data values. If NULL, the name will not be created.
hemi	character string, one of 'lh' or 'rh'. The name to use for the data in the returned list.
hemilist	optional hemilist, an existing hemilist to add the entry to. If left at the default value 'NULL', a new list will be created.

Value

named list, with the 'data' in the name given by parameter 'hemi'

is.fs.coloredmesh *Check whether object is an fs.coloredmesh (S3)*

Description

Check whether object is an fs.coloredmesh (S3)

Usage

```
is.fs.coloredmesh(x)
```

Arguments

x any 'R' object

Value

TRUE if its argument is a coloredmesh (that is, has "fs.coloredmesh" amongst its classes) and FALSE otherwise.

is.fs.coloredvoxels *Check whether object is an fs.coloredvoxels instance (S3)*

Description

Check whether object is an fs.coloredvoxels instance (S3)

Usage

```
is.fs.coloredvoxels(x)
```

Arguments

x any 'R' object

Value

TRUE if its argument is a fs.coloredvoxels instance (that is, has "fs.coloredvoxels" among its classes) and FALSE otherwise.

is.hemilist	<i>Check whether x is a hemilist</i>
-------------	--------------------------------------

Description

A hemilist is a named list with entries 'lh' and/or 'rh'.

Usage

```
is.hemilist(x)
```

Arguments

x any R object

Value

whether 'x' is a hemilist

label.border	<i>Compute border of a label.</i>
--------------	-----------------------------------

Description

Compute the border of a label (i.e., a subset of the vertices of a mesh). The border thickness can be specified. Useful to draw the outline of a region, e.g., a significant cluster on the surface or a part of a ROI from a brain parcellation.

Usage

```
label.border(  
  surface_mesh,  
  label,  
  inner_only = TRUE,  
  expand_inwards = 0L,  
  derive = FALSE  
)
```

Arguments

surface_mesh surface mesh, as loaded by [subject.surface](#) or [read.fs.surface](#).
label instance of class 'fs.label' or an integer vector, the vertex indices. This function only makes sense if they form a patch on the surface, but that is not checked.

inner_only	logical, whether only faces consisting only of label_vertices should be considered to be label faces. If FALSE, faces containing at least one label vertex will be used. Defaults to TRUE. Leave this alone if in doubt, especially if you want to draw several label borders which are directly adjacent on the surface.
expand_inwards	integer, border thickness extension. If given, once the border has been computed, it is extended by the given graph distance. It is guaranteed that the border only extends inwards, i.e., it will never extend to vertices which are not part of the label.
derive	logical, whether the returned result should also include the border edges and faces in addition to the border vertices. Takes longer if requested, defaults to FALSE.

Value

the border as a list with the following entries: 'vertices': integer vector, the vertex indices of the border. Iff the parameter 'derive' is TRUE, the following two additional fields are included: 'edges': integer matrix of size (n, 2) for n edges. Each row defines an edge by its start and target vertex. 'faces': integer vector, the face indices of the border.

See Also

Other surface mesh functions: [face.edges\(\)](#), [mesh.vertex.included.faces\(\)](#), [mesh.vertex.neighbors\(\)](#), [subject.surface\(\)](#), [vis.path.along.verts\(\)](#)

label.colFn	<i>A simple colormap function for binary colors.</i>
-------------	--

Description

Useful for plotting labels.

Usage

```
label.colFn(n = 2L, col_a = "#228B22", col_b = "#FFFFFF")
```

Arguments

n	positive integer, the number of colors. Must be 1 or 2 for this function.
col_a	color string, the foreground color
col_b	color string, the background color

Value

vector of 'n' RGB colorstrings

label.colFn.inv *A simple colormap function for binary colors.*

Description

Useful for plotting labels.

Usage

```
label.colFn.inv(n = 2L, col_a = "#228B22", col_b = "#FFFFFF")
```

Arguments

n positive integer, the number of colors. Must be 1 or 2 for this function.
col_a color string, the foreground color
col_b color string, the background color

Value

vector of 'n' RGB colorstrings

label.from.annotdata *Extract a region from an annotation as a label.*

Description

The returned label can be used to mask morphometry data, e.g., to set the values of a certain region to 'NaN' or to extract only values from a certain region.

Usage

```
label.from.annotdata(  
  annotdata,  
  region,  
  return_one_based_indices = TRUE,  
  invert = FALSE,  
  error_on_invalid_region = TRUE  
)
```

Arguments

- annotdata, annotation. An annotation for one hemisphere, as returned by `subject.annot`. This must be the loaded data, not a path to a file.
- region, string. A valid region name for the annotation, i.e., one of the regions of the atlas used to create the annotation.
- return_one_based_indices, logical. Whether the indices should be 1-based. Indices are stored zero-based in label files, but R uses 1-based indices. Defaults to TRUE.
- invert, logical. If TRUE, return the indices of all vertices which are NOT part of the region. Defaults to FALSE.
- error_on_invalid_region, logical. Whether to throw an error if the given region does not appear in the region list of the annotation. If set to FALSE, this will be ignored and an empty vertex list will be returned. Defaults to TRUE.

Value

integer vector with label data: the list of vertex indices in the label. See `'return_one_based_indices'` for important information.

See Also

Other atlas functions: `get.atlas.region.names()`, `group.agg.atlas.native()`, `group.agg.atlas.standard()`, `group.annot()`, `group.label.from.annot()`, `label.to.annot()`, `regions.to.ignore()`, `spread.values.over.annot()`, `spread.values.over.hemi()`, `spread.values.over.subject()`, `subject.annot()`, `subject.atlas.agg()`, `subject.label.from.annot()`, `subject.lobes()`

label.to.annot	<i>Merge several labels into an annotation</i>
----------------	--

Description

Merge several labels and a colortable into an annotation.

Usage

```
label.to.annot(
  label_vertices_by_region,
  num_vertices_in_surface,
  colortable_df = NULL,
  index_of_unknown_region = 1L
)
```

Arguments

- `label_vertices_by_region`
named list of integer vectors, the keys are strings which define region names, and the values are integer vectors: the vertex indices of the region.
- `num_vertices_in_surface`
integer, total number of vertices in the surface mesh
- `colortable_df` NULL or dataframe, a colortable. It must contain the columns 'struct_name', 'r', 'g', 'b', and 'a'. All other columns will be derived if missing. The entries in 'struct_name' must match keys from the 'label_vertices_by_region' parameter. There must be one more row in here than there are labels. This row identifies the 'unknown' region (see also parameter 'index_of_unknown_region'). If NULL, a colortable will be auto-generated.
- `index_of_unknown_region`
positive integer, the index of the row in 'colortable_df' that defines the 'unknown' or 'background' region to which all vertices will be assigned which are *not* part of any of the given labels.

Value

an annotation, see [read.fs.annot](#) for details.

See Also

Other atlas functions: [get.atlas.region.names\(\)](#), [group.agg.atlas.native\(\)](#), [group.agg.atlas.standard\(\)](#), [group.annot\(\)](#), [group.label.from.annot\(\)](#), [label.from.annotdata\(\)](#), [regions.to.ignore\(\)](#), [spread.values.over.annot\(\)](#), [spread.values.over.hemi\(\)](#), [spread.values.over.subject\(\)](#), [subject.annot\(\)](#), [subject.atlas.agg\(\)](#), [subject.label.from.annot\(\)](#), [subject.lobes\(\)](#)

Examples

```
# Create two labels. Real-world labels would have more vertices, of course.
label1 = c(46666, 46777);
label2 = c(99888, 99889);
label_vertices = list("region1"=label1, "region2"=label2);
colortable_df = data.frame("struct_index"=seq(0, 2),
  "struct_name"=c("unknown", "region1", "region2"),
  "r"=c(255L, 255L, 0L), "g"=c(255L, 0L, 255L), "b"=c(255L, 0L, 0L), "a"=c(0L, 0L, 0L));
annot = label.to.annot(label_vertices, 100000, colortable_df);
```

labeldata.from.mask *Create labeldata from a mask.*

Description

Create labeldata from a mask. This function is trivial and only calls [which](#) after performing basic sanity checks.

Usage

```
labeldata.from.mask(mask, invert = FALSE)
```

Arguments

mask	a logical vector
invert	Whether to report the inverse the mask before determining the indices. Defaults to FALSE.

Value

labeldata. The list of indices which are TRUE in the mask (or the ones which FALSE if 'invert' is TRUE).

See Also

Other label data functions: [group.label\(\)](#), [mask.from.labeldata.for.hemi\(\)](#), [subject.label\(\)](#)

limit_fun	<i>Get data limiting function.</i>
-----------	------------------------------------

Description

Get data limiting function to use in rglactions as 'trans_fun' to transform data. This is typically used to limit the colorbar in a plot to a certain range. This is similar to [clip.data](#), but uses absolute values instead of percentiles to clip.

Usage

```
limit_fun(vmin, vmax)
```

Arguments

vmin	numerical scalar, the lower border. Data values below this will be set to vmin in the return value.
vmax	numerical scalar, the upper border. Data values above this will be set to vmax in the return value.

Value

a function that takes as argument the data, and clips it to the requested range. I.e., values outside the range will be set to the closest border value ('vmin' or 'vmax'). Designed to be used as `rglactions$trans_fun` in vis functions, to limit the colorbar and data range.

Examples

```
rglactions = list("trans_fun"=limit_fun(2,3));
```

limit_fun_na	<i>Get data limiting function to NA.</i>
--------------	--

Description

Get data limiting function to use in `rglactions` as `'trans_fun'` to transform data. This is typically used to limit the colorbar in a plot to a certain range. This is similar to `clip.data`, but uses absolute values instead of percentiles to clip.

Usage

```
limit_fun_na(vmin, vmax)
```

Arguments

vmin	numerical scalar, the lower border. Data values below this will be set to vmin in the return value.
vmax	numerical scalar, the upper border. Data values above this will be set to vmax in the return value.

Value

a function that takes as argument the data, and clips it to the requested range. I.e., values outside the range will be set to 'NA'. Designed to be used as `rglactions$trans_fun` in vis functions, to limit the colorbar and data range.

Note

This is useful for thresholding stuff like t-value maps. All values outside the range will be displayed as the background color.

Examples

```
rglactions = list("trans_fun"=limit_fun_na(2,3));
```

list_optional_data	<i>Get file names available in package cache.</i>
--------------------	---

Description

Get file names of optional data files which are available in the local package cache. You can access these files with `get_optional_data_file()`.

Usage

```
list_optional_data()
```

Value

vector of strings. The file names available, relative to the package cache.

```
mask.from.labeldata.for.hemi
```

Create a binary mask from labels.

Description

Create a binary mask for the data of a single hemisphere from one or more labels. A label contains the vertex indices which are part of it, but often having a mask in more convenient.

Usage

```
mask.from.labeldata.for.hemi(
  labels,
  num_vertices_in_hemi,
  invert_labels = FALSE,
  existing_mask = NULL
)
```

Arguments

labels	list of labels. A label is just a vector of vertex indices. It can be created manually, but is typically loaded from a label file using subject.label .
num_vertices_in_hemi	integer. The number of vertices of the surface for which the mask is created. This must be for a single hemisphere.
invert_labels	logical, whether to invert the label data.
existing_mask	an existing mask to modify or NULL. If it is NULL, a new mask will be created before applying any labels, and the values set during initialization of this new mask are the negation of the 'invert_label' parameter. Defaults to NULL.

Value

logical vector. The mask. It contains a logical value for each vertex. By default, the vertex indices from the labels are FALSE and the rest are TRUE, but this can be changed with the parameter 'invert_labels'.

See Also

Other label data functions: [group.label\(\)](#), [labeldata.from.mask\(\)](#), [subject.label\(\)](#)

Other mask functions: [coloredmesh.from.mask\(\)](#), [vis.mask.on.subject\(\)](#)

Examples

```

fsbrain::download_optional_data();

# Define the data to use:
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
subject_id = 'subject1';
surface = 'white';
hemi = 'both';
atlas = 'aparc';
region = 'bankssts';

# Create a mask from a region of an annotation:
lh_annot = subject.annot(subjects_dir, subject_id, 'lh', atlas);
rh_annot = subject.annot(subjects_dir, subject_id, 'rh', atlas);
lh_label = label.from.annotdata(lh_annot, region);
rh_label = label.from.annotdata(rh_annot, region);
lh_mask = mask.from.labeldata.for.hemi(lh_label, length(lh_annot$vertices));
rh_mask = mask.from.labeldata.for.hemi(rh_label, length(rh_annot$vertices));

# Edit the mask: add the vertices from another region to it:
region2 = 'medialorbitofrontal';
lh_label2 = label.from.annotdata(lh_annot, region2);
rh_label2 = label.from.annotdata(rh_annot, region2);
lh_mask2 = mask.from.labeldata.for.hemi(lh_label2, length(lh_annot$vertices),
existing_mask = lh_mask);
rh_mask2 = mask.from.labeldata.for.hemi(rh_label2, length(rh_annot$vertices),
existing_mask = rh_mask);

```

mesh.vertex.neighbors *Compute neighborhood of a vertex*

Description

Given a set of query vertex indices and a mesh **m**, compute all vertices which are adjacent to the query vertices in the mesh. A vertex **u** is **adjacent** to another vertex **v** iff there exists an edge **e = (u, v)** in **m**. While you could call this function repeatedly with the old output as its new input to extend the neighborhood, you should maybe use a proper graph library for this.

Usage

```

mesh.vertex.neighbors(
  surface,
  source_vertices,
  k = 1L,
  restrict_to_vertices = NULL
)

```

Arguments

surface	a surface as returned by functions like <code>subject.surface</code> or <code>read.fs.surface</code> .
source_vertices	Vector of source vertex indices.
k	positive integer, how often to repeat the procedure and grow the neighborhood, using the output 'vertices' as the 'source_vertices' for the next iteration. Warning: settings this to high values will be very slow for large meshes.
restrict_to_vertices	integer vector of vertex indices. If given, the neighborhood growth will be limited to the given vertex indices. Defaults to NULL, which means the neighborhood is not restricted.

Value

the neighborhood as a list with two entries: "faces": integer vector, the face indices of all faces the source_vertices are a part of. "vertices": integer vector, the unique vertex indices of all vertices of the faces in the 'faces' property. These vertex indices include the indices of the source_vertices themselves.

See Also

Other surface mesh functions: `face.edges()`, `label.border()`, `mesh.vertex.included.faces()`, `subject.surface()`, `vis.path.along.verts()`

mkco.cluster

Return recommended 'makecmap_options' for diverging cluster data.

Description

This function returns recommended visualization settings (a colormap function and suitable other settings) for the given type of data. The return value is meant to be passed as parameter 'makecmap_options' to the vis.* functions, e.g., `vis.subject.morph.native`.

Usage

```
mkco.cluster()
```

Value

named list, visualization settings to be used as 'makecmap_options' for diverging data.

Note

This uses a cyan blue red yellow colormap, which is popular for displaying clusters in neuroscience.

mkco.div	<i>Return recommended 'makecmap_options' for diverging data.</i>
----------	--

Description

This function returns recommended visualization settings (a colormap function and suitable other settings) for the given type of data. The return value is meant to be passed as parameter 'makecmap_options' to the vis.* functions, e.g., [vis.subject.morph.native](#).

Usage

```
mkco.div()
```

Value

named list, visualization settings to be used as 'makecmap_options' for diverging data.

mkco.heat	<i>Return recommended 'makecmap_options' for sequential data with heatmap style.</i>
-----------	--

Description

This function returns recommended visualization settings (a colormap function and suitable other settings) for the given type of data. The return value is meant to be passed as parameter 'makecmap_options' to the vis.* functions, e.g., [vis.subject.morph.native](#).

Usage

```
mkco.heat()
```

Value

named list, visualization settings to be used as 'makecmap_options' for sequential data with heatmap style.

mkco.seq	<i>Return recommended 'makecmap_options' for sequential data.</i>
----------	---

Description

This function returns recommended visualization settings (a colormap function and suitable other settings) for the given type of data. The return value is meant to be passed as parameter 'makecmap_options' to the vis.* functions, e.g., [vis.subject.morph.native](#).

Usage

```
mkco.seq()
```

Value

named list, visualization settings to be used as 'makecmap_options' for sequential data.

principal.curvatures	<i>Computes principal curvatures according to 2 definitions from raw k1 and k2 values.</i>
----------------------	--

Description

Computes principal curvatures according to 2 definitions from raw k1 and k2 values.

Usage

```
principal.curvatures(k1_raw, k2_raw)
```

Arguments

k1_raw	numerical vector, one of the two principal curvatures, one value per vertex
k2_raw	numerical vector, the other one of the two principal curvatures, one value per vertex

Value

a named 'principal_curvatures' list, with entries 'principal_curvature_k1': larger value of k1_raw, k2_raw. 'principal_curvature_k2': smaller value of k1_raw, k2_raw. 'principal_curvature_k_major': larger value of abs(k1_raw), abs(k2_raw). 'principal_curvature_k_minor': smaller value of abs(k1_raw), abs(k2_raw).

print.fs.coloredmesh *Print description of a brain coloredmesh (S3).*

Description

Print description of a brain coloredmesh (S3).

Usage

```
## S3 method for class 'fs.coloredmesh'  
print(x, ...)
```

Arguments

x	brain surface with class 'fs.coloredmesh'.
...	further arguments passed to or from other methods

print.fs.coloredvoxels
Print description of fs.coloredvoxels (S3).

Description

Print description of fs.coloredvoxels (S3).

Usage

```
## S3 method for class 'fs.coloredvoxels'  
print(x, ...)
```

Arguments

x	brain voxel tris with class 'fs.coloredvoxels'.
...	further arguments passed to or from other methods

 qc.for.group

Perform data quality check based on computed region stats.

Description

Determine subjects that potentially failed segmentation, based on region-wise morphometry data. The stats can be computed from any kind of data, but something like area or volume most likely works best. The stats are based on the mean of the region values, so the measure should at least roughly follow a normal distribution.

Usage

```
qc.for.group(subjects_dir, subjects_list, measure, atlas, hemi = "both", ...)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subjects_list	string vector. A vector of subject identifiers that match the directory names within subjects_dir.
measure	string. Name of the vertex-wise measure of morphometry data file. E.g., "area" or "thickness". Used to construct the name of the morphometry file to be loaded.
atlas	string. The atlas name. E.g., "aparc", "aparc.2009s", or "aparc.DKTatlas". Used to construct the name of the annotation file to be loaded.
hemi	string, one of 'lh', 'rh', or 'both'. The hemisphere name. Used to construct the names of the annotation and morphometry data files to be loaded. If set to 'both', combined data for 'lh' and 'rh' will be used.
...	parameters passed to qc.from.regionwise.df .

Value

qc result as a hemilist, each entry contains a named list as returned by [qc.from.regionwise.df](#).

See Also

Other quality check functions: [qc.from.regionwise.df\(\)](#), [qc.from.segstats.table\(\)](#)

qc.from.regionwise.df *Perform data quality check based on a dataframe containing aggregated region-wise data.*

Description

Determine subjects that potentially failed segmentation, based on region-wise data. The data can be anything, but there must be one numerical value per subject per region.

Usage

```
qc.from.regionwise.df(  
  rdf,  
  z_threshold = 2.8,  
  verbosity = 0L,  
  num_bad_regions_allowed = 1L  
)
```

Arguments

rdf	data.frame, the region data. The first column must contain the subject identifier, all other columns should contain numerical data for a single region. (Each row represents a subject.) This can be produced by calling group.agg.atlas.native .
z_threshold	numerical, the cutoff value for considering a subject an outlier (in standard deviations).
verbosity	integer, controls the output to stdout. 0=off, 1=normal, 2=verbose.
num_bad_regions_allowed	integer, the number of regions in which subjects are allowed to be outliers without being reported as potentially failed segmentation

Value

named list with entries: 'failed_subjects': vector of character strings, the subject identifiers which potentially failed segmentation. 'mean_dists_z': distance to mean, in standard deviations, per subject per region. 'num_outlier_subjects_per_region': number of outlier subjects by region. 'meta-data': named list of metadata, e.g., hemi, atlas and measure used to compute these QC results.

See Also

Other quality check functions: [qc.for.group\(\)](#), [qc.from.segstats.table\(\)](#)

```
qc.from.segstats.tables
```

Perform data quality check based on a segstats table.

Description

Determine subjects that potentially failed segmentation, based on segstats table data. The input table file must be a segmentation or parcellation table, generated by running the FreeSurfer tools 'aparcstats2table' or 'asegstats2table' for your subjects.

Usage

```
qc.from.segstats.tables(filepath_lh, filepath_rh, ...)
```

Arguments

filepath_lh	path to left hemisphere input file, a tab-separated file generated by running the FreeSurfer tools 'aparcstats2table' or 'asegstats2table'. The command line in the system shell would be something like 'aparcstats2table_bin -subjectsfile \$subjects_file -meas \$measure -hemi \$hemi -t \$aparc_output_table'.
filepath_rh	path to equivalent right hemisphere input file.
...	parameters passed to qc.from.regionwise.df .

Value

qc result as a hemilist, each entry contains a named list as returned by [qc.from.regionwise.df](#).

```
qc.vis.failcount.by.region
```

Visualize the number of outlier subjects per region in your dataset.

Description

The function helps you to see which regions are affected the most by QC issues: for each region, it plots the number of subjects which are outliers in the region.

Usage

```
qc.vis.failcount.by.region(
  qc_res,
  atlas,
  subjects_dir = fsaverage.path(),
  subject_id = "fsaverage",
  ...
)
```

Arguments

qc_res	hemilist of QC results, as returned by functions like qc.for.group or qc.from.segstats.tables .
atlas	string. The brain atlas to use. E.g., 'aparc' or 'aparc.a2009s'.
subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	string. The subject identifier.
...	extra parameters passed to vis.region.values.on.subject . E.g., to change to interactive view, get a colorbar and better resolution, try: <code>draw_colorbar=T,rgloptions = rglo(),views='si'</code> .

Note

You can visualize this on any subject you like, 'fsaverage' is a typical choice. The atlas must be the one used during the QC step.

ras2vox_tkr	<i>The FreeSurfer default ras2vox_tkr matrix.</i>
-------------	---

Description

Applying this matrix to a FreeSurfer surface RAS coordinate (from a surface file like 'lh.white') will give you the voxel index (CRS) in a conformed FreeSurfer volume. The returned matrix is the inverse of the 'vox2ras_tkr' matrix.

Usage

```
ras2vox_tkr()
```

Value

numeric 4x4 matrix, the FreeSurfer ras2vox_tkr matrix.

See Also

Other surface and volume coordinates: [vox2ras_tkr\(\)](#)

Examples

```
# Compute the FreeSurfer CRS voxel index of surface RAS coordinate (0.0, 0.0, 0.0):
ras2vox_tkr() %*% c(0, 0, 0, 1);
# Show that the voxel at surface RAS corrd (0.0, 0.0, 0.0) is the one with CRS (128, 128, 128):
ras2vox_tkr() %*% c(0.0, 0.0, 0.0, 1);
```

read.colorcsv	<i>Read colors from CSV file.</i>
---------------	-----------------------------------

Description

Read colors from CSV file.

Usage

```
read.colorcsv(filepath)
```

Arguments

filepath character string, path to a CSV file containing colors

Value

vector of hex color strings

read.md.demographics	<i>Read demographics file</i>
----------------------	-------------------------------

Description

Load a list of subjects and metadata from a demographics file, i.e., a tab-separated file containing an arbitrary number of columns, one of which must be the subject id.

Usage

```
read.md.demographics(  
  demographics_file,  
  column_names = NULL,  
  header = FALSE,  
  scale_and_center = FALSE,  
  sep = "",  
  report = FALSE,  
  stringsAsFactors = TRUE,  
  group_column_name = NULL  
)
```


Arguments

demographics_file,	string. The path to the file.
column_names,	vector of strings. The column names to set in the returned dataframe. The length must match the number of columns in the file.
header,	logical. Whether the file starts with a header line.
scale_and_center,	logical. Whether to center and scale the data. Defaults to FALSE.
sep,	string. Separator passed to read.table , defaults to tabulator.
report,	logical. Whether to write an overview, i.e., some descriptive statistics for each column, to STDOUT. Defaults to FALSE. See report.on.demographics .
stringsAsFactors,	logical. Whether to convert strings in the input data to factors. Defaults to TRUE.
group_column_name,	string or NULL. If given, the column name of the group column. It must be a factor column with 2 levels. Enables group-comparison tests. Defaults to NULL.

Value

a dataframe. The data in the file. String columns will be returned as factors, which you may want to adapt afterwards for the subject identifier column.

See Also

Other metadata functions: [demographics.to.fsgd.file\(\)](#), [read.md.subjects\(\)](#), [report.on.demographics\(\)](#)

Examples

```
demographics_file =
  system.file("extdata", "demographics.tsv", package = "fsbrain", mustWork = TRUE);
column_names = c("subject_id", "group", "age");
demographics = read.md.demographics(demographics_file,
  header = TRUE, column_names = column_names, report = FALSE);
```

read.md.subjects	<i>Read subjects file</i>
------------------	---------------------------

Description

Load a list of subjects from a subjects file, i.e., a simple text file containing one subject name per line.

Usage

```
read.md.subjects(subjects_file, header)
```

Arguments

`subjects_file` character string, the path to the subjects file.
`header` logical, whether the file starts with a header line.

Value

vector of strings, the subject identifiers.

See Also

Other metadata functions: [demographics.to.fsgd.file\(\)](#), [read.md.demographics\(\)](#), [report.on.demographics\(\)](#)

Examples

```
subjects_file = system.file("extdata", "subjects.txt", package = "fsbrain", mustWork = TRUE);  
subjects_list = read.md.subjects(subjects_file, header = FALSE);
```

```
read.md.subjects.from.fsgd
```

Read subjects list from an FSGD file.

Description

Read subjects list from an FSGD file.

Usage

```
read.md.subjects.from.fsgd(filepath)
```

Arguments

`filepath` character string, path to a FreeSurfer Group Descriptor (FSGD) file.

Value

vector of character strings, the subject identifiers

Note

This is not a parser for all data in an FSGD file.

See Also

[demographics.to.fsgd.file](#)

regions.to.ignore *Give suggestions for regions to ignore for an atlas.*

Description

Give suggestions for regions to ignore for an atlas. These are regions for which many subjects do not have any vertices in them, or the Medial Wall and Unknown regions.

Usage

```
regions.to.ignore(atlas)
```

Arguments

atlas, string. The name of an atlas. Supported strings are 'aparc' and 'aparc.a2009s'.

Value

vector of strings, the region names.

See Also

Other atlas functions: [get.atlas.region.names\(\)](#), [group.agg.atlas.native\(\)](#), [group.agg.atlas.standard\(\)](#), [group.annot\(\)](#), [group.label.from.annot\(\)](#), [label.from.annotdata\(\)](#), [label.to.annot\(\)](#), [spread.values.over.annot\(\)](#), [spread.values.over.hemi\(\)](#), [spread.values.over.subject\(\)](#), [subject.annot\(\)](#), [subject.atlas.agg\(\)](#), [subject.label.from.annot\(\)](#), [subject.lobes\(\)](#)

Examples

```
aparc_regions_ign = regions.to.ignore('aparc');
aparc_a2009s_regions_ign = regions.to.ignore('aparc.a2009s');
```

report.on.demographics

Print a demographics report

Description

Print a demographics report

Usage

```
report.on.demographics(
  demographics_df,
  group_column_name = NULL,
  paired = FALSE
)
```

Arguments

demographics_df	a demographics data.frame, as returned by read.md.demographics .
group_column_name,	string or NULL. If given, the column name of the group column. It must be a factor column with 2 levels. Enables group-comparison tests. Defaults to 'NULL'.
paired	Whether the data of the two groups is paired (repeated measurements). Only relevant if group_column_name is given and tests for group differences are included in the report. Defaults to 'FALSE'.

Value

vector of character strings, the lines of the demographics report.

See Also

Other metadata functions: [demographics.to.fsgd.file\(\)](#), [read.md.demographics\(\)](#), [read.md.subjects\(\)](#)

rglactions	<i>Create rglactions list, suitable to be passed as parameter to vis functions.</i>
------------	---

Description

Create rglactions list, suitable to be passed as parameter to vis functions.

Usage

```
rglactions()
```

Value

named list, an example 'rglactions' instance that will save a screenshot of the plot produced by the vis function in the current working directory (see `getwd`), under the name 'fsbrain_out.png'.

Note

List of all available rglactions: (1) 'snapshot_png=filepath' takes a screenshot in PNG format and saves it in at 'filepath'. (2) 'trans_fun=function' uses the transformation function trans_fun to the data before mapping data values to colors and plotting. Popular transformation functions are [limit_fun](#), [limit_fun_na](#), and [clip.data](#). (3) 'text=arglist' calls [text3d](#) with the given args after plotting.

Examples

```

rgla_screenie = list('snapshot_png'='fsbain_out.png');
rgla_screenie = rglactions(); # same as above
rgla_clamp = list('trans_fun'=clip.data);
rgla_limit = list('trans_fun'=limit_fun(2,5));
rgla_ls = list('trans_fun'=limit_fun_na(2,5), 'snapshot_png'='~/fig1.png');

```

rglo

Get rgloptions and consider global options.

Description

This function retrieves the global rgloptions defined in `getOption('fsbrain.rgloptions')`, or, if this is not set, returns the value from `rglot`.

Usage

```
rglo()
```

Value

named list, usable as 'rgloptions' parameter for vis functions like `vis.subject.morph.native`.

Note

You can set the default size for all fsbrain figures to 1200x1200 pixels like this: `options("fsbrain.rgloptions"=list("wi`

rglot

Get rgloptions for testing.

Description

This function defines the figure size that is used during the unit tests. Currently `list('windowRect' = c(50, 50, 800, 800))`.

Usage

```
rglot()
```

Value

named list, usable as 'rgloptions' parameter for vis functions like `vis.subject.morph.native`.

 rglvoxels

Draw 3D boxes at locations using rgl.

Description

Draw 3D boxes at all given coordinates using `rgl`, analogous to `rgl.spheres`. Constructs the coordinates for triangles making up the boxes, then uses `triangles3d` to render them.

Usage

```
rglvoxels(centers, r = 1, voxelcol = NULL, do_show = TRUE, ...)
```

Arguments

<code>centers</code>	numerical matrix with 3 columns. Each column represents the x, y, z coordinates of a center at which to create a cube.
<code>r</code>	numerical vector or scalar, the cube edge length. This is the length of the axis-parallel edges of the cube. The vector must have length 1 (same edge length for all cubes), or the length must be identical to the number of rows in parameter ‘centers’.
<code>voxelcol</code>	vector of rgb color strings for the individual voxels. Its length must be identical to <code>nrow(centers)</code> if given.
<code>do_show</code>	logical, whether to visualize the result in the current rgl scene
<code>...</code>	material properties, passed to <code>triangles3d</code> . Example: <code>color = "#0000ff", lit=FALSE</code> .

Value

list of ‘fs.coloredvoxels’ instances, invisible. The function is called for the side effect of visualizing the data, and usually you can ignore the return value.

Examples

```
# Plot a 3D cloud of 500 red voxels:
centers = matrix(rnorm(500*3)*100, ncol=3);
rglvoxels(centers, voxelcol="red");
```

`shape.descriptor.names`*Get all shape descriptor names.*

Description

Get all shape descriptor names.

Usage

```
shape.descriptor.names()
```

Value

vector of character strings, the names

`shape.descriptors`*Computes geometric curvature-based descriptors.*

Description

Computes geometric curvature-based descriptors.

Usage

```
shape.descriptors(pc, descriptors = shape.descriptor.names())
```

Arguments

<code>pc</code>	a 'principal_curvatures' data list, see principal.curvatures for details.
<code>descriptors</code>	vector of character strings, the descriptors you want. See shape.descriptor.names for all available names.

Value

dataframe of descriptor values, each columns contains one descriptor.

References

Shimony et al. (2016). Comparison of cortical folding measures for evaluation of developing human brain. *Neuroimage*, 125, 780-790.

shift.hemis.apart *Shift hemispheres apart.*

Description

Modify mesh coordinates of a hemilist of coloredmeshes to introduce a gap between the two hemispheres.

Usage

```
shift.hemis.apart(
    coloredmeshes_hl,
    shift_by = NULL,
    axis = 1L,
    hemi_order_on_axis = "lr",
    min_dist = 0
)
```

Arguments

coloredmeshes_hl	hemilist of coloredmeshes
shift_by	numerical vector of length 2, the amount by which to shift the hemis. The first value is for the left hemi, the second for the right hemi (values can be negative). Pass 'NULL' to determine the shift automatically from the mesh coordinates, and adapt 'hemi_order_on_axis' to define how that happens.
axis	positive integer, one of 1L, 2L or 3L. The axis on which to shift (x,y,z).
hemi_order_on_axis	character string, one of 'auto', 'auto_flipped', 'lr' or 'rl'. Defines how to determine the order of the hemis on the axes. This is ignored unless 'shift_by' is 'NULL'. The 'auto' setting assumes that the hemisphere with the smaller minimal vertex coordinate (on the given axis) comes first. Note that if the overlap (or shift) is extreme, this may not hold anymore. Therefore, the default value is 'lr', which works for FreeSurfer data. The 'auto_flipped' setting will always return the inverse of 'auto', so if 'auto' did not work, 'auto_flipped' will.
min_dist	numerical scalar, the minimal distance of the hemispheres. Ignored unless 'shift_by' is 'NULL'.

Value

hemilist of coloredmeshes, the shifted meshes

```
spread.values.over.annot
```

Spread a single value for a region to all region vertices.

Description

Given an annotation and a list of values (one per brain region), return data that has the values for each region mapped to all region vertices.

Usage

```
spread.values.over.annot(
  annot,
  region_value_list,
  value_for_unlisted_regions = NaN,
  warn_on_unmatched_list_regions = FALSE,
  warn_on_unmatched_atlas_regions = FALSE
)
```

Arguments

`annot`, annotation. The result of calling `fs.read.annot`.

`region_value_list`, named list of strings. Each name must be a region name from the annotation, and the value must be the value to spread to all region vertices.

`value_for_unlisted_regions`, numeric scalar. The value to assign to vertices which are part of atlas regions that are not listed in `region_value_list`. Defaults to `NaN`.

`warn_on_unmatched_list_regions`, logical. Whether to print a warning when a region occurs in the `region_value_list` that is not part of the given atlas (and the value assigned to this region is thus ignored in the output file and data). Defaults to `FALSE`.

`warn_on_unmatched_atlas_regions`, logical. Whether to print a warning when a region occurs in the atlas that is not part of the given `region_value_list` (and thus the vertices of the region will be assigned the value 'value_for_unlisted_regions' in the output file and data). Defaults to `FALSE`.

Value

named list with following entries: "spread_data": a vector of length `n`, where `n` is the number of vertices in the annotation. One could write this to an MGH or curv file for visualization. "regions_not_in_annot": list of regions which are not in the annotation, but in the `region_value_list`. Their values were ignored.

See Also

Other atlas functions: `get.atlas.region.names()`, `group.agg.atlas.native()`, `group.agg.atlas.standard()`, `group.annot()`, `group.label.from.annot()`, `label.from.annotdata()`, `label.to.annot()`, `regions.to.ignore()`, `spread.values.over.hemi()`, `spread.values.over.subject()`, `subject.annot()`, `subject.atlas.agg()`, `subject.label.from.annot()`, `subject.lobes()`

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
annot = subject.annot(subjects_dir, 'subject1', 'lh', 'aparc');
region_value_list = list("bankssts"=0.9, "precuneus"=0.7);
morph_like_data =
spread.values.over.annot(annot, region_value_list, value_for_unlisted_regions=0.0);
```

spread.values.over.hemi

Spread the values in the region_value_list and return them for one hemisphere.

Description

Given an atlas and a list that contains one value for each atlas region, create morphometry data in which all region vertices are assigned the value. Can be used to plot stuff like p-values or effect sizes onto brain regions.

Usage

```
spread.values.over.hemi(
  subjects_dir,
  subject_id,
  hemi,
  atlas,
  region_value_list,
  value_for_unlisted_regions = NA,
  silent = FALSE
)
```

Arguments

`subjects_dir`, string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.

`subject_id`, string. The subject identifier

hemi,	string, one of 'lh' or 'rh'. The hemisphere name. Used to construct the names of the annotation and morphometry data files to be loaded.
atlas,	string. The atlas name. E.g., "aparc", "aparc.2009s", or "aparc.DKAtlas". Used to construct the name of the annotation file to be loaded.
region_value_list,	named list. A list in which the names are atlas regions, and the values are the value to write to all vertices of that region. You can pass an unnamed list or vector, but then the length must exactly match the number of regions in the atlas, and the order must match the annotation file of the subject and hemisphere. Use with care, and keep in mind that some subjects do not have all regions.
value_for_unlisted_regions,	numeric scalar. The value to assign to vertices which are part of atlas regions that are not listed in region_value_list. Defaults to NaN.
silent	logical, whether to suppress mapping info in case of unnamed region value lists (see 'lh_region_value_list' description).

Value

numeric vector containing the data.

See Also

Other atlas functions: [get.atlas.region.names\(\)](#), [group.agg.atlas.native\(\)](#), [group.agg.atlas.standard\(\)](#), [group.annot\(\)](#), [group.label.from.annot\(\)](#), [label.from.annotdata\(\)](#), [label.to.annot\(\)](#), [regions.to.ignore\(\)](#), [spread.values.over.annot\(\)](#), [spread.values.over.subject\(\)](#), [subject.annot\(\)](#), [subject.atlas.agg\(\)](#), [subject.label.from.annot\(\)](#), [subject.lobes\(\)](#)

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
region_value_list = list("bankssts"=0.9, "precuneus"=0.7);
morph_like_data =
  spread.values.over.hemi(subjects_dir, 'subject1', 'lh', 'aparc', region_value_list);
```

spread.values.over.subject

Spread the values in the region_value_list and return them for one hemisphere.

Description

Given an atlas and a list that contains one value for each atlas region, create morphometry data in which all region vertices are assigned the value. Can be used to plot stuff like p-values or effect sizes onto brain regions.

Usage

```
spread.values.over.subject(
  subjects_dir,
  subject_id,
  atlas,
  lh_region_value_list,
  rh_region_value_list,
  value_for_unlisted_regions = NaN,
  silent = FALSE
)
```

Arguments

`subjects_dir`, string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.

`subject_id`, string. The subject identifier

`atlas`, string. The atlas name. E.g., "aparc", "aparc.2009s", or "aparc.DKTatlas". Used to construct the name of the annotation file to be loaded.

`lh_region_value_list`,
named list. A list in which the names are atlas regions, and the values are the value to write to all vertices of that region. Applied to the left hemisphere.

`rh_region_value_list`,
named list. A list in which the names are atlas regions, and the values are the value to write to all vertices of that region. Applied to the right hemisphere.

`value_for_unlisted_regions`,
numeric scalar. The value to assign to vertices which are part of atlas regions that are not listed in `region_value_list`. Defaults to NaN.

`silent` logical, whether to suppress mapping info in case of unnamed region value lists (see 'lh_region_value_list' description).

Value

named list with entries 'lh' and 'rh'. Each value is a numeric vector containing the data for the respective hemisphere.

See Also

Other atlas functions: [get.atlas.region.names\(\)](#), [group.agg.atlas.native\(\)](#), [group.agg.atlas.standard\(\)](#), [group.annot\(\)](#), [group.label.from.annot\(\)](#), [label.from.annotdata\(\)](#), [label.to.annot\(\)](#), [regions.to.ignore\(\)](#), [spread.values.over.annot\(\)](#), [spread.values.over.hemi\(\)](#), [subject.annot\(\)](#), [subject.atlas.agg\(\)](#), [subject.label.from.annot\(\)](#), [subject.lobes\(\)](#)

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
lh_region_value_list = list("bankssts"=0.9, "precuneus"=0.7);
```

```
rh_region_value_list = list("bankssts"=0.5);
morph_like_data =
spread.values.over.subject(subjects_dir, 'subject1', 'aparc',
lh_region_value_list, rh_region_value_list);
```

subject.annot	<i>Load an annotation for a subject.</i>
---------------	--

Description

Load a brain surface annotation, i.e., a cortical parcellation based on an atlas, for a subject.

Usage

```
subject.annot(subjects_dir, subject_id, hemi, atlas)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	string. The subject identifier
hemi	string, one of 'lh' or 'rh'. The hemisphere name. Used to construct the names of the annotation and morphometry data files to be loaded.
atlas	string. The atlas name. E.g., "aparc", "aparc.2009s", or "aparc.DKAtlas". Used to construct the name of the annotation file to be loaded.

Value

the annotation, as returned by [read.fs.annot](#). It is a named list, entries are: "vertices" vector of n vertex indices, starting with 0. "label_codes": vector of n integers, each entry is a color code, i.e., a value from the 5th column in the table structure included in the "colortable" entry (see below). "label_names": the n brain structure names for the vertices, already retrieved from the colortable using the code. "hex_colors_rgb": Vector of hex color for each vertex. The "colortable" is another named list with 3 entries: "num_entries": int, number of brain structures. "struct_names": vector of strings, the brain structure names. "table": numeric matrix with num_entries rows and 5 columns. The 5 columns are: 1 = color red channel, 2=color blue channel, 3=color green channel, 4=color alpha channel, 5=unique color code. "colortable_df": The same information as a dataframe. Contains the extra columns "hex_color_string_rgb" and "hex_color_string_rgba" that hold the color as an RGB(A) hex string, like "#rrggbbaa".

See Also

Other atlas functions: [get.atlas.region.names\(\)](#), [group.agg.atlas.native\(\)](#), [group.agg.atlas.standard\(\)](#), [group.annot\(\)](#), [group.label.from.annot\(\)](#), [label.from.annotdata\(\)](#), [label.to.annot\(\)](#), [regions.to.ignore\(\)](#), [spread.values.over.annot\(\)](#), [spread.values.over.hemi\(\)](#), [spread.values.over.subject\(\)](#), [subject.atlas.agg\(\)](#), [subject.label.from.annot\(\)](#), [subject.lobes\(\)](#)

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
annot_lh = subject.annot(subjects_dir, "subject1", "lh", "aparc");
```

subject.atlas.agg *Aggregate morphometry data over brain atlas regions for a subject.*

Description

Aggregate morphometry data over brain atlas regions, e.g., compute the mean thickness value over all regions in an atlas.

Usage

```
subject.atlas.agg(
  vertex_morph_data,
  vertex_label_names,
  agg_fun = base::mean,
  requested_label_names = c()
)
```

Arguments

vertex_morph_data, numeric vector. The morphometry data, one value per vertex. The morphometry data are typically loaded from an MGZ or curv format file with the read.fs.curv or read.fs.mgh functions.

vertex_label_names, string vector. The region names for the vertices, one string per vertex. The region names are typically loaded from an annotation file with the read.fs.annot function.

agg_fun, function. An R function that aggregates data, typically max, mean, min or something similar. Note: this is NOT a string, put the function name without quotes. Defaults to base::mean.

requested_label_names, string vector. The label (or region) names that you want to occur in the output. If not specified, all region names which occur in the data are used. If given, and one of the requested names does NOT occur in the data, it will occur in the output with aggregation value NaN. If given, and one of the names from the data does NOT occur in the requested list, it will NOT occur in the output. So if you specify this, the output dataframe will contain a row for a region if and only if it is in the requested list.

Value

dataframe with aggregated values for all regions, with 2 columns and n rows, where n is the number of effective regions. The columns are: "region": string, contains the region name. "aggregated": numeric, contains the result of applying `agg_fun` to the morphometry data in that region.

See Also

Other aggregation functions: `group.agg.atlas.native()`, `group.agg.atlas.standard()`, `group.morph.agg.standard()`

Other atlas functions: `get.atlas.region.names()`, `group.agg.atlas.native()`, `group.agg.atlas.standard()`, `group.annot()`, `group.label.from.annot()`, `label.from.annotdata()`, `label.to.annot()`, `regions.to.ignore()`, `spread.values.over.annot()`, `spread.values.over.hemi()`, `spread.values.over.subject()`, `subject.annot()`, `subject.label.from.annot()`, `subject.lobes()`

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
morph_data = subject.morph.native(subjects_dir, 'subject1', 'thickness', 'lh');
annot = subject.annot(subjects_dir, 'subject1', 'lh', 'aparc');
agg = subject.atlas.agg(morph_data, annot$label_names);
```

subject.filepath.morph.native

Construct filepath of native space morphometry data file.

Description

Construct filepath of native space morphometry data file.

Usage

```
subject.filepath.morph.native(
  subjects_dir,
  subject_id,
  measure,
  hemi,
  format = "curv",
  warn_if_nonexistent = FALSE,
  error_if_nonexistent = FALSE
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	string. The subject identifier
measure	string. Name of the vertex-wise measure of morphometry data file. E.g., "area" or "thickness". Used to construct the name of the morphometry file to be loaded.
hemi	string, one of 'lh' or 'rh'. The hemisphere name.
format,	string. One of 'mgh', 'mgz', 'curv'. Defaults to 'curv'.
warn_if_nonexistent	logical. Whether to print a warning if the file does not exist or cannot be accessed. Defaults to FALSE.
error_if_nonexistent	logical. Whether to raise an error if the file does not exist or cannot be accessed. Defaults to FALSE.

Value

string, the file path.

subject.filepath.morph.standard

Construct filepath of standard space morphometry data file.

Description

Construct filepath of standard space morphometry data file.

Usage

```
subject.filepath.morph.standard(
  subjects_dir,
  subject_id,
  measure,
  hemi,
  fwhm = "10",
  template_subject = "fsaverage",
  format = "auto",
  warn_if_nonexistent = FALSE,
  error_if_nonexistent = FALSE
)
```


Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	string. The subject identifier
measure	string. Name of the vertex-wise measure of morphometry data file. E.g., "area" or "thickness". Used to construct the name of the morphometry file to be loaded.
hemi	string, one of 'lh' or 'rh'. The hemisphere name.
fwhm	string. Smoothing as string, e.g. '10' or '25'. Defaults to '10'.
template_subject	string. Template subject name, defaults to 'fsaverage'.
format	string. One of 'mgh', 'mgz', 'curv'. Defaults to 'mgh'.
warn_if_nonexistent	logical. Whether to print a warning if the file does not exist or cannot be accessed. Defaults to FALSE.
error_if_nonexistent	logical. Whether to raise an error if the file does not exist or cannot be accessed. Defaults to FALSE.

Value

string, the file path.

subject.label	<i>Retrieve label data for a single subject.</i>
---------------	--

Description

Load a label (like 'label/lh.cortex.label') for a subject from disk. Uses knowledge about the FreeSurfer directory structure to load the correct file.

Usage

```
subject.label(
  subjects_dir,
  subject_id,
  label,
  hemi,
  return_one_based_indices = TRUE
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	string. The subject identifier
label	string. Name of the label file, without the hemi part. You can include the '.label' suffix. E.g., 'cortex.label' for '?h.cortex.label'. You can also pass just the label (e.g., 'cortex'): if the string does not end with the suffix '.label', that suffix gets added automatically.
hemi	string, one of 'lh', 'rh', or 'both'. The hemisphere name. Used to construct the names of the label data files to be loaded. For 'both', see the information on the return value.
return_one_based_indices	logical. Whether the indices should be 1-based. Indices are stored zero-based in the file, but R uses 1-based indices. Defaults to TRUE, which means that 1 will be added to all indices read from the file before returning them.

Value

integer vector with label data: the list of vertex indices in the label. See 'return_one_based_indices' for important information. If parameter 'hemi' is set to 'both', a named list with entries 'lh' and 'rh' is returned, and the values of are the respective labels.

See Also

Other label data functions: [group.label\(\)](#), [labeldata.from.mask\(\)](#), [mask.from.labeldata.for.hemi\(\)](#)

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
cortex_lh = subject.label(subjects_dir, "subject1", "cortex.label", "lh");
```

subject.label.from.annot

Extract a region from an atlas annotation as a label for a subject.

Description

The returned label can be used to mask morphometry data, e.g., to set the values of a certain region to NaN or to extract only values from a certain region.

Usage

```
subject.label.from.annot(
  subjects_dir,
  subject_id,
  hemi,
  atlas,
  region,
  return_one_based_indices = TRUE,
  invert = FALSE,
  error_on_invalid_region = TRUE
)
```

Arguments

`subjects_dir`, string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.

`subject_id`, string. The subject identifier.

`hemi`, string, one of 'lh' or 'rh'. The hemisphere name. Used to construct the names of the label data files to be loaded.

`atlas`, string. The atlas name. E.g., "aparc", "aparc.2009s", or "aparc.DKTatlas". Used to construct the name of the annotation file to be loaded.

`region`, string. A valid region name for the annotation, i.e., one of the regions of the atlas.

`return_one_based_indices`, logical. Whether the indices should be 1-based. Indices are stored zero-based in label files, but R uses 1-based indices. Defaults to TRUE.

`invert`, logical. If TRUE, return the indices of all vertices which are NOT part of the region. Defaults to FALSE.

`error_on_invalid_region`, logical. Whether to throw an error if the given region does not appear in the region list of the annotation. If set to FALSE, this will be ignored and an empty vertex list will be returned. Defaults to TRUE.

Value

integer vector with label data: the list of vertex indices in the label.

See Also

Other atlas functions: [get.atlas.region.names\(\)](#), [group.agg.atlas.native\(\)](#), [group.agg.atlas.standard\(\)](#), [group.annot\(\)](#), [group.label.from.annot\(\)](#), [label.from.annotdata\(\)](#), [label.to.annot\(\)](#), [regions.to.ignore\(\)](#), [spread.values.over.annot\(\)](#), [spread.values.over.hemi\(\)](#), [spread.values.over.subject](#), [subject.annot\(\)](#), [subject.atlas.agg\(\)](#), [subject.lobes\(\)](#)

subject.lobes	<i>Load labels representing brain lobes.</i>
---------------	--

Description

This gives you labels that represent brain lobes for a subject. The lobe definition is based on the Desikan-Killiany atlas (Desikan *et al.*, 2010) as suggested on the FreeSurfer website at <https://surfer.nmr.mgh.harvard.edu/fswiki/CorticalParcellation>.

Usage

```
subject.lobes(
  subjects_dir,
  subject_id,
  hemi = "both",
  include_cingulate = TRUE,
  as_annot = FALSE
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	string. The subject identifier
hemi	string, one of 'lh', 'rh', or 'both'. The hemisphere name. Used to construct the names of the surface file to be loaded. For 'both', see the information on the return value.
include_cingulate	logical, whether to include the vertices of the cingulate in the lobes
as_annot	return a hemilist of annotations instead of the return value described in the *value* section

Value

hemilist of integer vectors, the vectors represent vertex indices of the hemispheres, and each vertex is assigned one of the following values: '0'=no_lobe, '1'=frontal, '2'=parietal, '3'=temporal, '4'=occipital.

See Also

Other atlas functions: [get.atlas.region.names\(\)](#), [group.agg.atlas.native\(\)](#), [group.agg.atlas.standard\(\)](#), [group.annot\(\)](#), [group.label.from.annot\(\)](#), [label.from.annotdata\(\)](#), [label.to.annot\(\)](#), [regions.to.ignore\(\)](#), [spread.values.over.annot\(\)](#), [spread.values.over.hemi\(\)](#), [spread.values.over.subject\(\)](#), [subject.annot\(\)](#), [subject.atlas.agg\(\)](#), [subject.label.from.annot\(\)](#)

Other label functions: [apply.label.to.morphdata\(\)](#), [apply.labeldata.to.morphdata\(\)](#), [subject.mask\(\)](#), [vis.labeldata.on.subject\(\)](#), [vis.subject.label\(\)](#)

subject.mask *Compute a mask for a subject.*

Description

Compute a binary vertex mask for the surface vertices of a subject. By defaults, the medial wall is masked.

Usage

```
subject.mask(
    subjects_dir,
    subject_id,
    hemi = "both",
    from_label = "cortex",
    surf_num_verts = "white",
    invert_mask = TRUE
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	string. The subject identifier
hemi	string, one of 'lh', 'rh' or 'both'. The hemisphere name. Used to construct the names of the annotation and morphometry data files to be loaded.
from_label	string, the label file to use. Defaults to 'cortex', which will result in a mask of the medial wall versus cortex vertices.
surf_num_verts	string or integer. If an integer, interpreted as the number of vertices in the respective surface (lh or rh). If a character string, interpreted as a surface name, (e.g., 'white' or 'pial'), and the respective surface will be loaded to determine the number of vertices in it. If parameter 'hemi' is set to 'both' and you supply the vertex count as an integer, this can be a vector of length 2 if the surfaces have different vertex counts (the first entry for 'lh', the second for 'rh').
invert_mask	logical, whether to invert the mask. E.g., when the mask is loaded from the cortex labels, if this is set to FALSE, the cortex would be masked (set to 0 in the final mask). If you want everything but the cortex to be masked (set to 0), you should set this to 'TRUE'. Defaults to 'TRUE'.

Value

the mask, a logical vector with the length of the vertices in the surface. If parameter 'hemi' is set to 'both', a named list with entries 'lh' and 'rh' is returned, and the values of are the respective masks.

See Also

Other label functions: [apply.label.to.morphdata\(\)](#), [apply.labeldata.to.morphdata\(\)](#), [subject.lobes\(\)](#), [vis.labeldata.on.subject\(\)](#), [vis.subject.label\(\)](#)

Examples

```
# Generate a binary mask of the medial wall. Wall vertices will
# be set to 0, cortex vertices will be set to 1.
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
mask = subject.mask(subjects_dir, "subject1");
# Print some information on the mask:
#cat(sprintf("lh: %d verts, %d in cortex, %d medial wall.\n", length(mask$lh),
# sum(mask$lh), (length(mask$lh)- sum(mask$lh))))
# Output: lh: 149244 verts, 140891 in cortex, 8353 medial wall.
# Now visualize the mask to illustrate that it is correct:
vis.mask.on.subject(subjects_dir, "subject1", mask$lh, mask$rh);
```

subject.morph.native *Retrieve native space morphometry data for a single subject.*

Description

Load native space morphometry data (like 'surf/lh.area') for a subject from disk. Uses knowledge about the FreeSurfer directory structure to load the correct file.

Usage

```
subject.morph.native(
  subjects_dir,
  subject_id,
  measure,
  hemi,
  format = "curv",
  cortex_only = FALSE,
  split_by_hemi = FALSE
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	string. The subject identifier
measure	string. Name of the vertex-wise measure of morphometry data file. E.g., "area" or "thickness". Used to construct the name of the morphometry file to be loaded.

hemi	string, one of 'lh', 'rh' or 'both'. The hemisphere name. Used to construct the names of the annotation and morphometry data files to be loaded.
format	string. One of 'mgh', 'mgz', 'curv'. Defaults to 'curv'.
cortex_only	logical, whether to mask the medial wall, i.e., whether the morphometry data for all vertices which are <i>*not*</i> part of the cortex (as defined by the label file 'label/?h.cortex.label') should be replaced with NA values. In other words, setting this to TRUE will ignore the values of the medial wall between the two hemispheres. If set to true, the mentioned label file needs to exist for the subject. Defaults to FALSE.
split_by_hemi	logical, whether the returned data should be encapsulated in a named list, where the names are from 'lh' and 'rh', and the values are the respective data.

Value

vector with native space morph data, as returned by `read.fs.morph`.

See Also

Other morphometry data functions: `apply.label.to.morphdata()`, `apply.labeldata.to.morphdata()`, `group.morph.native()`, `group.morph.standard()`, `subject.morph.standard()`

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");

# Load the full data:
thickness_lh = subject.morph.native(subjects_dir, "subject1", "thickness", "lh");
mean(thickness_lh); # prints 2.437466

# Load the data again, but this time exclude the medial wall:
thickness_lh_cortex = subject.morph.native(subjects_dir, "subject1", "thickness",
  "lh", cortex_only=TRUE);
mean(thickness_lh_cortex, na.rm=TRUE); # prints 2.544132
vis.data.on.subject(subjects_dir, "subject1", thickness_lh_cortex, NULL);
```

subject.morph.standard

Retrieve standard space morphometry data for a single subject.

Description

Load standard space morphometry data (like 'surf/lh.area.fwhm10.fsaverage.mgh') for a subject from disk. Uses knowledge about the FreeSurfer directory structure to load the correct file.

Usage

```
subject.morph.standard(
  subjects_dir,
  subject_id,
  measure,
  hemi,
  fwhm = "10",
  template_subject = "fsaverage",
  format = "mgh",
  cortex_only = FALSE,
  split_by_hemi = FALSE
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	string. The subject identifier
measure	string. Name of the vertex-wise measure of morphometry data file. E.g., "area" or "thickness". Used to construct the name of the morphometry file to be loaded.
hemi	string, one of 'lh', 'rh', or 'both'. The hemisphere name. Used to construct the names of the annotation and morphometry data files to be loaded.
fwhm	string. Smoothing as string, e.g. '10' or '25'.
template_subject	string. Template subject name, defaults to 'fsaverage'.
format	string. One of 'mgh', 'mgz', 'curv'. Defaults to 'mgh'.
cortex_only	logical, whether to mask the medial wall, i.e., whether the morphometry data for all vertices which are <i>*not*</i> part of the cortex (as defined by the label file 'label/?h.cortex.label') should be replaced with NA values. In other words, setting this to TRUE will ignore the values of the medial wall between the two hemispheres. If set to true, the mentioned label file needs to exist for the template subject. Defaults to FALSE.
split_by_hemi	logical, whether the returned data should be encapsulated in a named list, where the names are from 'lh' and 'rh', and the values are the respective data.

Value

vector with standard space morph data

See Also

Other morphometry data functions: [apply.label.to.morphdata\(\)](#), [apply.labeldata.to.morphdata\(\)](#), [group.morph.native\(\)](#), [group.morph.standard\(\)](#), [subject.morph.native\(\)](#)

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
thickness_lh = subject.morph.standard(subjects_dir, "subject1", "thickness", "lh", fwhm='10');
```

subject.num.verts	<i>Get subjects vertex count.</i>
-------------------	-----------------------------------

Description

Determine vertex counts for the brain meshes of a subject.

Usage

```
subject.num.verts(
  subjects_dir,
  subject_id,
  surface = "white",
  hemi = "both",
  do_sum = FALSE
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	string. The subject identifier
surface	string. The surface name. E.g., "white", or "pial". Used to construct the name of the surface file to be loaded.
hemi	string, one of 'lh', 'rh', or 'both'. The hemisphere name. Used to construct the names of the surface file to be loaded. For 'both', see the information on the return value.
do_sum	logical, whether to return the sum of the vertex counts for lh and rh. Ignored unless 'hemi' is 'both'. If set, a single scalar will be returned.

Value

integer of hemilist of integers, the vertex count. If hemi is 'both' and 'do_sum' is 'FALSE', a hemilist of integers is returned. Otherwise, a single integer.

subject.surface *Load a surface for a subject.*

Description

Load a brain surface mesh for a subject.

Usage

```
subject.surface(
    subjects_dir,
    subject_id,
    surface,
    hemi,
    force_hemilist = FALSE
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	string. The subject identifier
surface	string. The surface name. E.g., "white", or "pial". Used to construct the name of the surface file to be loaded.
hemi	string, one of 'lh', 'rh', or 'both'. The hemisphere name. Used to construct the names of the surface file to be loaded. For 'both', see the information on the return value.
force_hemilist	logical, whether to return a hemilist even if the 'hemi' parameter is not set to 'both'

Value

the 'fs.surface' instance, as returned by [read.fs.surface](#). If parameter 'hemi' is set to 'both', a named list with entries 'lh' and 'rh' is returned, and the values of are the respective surfaces. The mesh data structure used in 'fs.surface' is a *face index set*.

See Also

Other surface mesh functions: [face.edges\(\)](#), [label.border\(\)](#), [mesh.vertex.included.faces\(\)](#), [mesh.vertex.neighbors\(\)](#), [vis.path.along.verts\(\)](#)

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
lh_white = subject.surface(subjects_dir, "subject1", "white", "lh");
```

subject.volume	<i>Read a brain volume.</i>
----------------	-----------------------------

Description

Load a brain volume (like 'mri/brain.mgz') for a subject from disk. Uses knowledge about the FreeSurfer directory structure to load the correct file.

Usage

```
subject.volume(
  subjects_dir,
  subject_id,
  volume,
  format = "auto",
  drop_empty_dims = TRUE,
  with_header = FALSE,
  mri_subdir = NULL
)
```

Arguments

subjects_dir	character string, the FreeSurfer 'SUBJECTS_DIR', i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	character string, the subject identifier.
volume	character string, name of the volume file without file extension. Examples: 'brain' or 'aseg'.
format	string. One of 'mgh', 'mgz', 'AUTO'. If left at the default value 'AUTO', the function will look for files with extensions 'mgh' and 'mgz' (in that order) and use the first one that exists.
drop_empty_dims	logical, whether to drop empty dimensions of the returned data. Passed to read.fs.mgh .
with_header	logical. Whether to return the header as well. If TRUE, return a named list with entries "data" and "header". The latter is another named list which contains the header data. These header entries exist: "dtype": int, one of: 0=MRI_UCHAR; 1=MRI_INT; 3=MRI_FLOAT; 4=MRI_SHORT. "voldim": integer vector. The volume (=data) dimensions. E.g., c(256, 256, 256, 1). These header entries may exist: "vox2ras_matrix" (exists if "ras_good_flag" is 1), "mr_params" (exists if "has_mr_params" is 1). Passed to read.fs.mgh .
mri_subdir	character string or NULL, the subdir to use within the 'mri' directory. Defaults to 'NULL', which means to read directly from the 'mri' dir. You could use this to read volumes from the 'mri/orig/' directory by setting it to 'orig'.

Value

numerical array, the voxel data. If 'with_header', the full volume datastructure (see above).

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
brain = subject.volume(subjects_dir, 'subject1', 'brain', with_header = TRUE);
# Use the vox2ras matrix from the header to compute RAS coordinates at CRS voxel (0, 0, 0):
brain$header$vox2ras_matrix %*% c(0,0,0,1);
```

vdata.split.by.hemi *Split morph data vector at hemisphere boundary.*

Description

Given a single vector of per-vertex data for a mesh, split it at the hemi boundary. This is achieved by loading the respective surface and checking the number of vertices for the 2 hemispheres.

Usage

```
vdata.split.by.hemi(
  subjects_dir,
  subject_id,
  vdata,
  surface = "white",
  expand = TRUE
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	string. The subject identifier
vdata	numerical vector of data for both hemispheres, one value per vertex
surface	the surface to load to determine the vertex counts
expand	logical, whether to allow input of length 1, and expand (repeat) it to the length of the hemispheres.

Value

a hemilist, each entry contains the data part of the respective hemi.

Note

Instead of calling this function to split the data, you could use the 'split_by_hemi' parameter of [subject.morph.native](#).

vis.color.on.subject *Visualize pre-defined vertex colors on a subject.*

Description

Visualize pre-defined vertex colors on a subject.

Usage

```
vis.color.on.subject(
  subjects_dir,
  vis_subject_id,
  color_lh = NULL,
  color_rh = NULL,
  surface = "white",
  views = c("t4"),
  rgloptions = rglo(),
  rglactions = list(),
  color_both = NULL
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
vis_subject_id	string. The subject identifier from which to obtain the surface for data visualization. Example: 'fsaverage'.
color_lh	vector of colors to visualize on the left hemisphere surface. Length must match number of vertices in hemi surface, or be a single color.
color_rh	vector of colors to visualize on the right hemisphere surface. Length must match number of vertices in hemi surface, or be a single color.
surface	string. The display surface. E.g., "white", "pial", or "inflated". Defaults to "white".
views	list of strings. Valid entries include: 'si': single interactive view. 't4': tiled view showing the brain from 4 angles. 't9': tiled view showing the brain from 9 angles.
rgloptions	option list passed to par3d . Example: rgloptions = list("windowRect"=c(50,50,1000,1000)).
rglactions	named list. A list in which the names are from a set of pre-defined actions. The values can be used to specify parameters for the action. The following example clips outliers in the data before plotting and writes a screenshot in PNG format: rglactions = list("snapshot_png"=~"/fsbrain.png", "clip_data"=c(0.05,0.95)). See rglactions .

color_both vector of colors to visualize on the left and right hemispheres. Alternative to 'color_lh' and 'color_rh'. Length must match sum of vertices in both hemis.

Value

list of coloredmeshes. The coloredmeshes used for the visualization.

See Also

Other visualization functions: [vis.data.on.fsaverage\(\)](#), [vis.data.on.subject\(\)](#), [vis.labeldata.on.subject\(\)](#), [vis.mask.on.subject\(\)](#), [vis.region.values.on.subject\(\)](#), [vis.subject.annot\(\)](#), [vis.subject.label\(\)](#), [vis.subject.morph.native\(\)](#), [vis.subject.morph.standard\(\)](#), [vis.symmetric.data.on.subject\(\)](#), [vislayout.from.coloredmeshes\(\)](#)

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
color_lh = '#ff0000';
num_verts_subject1_rh = 153333;
color_rh = rep('#333333', num_verts_subject1_rh);
color_rh[1:30000] = '#00ff00';
color_rh[30001:60000] = '#ff0000';
color_rh[60001:90000] = '#0000ff';
color_rh[90001:120000] = '#ffff00';
color_rh[120001:150000] = '#00ffff';
vis.color.on.subject(subjects_dir, 'subject1', color_lh, color_rh);
```

vis.coloredmeshes *Visualize a list of colored meshes in a single scene.*

Description

Visualize a list of colored meshes in a single scene.

Usage

```
vis.coloredmeshes(
  coloredmeshes,
  background = "white",
  skip_all_na = TRUE,
  style = "default",
  rgloptions = rglo(),
  rglactions = list(),
  draw_colorbar = FALSE
)
```

Arguments

coloredmeshes	list of coloredmesh. A coloredmesh is a named list as returned by the coloredmesh.from.* functions. It has the entries 'mesh' of type tmesh3d, a 'col', which is a color specification for such a mesh.
background	string, background color passed to rgl::bg3d()
skip_all_na	logical, whether to skip (i.e., not render) meshes in the list that have the property 'render' set to FALSE. Defaults to TRUE. Practically, this means that a hemisphere for which the data was not given is not rendered, instead of being rendered in a single color.
style,	a named list of style parameters or a string specifying an available style by name (e.g., 'shiny'). Defaults to 'default', the default style.
rgloptions	option list passed to par3d. Example: rgloptions = list("windowRect"=c(50,50,1000,1000));
rglactions	named list. A list in which the names are from a set of pre-defined actions. Defaults to the empty list.
draw_colorbar	logical. Whether to draw a colorbar. WARNING: Will only show up if there is enough space in the plot area and does not resize properly. Defaults to FALSE. See coloredmesh.plot.colorbar.separate for an alternative.

Value

the list of visualized coloredmeshes

Note

To change or adapt the colorbar, you should use the makecmap_options parameter when constructing them in a vis function. See the example.

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
cm = vis.subject.morph.native(subjects_dir, 'subject1', 'thickness',
  makecmap_options=list('n'=100));
# You could mess with the meshes here.
vis.coloredmeshes(cm);
```

vis.coloredmeshes.rotating

Visualize a list of colored meshes in a single scene and rotate them, movie-style.

Description

Visualize a list of colored meshes in a single scene and rotate them, movie-style.

Usage

```
vis.coloredmeshes.rotating(
    coloredmeshes,
    background = "white",
    skip_all_na = TRUE,
    style = "default",
    x = 0,
    y = 0,
    z = 1,
    rpm = 6,
    duration = 10,
    rgloptions = rglo(),
    rglactions = list()
)
```

Arguments

coloredmeshes	list of coloredmesh. A coloredmesh is a named list as returned by the coloredmesh.from.* functions. It has the entries 'mesh' of type tmesh3d, a 'col', which is a color specification for such a mesh.
background	string, background color passed to rgl::bg3d()
skip_all_na	logical, whether to skip (i.e., not render) meshes in the list that have the property 'render' set to FALSE. Defaults to TRUE. Practically, this means that a hemisphere for which the data was not given is not rendered, instead of being rendered in a single color.
style	a named list of style parameters or a string specifying an available style by name (e.g., 'shiny'). Defaults to 'default', the default style.
x	rotation x axis value, passed to spin3d. Defaults to 0.
y	rotation y axis value, passed to spin3d. Defaults to 1.
z	rotation z axis value, passed to spin3d. Defaults to 0.
rpm	rotation rpm value, passed to spin3d. Defaults to 15.
duration	rotation duration value, passed to spin3d. Defaults to 20.
rgloptions	option list passed to par3d. Example: rgloptions = list("windowRect"=c(50,50,1000,1000));
rglactions	named list. A list in which the names are from a set of pre-defined actions. Defaults to the empty list.

Value

the list of visualized coloredmeshes

vis.colortable.legend *Create a separate legend plot for a colortable or an annotation.*

Description

This plots a legend for a colortable or an atlas (annotation), showing the region names and their assigned colors. This function creates a new plot.

Usage

```
vis.colortable.legend(colortable, ncols = 1L, plot_struct_index = TRUE)
```

Arguments

colortable dataframe, a colortable as returned by [read.fs.colortable](#) or the inner 'colortable_df' returned by [subject.annot](#). One can also pass an annotation (*fs.annot* instance).

ncols positive integer, the number of columns to use when plotting

plot_struct_index logical, whether to plot the region index from tge 'struct_index' field. If there is no such field, this is silently ignored.

Note

This function plots one or more legends (see [legend](#)). You may have to adapt the device size before calling this function if you inted to plot a large colortable.

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
annot = subject.annot(subjects_dir, 'subject1', 'lh', 'aparc');
vis.colortable.legend(annot$colortable_df, ncols=3);
```

vis.data.on.fsaverage *Visualize arbitrary data on the fsaverage template subject, if available.*

Description

Creates a surface mesh, applies a colormap transform the morphometry data values into colors, and renders the resulting colored mesh in an interactive window. If hemi is 'both', the data is rendered for the whole brain. This function tries to automatically retrieve the subjects dir of the fsaverage template subject by checking the environment variables SUBJECTS_DIR and FREESURFER_HOME for the subject. The subject is required for its surfaces, which are not shipped with this package for licensing reasons.

Usage

```
vis.data.on.fsaverage(
    subjects_dir = NULL,
    vis_subject_id = "fsaverage",
    morph_data_lh = NULL,
    morph_data_rh = NULL,
    surface = "white",
    colormap = NULL,
    views = c("t4"),
    rgloptions = rglo(),
    rglactions = list(),
    draw_colorbar = FALSE,
    makecmap_options = mkco.seq(),
    bg = NULL,
    morph_data_both = NULL
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
vis_subject_id	string. The subject identifier from which to obtain the surface for data visualization. Defaults to 'fsaverage'.
morph_data_lh	numeric vector or character string or NULL, the data to visualize on the left hemisphere surface. If a string, it is treated as a filename and data is loaded from it first. When it is a numerical vector, this is assumed to be the data already. The data must have the same length as the surface of the vis_subject_id has vertices. If NULL, this surface will not be rendered. Only one of morph_data_lh or morph_data_rh is allowed to be NULL.
morph_data_rh	numeric vector or character string or NULL, the data to visualize on the right hemisphere surface. If a string, it is treated as a filename and data is loaded from it first. When it is a numerical vector, this is assumed to be the data already. The data must have the same length as the surface of the vis_subject_id has vertices. If NULL, this surface will not be rendered. Only one of morph_data_lh or morph_data_rh is allowed to be NULL.
surface	string. The display surface. E.g., "white", "pial", or "inflated". Defaults to "white".
colormap	a colormap function. **DEPRECATED** : use parameter 'makecmap_options' instead, like this: makecmap_options=list('colFn'=terrain.colors).
views	list of strings. Valid entries include: 'si': single interactive view. 't4': tiled view showing the brain from 4 angles. 't9': tiled view showing the brain from 9 angles.
rgloptions	option list passed to par3d . Example: rgloptions = list("windowRect"=c(50,50,1000,1000)).
rglactions	named list. A list in which the names are from a set of pre-defined actions. The values can be used to specify parameters for the action. The following example clips outliers in the data before plotting and writes a screenshot in PNG format:

- `rglactions = list("snapshot_png"=~"/fsbrain.png", "clip_data"=c(0.05,0.95)).`
See [rglactions](#).
- `draw_colorbar` logical or one of the character strings 'vertical' or 'horizontal', whether to draw a colorbar. Notice: the colorbar is drawn to a separate subplot, and this only works if there is enough space for it, i.e., the plot resolution must be high enough. You may have to increase the plot size for the colorbar to show up, see the vignette for instructions. Defaults to 'FALSE'. See [coloredmesh.plot.colorbar.separate](#) for an alternative.
- `makecmap_options` named list of parameters to pass to [makecmap](#). Must not include the unnamed first parameter, which is derived from 'measure'. Should include at least a colormap function as name 'colFn'.
- `bg` a background definition. Can be a surface color layer or a character string like 'curv_light' to select a pre-defined layer, see [collayer.bg](#) for valid strings.
- `morph_data_both` numeric vector or NULL, the data to visualize on both hemispheres. This must be a single vector with length equal to the sum of the vertex counts of the left and the right hemisphere. The data for the left hemisphere must come first. If this is given, 'morph_data_lh' and 'morph_data_rh' must be NULL.

Value

list of coloredmeshes. The coloredmeshes used for the visualization.

See Also

Other visualization functions: [vis.color.on.subject\(\)](#), [vis.data.on.subject\(\)](#), [vis.labeldata.on.subject\(\)](#), [vis.mask.on.subject\(\)](#), [vis.region.values.on.subject\(\)](#), [vis.subject.annot\(\)](#), [vis.subject.label\(\)](#), [vis.subject.morph.native\(\)](#), [vis.subject.morph.standard\(\)](#), [vis.symmetric.data.on.subject\(\)](#), [vislayout.from.coloredmeshes\(\)](#)

Other morphometry visualization functions: [vis.data.on.subject\(\)](#), [vis.subject.morph.native\(\)](#), [vis.subject.morph.standard\(\)](#), [vis.symmetric.data.on.subject\(\)](#)

vis.data.on.group.native

Visualize native space data on a group of subjects.

Description

Plot surface data on the native space surfaces of a group of subjects and combine the tiles into a single large image.

Usage

```
vis.data.on.group.native(
    subjects_dir,
    subject_id,
    morph_data_both,
    view_angles = "sd_dorsal",
    output_img = "fsbrain_group_morph.png",
    num_per_row = 5L,
    captions = subject_id,
    rglactions = list(no_vis = TRUE),
    ...
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	vector of character strings, the subject identifiers
morph_data_both	named list of numerical vectors, the morph data for both hemispheres of all subjects. Can be loaded with group.morph.native .
view_angles	see get.view.angle.names .
output_img	character string, the file path for the output image. Should end with '.png'.
num_per_row	positive integer, the number of tiles per row.
captions	optional vector of character strings, the short text annotations for the individual tiles. Typically used to plot the subject identifier.
rglactions	named list. A list in which the names are from a set of pre-defined actions. The values can be used to specify parameters for the action. The following example clips outliers in the data before plotting and writes a screenshot in PNG format: <code>rglactions = list("snapshot_png"="~/fsbrain.png", "clip_data"=c(0.05,0.95))</code> . See rglactions .
...	extra parameters passed to the subject level visualization function. Not all may make sense in this context. Example: <code>surface='pial'</code> .

Value

named list, see the return value of [arrange.brainview.images.grid](#) for details.

Note

The subjects are plotted row-wise, in the order in which they appear in the 'morph_data_both' parameter. The surfaces are loaded in the order of the 'subject_id' parameter, so the order in both must match.

You can force an identical plot range for all subjects, so that one color represents identical values across subjects, via 'makecmap_options'. E.g., for the ... parameter, pass `makecmap_options=list('colFn'=viridis::vi`

See Also

Other group visualization functions: [vis.data.on.group.standard\(\)](#), [vis.group.annot\(\)](#), [vis.group.coloredmeshes](#), [vis.group.morph.native\(\)](#), [vis.group.morph.standard\(\)](#)

vis.data.on.group.standard

Visualize standard space data for a group on template.

Description

Plot standard space data for a group of subjects onto a template brain and combine the tiles into a single large image.

Usage

```
vis.data.on.group.standard(
    subjects_dir,
    vis_subject_id,
    morph_data_both,
    captions = NULL,
    view_angles = "sd_dorsal",
    output_img = "fsbrain_group_morph.png",
    num_per_row = 5L,
    rglactions = list(no_vis = TRUE),
    ...
)
```

Arguments

subjects_dir	character string, the path to the SUBJECTS_DIR containing the template subject
vis_subject_id	character string, the template subject name. A typical choice is 'fsaverage'.
morph_data_both	named list of numerical vectors, 4D array or dataframe, the morph data for both hemispheres of all subjects. Can be loaded with group.morph.standard or group.morph.standard.sf .
captions	optional vector of character strings, the short text annotations for the individual tiles. Typically used to plot the subject identifier.
view_angles	see get.view.angle.names .
output_img	character string, the file path for the output image. Should end with '.png'.
num_per_row	positive integer, the number of tiles per row.
rglactions	named list. A list in which the names are from a set of pre-defined actions. The values can be used to specify parameters for the action. The following example clips outliers in the data before plotting and writes a screenshot in PNG format: <code>rglactions = list("snapshot_png"=~"/fsbrain.png", "clip_data"=c(0.05,0.95))</code> . See rglactions .
...	extra parameters passed to the subject level visualization function. Not all may make sense in this context. Example: <code>surface='pial'</code> .

Value

named list, see the return value of [arrange.brainview.images.grid](#) for details.

Note

The subject data are plotted row-wise, in the order in which they appear in the 'morph_data_both' parameter.

You can force an identical plot range for all subjects, so that one color represents identical values across subjects, via 'makecmap_options'. E.g., for the ... parameter, pass `makecmap_options=list('colFn'=viridis::vi`

See Also

Other group visualization functions: [vis.data.on.group.native\(\)](#), [vis.group.annot\(\)](#), [vis.group.coloredmeshes\(\)](#), [vis.group.morph.native\(\)](#), [vis.group.morph.standard\(\)](#)

`vis.data.on.subject` *Visualize arbitrary data on the surface of any subject.*

Description

Creates a surface mesh, applies a colormap transform the morphometry data values into colors, and renders the resulting colored mesh in an interactive window. If hemi is 'both', the data is rendered for the whole brain.

Usage

```
vis.data.on.subject(
  subjects_dir,
  vis_subject_id,
  morph_data_lh = NULL,
  morph_data_rh = NULL,
  surface = "white",
  colormap = NULL,
  views = c("t4"),
  rgloptions = rglo(),
  rglactions = list(),
  draw_colorbar = FALSE,
  makecmap_options = mkco.seq(),
  bg = NULL,
  morph_data_both = NULL
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
vis_subject_id	string. The subject identifier from which to obtain the surface for data visualization. Example: 'fsaverage'.
morph_data_lh	numeric vector or character string or NULL, the data to visualize on the left hemisphere surface. If a string, it is treated as a filename and data is loaded from it first. When it is a numerical vector, this is assumed to be the data already. The data must have the same length as the surface of the vis_subject_id has vertices. If NULL, this surface will not be rendered. Only one of morph_data_lh or morph_data_rh is allowed to be NULL.
morph_data_rh	numeric vector or character string or NULL, the data to visualize on the right hemisphere surface. If a string, it is treated as a filename and data is loaded from it first. When it is a numerical vector, this is assumed to be the data already. The data must have the same length as the surface of the vis_subject_id has vertices. If NULL, this surface will not be rendered. Only one of morph_data_lh or morph_data_rh is allowed to be NULL.
surface	string. The display surface. E.g., "white", "pial", or "inflated". Defaults to "white".
colormap	a colormap function. **DEPRECATED** : use parameter 'makecmap_options' instead, like this: makecmap_options=list('colFn'=terrain.colors).
views	list of strings. Valid entries include: 'si': single interactive view. 't4': tiled view showing the brain from 4 angles. 't9': tiled view showing the brain from 9 angles.
rgloptions	option list passed to par3d . Example: rgloptions = list("windowRect"=c(50,50,1000,1000)).
rglactions	named list. A list in which the names are from a set of pre-defined actions. The values can be used to specify parameters for the action. The following example clips outliers in the data before plotting and writes a screenshot in PNG format: rglactions = list("snapshot_png"=~"/fsbrain.png", "clip_data"=c(0.05,0.95)). See rglactions .
draw_colorbar	logical or one of the character strings 'vertical' or 'horizontal', whether to draw a colorbar. Notice: the colorbar is drawn to a separate subplot, and this only works if there is enough space for it, i.e., the plot resolution must be high enough. You may have to increase the plot size for the colorbar to show up, see the vignette for instructions. Defaults to 'FALSE'. See coloredmesh.plot.colorbar.separate for an alternative.
makecmap_options	named list of parameters to pass to makecmap . Must not include the unnamed first parameter, which is derived from 'measure'. Should include at least a colormap function as name 'colFn'.
bg	a background definition. Can be a surface color layer or a character string like 'curv_light' to select a pre-defined layer, see collayer.bg for valid strings.
morph_data_both	numeric vector or NULL, the data to visualize on both hemispheres. This must be a single vector with length equal to the sum of the vertex counts of the left

and the right hemisphere. The data for the left hemisphere must come first. If this is given, 'morph_data_lh' and 'morph_data_rh' must be NULL.

Value

list of coloredmeshes. The coloredmeshes used for the visualization.

See Also

Other visualization functions: `vis.color.on.subject()`, `vis.data.on.fsaverage()`, `vis.labeldata.on.subject()`, `vis.mask.on.subject()`, `vis.region.values.on.subject()`, `vis.subject.annot()`, `vis.subject.label()`, `vis.subject.morph.native()`, `vis.subject.morph.standard()`, `vis.symmetric.data.on.subject()`, `vislayout.from.coloredmeshes()`

Other morphometry visualization functions: `vis.data.on.fsaverage()`, `vis.subject.morph.native()`, `vis.subject.morph.standard()`, `vis.symmetric.data.on.subject()`

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
morph_data_lh = subject.morph.native(subjects_dir, 'subject1', 'thickness', 'lh');
morph_data_rh = NULL;
vis.data.on.subject(subjects_dir, 'subject1', morph_data_lh, morph_data_rh);
```

vis.export.from.coloredmeshes

Export high-quality brainview image with horizontal colorbar.

Description

This function serves as an easy (but slightly inflexible) way to export a high-quality, tight-layout, horizontal colorbar figure to disk.

Usage

```
vis.export.from.coloredmeshes(
  coloredmeshes,
  colorbar_legend = NULL,
  img_only = TRUE,
  horizontal = TRUE,
  silent = TRUE,
  quality = 1L,
  output_img = "fsbrain_arranged.png",
  image.plot_extra_options = NULL,
  large_legend = TRUE,
```



```

    view_angles = get.view.angle.names(angle_set = "t4"),
    style = "default"
)

```

Arguments

coloredmeshes	list of coloredmesh. A coloredmesh is a named list as returned by the ‘coloredmesh.from*’ functions (like coloredmesh.from.morph.native). It has the entries ‘mesh’ of type tmesh3d, a ‘col’, which is a color specification for such a mesh. The ‘vis*’ functions (like vis.subject.morph.native) all return a list of coloredmeshes.
colorbar_legend	character string or NULL, the title for the colorbar.
img_only	logical, whether to return only the resulting image
horizontal	logical, whether to plot the colorbar horizontally
silent	logical, whether to suppress messages
quality	integer, an arbitrary quality. This is the resolution per tile before trimming, divided by 1000, in pixels. Example: 1L means 1000x1000 pixels per tile before trimming. Currently supported values: 1L . . 2L. Note that the resolution you can get is also limited by your screen resolution.
output_img	string, path to the output file. Defaults to "fsbrain_arranged.png"
image.plot_extra_options	named list, custom options for fields::image.plot. Overwrites those derived from the quality setting. If in doubt, leave this alone.
large_legend	logical, whether to plot extra large legend text, affects the font size of the colorbar_legend and the tick labels.
view_angles	list of strings. See get.view.angle.names for all valid strings.
style	the rendering style, see material3d or use a predefined style like ‘default’ or ‘shiny’.

Value

magick image instance or named list, depending on the value of ‘img_only’. If the latter, the list contains the fields ‘rev_vl’, ‘rev_cb’, and ‘rev_ex’, which are the return values of the functions [vislayout.from.coloredmeshes](#), [coloredmesh.plot.colorbar.separate](#), and [combine.colorbar.with.brainview.image](#), respectively.

Note

This function also exports the resulting image to disk in PNG format, in the current working directory, named ‘fsbrain_merged.png’. Note that your screen resolution has to be high enough to generate the final image.

Examples

```
## Not run:
  rand_data = rnorm(327684, 5, 1.5);
  cm = vis.data.on.fsaverage(morph_data_both=rand_data, rglactions=list('no_vis'=T));
  vis.export.from.coloredmeshes(cm, colorbar_legend='Random data');

## End(Not run)
```

vis.fs.surface	<i>Visualize fs.surface mesh</i>
----------------	----------------------------------

Description

Render a mesh. All mesh formats supported by the **freesurferformats** package are supported, including OFF, PLY, OBJ, STL, and many more.

Usage

```
vis.fs.surface(
  fs_surface,
  col = "white",
  per_vertex_data = NULL,
  hemi = "lh",
  makecmap_options = mkco.seq(),
  ...
)
```

Arguments

fs_surface	an fs.surface instance, as returned by function like subject.surface or read.fs.surface . If a character string, it is assumed to be the full path of a surface file, and the respective file is loaded with read.fs.surface . If parameter 'hemi' is 'both', this must be a hemilist.
col	vector of colors, the per-vertex-colors. Defaults to white. Must be a single color or one color per vertex. If parameter 'hemi' is 'both', this must be a hemilist.
per_vertex_data	numerical vector, per-vertex data. If given, takes precedence over 'col'. Used to color the mesh using the colormap options in parameter 'makecmap_options'. If a character string, it is assumed to be the full path of a morphometry data file, and the respective file is loaded with read.fs.morph . If parameter 'hemi' is 'both', this must be a hemilist.
hemi	character string, one of 'lh' or 'rh'. This may be used by visualization functions to decide whether or not to show this mesh in a certain view.

makecmap_options named list of parameters to pass to [makecmap](#). Must not include the unnamed first parameter, which is derived from 'measure'. Should include at least a colormap function as name 'colFn'.

... extra parameters to pass to [vis.coloredmeshes](#).

Value

see [vis.coloredmeshes](#)

Note

This function can be used to visualize arbitrary triangular meshes in R. Despite its name, it is not limited to brain surface meshes.

vis.group.annot *Plot atlas annotations for a group of subjects.*

Description

Plot atlas annotations for a group of subjects and combine them into a single large image.

Usage

```
vis.group.annot(
  subjects_dir,
  subject_id,
  atlas,
  view_angles = "sd_dorsal",
  output_img = "fsbrain_group_annot.png",
  num_per_row = 5L,
  captions = subject_id,
  rglactions = list(no_vis = TRUE),
  ...
)
```

Arguments

subjects_dir string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.

subject_id vector of character strings, the subject identifiers

atlas vector of character strings, the atlas names. Example: c('aparc', 'aparc.a2009s')

view_angles see [get.view.angle.names](#).

output_img character string, the file path for the output image. Should end with '.png'.

num_per_row positive integer, the number of tiles per row.

captions	optional vector of character strings, the short text annotations for the individual tiles. Typically used to plot the subject identifier.
rglactions	named list. A list in which the names are from a set of pre-defined actions. The values can be used to specify parameters for the action. The following example clips outliers in the data before plotting and writes a screenshot in PNG format: <code>rglactions = list("snapshot_png"=~"/fsbrain.png", "clip_data"=c(0.05,0.95))</code> . See rglactions .
...	extra parameters passed to the subject level visualization function. Not all may make sense in this context. Example: <code>surface='pial'</code> .

Value

named list, see the return value of [arrange.brainview.images.grid](#) for details.

Note

The subjects are plotted row-wise, in the order in which they appear in the 'subject_id' parameter. This function is vectorized over 'subject_id' and 'atlas'.

See Also

Other group visualization functions: [vis.data.on.group.native\(\)](#), [vis.data.on.group.standard\(\)](#), [vis.group.coloredmeshes\(\)](#), [vis.group.morph.native\(\)](#), [vis.group.morph.standard\(\)](#)

vis.group.coloredmeshes

Plot coloredmeshes for a group of subjects.

Description

Plot coloredmeshes for a group of subjects into a single image.

Usage

```
vis.group.coloredmeshes(
  coloredmeshes,
  view_angles = "sd_dorsal",
  output_img = "fsbrain_group_annot.png",
  num_per_row = 5L,
  captions = NULL
)
```

Arguments

coloredmeshes	a list of coloredmeshes lists, each entry in the outer list contains the hemilist of coloredmeshes (left and right hemisphere mesh) for one subject.
view_angles	see get.view.angle.names .
output_img	character string, the file path for the output image. Should end with '.png'.
num_per_row	positive integer, the number of tiles per row.
captions	optional vector of character strings, the short text annotations for the individual tiles. Typically used to plot the subject identifier.

Value

named list, see the return value of [arrange.brainview.images.grid](#) for details.

Note

This is a mid-level function, end users may want to call high-level functions like [vis.group.annot](#) instead.

See Also

Other group visualization functions: [vis.data.on.group.native\(\)](#), [vis.data.on.group.standard\(\)](#), [vis.group.annot\(\)](#), [vis.group.morph.native\(\)](#), [vis.group.morph.standard\(\)](#)

vis.group.morph.native

Plot native space morphometry data for a group of subjects.

Description

Plot native space morphometry data for a group of subjects and combine them into a single large image.

Usage

```
vis.group.morph.native(
  subjects_dir,
  subject_id,
  measure,
  view_angles = "sd_dorsal",
  output_img = "fsbrain_group_morph.png",
  num_per_row = 5L,
  captions = subject_id,
  rglactions = list(no_vis = TRUE),
  ...
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	vector of character strings, the subject identifiers
measure	vector of character strings, the morphometry measures, e.g., c('thickness', 'area')
view_angles	see get.view.angle.names .
output_img	character string, the file path for the output image. Should end with '.png'.
num_per_row	positive integer, the number of tiles per row.
captions	optional vector of character strings, the short text annotations for the individual tiles. Typically used to plot the subject identifier.
rglactions	named list. A list in which the names are from a set of pre-defined actions. The values can be used to specify parameters for the action. The following example clips outliers in the data before plotting and writes a screenshot in PNG format: <code>rglactions = list("snapshot_png"="~/fsbrain.png", "clip_data"=c(0.05,0.95))</code> . See rglactions .
...	extra parameters passed to the subject level visualization function. Not all may make sense in this context. Example: <code>surface='pial'</code> .

Value

named list, see the return value of [arrange.brainview.images.grid](#) for details.

Note

The subjects are plotted row-wise, in the order in which they appear in the 'subject_id' parameter. This function is vectorized over 'subject_id' and 'measure'.

You can force an identical plot range for all subjects, so that one color represents identical values across subjects, via 'makecmap_options'. E.g., for the ... parameter, pass `makecmap_options=list('colFn'=viridis::vi`

See Also

Other group visualization functions: [vis.data.on.group.native\(\)](#), [vis.data.on.group.standard\(\)](#), [vis.group.annot\(\)](#), [vis.group.coloredmeshes\(\)](#), [vis.group.morph.standard\(\)](#)

vis.group.morph.standard

Plot standard space morphometry data for a group of subjects.

Description

Plot standard space morphometry data for a group of subjects and combine them into a single large image.

Usage

```
vis.group.morph.standard(
    subjects_dir,
    subject_id,
    measure,
    fwhm = "10",
    view_angles = "sd_dorsal",
    output_img = "fsbrain_group_morph.png",
    num_per_row = 5L,
    captions = subject_id,
    rglactions = list(no_vis = TRUE),
    ...
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	vector of character strings, the subject identifiers
measure	vector of character strings, the morphometry measures, e.g., c('thickness', 'area')
fwhm	vector of character strings, the smoothing kernel FWHM strings, e.g., c('0', '10', '15')
view_angles	see get.view.angle.names .
output_img	character string, the file path for the output image. Should end with '.png'.
num_per_row	positive integer, the number of tiles per row.
captions	optional vector of character strings, the short text annotations for the individual tiles. Typically used to plot the subject identifier.
rglactions	named list. A list in which the names are from a set of pre-defined actions. The values can be used to specify parameters for the action. The following example clips outliers in the data before plotting and writes a screenshot in PNG format: <code>rglactions = list("snapshot_png"="~/fsbrain.png", "clip_data"=c(0.05, 0.95))</code> . See rglactions .
...	extra parameters passed to the subject level visualization function. Not all may make sense in this context. Example: <code>surface='pial'</code> .

Value

named list, see the return value of [arrange.brainview.images.grid](#) for details.

Note

The subjects are plotted row-wise, in the order in which they appear in the 'subject_id' parameter. This function is vectorized over 'subject_id', 'measure' and 'fwhm'.

You can force an identical plot range for all subjects, so that one color represents identical values across subjects, via 'makecmap_options'. E.g., for the ... parameter, pass `makecmap_options=list('colFn'=viridis::vi`

See Also

Other group visualization functions: [vis.data.on.group.native\(\)](#), [vis.data.on.group.standard\(\)](#), [vis.group.annot\(\)](#), [vis.group.coloredmeshes\(\)](#), [vis.group.morph.native\(\)](#)

```
vis.labeldata.on.subject
```

Visualize a label on the surface of a subject.

Description

Visualizes a label. Note that a label is just a set of vertices, and that you can use this function to visualize sets of vertices, e.g., to see where on the mesh a certain vertex lies. It may be helpful the visualize the vertex with its neighbors, because otherwise it may be too small to spot. Use the function `[fsbrain::mesh.vertex.neighbors]` to get them. It is advisable to set the view to the interactive 'si' mode and use the 'inflated' surface to identify single vertices.

Usage

```
vis.labeldata.on.subject(  
  subjects_dir,  
  vis_subject_id,  
  lh_labeldata,  
  rh_labeldata,  
  surface = "white",  
  colormap = NULL,  
  views = c("t4"),  
  rgloptions = rglo(),  
  rglactions = list(),  
  draw_colorbar = FALSE,  
  makecmap_options = list(colFn = label.colFn.inv)  
)
```

Arguments

<code>subjects_dir</code>	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
<code>vis_subject_id</code>	string. The subject identifier from which to obtain the surface for data visualization. Example: 'fsaverage'.
<code>lh_labeldata</code>	integer vector of vertex indices for the left hemisphere
<code>rh_labeldata</code>	integer vector of vertex indices for the right hemisphere
<code>surface</code>	string. The display surface. E.g., "white", "pial", or "inflated". Defaults to "white".
<code>colormap</code>	a colormap function. **DEPRECATED** : use parameter 'makecmap_options' instead, like this: <code>makecmap_options=list('colFn'=terrain.colors)</code> .

views	list of strings. Valid entries include: 'si': single interactive view. 't4': tiled view showing the brain from 4 angles. 't9': tiled view showing the brain from 9 angles.
rgloptions	option list passed to par3d . Example: <code>rgloptions = list("windowRect"=c(50,50,1000,1000))</code> .
rglactions	named list. A list in which the names are from a set of pre-defined actions. The values can be used to specify parameters for the action. The following example clips outliers in the data before plotting and writes a screenshot in PNG format: <code>rglactions = list("snapshot_png"=~"/fsbrain.png", "clip_data"=c(0.05,0.95))</code> . See rglactions .
draw_colorbar	logical or one of the character strings 'vertical' or 'horizontal', whether to draw a colorbar. Notice: the colorbar is drawn to a separate subplot, and this only works if there is enough space for it, i.e., the plot resolution must be high enough. You may have to increase the plot size for the colorbar to show up, see the vignette for instructions. Defaults to 'FALSE'. See coloredmesh.plot.colorbar.separate for an alternative.
makecmap_options	named list of parameters to pass to makecmap . Must not include the unnamed first parameter, which is derived from 'measure'. Should include at least a colormap function as name 'colFn'.

Value

list of coloredmeshes. The coloredmeshes used for the visualization.

Note

Drawing a colorbar for label data makes limited sense, use a legend instead. The colorbar can give a rough overview of the relative number of label and non-label vertices though, so it is possible to request one.

See Also

Other label functions: [apply.label.to.morphdata\(\)](#), [apply.labeldata.to.morphdata\(\)](#), [subject.lobes\(\)](#), [subject.mask\(\)](#), [vis.subject.label\(\)](#)

Other visualization functions: [vis.color.on.subject\(\)](#), [vis.data.on.fsaverage\(\)](#), [vis.data.on.subject\(\)](#), [vis.mask.on.subject\(\)](#), [vis.region.values.on.subject\(\)](#), [vis.subject.annot\(\)](#), [vis.subject.label\(\)](#), [vis.subject.morph.native\(\)](#), [vis.subject.morph.standard\(\)](#), [vis.symmetric.data.on.subject\(\)](#), [vislayout.from.coloredmeshes\(\)](#)

Examples

```
fsbrain::download_optional_data();

# Define the data to use:
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
lh_labeldata = c(1000, 1001, 1002); # only the vertices, will be tiny.
subject_id = 'subject1';
surface = 'white'; # Should use 'inflated', but we do not currently
```

```

# ship it for the example subject to reduce download size.

# For the right hemi, extend them to neighborhood for better visibility:
rh_labeldata = c(500, 5000);
rh_surface = subject.surface(subjects_dir, subject_id, surface, 'rh');
rh_labeldata_neighborhood = mesh.vertex.neighbors(rh_surface, rh_labeldata);
vis.labeldata.on.subject(subjects_dir, subject_id, lh_labeldata,
  rh_labeldata_neighborhood$vertices, surface=surface, views=c('si'));

```

vis.mask.on.subject *Visualize a vertex mask on the surface of a subject.*

Description

A mask is a logical vector that contains one value per vertex. You can create it manually, or use functions like [mask.from.labeldata.for.hemi](#) to create and modify it. Check the example for this function.

Usage

```

vis.mask.on.subject(
  subjects_dir,
  vis_subject_id,
  mask_lh,
  mask_rh,
  surface = "white",
  colormap = NULL,
  views = c("t4"),
  rgloptions = rglo(),
  rglactions = list(),
  draw_colorbar = FALSE,
  makecmap_options = list(colFn = label.colFn.inv)
)

```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
vis_subject_id	string. The subject identifier from which to obtain the surface for data visualization. Example: 'fsaverage'.
mask_lh	logical vector or NULL, the mask to visualize on the left hemisphere surface. Must have the same length as the lh surface of the vis_subject_id has vertices. If NULL, this surface will not be rendered. Only one of mask_lh or mask_rh is allowed to be NULL.

mask_rh	logical vector or NULL, the mask to visualize on the right hemisphere surface. Must have the same length as the rh surface of the vis_subject_id has vertices. If NULL, this surface will not be rendered. Only one of mask_lh or mask_rh is allowed to be NULL.
surface	string. The display surface. E.g., "white", "pial", or "inflated". Defaults to "white".
colormap	a colormap function. **DEPRECATED** : use parameter 'makecmap_options' instead, like this: makecmap_options=list('colFn'=terrain.colors).
views	list of strings. Valid entries include: 'si': single interactive view. 't4': tiled view showing the brain from 4 angles. 't9': tiled view showing the brain from 9 angles.
rgloptions	option list passed to par3d . Example: rgloptions = list("windowRect"=c(50,50,1000,1000)).
rglactions	named list. A list in which the names are from a set of pre-defined actions. The values can be used to specify parameters for the action. The following example clips outliers in the data before plotting and writes a screenshot in PNG format: rglactions = list("snapshot_png"=~"/fsbrain.png", "clip_data"=c(0.05,0.95)). See rglactions .
draw_colorbar	logical or one of the character strings 'vertical' or 'horizontal', whether to draw a colorbar. Notice: the colorbar is drawn to a separate subplot, and this only works if there is enough space for it, i.e., the plot resolution must be high enough. You may have to increase the plot size for the colorbar to show up, see the vignette for instructions. Defaults to 'FALSE'. See coloredmesh.plot.colorbar.separate for an alternative.
makecmap_options	named list of parameters to pass to makecmap . Must not include the unnamed first parameter, which is derived from 'measure'. Should include at least a colormap function as name 'colFn'.

Value

list of coloredmeshes. The coloredmeshes used for the visualization.

Note

Drawing a colorbar for label data makes limited sense, use a legend instead. The colorbar can give a rough overview of the relative number of label and non-label vertices though, so it is possible to request one.

See Also

Other mask functions: [coloredmesh.from.mask\(\)](#), [mask.from.labeldata.for.hemi\(\)](#)

Other visualization functions: [vis.color.on.subject\(\)](#), [vis.data.on.fsaverage\(\)](#), [vis.data.on.subject\(\)](#), [vis.labeldata.on.subject\(\)](#), [vis.region.values.on.subject\(\)](#), [vis.subject.annot\(\)](#), [vis.subject.label\(\)](#), [vis.subject.morph.native\(\)](#), [vis.subject.morph.standard\(\)](#), [vis.symmetric.data.on.subject\(\)](#), [vislayout.from.coloredmeshes\(\)](#)

Examples

```

fsbrain::download_optional_data();

# Define the data to use:
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
subject_id = 'subject1';
surface = 'white';
hemi = 'both';
atlas = 'aparc';
region = 'bankssts';

# Create a mask from a region of an annotation:
lh_annot = subject.annot(subjects_dir, subject_id, 'lh', atlas);
rh_annot = subject.annot(subjects_dir, subject_id, 'rh', atlas);
lh_label = label.from.annotdata(lh_annot, region);
rh_label = label.from.annotdata(rh_annot, region);
lh_mask = mask.from.labeldata.for.hemi(lh_label, length(lh_annot$vertices));
rh_mask = mask.from.labeldata.for.hemi(rh_label, length(rh_annot$vertices));

# Edit the mask: add the vertices from another region to it:
region2 = 'medialorbitofrontal';
lh_label2 = label.from.annotdata(lh_annot, region2);
rh_label2 = label.from.annotdata(rh_annot, region2);
lh_mask2 = mask.from.labeldata.for.hemi(lh_label2, length(lh_annot$vertices),
existing_mask = lh_mask);
rh_mask2 = mask.from.labeldata.for.hemi(rh_label2, length(rh_annot$vertices),
existing_mask = rh_mask);
# Visualize the mask:
vis.mask.on.subject(subjects_dir, subject_id, lh_mask2, rh_mask2);

```

vis.path.along.verts *Draw a 3D line from vertex to vertex*

Description

To get a nice path along the surface, pass the vertex indices along a geodesic path. Note: You can first open an interactive brain view ('view='si') with a vis* function like [vis.subject.morph.native](#), then run this function to draw into the active plot.

Usage

```
vis.path.along.verts(surface_vertices, path_vertex_indices)
```

Arguments

- surface_vertices
float matrix of size (n, 3), the surface vertex coordinates, as returned as part of `subject.surface` or `read.fs.surface`, in the member "vertices".
- path_vertex_indices
vector of vertex indices, the path. You will need to have it computed already. (This function does **not** compute geodesic paths. You can use it to visualize such a path though.)

See Also

Other surface mesh functions: `face.edges()`, `label.border()`, `mesh.vertex.included.faces()`, `mesh.vertex.neighbors()`, `subject.surface()`

vis.region.values.on.subject

Visualize arbitrary data, one value per atlas region, on the surface of any subject (including template subjects).

Description

This function can be used for rendering a single value (color) for all vertices of an atlas region. The typical usecase is the visualization of results of atlas-based analyses, e.g., p-value, means or other aggregated values over all vertices of a region.

Usage

```
vis.region.values.on.subject(
  subjects_dir,
  subject_id,
  atlas,
  lh_region_value_list,
  rh_region_value_list,
  surface = "white",
  colormap = NULL,
  views = c("t4"),
  rgloptions = rglo(),
  rglactions = list(),
  value_for_unlisted_regions = NA,
  draw_colorbar = FALSE,
  makecmap_options = mkco.heat(),
  bg = NULL,
  silent = FALSE
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	string. The subject identifier.
atlas	string. The brain atlas to use. E.g., 'aparc' or 'aparc.a2009s'.
lh_region_value_list	named list. A list for the left hemisphere in which the names are atlas regions, and the values are the value to write to all vertices of that region. You can pass an unnamed list, but then the its length must exactly match the number of atlas regions. The order of values must also match the order of regions in the annotation, of course. The resulting mapping will be printed so you can check it (unless 'silent' is set).
rh_region_value_list	named list. A list for the right hemisphere in which the names are atlas regions, and the values are the value to write to all vertices of that region.
surface	string. The display surface. E.g., "white", "pial", or "inflated". Defaults to "white".
colormap	a colormap function. **DEPRECATED** : use parameter 'makecmap_options' instead, like this: makecmap_options=list('colFn'=terrain.colors).
views	list of strings. Valid entries include: 'si': single interactive view. 't4': tiled view showing the brain from 4 angles. 't9': tiled view showing the brain from 9 angles.
rgloptions	option list passed to par3d . Example: rgloptions = list("windowRect"=c(50,50,1000,1000)).
rglactions	named list. A list in which the names are from a set of pre-defined actions. The values can be used to specify parameters for the action. The following example clips outliers in the data before plotting and writes a screenshot in PNG format: rglactions = list("snapshot_png"="~/fsbrain.png", "clip_data"=c(0.05,0.95)). See rglactions .
value_for_unlisted_regions	numerical scalar or 'NaN', the value to assign to regions which do not occur in the region_value_lists. Defaults to 'NaN'.
draw_colorbar	logical or one of the character strings 'vertical' or 'horizontal', whether to draw a colorbar. Notice: the colorbar is drawn to a separate subplot, and this only works if there is enough space for it, i.e., the plot resolution must be high enough. You may have to increase the plot size for the colorbar to show up, see the vignette for instructions. Defaults to 'FALSE'. See coloredmesh.plot.colorbar.separate for an alternative.
makecmap_options	named list of parameters to pass to makecmap . Must not include the unnamed first parameter, which is derived from 'measure'. Should include at least a colormap function as name 'colFn'.
bg	a background definition. Can be a surface color layer or a character string like 'curv_light' to select a pre-defined layer, see collayer.bg for valid strings.
silent	logical, whether to suppress mapping info in case of unnamed region value lists (see 'lh_region_value_list' description).

Value

list of coloredmeshes. The coloredmeshes used for the visualization.

See Also

Other visualization functions: [vis.color.on.subject\(\)](#), [vis.data.on.fsaverage\(\)](#), [vis.data.on.subject\(\)](#), [vis.labeldata.on.subject\(\)](#), [vis.mask.on.subject\(\)](#), [vis.subject.annot\(\)](#), [vis.subject.label\(\)](#), [vis.subject.morph.native\(\)](#), [vis.subject.morph.standard\(\)](#), [vis.symmetric.data.on.subject\(\)](#), [vislayout.from.coloredmeshes\(\)](#)

Other region-based visualization functions: [vis.subject.annot\(\)](#)

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
atlas = 'aparc'; # Desikan atlas
# For the left hemisphere, we just assign a subset of the
# atlas regions. The others will get the default value.
lh_region_value_list = list("bankssts"=0.9, "precuneus"=0.7, "postcentral"=0.8, "lingual"=0.6);
# For the right hemisphere, we retrieve the full list of regions for
# the atlas, and assign random values to all of them.
atlas_region_names = get.atlas.region.names(atlas, template_subjects_dir = subjects_dir,
  template_subject='subject1');
rh_region_value_list = rnorm(length(atlas_region_names), 3.0, 1.0);
names(rh_region_value_list) = atlas_region_names;
vis.region.values.on.subject(subjects_dir, 'subject1', atlas,
  lh_region_value_list, rh_region_value_list);
```

vis.subject.annot

Visualize an annotation for a subject.

Description

Creates a surface mesh, loads the colors from the annotation, and renders the resulting colored mesh in an interactive window. If hemi is 'both', the data is rendered for the whole brain.

Usage

```
vis.subject.annot(
  subjects_dir,
  subject_id,
  atlas,
  hemi = "both",
  surface = "white",
  views = c("t4"),
```

```

    rgloptions = rglo(),
    rglactions = list(),
    outline = FALSE
)

```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	string. The subject identifier.
atlas	string. The atlas name. E.g., "aparc", "aparc.2009s", or "aparc.DKTatlas". Used to construct the name of the annotation file to be loaded. Can also be a hemilist of already loaded annotations.
hemi	string, one of 'lh', 'rh', or 'both'. The hemisphere name. Used to construct the names of the label data files to be loaded.
surface	string. The display surface. E.g., "white", "pial", or "inflated". Defaults to "white".
views	list of strings. Valid entries include: 'si': single interactive view. 't4': tiled view showing the brain from 4 angles. 't9': tiled view showing the brain from 9 angles.
rgloptions	option list passed to par3d . Example: <code>rgloptions = list("windowRect"=c(50,50,1000,1000))</code> .
rglactions	named list. A list in which the names are from a set of pre-defined actions. The values can be used to specify parameters for the action. The following example clips outliers in the data before plotting and writes a screenshot in PNG format: <code>rglactions = list("snapshot_png"=~"/fsbrain.png", "clip_data"=c(0.05,0.95))</code> . See rglactions .
outline	logical, whether to draw an outline only instead of filling the regions. Defaults to 'FALSE'. Instead of passing 'TRUE', one can also pass a list of extra parameters to pass to annot.outline , e.g., <code>outline=list('outline_color'='#000000')</code> . Using this increases computation time dramatically, sorry for the performance.

Value

list of coloredmeshes. The coloredmeshes used for the visualization.

See Also

Other visualization functions: [vis.color.on.subject\(\)](#), [vis.data.on.fsaverage\(\)](#), [vis.data.on.subject\(\)](#), [vis.labeldata.on.subject\(\)](#), [vis.mask.on.subject\(\)](#), [vis.region.values.on.subject\(\)](#), [vis.subject.label\(\)](#), [vis.subject.morph.native\(\)](#), [vis.subject.morph.standard\(\)](#), [vis.symmetric.data.on.subject\(\)](#), [vislayout.from.coloredmeshes\(\)](#)

Other region-based visualization functions: [vis.region.values.on.subject\(\)](#)

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
vis.subject.annot(subjects_dir, 'subject1', 'aparc', 'both');
```

vis.subject.label *Visualize a binary label for a subject.*

Description

Visualize a label for a subject. A label is just a logical vector with one entry for each vertex in the mesh. Each vertex may additionally be associated with a scalar value, but this function ignored that.

Usage

```
vis.subject.label(
  subjects_dir,
  subject_id,
  label,
  hemi,
  surface = "white",
  colormap = NULL,
  views = c("t4"),
  rgloptions = rglo(),
  rglactions = list(),
  draw_colorbar = FALSE,
  makecmap_options = list(colFn = label.colFn.inv, col.na = "#FFFFFF00"),
  map_to_NA = 0L,
  bg = NULL
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	string. The subject identifier.
label	string. Name of the label file, without the hemi part (if any), but including the '.label' suffix. E.g., 'cortex.label' for '?h.cortex.label'.
hemi	string, one of 'lh', 'rh', or 'both'. The hemisphere name. Used to construct the names of the label data files to be loaded.
surface	string. The display surface. E.g., "white", "pial", or "inflated". Defaults to "white".

colormap	a colormap function. **DEPRECATED** : use parameter 'makecmap_options' instead, like this: <code>makecmap_options=list('colFn'=terrain.colors)</code> .
views	list of strings. Valid entries include: 'si': single interactive view. 't4': tiled view showing the brain from 4 angles. 't9': tiled view showing the brain from 9 angles.
rgloptions	option list passed to par3d . Example: <code>rgloptions = list("windowRect"=c(50,50,1000,1000))</code> .
rglactions	named list. A list in which the names are from a set of pre-defined actions. The values can be used to specify parameters for the action. The following example clips outliers in the data before plotting and writes a screenshot in PNG format: <code>rglactions = list("snapshot_png"=~"/fsbrain.png", "clip_data"=c(0.05,0.95))</code> . See rglactions .
draw_colorbar	logical or one of the character strings 'vertical' or 'horizontal', whether to draw a colorbar. Notice: the colorbar is drawn to a separate subplot, and this only works if there is enough space for it, i.e., the plot resolution must be high enough. You may have to increase the plot size for the colorbar to show up, see the vignette for instructions. Defaults to 'FALSE'. See coloredmesh.plot.colorbar.separate for an alternative.
makecmap_options	named list of parameters to pass to makecmap . Must not include the unnamed first parameter, which is derived from 'measure'. Should include at least a colormap function as name 'colFn'.
map_to_NA	the value or value range that should **not** be considered part of the label, and should thus be plotted as background color. Only used if 'bg' is not 'NULL'. If a single value, only exactly this value is used (typically 0). If two values, they are interpreted as a range, and a values between them are mapped to NA. If you prefer to map the data to NA yourself before using this function, pass 'NULL'.
bg	a background definition. Can be a surface color layer or a character string like 'curv_light' to select a pre-defined layer, see collayer.bg for valid strings.

Value

list of coloredmeshes. The coloredmeshes used for the visualization.

Note

Drawing a colorbar for label data makes limited sense, use a legend instead. The colorbar can give a rough overview of the relative number of label and non-label vertices though, so it is possible to request one.

See Also

Other visualization functions: [vis.color.on.subject\(\)](#), [vis.data.on.fsaverage\(\)](#), [vis.data.on.subject\(\)](#), [vis.labeldata.on.subject\(\)](#), [vis.mask.on.subject\(\)](#), [vis.region.values.on.subject\(\)](#), [vis.subject.annot\(\)](#), [vis.subject.morph.native\(\)](#), [vis.subject.morph.standard\(\)](#), [vis.symmetric.data.on.subject\(\)](#), [vislayout.from.coloredmeshes\(\)](#)

Other label functions: [apply.label.to.morphdata\(\)](#), [apply.labeldata.to.morphdata\(\)](#), [subject.lobes\(\)](#), [subject.mask\(\)](#), [vis.labeldata.on.subject\(\)](#)

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
subject_id = 'subject1';
surface = 'white';
hemi = 'both';
label = 'cortex.label';
vis.subject.label(subjects_dir, subject_id, label, hemi, views="si");
```

vis.subject.morph.native

Visualize native space morphometry data for a subject.

Description

Creates a surface mesh, applies a colormap transform the morphometry data values into colors, and renders the resulting colored mesh in an interactive window. If hemi is 'both', the data is rendered for the whole brain.

Usage

```
vis.subject.morph.native(  
  subjects_dir,  
  subject_id,  
  measure,  
  hemi = "both",  
  surface = "white",  
  colormap = NULL,  
  views = c("t4"),  
  rgloptions = rglo(),  
  rglactions = list(),  
  draw_colorbar = FALSE,  
  cortex_only = FALSE,  
  style = "default",  
  makecmap_options = mkco.seq(),  
  bg = NULL  
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	string. The subject identifier.

measure	string. The morphometry data to use. E.g., 'area' or 'thickness'. Pass NULL to render just the surface in white, without any data.
hemi	string, one of 'lh', 'rh', or 'both'. The hemisphere name. Used to construct the names of the label data files to be loaded.
surface	string. The display surface. E.g., "white", "pial", or "inflated". Defaults to "white".
colormap	a colormap function. **DEPRECATED** : use parameter 'makecmap_options' instead, like this: <code>makecmap_options=list('colFn'=terrain.colors)</code> .
views	list of strings. Valid entries include: 'si': single interactive view. 't4': tiled view showing the brain from 4 angles. 't9': tiled view showing the brain from 9 angles.
rgloptions	option list passed to <code>par3d</code> . Example: <code>rgloptions = list("windowRect"=c(50,50,1000,1000))</code> .
rglactions	named list. A list in which the names are from a set of pre-defined actions. The values can be used to specify parameters for the action. The following example clips outliers in the data before plotting and writes a screenshot in PNG format: <code>rglactions = list("snapshot_png"=~"/fsbrain.png", "clip_data"=c(0.05,0.95))</code> . See rglactions .
draw_colorbar	logical or one of the character strings 'vertical' or 'horizontal', whether to draw a colorbar. Notice: the colorbar is drawn to a separate subplot, and this only works if there is enough space for it, i.e., the plot resolution must be high enough. You may have to increase the plot size for the colorbar to show up, see the vignette for instructions. Defaults to 'FALSE'. See coloredmesh.plot.colorbar.separate for an alternative.
cortex_only	logical, whether to mask the medial wall, i.e., whether the morphometry data for all vertices which are <i>not</i> part of the cortex (as defined by the label file 'label/?h.cortex.label') should be replaced with NA values. In other words, setting this to TRUE will ignore the values of the medial wall between the two hemispheres. If set to true, the mentioned label file needs to exist for the subject. Defaults to FALSE.
style	character string, a rendering style, e.g., 'default', 'shiny' or 'semitransparent'.
makecmap_options	named list of parameters to pass to <code>makecmap</code> . Must not include the unnamed first parameter, which is derived from 'measure'. Should include at least a colormap function as name 'colFn'.
bg	a background definition. Can be a surface color layer or a character string like 'curv_light' to select a pre-defined layer, see collayer.bg for valid strings.

Value

list of coloredmeshes. The coloredmeshes used for the visualization.

See Also

Other visualization functions: [vis.color.on.subject\(\)](#), [vis.data.on.fsaverage\(\)](#), [vis.data.on.subject\(\)](#), [vis.labeldata.on.subject\(\)](#), [vis.mask.on.subject\(\)](#), [vis.region.values.on.subject\(\)](#),

```
vis.subject.annot(), vis.subject.label(), vis.subject.morph.standard(), vis.symmetric.data.on.subject(),
vislayout.from.coloredmeshes()
```

Other morphometry visualization functions: `vis.data.on.fsaverage()`, `vis.data.on.subject()`,
`vis.subject.morph.standard()`, `vis.symmetric.data.on.subject()`

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
vis.subject.morph.native(subjects_dir, 'subject1', 'thickness', 'lh', views=c("t9"));
```

```
vis.subject.morph.standard
```

Visualize native space morphometry data for a subject or a group.

Description

Renders standard space morphometry data for a single subject, or the group mean for a group of subjects. The default template subject is fsaverage.

Usage

```
vis.subject.morph.standard(
  subjects_dir,
  subject_id,
  measure,
  hemi = "both",
  fwhm = "10",
  surface = "white",
  template_subject = "fsaverage",
  template_subjects_dir = NULL,
  colormap = NULL,
  views = c("t4"),
  rgloptions = rglo(),
  rglactions = list(),
  draw_colorbar = FALSE,
  cortex_only = FALSE,
  makecmap_options = mkco.seq(),
  bg = NULL,
  style = "default"
)
```

Arguments

subjects_dir	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id	character string or vector of character strings, the subject or subjects. For a single subjects, its data will be plotted. If a group of subjects is given instead, at each vertex the mean value over all the subjects will be plotted.
measure	string. The morphometry data to use. E.g., 'area' or 'thickness'. Pass NULL to render just the surface in white, without any data.
hemi	string, one of 'lh', 'rh', or 'both'. The hemisphere name. Used to construct the names of the label data files to be loaded.
fwhm	string, smoothing setting (full width at half maximum of the kernel). The smoothing part of the filename, typically something like '0', '5', '10', ..., or '25'.
surface	string. The display surface. E.g., "white", "pial", or "inflated". Defaults to "white".
template_subject	The template subject used. This will be used as part of the filename, and its surfaces are loaded for data visualization. Defaults to 'fsaverage'.
template_subjects_dir	The template subjects dir. If NULL, the value of the parameter 'subjects_dir' is used. If you have FreeSurfer installed and configured, and are using the standard fsaverage subject, try passing the result of calling 'file.path(Sys.getenv('FREESURFER_HOME'), 'subjects')'.
colormap	a colormap function. **DEPRECATED** : use parameter 'makecmap_options' instead, like this: makecmap_options=list('colFn'=terrain.colors).
views	list of strings. Valid entries include: 'si': single interactive view. 't4': tiled view showing the brain from 4 angles. 't9': tiled view showing the brain from 9 angles.
rgloptions	option list passed to par3d . Example: rgloptions = list("windowRect"=c(50,50,1000,1000)).
rglactions	named list. A list in which the names are from a set of pre-defined actions. The values can be used to specify parameters for the action. The following example clips outliers in the data before plotting and writes a screenshot in PNG format: rglactions = list("snapshot_png"=~"/fsbrain.png", "clip_data"=c(0.05,0.95)). See rglactions .
draw_colorbar	logical or one of the character strings 'vertical' or 'horizontal', whether to draw a colorbar. Notice: the colorbar is drawn to a separate subplot, and this only works if there is enough space for it, i.e., the plot resolution must be high enough. You may have to increase the plot size for the colorbar to show up, see the vignette for instructions. Defaults to 'FALSE'. See coloredmesh.plot.colorbar.separate for an alternative.
cortex_only	logical, whether to mask the medial wall, i.e., whether the morphometry data for all vertices which are <i>not</i> part of the cortex (as defined by the label file 'label/?h.cortex.label') should be replaced with NA values. In other words, setting this to TRUE will ignore the values of the medial wall between the two hemispheres. If set to true, the mentioned label file needs to exist for the subject. Defaults to FALSE.

makecmap_options	named list of parameters to pass to makecmap . Must not include the unnamed first parameter, which is derived from 'measure'. Should include at least a colormap function as name 'colFn'.
bg	a background definition. Can be a surface color layer or a character string like 'curv_light' to select a pre-defined layer, see collayer.bg for valid strings.
style	character string, a rendering style, e.g., 'default', 'shiny' or 'semitransparent'.

Value

list of coloredmeshes. The coloredmeshes used for the visualization.

See Also

Other visualization functions: [vis.color.on.subject\(\)](#), [vis.data.on.fsaverage\(\)](#), [vis.data.on.subject\(\)](#), [vis.labeldata.on.subject\(\)](#), [vis.mask.on.subject\(\)](#), [vis.region.values.on.subject\(\)](#), [vis.subject.annot\(\)](#), [vis.subject.label\(\)](#), [vis.subject.morph.native\(\)](#), [vis.symmetric.data.on.subject\(\)](#), [vislayout.from.coloredmeshes\(\)](#)

Other morphometry visualization functions: [vis.data.on.fsaverage\(\)](#), [vis.data.on.subject\(\)](#), [vis.subject.morph.native\(\)](#), [vis.symmetric.data.on.subject\(\)](#)

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
fsaverage_dir = file.path(Sys.getenv('FREESURFER_HOME'), 'subjects');
if(dir.exists(fsaverage_dir)) {
  vis.subject.morph.standard(subjects_dir, 'subject1', 'thickness', 'lh',
    '10', template_subjects_dir=fsaverage_dir);
}
# The last command will load the file
# *<subjects_dir>/subject1/surf/lh.thickness.fwhm10.fsaverage.mgh* and
# visualize the data on *$FREESURFER_HOME/subjects/fsaverage/surf/lh.white*.
```

vis.symmetric.data.on.subject

Visualize clusters or activation data on the surface of any subject.

Description

This function is intended to plot symmetric data around zero (like positive and negative activation data, signed p-values, etc.) on a subject's surface. It is a thin wrapper around [vis.data.on.subject](#).

Usage

```
vis.symmetric.data.on.subject(
  subjects_dir,
  vis_subject_id,
  morph_data_lh = NULL,
  morph_data_rh = NULL,
  surface = "white",
  colormap = NULL,
  views = c("t4"),
  rgloptions = rglo(),
  rglactions = list(),
  draw_colorbar = TRUE,
  makecmap_options = list(colFn = cm.cbry(), symm = TRUE, col.na = "#FFFFFF00", n =
    200),
  map_to_NA = c(0),
  bg = NULL,
  morph_data_both = NULL
)
```

Arguments

- | | |
|----------------|--|
| subjects_dir | string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier. |
| vis_subject_id | string. The subject identifier from which to obtain the surface for data visualization. Example: 'fsaverage'. |
| morph_data_lh | numeric vector or character string or NULL, the data to visualize on the left hemisphere surface. If a string, it is treated as a filename and data is loaded from it first. When it is a numerical vector, this is assumed to be the data already. The data must have the same length as the surface of the vis_subject_id has vertices. If NULL, this surface will not be rendered. Only one of morph_data_lh or morph_data_rh is allowed to be NULL. |
| morph_data_rh | numeric vector or character string or NULL, the data to visualize on the right hemisphere surface. If a string, it is treated as a filename and data is loaded from it first. When it is a numerical vector, this is assumed to be the data already. The data must have the same length as the surface of the vis_subject_id has vertices. If NULL, this surface will not be rendered. Only one of morph_data_lh or morph_data_rh is allowed to be NULL. |
| surface | string. The display surface. E.g., "white", "pial", or "inflated". Defaults to "white". |
| colormap | a colormap function. **DEPRECATED** : use parameter 'makecmap_options' instead, like this: makecmap_options=list('colFn'=terrain.colors). |
| views | list of strings. Valid entries include: 'si': single interactive view. 't4': tiled view showing the brain from 4 angles. 't9': tiled view showing the brain from 9 angles. |
| rgloptions | option list passed to par3d . Example: rgloptions = list("windowRect"=c(50, 50, 1000, 1000)). |

<code>rglactions</code>	named list. A list in which the names are from a set of pre-defined actions. The values can be used to specify parameters for the action. The following example clips outliers in the data before plotting and writes a screenshot in PNG format: <code>rglactions = list("snapshot_png"=~"/fsbrain.png", "clip_data"=c(0.05,0.95)).</code> See rglactions .
<code>draw_colorbar</code>	logical or one of the character strings 'vertical' or 'horizontal', whether to draw a colorbar. Notice: the colorbar is drawn to a separate subplot, and this only works if there is enough space for it, i.e., the plot resolution must be high enough. You may have to increase the plot size for the colorbar to show up, see the vignette for instructions. Defaults to 'FALSE'. See coloredmesh.plot.colorbar.separate for an alternative.
<code>makecmap_options</code>	named list of parameters to pass to makecmap . Must not include the unnamed first parameter, which is derived from 'measure'. Should include at least a colormap function as name 'colFn'.
<code>map_to_NA</code>	the value or value range that should not be considered a cluster, and should thus be plotted as background color. These values will be set to NA, leading to transparent rendering, so the background will be visible instead. If a single value, only exactly this value is used (typically 0). If two values, they are interpreted as a range, and a values between them are mapped to NA. If you prefer to map the data to NA yourself before using this function or do not want to use a , pass 'NULL'.
<code>bg</code>	a background definition. Can be a surface color layer or a character string like 'curv_light' to select a pre-defined layer, see collayer.bg for valid strings.
<code>morph_data_both</code>	numeric vector or NULL, the data to visualize on both hemispheres. This must be a single vector with length equal to the sum of the vertex counts of the left and the right hemisphere. The data for the left hemisphere must come first. If this is given, 'morph_data_lh' and 'morph_data_rh' must be NULL.

Value

list of coloredmeshes. The coloredmeshes used for the visualization.

See Also

Other visualization functions: [vis.color.on.subject\(\)](#), [vis.data.on.fsaverage\(\)](#), [vis.data.on.subject\(\)](#), [vis.labeldata.on.subject\(\)](#), [vis.mask.on.subject\(\)](#), [vis.region.values.on.subject\(\)](#), [vis.subject.annot\(\)](#), [vis.subject.label\(\)](#), [vis.subject.morph.native\(\)](#), [vis.subject.morph.standard\(\)](#), [vislayout.from.coloredmeshes\(\)](#)

Other morphometry visualization functions: [vis.data.on.fsaverage\(\)](#), [vis.data.on.subject\(\)](#), [vis.subject.morph.native\(\)](#), [vis.subject.morph.standard\(\)](#)

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
```

```
morph_data_lh = subject.morph.native(subjects_dir, 'subject1', 'thickness', 'lh');
morph_data_rh = NULL;
vis.symmetric.data.on.subject(subjects_dir, 'subject1', morph_data_lh, morph_data_rh);
```

vislayout.from.coloredmeshes

Visualize coloredmeshes from several angles and combine the images into a new figure.

Description

Create a tight layout view of coloredmeshes from several angles. Creates separate ‘sd_<angle>’ images, then crops and finally merges them into a single output image with image magick. The ‘coloredmeshes’ to pass to this function are usually obtained by running any ‘vis*’ function (like [vis.subject.morph.native](#), [vis.subject.morph.standard](#), [vis.subject.label](#), [vis.subject.annot](#), and others). That means you can use this function to visualize all kinds of data, e.g., morphometry data in native and standard space, labels, and brain atlases.

Usage

```
vislayout.from.coloredmeshes(
  coloredmeshes,
  view_angles = get.view.angle.names(angle_set = "t4"),
  rgloptions = rglo(),
  rglactions = list(),
  style = "default",
  output_img = "fsbrain_arranged.png",
  silent = FALSE,
  grid_like = TRUE
)
```

Arguments

coloredmeshes,	list of coloredmesh. A coloredmesh is a named list as returned by the ‘coloredmesh.from*’ functions (like coloredmesh.from.morph.native). It has the entries ‘mesh’ of type tmesh3d, a ‘col’, which is a color specification for such a mesh. The ‘vis*’ functions (like vis.subject.morph.native) all return a list of coloredmeshes.
view_angles	list of strings. See get.view.angle.names for all valid strings.
rgloptions	option list passed to par3d . Example: <code>rgloptions = list("windowRect"=c(50, 50, 1000, 1000))</code> .
rglactions	named list. A list in which the names are from a set of pre-defined actions. The values can be used to specify parameters for the action.

style	character string, a rendering style, e.g., 'default', 'shiny' or 'semitransparent'. Alternatively, a named list of style parameters (see material3d), e.g., <code>list("shininess"=50, specular=...)</code> . Use the magic word 'from_mesh' to use the 'style' field of each <code>coloredmesh</code> instead of a single, global style. In that case, you will have to make sure your meshes have such a field, if not, the style 'default' is used as a fallback for those which don't.
output_img	string, path to the output file. Defaults to "fsbrain_arranged.png"
silent	logical, whether to suppress all messages
grid_like	logical, whether to arrange the images in a grid-like fashion. If FALSE, they will all be merged horizontally. Passed to arrange.brainview.images .

Value

named list, see [arrange.brainview.images](#) for details

See Also

Other visualization functions: [vis.color.on.subject\(\)](#), [vis.data.on.fsaverage\(\)](#), [vis.data.on.subject\(\)](#), [vis.labeldata.on.subject\(\)](#), [vis.mask.on.subject\(\)](#), [vis.region.values.on.subject\(\)](#), [vis.subject.annot\(\)](#), [vis.subject.label\(\)](#), [vis.subject.morph.native\(\)](#), [vis.subject.morph.standard\(\)](#), [vis.symmetric.data.on.subject\(\)](#)

Examples

```
## Not run:
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
# Use any vis function to get coloredmeshes. You can visualize morphometry,
# labels, an atlas, whatever. You can suppress the view unless you need it.
coloredmeshes = vis.subject.morph.native(subjects_dir, "subject1", "thickness",
  cortex_only=TRUE, rglactions=list("clip_data"=c(0.05, 0.95)),
  views=NULL);
# The meshes contain the surface, data, and color information and can be
# visualized. You could adapt the rendering style while doing so:
vislayout.from.coloredmeshes(coloredmeshes, style='shiny');
# You could change the rendering style on a per-mesh basis.
coloredmeshes[[1]]$style = list("shininess"=50, alpha=0.5);
vislayout.from.coloredmeshes(coloredmeshes, style='from_mesh');

## End(Not run)
```

Description

Compute the axis-aligned foreground bounding box of a 3D volume, i.e. the inner foreground area that must be retained if you want to remove all background from the corners of the volume. The foreground is determined by thresholding, such that all values greater than 0 are considered foreground. See [vol.boundary.mask](#) for details.

Usage

```
vol.boundary.box(volume, threshold = 0L, apply = FALSE)
```

Arguments

volume	a 3D image volume
threshold	numerical, the threshold intensity used to separate background and foreground. All voxels with intensity values greater than this value will be considered ‘foreground’ voxels.
apply	logical, whether to directly apply the bounding box and return the resulting volume instead.

Value

named list with 2 entries: ‘from’ is an integer vector of length 3, defining the minimal (x,y,z) foreground indices. ‘to’ is an integer vector of length 3, defining the maximal (x,y,z) foreground indices.

See Also

Other volume utility: [vol.imagestack\(\)](#), [vol.merge\(\)](#), [vol.overlay.colors.from.activation\(\)](#), [vol.planes\(\)](#), [vol.slice\(\)](#)

```
vol.boundary.box.apply
```

Apply a boundary box to a volume, returning the inner volume part

Description

Apply a boundary box to a volume, returning the inner volume part

Usage

```
vol.boundary.box.apply(volume, bbox)
```

Arguments

volume	a 3D image volume
bbox	the boundary box

Value

a 3D image volume, the inner volume part, resulting from the application of the boundary box

vol.hull	<i>Retain only the outer hull voxels of the foreground.</i>
----------	---

Description

Filters the *foreground* voxel in the volume by keeping only an outer border of voxels, and setting the inner core voxels to 'NA'. This is a utility function for voxel-based visualization. The goal is to remove the inner voxels, which will not be visible anyways, and thus to dramatically reduce the number of triangles that will need to be computed for the mesh.

Usage

```
vol.hull(volume, thickness = 1L, axes = c(2L))
```

Arguments

volume	numeric 3d array, must contain foreground voxel and background voxels. The latter must have value 'NA'. This function assumes that a solid foreground object surrounded by background exists in the volume.
thickness	integer, the width of the border in voxels, i.e., how many of the voxels in each upright column to keep at the top and at the bottom.
axes	integer vector, the axes to use. Valid values in the vector are 1L, 2L and 3L. You will have to use all 3 axes if you do not want any holes in the object. (Obviously, having noise around the object can still lead to holes.)

Value

numeric 3d array, a filtered version of the input. It contains at least as many 'NA' voxels as the input. If the function had any effect, it contains a lot more 'NA' values. The other values and the volume dimensions are left unchanged.

vol.imagestack	<i>Turn volume into an ImageMagick image stack.</i>
----------------	---

Description

Create an image from each slice along the axis, then stack those into an ImageMagick image stack.

Usage

```
vol.imagestack(volume, axis = 1L, intensity_scale = 255)
```

Arguments

volume	a 3D image volume. Can be numeric, or something that can be read directly by <code>magick::image_read</code> in 2D matrices (slices along the axis), e.g., a 3D array of color strings. If a 2D matrix is passed, the resulting stack will contain a single image.
axis	positive integer in range 1L..3L or an axis name, the axis to use.
intensity_scale	integer, value by which to scale the intensities in the volume to the range <code>'[0, 1]'</code> . Only used for numeric volumes. Set to <code>NULL</code> for data that can be read directly by <code>magick::image_read</code> , and to 1 for intensity data that requires no scaling. Defaults to 255, which is suitable for 8 bit image data.

Value

a vectorized ImageMagick image, containing one subimage per slice. This can be interpreted as an animation or whatever.

See Also

Other volume utility: [vol.boundary.box\(\)](#), [vol.merge\(\)](#), [vol.overlay.colors.from.activation\(\)](#), [vol.planes\(\)](#), [vol.slice\(\)](#)

vol.intensity.to.color

Convert integer intensity image to RGB color string form.

Description

Convert a gray-scale image defined by intensity values in range `'[0, 1]'` to an image with identical dimensions that contains an R color string (like `'#222222'`) at each position. The color strings are computed from the intensities, by taking the intensity value as the value for all three RGB channels. I.e., the output is still gray-scale, but defined in RGB space. To make it clear, this function does **not** apply a colormap. It only changes the representation of the data, not the resulting colors.

Usage

```
vol.intensity.to.color(volume, scale = NULL)
```

Arguments

volume	numeric array, typically a 3D image with intensities in range <code>'[0, 1]'</code> . This function now also supports numeric matrices (2D images, slices) and numeric vectors (1D).
--------	--

`scale` numeric or character string, a scaling to apply to the values. Defaults to `NULL`, which means *no scaling* and requires the values in `'volume'` to be in range `'[0, 1]'`. You can pass a number like 255 or the string `'normalize'` to scale based on the data. You can pass the string `'normalize_if_needed'` to scale only if the data is *outside* the range `'[0, 1]'`, so that data in range `'[0.3, 0.5]'` would *not* be rescaled to `'[0, 1]'`.

Value

array (or matrix, or vector) of RGB color strings. All of them will represent gray values.

Examples

```
vol.intensity.to.color(c(0.0, 0.5, 1.0));
# output: "#000000" "#808080" "#FFFFFF"
vol.intensity.to.color(c(20, 186, 240), scale="normalize");
vol.intensity.to.color(c(20, 186, 240), scale=255);
vol.intensity.to.color(c(0.0, 0.5, 0.8), scale="normalize");
vol.intensity.to.color(c(0.0, 0.5, 0.8), scale="normalize_if_needed");
```

`vol.mask.from.segmentation`

Extract subset from a volume by value.

Description

Extract subset from a volume by value, set all other voxel values to `'NA'`. Typically used to extract a brain structure (like corpus callosum) from a volume segmentation (like the `'mri/aseg.mgz'` file of a subject). You should consider passing the volume and the include values as integers.

Usage

```
vol.mask.from.segmentation(volume, include_values)
```

Arguments

`volume` numeric 3D array

`include_values` numerical vector, the intensity values which qualify a voxel to be part of the result (without being set to `NA`)

Value

numerical array with same dimensions as the input volume. All values which are not part of `'include_values'` replaced with `'NA'`.

`vol.merge`*Merge background volume and overlay to new colors.*

Description

Merge background volume and overlay to new colors.

Usage

```
vol.merge(  
    volume,  
    overlay_colors,  
    bbox_threshold = 0L,  
    forced_overlay_color = NULL  
)
```

Arguments

`volume` 3D array, can be numeric (gray-scale intensity values) or color strings. If numeric, the intensity values must be in range `[0, 1]`.

`overlay_colors` 3D array of color strings, values which are not part of the overlay (and should display background in the result) must have `'NA'` instead of a color string. Must have same dimensions as the `'volume'`.

`bbox_threshold` numerical, the threshold intensity used to separate background and foreground. All voxels with intensity values greater than this value in the background `'volume'` will be considered `'foreground'` voxels. Background-only slices at the borders of the volume will be discarded. Pass `'NULL'` to use the full image without applying any bounding box.

`forced_overlay_color` NULL or an rgb color string, like `'#FF0000'` for red. If NULL, the activation colors will be used as foreground colors. Otherwise, the given color will be for all foreground vertices.

Value

3D array of color strings, the merged colors

See Also

Other volume utility: [vol.boundary.box\(\)](#), [vol.imagystack\(\)](#), [vol.overlay.colors.from.activation\(\)](#), [vol.planes\(\)](#), [vol.slice\(\)](#)

```
vol.overlay.colors.from.activation
```

Generate colors for a 3D volume, based on the activation data and a colormap.

Description

Applies the colormap function to the data, then sets the alpha value (transparency) to full in all areas without any activation. Feel free to clip data or whatever before passing it, so that all your no-activation data has the same value.

Usage

```
vol.overlay.colors.from.activation(  
    volume,  
    colormap_fn = squash::blueorange,  
    no_act_source_value = 0  
)
```

Arguments

`volume` a 3D array, the activation data (or p-values, effect sizes, or whatever)
`colormap_fn` function, a colormap function
`no_act_source_value` numerical scalar, the value from the data in 'volume' that means no activation. The output colors for this value will be set to 'NA'.

Value

a 3D matrix of color strings, with the same dimensions as the input volume

See Also

Other volume utility: [vol.boundary.box\(\)](#), [vol.imagestack\(\)](#), [vol.merge\(\)](#), [vol.planes\(\)](#), [vol.slice\(\)](#)

```
vol.overlay.colors.from.colortable
```

Compute voxel colors based on colortable.

Description

Use the intensity values of the voxels in volume and lookup the respective colors in a colortable.

Usage

```
vol.overlay.colors.from.colortable(
  volume,
  colortable,
  ignored_struct_indices = c(),
  ignored_struct_names = c("unknown", "Unknown")
)
```

Arguments

volume numeric 3D array, the values should be integers present in the 'struct_index' column of the colortable. All other values will be assigned 'NA' as a color.

colortable a colortable, as returned by [read.fs.colortable](#).

ignored_struct_indices integer vector, 'struct_index' entries in the colortable that should be ignored

ignored_struct_names vector of character strings, 'struct_name' entries in the colortable that should be ignored. Can be combined with 'ignored_struct_indices'.

Value

character string 3D array, the colors. Voxels in the volume which were not matched by the colortable are set to 'NA' in it.

vol.planes	<i>Translate names and indices of planes.</i>
------------	---

Description

Translate names and indices of 3D image planes. The names only make sense if the data in the volume is in the default FreeSurfer conformed orientation.

Usage

```
vol.planes(plane = NULL)
```

Arguments

plane NULL, a plane index, or a plane name.

Value

if 'plane' is NULL, all available planes and their indices as a named list. If 'plane' is an integer (a plane index), its name. If 'plane' is an characters string (a plane name), its index.

See Also

Other volume utility: [vol.boundary.box\(\)](#), [vol.imagestack\(\)](#), [vol.merge\(\)](#), [vol.overlay.colors.from.activation](#)
[vol.slice\(\)](#)

vol.slice	<i>Extract a slice of a 3D image stack.</i>
-----------	---

Description

Extracts one or more 2D slices from a 3D image (or a frame of a 4D image). To display the result, you can use [volvis.lightbox](#).

Usage

```
vol.slice(
    volume,
    slice_index = NULL,
    frame = 1L,
    axis = 1L,
    rotation = 0L,
    flip = NULL
)
```

Arguments

volume	a 3D or 4D image volume. Note that empty dimensions will be dropped before any processing, and the remaining volume must have 3 or 4 dimensions.
slice_index	positive integer or vector of positive integers, the index into the slices (for the axis). A <i>*slice*</i> in the sense of this function is any 2D image plane extracted from the 3D volume (no matter the axis). If NULL, the slice in the middle of the volume is used. One can pass the magic character string 'all' to use all slice indices along the axis.
frame	positive integer, optional. The frame (time point) to use, only relevant for 4D volumes. The last (i.e. 4th) dimension is assumed to be the time dimension in that case.
axis	positive integer, the axis to use when indexing the slices. Defaults to 1.
rotation	integer, rotation in degrees. Defaults to 0 (no rotation). Must be a multiple of 90L if given.
flip	NULL or one of the character strings 'vertically' or 'horizontally'. Note that flipping <i>*horizontally*</i> means that the image will be mirrored along the central <i>*vertical*</i> axis. If 'NULL' is passed, nothing is flipped. Flipping occurs after rotation.

Value

slice data. If 'slice_index' is a scalar, a numerical 2D matrix (a 2D image from the stack). Otherwise, a numerical 3D array that contains the selected 2D images.

See Also

Other volume utility: `vol.boundary.box()`, `vol.imagestack()`, `vol.merge()`, `vol.overlay.colors.from.activation`
`vol.planes()`

`vol.vox.from.crs` *Compute R voxel index for FreeSurfer CRS voxel index.*

Description

Performs a vox2vos transform from FreeSurfer to R indices.

Usage

```
vol.vox.from.crs(fs_crs, add_affine = FALSE)
```

Arguments

`fs_crs` integer vector of length 3, FreeSurfer indices for column, row, and slice (CRS).
`add_affine` logical, whether to add 1 to the output vector as the 4th value

Value

the R indices into the volume data for the given FreeSurfer CRS indices

Examples

```
# Get voxel intensity data on the command line, based
# on the FreeSurfer (zero-based) CRS voxel indices:
# `mri_info --voxel 127 100 100 ~/data/tim_only/tim/mri/brain.mgz`
# (the result is: 106.0)
#
# That should be identical to:
# our_crs = vol.vox.from.crs(c(127,100,100), add_affine = FALSE);
# brain$data[our_crs[1], our_crs[2], our_crs[3]]; # gives 106
```

`volvis.contour` *Visualize contour of a volume.*

Description

Compute a smoothed surface from the voxel intensities in the given volume and render it. Requires the ‘misc3d’ package to be installed, which is an optional dependency.

Usage

```
volvis.contour(volume, level = 80, show = TRUE, frame = 1L, color = "white")
```

Arguments

volume	a 3D brain volume
level	numeric, intensity threshold for the data. Voxels with intensity value smaller than 'level' will be ignored when creating the contour surface.
show	logical, whether to display the triangles. Defaults to 'TRUE'.
frame	integer, the frame to show in case of a 4D input volume. Can also be the character string 'all' to draw the contents of all frames at once. Useful to plot white matter tracts from DTI data, where each tract is stored in a different frame.
color	the color to use when plotting. Can be a vector of colors when plotting all frames of a 4D image (one color per frame).

Value

the rendered triangles (a 'Triangles3D' instance) with coordinates in surface RAS space if any, 'NULL' otherwise. This will be a list if you pass a 4D volume and select 'all' frames.

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
brain = subject.volume(subjects_dir, 'subject1', 'brain');
# Plot all voxels of the brain:
volvis.contour(brain);
```

volvis.lightbox *Draw a lightbox view from volume slices.*

Description

A lightbox is a single image that holds a set of subimages, arranged in a grid. The images can have a small border or spacing between them. Consecutive subimages will be appear the same row of the grid.

If `overlay_colors` are given, the volume will be used as the background, and it will only be visible where `overlay_colors` has transparency.

Usage

```
volvis.lightbox(
  volume,
  slices = -5,
  axis = 1L,
  per_row = 5L,
  per_col = NULL,
```

```

border_geometry = "5x5",
background_color = "#000000",
arrange_single_image = FALSE
)

```

Arguments

volume	3D array, can be numeric (gray-scale intensity values) or color strings. If numeric, the intensity values must be in range <code>[0, 1]</code> .
slices	slice index definition. If a vector of integers, interpreted as slice indices. If a single negative interger <code>-n</code> , interpreted as every <code>'nth'</code> slice, starting at slice 1. The character string <code>'all'</code> or the value <code>'NULL'</code> will be interpreted as <i>*all slices*</i> .
axis	positive integer in range 1L..3L, the axis to use.
per_row	positive integer, the number of subimages per row in the output image. If <code>'NULL'</code> , automatically computed from the number of slices and the <code>'per_col'</code> parameter.
per_col	positive integer, the number of subimages per column in the output image. If <code>'NULL'</code> , automatically computed from the number of slices and the <code>'per_row'</code> parameter.
border_geometry	string, a geometry string passed to <code>magick::image_border</code> to define the borders to add to each image tile. The default value adds 5 pixels, both horizontally and vertically.
background_color	string, a valid ImageMagick color string such as <code>"white"</code> or <code>"#000080"</code> . The color to use when extending images (e.g., when creating the border). Defaults to black.
arrange_single_image	logical, whether to apply the given arrangement (from parameters <code>'per_row'</code> and <code>'per_column'</code>) even if a single slice (a 2D image) is passed as <code>'volume'</code> . Defaults to <code>FALSE</code> , which prevents that background tiles are added to fill the row up to <code>'per_row'</code> images. This also prevents the border from getting added to a single image, so all you see is the raw image. Set to <code>'TRUE'</code> if you want to arrange even a single image in a row with a border.

Value

a magick image instance

Description

Plots a 3D box at every **foreground** voxel in the given volume. All voxels which do not have their intensity value set to 'NA' are considered **foreground** voxels. The locations at which to plot the voxels is computed from the voxel CRS indices using the FreeSurfer `vox2ras_tkr` matrix. This means that the position of the rendered data fits to the surface coordinates (in files like 'surf/lh.white'), and that you can call this function while an active surface rendering window is open (e.g., from calling `vis.subject.morph.native`), to superimpose the surface and volume data. ****On coloring the voxels**** (using **rgl materials**): Note that you can call this function several times for the active plot, and color the voxels differently by passing different material properties in each call. Alternatively, check the 'voxelcol' parameter.

Usage

```
volvis.voxels(volume, render_every = 1, voxelcol = NULL, ...)
```

Arguments

volume	numeric 3d array, voxels which should not be plotted must have value 'NA'. Take care not to plot too many.
render_every	integer, how many to skip before rendering the next one (to improve performance and/or see deeper structures). Use higher values to see a less dense representation of your data that usually still allows you to see the general shape, but at lower computational burden. Set to 1 to render every (foreground) voxel.
voxelcol	character string or a <i>*voxel coloring*</i> . A <i>*voxel coloring*</i> can be specified in three ways: 1) the string 'from_intensity' will compute colors based on the intensity values of the foreground voxels in the volume, applying normalization of the intensity values if needed. 2) an array of RGB color strings: will be used to retrieve the colors for all foreground vertices, at their CRS indices. 3) A vector with length identical to the number of foreground voxels in the volume: will be applied directly. Obviously, you should not pass a color material parameter (see '...') when using this.
...	material properties, passed to <code>triangles3d</code> . Example: <code>color = "#0000ff", lit=FALSE</code> .

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
brain = subject.volume(subjects_dir, 'subject1', 'brain');
# Plot all voxels of the brain:
brain[which(brain==0L, arr.ind = TRUE)] = NA; # mark background
brain = vol.hull(brain); # remove inner triangles
volvis.voxels(brain);
```

 vox2ras_tkr

The FreeSurfer default vox2ras_tkr matrix.

Description

Applying this matrix to a FreeSurfer CRS index of a conformed volume will give you the RAS coordinates of the voxel in surface coordinates, i.e., in the coordinates used in surface file like 'lh.white'. The central voxel is 127,127,127 when using zero-based indices (or 128,128,128 when using one-based indices), meaning its surface RAS coordinates are 0.0, 0.0, 0.0. The returned matrix is the inverse of the 'ras2vox_tkr' matrix.

Usage

```
vox2ras_tkr()
```

Value

numeric 4x4 matrix, the FreeSurfer vox2ras_tkr matrix.

See Also

Other surface and volume coordinates: [ras2vox_tkr\(\)](#)

Examples

```
# Compute surface RAS coordinate of voxel with CRS (0L, 0L, 0L):
vox2ras_tkr() %% c(0, 0, 0, 1);
# Show that voxel with CRS (128,128,128) is at the
# origin (0.0, 0.0, 0.0) of the surface RAS coordinate system:
(vox2ras_tkr() %% c(128, 128, 128, 1))[1:3];
```

 write.group.morph.standard.sf

Write combined data for a group to a single MGH file.

Description

Write morphometry data for a group into a single MGH or MGZ file.

Usage

```
write.group.morph.standard.sf(filepath, data)
```


Arguments

filepath	character string, path to the target file, should end with '.mgh' or '.mgz'.
data	numerical 4d matrix, with first dimension holding 1D data for all vertices of a subject, 2nd and 3rd dimension are of length 1, and 4th dimension is for each subject.

Note

The file will contain no information on the subject identifiers. The data can be for one or both hemispheres. See [group.morph.standard.sf](#) to read the data back into R.

```
write.region.aggregated
```

Write data aggregated over regions to morphometry file for group.

Description

Given an atlas, a subjects list and a measure, aggregate the measure over each region (e.g., mean) and write an output morphometry file in which the value for all region vertices is set to the aggregated value.

Usage

```
write.region.aggregated(
  subjects_dir,
  subjects_list,
  measure,
  hemi,
  atlas,
  agg_fun = mean,
  outfile_morph_name = "",
  format = "mgz"
)
```

Arguments

subjects_dir,	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subjects_list,	string vector. A vector of subject identifiers that match the directory names within subjects_dir.
measure,	string. Name of the vertex-wise measure of morphometry data file. E.g., "area" or "thickness". Used to construct the name of the morphometry file to be loaded.
hemi,	string, one of 'lh' or 'rh'. The hemisphere name. Used to construct the names of the annotation and morphometry data files to be loaded.

atlas,	string. The atlas name. E.g., "aparc", "aparc.2009s", or "aparc.DKTatlas". Used to construct the name of the annotation file to be loaded.
agg_fun,	function. An R function that aggregates data, typically max, mean, min or something similar. Note: this is NOT a string, put the function name without quotes. Defaults to mean.
outfile_morph_name,	string. The measure part of the output file name. E.g., 'agg_thickness' will write the file '<subject>/surf/<hemi>.agg_thickness.mgh'. Defaults to 'agg_<measure>'.
format,	string. A morphometry file format. One of 'mgh', 'mgz' or 'curv.' The output file name extension will be set accordingly. Defaults to 'mgz'.

See Also

Other output functions: [write.region.values.fsaverage\(\)](#), [write.region.values\(\)](#)

write.region.values *Write one value per atlas region for a subject.*

Description

Given an atlas and a list that contains one value for each atlas region, write a morphometry file in which all region vertices are assigned the value. Can be used to plot stuff like p-values or effect sizes onto brain regions.

Usage

```
write.region.values(
  subjects_dir,
  subject_id,
  hemi,
  atlas,
  region_value_list,
  outfile_morph_name,
  format = "mgz",
  do_write_file = TRUE,
  output_path = NULL,
  value_for_unlisted_regions = NaN
)
```

Arguments

subjects_dir,	string. The FreeSurfer SUBJECTS_DIR, i.e., a directory containing the data for all your subjects, each in a subdir named after the subject identifier.
subject_id,	string. The subject identifier
hemi,	string, one of 'lh' or 'rh'. The hemisphere name. Used to construct the names of the annotation and morphometry data files to be loaded.

atlas,	string. The atlas name. E.g., "aparc", "aparc.2009s", or "aparc.DKTatlas". Used to construct the name of the annotation file to be loaded.
region_value_list,	named list. A list in which the names are atlas regions, and the values are the value to write to all vertices of that region.
outfile_morph_name,	string. The measure part of the output file name. E.g., 'agg_thickness' will write the file '<subject>/surf/<hemi>.agg_thickness.mgh'.
format,	string. A morphometry file format. One of 'mgh', 'mgz' or 'curv.' The output file name extension will be set accordingly. Defaults to 'mgz'.
do_write_file,	logical. Whether to write the data to a file on the disk. If FALSE, the data are only returned (and the outfile_morph_name parameter gets ignored). Default to TRUE.
output_path	string, path to the output directory. If omitted, defaults to the 'surf' directory of the subject (i.e., '<subjects_dir>/<subject_id>/surf/').
value_for_unlisted_regions,	numeric scalar. The value to assign to vertices which are part of atlas regions that are not listed in region_value_list. Defaults to NaN.

Value

a named list with the following entries: "data": a vector containing the data. "file_written": string, path to the file that was written, only exists if do_write = TRUE.

See Also

Other output functions: [write.region.aggregated\(\)](#), [write.region.values.fsaverage\(\)](#)

Examples

```
fsbrain::download_optional_data();
subjects_dir = fsbrain::get_optional_data_filepath("subjects_dir");
region_value_list = list("bankssts"=0.9, "precuneus"=0.7);
write.region.values(subjects_dir, 'subject1', 'lh', 'aparc',
  region_value_list, 'pvalues.mgz', do_write_file = FALSE);
```

write.region.values.fsaverage

Write one value per atlas region for a template subject.

Description

Given an atlas and a list that contains one value for each atlas region, write a morphometry file in which all region vertices are assigned the value. Can be used to plot stuff like p-values or effect sizes onto brain regions.

Usage

```
write.region.values.fsaverage(
    hemi,
    atlas,
    region_value_list,
    output_file,
    template_subject = "fsaverage",
    template_subjects_dir = NULL,
    show_freeview_tip = FALSE,
    value_for_unlisted_regions = NaN
)
```

Arguments

hemi,	string, one of 'lh' or 'rh'. The hemisphere name. Used to construct the names of the annotation and morphometry data files to be loaded.
atlas,	string. The atlas name. E.g., "aparc", "aparc.2009s", or "aparc.DKTatlas". Used to construct the name of the annotation file to be loaded.
region_value_list,	named list. A list in which the names are atlas regions, and the values are the value to write to all vertices of that region.
output_file,	string or 'NULL'. Path of the output file, including file name and extension. The format is determined from the (absence of a) file extension. If NULL, no file will be written.
template_subject	string, template subject name. Defaults to 'fsaverage'.
template_subjects_dir	string, the path to the subjects directory containing the template subject directory. If this is 'NULL', the function will try to find it using the environment, see the function find.subjectsdir.of for details. Defaults to NULL.
show_freeview_tip	logical, whether to print the freeview command on howto use the overlay to the console. (Only happens if the output_file is not 'NULL'.)
value_for_unlisted_regions,	numeric scalar. The value to assign to vertices which are part of atlas regions that are not listed in region_value_list. Defaults to NaN.

Value

a named list with the following entries: "data": a vector containing the data. "file_written": string, path to the file that was written, only exists if do_write = TRUE.

See Also

Other output functions: [write.region.aggregated\(\)](#), [write.region.values\(\)](#)

Index

- * **aggregation functions**
 - group.agg.atlas.native, 53
 - group.agg.atlas.standard, 55
 - group.morph.agg.standard.vertex, 66
 - subject.atlas.agg, 110
- * **atlas functions**
 - get.atlas.region.names, 50
 - group.agg.atlas.native, 53
 - group.agg.atlas.standard, 55
 - group.annot, 57
 - group.label.from.annot, 61
 - label.from.annotdata, 81
 - label.to.annot, 82
 - regions.to.ignore, 99
 - spread.values.over.annot, 105
 - spread.values.over.hemi, 106
 - spread.values.over.subject, 107
 - subject.annot, 109
 - subject.atlas.agg, 110
 - subject.label.from.annot, 114
 - subject.lobes, 116
- * **color functions**
 - alphablend, 5
 - desaturate, 40
- * **colorbar functions**
 - coloredmesh.plot.colorbar.separate, 31
 - combine.colorbar.with.brainview.animation, 35
 - combine.colorbar.with.brainview.image, 36
- * **coloredmesh functions**
 - coloredmesh.from.annot, 23
 - coloredmesh.from.label, 24
 - coloredmesh.from.mask, 26
 - coloredmesh.from.morph.native, 27
 - coloredmesh.from.morph.standard, 28
 - coloredmesh.from.morphdata, 30
 - coloredmeshes.from.color, 33
- * **concatination functions**
 - group.concat.measures.native, 58
 - group.concat.measures.standard, 59
- * **global aggregation functions**
 - group.morph.agg.native, 62
 - group.morph.agg.standard, 64
 - group.multimorph.agg.native, 71
 - group.multimorph.agg.standard, 72
- * **group visualization functions**
 - vis.data.on.group.native, 131
 - vis.data.on.group.standard, 133
 - vis.group.annot, 139
 - vis.group.coloredmeshes, 140
 - vis.group.morph.native, 141
 - vis.group.morph.standard, 142
- * **label data functions**
 - group.label, 60
 - labeldata.from.mask, 83
 - mask.from.labeldata.for.hemi, 86
 - subject.label, 113
- * **label functions**
 - apply.label.to.morphdata, 7
 - apply.labeldata.to.morphdata, 8
 - subject.lobes, 116
 - subject.mask, 117
 - vis.labeldata.on.subject, 144
 - vis.subject.label, 153
- * **mask functions**
 - coloredmesh.from.mask, 26
 - mask.from.labeldata.for.hemi, 86
 - vis.mask.on.subject, 146
- * **mesh data functions**
 - group.surface, 74
- * **metadata functions**
 - demographics.to.fsgd.file, 39
 - read.md.demographics, 96
 - read.md.subjects, 97

- report.on.demographics, 99
- * **morphometry data functions**
 - apply.label.to.morphdata, 7
 - apply.labeldata.to.morphdata, 8
 - group.morph.native, 67
 - group.morph.standard, 68
 - subject.morph.native, 118
 - subject.morph.standard, 119
- * **morphometry visualization functions**
 - vis.data.on.fsaverage, 129
 - vis.data.on.subject, 134
 - vis.subject.morph.native, 155
 - vis.subject.morph.standard, 157
 - vis.symmetric.data.on.subject, 159
- * **output functions**
 - write.region.aggregated, 177
 - write.region.values, 178
 - write.region.values.fsaverage, 179
- * **quality check functions**
 - qc.for.group, 92
 - qc.from.regionwise.df, 93
- * **region-based visualization functions**
 - vis.region.values.on.subject, 149
 - vis.subject.annot, 151
- * **surface and volume coordinates**
 - ras2vox_tkr, 95
 - vox2ras_tkr, 176
- * **surface color layer**
 - collayer.bg, 16
 - collayer.bg.atlas, 16
 - collayer.bg.meancurv, 18
 - collayer.bg.sulc, 19
 - collayer.from.annot, 20
 - collayer.from.annotdata, 20
 - collayer.from.mask.data, 21
 - collayer.from.morphlike.data, 22
 - collayers.merge, 23
- * **surface mesh functions**
 - face.edges, 44
 - label.border, 79
 - mesh.vertex.neighbors, 87
 - subject.surface, 122
 - vis.path.along.verts, 148
- * **surface visualization functions**
 - vis.color.on.subject, 125
- * **visualization functions**
 - vis.color.on.subject, 125
 - vis.data.on.fsaverage, 129
 - vis.data.on.subject, 134
 - vis.labeldata.on.subject, 144
 - vis.mask.on.subject, 146
 - vis.region.values.on.subject, 149
 - vis.subject.annot, 151
 - vis.subject.label, 153
 - vis.subject.morph.native, 155
 - vis.subject.morph.standard, 157
 - vis.symmetric.data.on.subject, 159
 - vislayout.from.coloredmeshes, 162
- * **volume utility**
 - vol.boundary.box, 163
 - vol.imagestack, 165
 - vol.merge, 168
 - vol.overlay.colors.from.activation, 169
 - vol.planes, 170
 - vol.slice, 171
- * **volume visualization**
 - volvis.lightbox, 173
- alphablend, 5, 41
- annot.outline, 6, 17, 152
- apply.label.to.morphdata, 7, 9, 68, 69, 116, 118–120, 145, 154
- apply.labeldata.to.morphdata, 8, 8, 68, 69, 116, 118–120, 145, 154
- apply.transform, 9
- arrange.brainview.images, 9, 163
- arrange.brainview.images.grid, 10, 132, 134, 140–143
- brainviews, 12
- c, 76
- clip.data, 13, 27, 29, 84, 85, 100
- cm.cbry, 13
- cm.div, 14
- cm.heat, 14
- cm.qual, 15
- cm.seq, 15
- collayer.bg, 16, 17–23, 131, 135, 150, 154, 156, 159, 161
- collayer.bg.atlas, 16, 16, 18–23
- collayer.bg.meancurv, 16, 17, 18, 19–23
- collayer.bg.sulc, 16–18, 19, 20–23
- collayer.from.annot, 16–19, 20, 21–23
- collayer.from.annotdata, 16–20, 20, 22, 23

- collayer.from.mask.data, [16–21](#), [21](#), [22](#), [23](#)
- collayer.from.morphlike.data, [16–22](#), [22](#), [23](#)
- collayers.merge, [16–22](#), [23](#)
- coloredmesh.from.annot, [23](#), [25](#), [27–30](#), [34](#)
- coloredmesh.from.label, [24](#), [24](#), [27–30](#), [34](#)
- coloredmesh.from.mask, [24](#), [25](#), [26](#), [28–30](#), [34](#), [86](#), [147](#)
- coloredmesh.from.morph.native, [24](#), [25](#), [27](#), [27](#), [29](#), [30](#), [34](#), [137](#), [162](#)
- coloredmesh.from.morph.standard, [24](#), [25](#), [27](#), [28](#), [28](#), [30](#), [34](#)
- coloredmesh.from.morphdata, [24](#), [25](#), [27–29](#), [30](#), [34](#)
- coloredmesh.from.preloaded.data, [31](#)
- coloredmesh.plot.colorbar.separate, [12](#), [31](#), [35](#), [37](#), [127](#), [131](#), [135](#), [145](#), [147](#), [150](#), [154](#), [156](#), [158](#), [161](#)
- coloredmeshes.from.color, [24](#), [25](#), [27–30](#), [33](#)
- colorlist.brain.clusters, [34](#)
- combine.colorbar.with.brainview.animation, [33](#), [35](#), [37](#)
- combine.colorbar.with.brainview.image, [33](#), [35](#), [36](#)
- cube3D.tris, [37](#), [38](#)
- cubes3D.tris, [38](#)
- delete_all_optional_data, [39](#)
- demographics.to.fsgd.file, [39](#), [97](#), [98](#), [100](#)
- desaturate, [6](#), [40](#)
- download_fsaverage, [41](#), [45](#)
- download_fsaverage3, [42](#)
- download_optional_data, [42](#)
- download_optional_paper_data, [43](#)
- export.coloredmesh.ply, [43](#)
- face.edges, [44](#), [80](#), [88](#), [122](#), [149](#)
- find.freesurferhome, [45](#)
- find.subjectsdir.of, [45](#), [180](#)
- fs.coloredmesh, [46](#)
- fs.home, [45](#), [47](#)
- fsaverage.path, [46](#), [47](#)
- fsbrain.set.default.figsize, [48](#)
- fup, [48](#)
- gen.test.volume, [49](#)
- get.atlas.region.names, [7](#), [50](#), [54](#), [56](#), [57](#), [62](#), [82](#), [83](#), [99](#), [106–109](#), [111](#), [115](#), [116](#)
- get.rglstyle, [51](#)
- get.view.angle.names, [12](#), [51](#), [132](#), [133](#), [137](#), [139](#), [141–143](#), [162](#)
- get_optional_data_filepath, [53](#)
- getIn, [52](#)
- group.agg.atlas.native, [50](#), [53](#), [56](#), [57](#), [62](#), [67](#), [82](#), [83](#), [93](#), [99](#), [106–109](#), [111](#), [115](#), [116](#)
- group.agg.atlas.standard, [50](#), [54](#), [55](#), [57](#), [62](#), [67](#), [82](#), [83](#), [99](#), [106–109](#), [111](#), [115](#), [116](#)
- group.annot, [50](#), [54](#), [56](#), [57](#), [62](#), [82](#), [83](#), [99](#), [106–109](#), [111](#), [115](#), [116](#)
- group.concat.measures.native, [58](#), [60](#)
- group.concat.measures.standard, [58](#), [59](#)
- group.label, [60](#), [84](#), [86](#), [114](#)
- group.label.from.annot, [50](#), [54](#), [56](#), [57](#), [61](#), [82](#), [83](#), [99](#), [106–109](#), [111](#), [115](#), [116](#)
- group.morph.agg.native, [62](#), [65](#), [72](#), [74](#)
- group.morph.agg.standard, [64](#), [64](#), [72](#), [74](#)
- group.morph.agg.standard.vertex, [54](#), [56](#), [66](#), [111](#)
- group.morph.native, [8](#), [9](#), [67](#), [69](#), [119](#), [120](#), [132](#)
- group.morph.standard, [8](#), [9](#), [68](#), [68](#), [119](#), [120](#), [133](#)
- group.morph.standard.sf, [70](#), [133](#), [177](#)
- group.multimorph.agg.native, [64](#), [65](#), [71](#), [74](#)
- group.multimorph.agg.standard, [64](#), [65](#), [72](#), [72](#)
- group.surface, [74](#)
- hasIn, [75](#)
- hemilist.derive.hemi, [76](#)
- hemilist.get.combined.data, [76](#)
- hemilist.unwrap, [77](#)
- hemilist.wrap, [77](#)
- image.plot, [32](#)
- is.fs.coloredmesh, [78](#)
- is.fs.coloredvoxels, [78](#)
- is.hemilist, [79](#)
- jet, [25–27](#), [29](#), [30](#)
- label.border, [44](#), [79](#), [88](#), [122](#), [149](#)

- label.colFn, 80
- label.colFn.inv, 81
- label.from.annotdata, 50, 54, 56, 57, 62, 81, 83, 99, 106–109, 111, 115, 116
- label.to.annot, 50, 54, 56, 57, 62, 82, 82, 99, 106–109, 111, 115, 116
- labeldata.from.mask, 61, 83, 86, 114
- legend, 129
- limit_fun, 84, 100
- limit_fun_na, 85, 100
- list_optional_data, 85
- makecmap, 21, 22, 25, 26, 28–31, 131, 135, 139, 145, 147, 150, 154, 156, 159, 161
- mask.from.labeldata.for.hemi, 27, 61, 84, 86, 114, 146, 147
- material3d, 51, 163
- max, 63, 65, 66, 71, 73
- mean, 63, 65, 67, 72, 73
- mesh.vertex.included.faces, 44, 80, 88, 122, 149
- mesh.vertex.neighbors, 44, 80, 87, 122, 149
- mkco.cluster, 88
- mkco.div, 89
- mkco.heat, 89
- mkco.seq, 90
- par3d, 12, 125, 127, 128, 130, 135, 145, 147, 150, 152, 154, 156, 158, 160, 162
- png, 32
- principal.curvatures, 90, 103
- print.fs.coloredmesh, 91
- print.fs.coloredvoxels, 91
- qc.for.group, 92, 93, 95
- qc.from.regionwise.df, 92, 93, 94
- qc.from.segstats.table, 92, 93
- qc.from.segstats.tables, 94, 95
- qc.vis.failcount.by.region, 94
- ras2vox_tkr, 95, 176
- read.colorcsv, 96
- read.fs.annot, 83, 109
- read.fs.colortable, 129, 170
- read.fs.mgh, 123
- read.fs.morph, 119, 138
- read.fs.surface, 6, 44, 79, 88, 122, 138, 149
- read.md.demographics, 40, 96, 98, 100
- read.md.subjects, 40, 97, 97, 100
- read.md.subjects.from.fsgd, 70, 98
- read.table, 97
- regions.to.ignore, 50, 54, 56, 57, 62, 82, 83, 99, 106–109, 111, 115, 116
- report.on.demographics, 40, 97, 98, 99
- rgb, 23
- rgl.spheres, 102
- rglactions, 100, 125, 131–133, 135, 140, 142, 143, 145, 147, 150, 152, 154, 156, 158, 161
- rglo, 101
- rglot, 101, 101
- rglvoxels, 102
- shade3d, 51
- shape.descriptor.names, 103, 103
- shape.descriptors, 103
- shift.hemis.apart, 104
- spin3d, 128
- spread.values.over.annot, 50, 54, 56, 57, 62, 82, 83, 99, 105, 107–109, 111, 115, 116
- spread.values.over.hemi, 50, 54, 56, 57, 62, 82, 83, 99, 106, 106, 108, 109, 111, 115, 116
- spread.values.over.subject, 50, 54, 56, 57, 62, 82, 83, 99, 106, 107, 107, 109, 111, 115, 116
- subject.annot, 6, 21, 50, 54, 56, 57, 62, 82, 83, 99, 106–108, 109, 111, 115, 116, 129
- subject.atlas.agg, 50, 54, 56, 57, 62, 67, 82, 83, 99, 106–109, 110, 115, 116
- subject.filepath.morph.native, 111
- subject.filepath.morph.standard, 112
- subject.label, 8, 61, 84, 86, 113
- subject.label.from.annot, 50, 54, 56, 57, 62, 82, 83, 99, 106–109, 111, 114, 116
- subject.lobes, 8, 9, 50, 54, 56, 57, 62, 82, 83, 99, 106–109, 111, 115, 116, 118, 145, 154
- subject.mask, 8, 9, 116, 117, 145, 154
- subject.morph.native, 8, 9, 68, 69, 118, 120, 125
- subject.morph.standard, 8, 9, 68, 69, 119, 119

- subject.num.verts, 121
- subject.surface, 6, 26, 44, 46, 79, 80, 88, 122, 138, 149
- subject.volume, 123
- text3d, 100
- tmesh3d, 24–26, 28–30, 34, 46
- toupper, 48
- triangles3d, 37, 102, 175
- vdata.split.by.hemi, 124
- vis.color.on.subject, 16–22, 125, 131, 136, 145, 147, 151, 152, 154, 156, 159, 161, 163
- vis.coloredmeshes, 126, 139
- vis.coloredmeshes.rotating, 127
- vis.colortable.legend, 129
- vis.data.on.fsaverage, 126, 129, 136, 145, 147, 151, 152, 154, 156, 157, 159, 161, 163
- vis.data.on.group.native, 131, 134, 140–142, 144
- vis.data.on.group.standard, 133, 133, 140–142, 144
- vis.data.on.subject, 126, 131, 134, 145, 147, 151, 152, 154, 156, 157, 159, 161, 163
- vis.export.from.coloredmeshes, 136
- vis.fs.surface, 138
- vis.group.annot, 133, 134, 139, 141, 142, 144
- vis.group.coloredmeshes, 133, 134, 140, 140, 142, 144
- vis.group.morph.native, 133, 134, 140, 141, 141, 144
- vis.group.morph.standard, 133, 134, 140–142, 142
- vis.labeldata.on.subject, 8, 9, 116, 118, 126, 131, 136, 144, 147, 151, 152, 154, 156, 159, 161, 163
- vis.mask.on.subject, 27, 86, 126, 131, 136, 145, 146, 151, 152, 154, 156, 159, 161, 163
- vis.path.along.verts, 44, 80, 88, 122, 148
- vis.region.values.on.subject, 95, 126, 131, 136, 145, 147, 149, 152, 154, 156, 159, 161, 163
- vis.subject.annot, 126, 131, 136, 145, 147, 151, 151, 154, 157, 159, 161–163
- vis.subject.label, 8, 9, 116, 118, 126, 131, 136, 145, 147, 151, 152, 153, 157, 159, 161–163
- vis.subject.morph.native, 32, 44, 51, 88–90, 101, 126, 131, 136, 137, 145, 147, 148, 151, 152, 154, 155, 159, 161–163, 175
- vis.subject.morph.standard, 126, 131, 136, 145, 147, 151, 152, 154, 157, 157, 161–163
- vis.symmetric.data.on.subject, 126, 131, 136, 145, 147, 151, 152, 154, 157, 159, 159, 163
- vislayout.from.coloredmeshes, 51, 126, 131, 136, 145, 147, 151, 152, 154, 157, 159, 161, 162
- vol.boundary.box, 163, 166, 168, 169, 171, 172
- vol.boundary.box.apply, 164
- vol.boundary.mask, 164
- vol.hull, 165
- vol.imagestack, 164, 165, 168, 169, 171, 172
- vol.intensity.to.color, 166
- vol.mask.from.segmentation, 167
- vol.merge, 164, 166, 168, 169, 171, 172
- vol.overlay.colors.from.activation, 164, 166, 168, 169, 171, 172
- vol.overlay.colors.from.colortable, 169
- vol.planes, 164, 166, 168, 169, 170, 172
- vol.slice, 164, 166, 168, 169, 171, 171
- vol.vox.from.crs, 172
- volvis.contour, 172
- volvis.lightbox, 171, 173
- volvis.voxels, 174
- vox2ras_tkr, 95, 175, 176
- which, 83
- write.group.morph.standard.sf, 70, 176
- write.region.aggregated, 177, 179, 181
- write.region.values, 178, 178, 181
- write.region.values.fsaverage, 178, 179, 179