

Package: freesurfer (via r-universe)

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Type Package

Title Wrapper Functions for 'Freesurfer'

Version 1.6.10

Description Wrapper functions that interface with 'Freesurfer' [<https://surfer.nmr.mgh.harvard.edu/>](https://surfer.nmr.mgh.harvard.edu/), a powerful and commonly-used 'neuroimaging' software, using system commands. The goal is to be able to interface with 'Freesurfer' completely in R, where you pass R objects of class 'nifti', implemented by package 'oro.nifti', and the function executes an 'Freesurfer' command and returns an R object of class 'nifti' or necessary output.

LazyData true

LazyLoad true

Imports methods, neurobase, tools, R.utils, reshape2, utils

Depends R (>= 3.2.0)

License GPL-3

Suggests magrittr, knitr, oro.nifti (>= 0.7), rgl, rmarkdown, pander, fslr (>= 2.9.2)

BugReports <https://github.com/muschellij2/freesurfer/issues>

SystemRequirements FSL, Freesurfer
(<https://surfer.nmr.mgh.harvard.edu/>)

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Repository <https://muschellij2.r-universe.dev>

RemoteUrl <https://github.com/muschellij2/freesurfer>

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aparcstats2table	<i>Parcellation Stats to Table</i>
------------------	------------------------------------

Description

This function calls `aparcstats2table` to convert parcellation statistics to a table

Usage

```

aparcstats2table(
  subjects,
  outfile = NULL,
  hemi = c("lh", "rh"),
  measure = c("area", "volume", "thickness", "thicknessstd", "meancurv", "gauscurv",
    "foldind", "curvind"),
  sep = c("tab", "space", "comma", "semicolon"),
  parc = c("aparc", "aparc.a2009s"),
  skip = FALSE,
  subj_dir = NULL,
  opts = "",
  verbose = TRUE
)

```

Arguments

subjects	(character) vector of subjects
outfile	(character) output filename
hemi	(character) hemisphere to run statistics
measure	(character) measure to be calculated
sep	(character) separator for the output file. This will be an attribute of outfile
parc	(character) parcellation to compute on
skip	(logical) if subject does not have parcellation, should the command skip that subject (TRUE) or error (FALSE)
subj_dir	(character path) if a different subjects directory is to be used other than SUBJECTS_DIR from shell, it can be specified here. Use with care as if the command fail, it may not reset the SUBJECTS_DIR back correctly after the error
opts	(character) additional options to aparcstats2table
verbose	(logical) print diagnostic messages

Value

Character filename of output file, with the attribute of the separator

Examples

```
if (have_fs()) {
  fs_subj_dir()
  outfile = aparcstats2table(subjects = "bert",
                             hemi = "lh",
                             meas = "thickness")
}
```

aparcstats2table.help *Parcellation Stats to Table Help*

Description

This calls Freesurfer's aparcstats2table help

Usage

```
aparcstats2table.help()
```

Value

Result of fs_help

aparcs_to_bg	<i>Convert Freesurfer aparcs Table to brainGraph</i>
--------------	--

Description

Converts Freesurfer aparcs table to brainGraph naming convention, relying on [aparcstats2table](#)

Usage

```
aparcs_to_bg(subjects, measure, ...)
```

Arguments

subjects	subjects to analyze, passed to aparcstats2table
measure	measure to be analyzed, passed to aparcstats2table
...	additional arguments passed to aparcstats2table

Value

Long data.frame

Examples

```
if (have_fs()) {
  fs_subj_dir()
  df = aparcs_to_bg(subjects = "bert", measure = "thickness")
  print(head(df))
}
```

asegstats2table	<i>Parcellation Stats to Table</i>
-----------------	------------------------------------

Description

This function calls `asegstats2table` to convert parcellation statistics to a table

Usage

```
asegstats2table(
  subjects = NULL,
  inputs = NULL,
  outfile = NULL,
  measure = c("volume", "mean", "std"),
  sep = c("tab", "space", "comma", "semicolon"),
  skip = FALSE,
  subj_dir = NULL,
```

```

    opts = "",
    verbose = TRUE
  )

```

Arguments

subjects	(character) vector of subjects
inputs	(character paths) vector of input filenames, e.g. aseg.stats.
outfile	(character) output filename
measure	(character) measure to be calculated
sep	(character) separator for the output file. This will be an attribute of outfile
skip	(logical) if subject does not have parcellation, should the command skip that subject (TRUE) or error (FALSE)
subj_dir	(character path) if a different subjects directory is to be used other than SUBJECTS_DIR from shell, it can be specified here. Use with care as if the command fail, it may not reset the SUBJECTS_DIR back correctly after the error
opts	(character) additional options to asegstats2table
verbose	(logical) print diagnostic messages

Value

Character filename of output file, with the attribute of the separator

Examples

```

if (have_fs()) {
  outfile = asegstats2table(subjects = "bert",
                           meas = "mean")
}

```

asegstats2table.help *Parcellation Stats to Table Help*

Description

This calls Freesurfer's asegstats2table help

Usage

```
asegstats2table.help()
```

Value

Result of fs_help

checkmnc-methods	<i>Force object to filename with .mnc extension</i>
------------------	---

Description

Ensures the output to be a character filename (or vector) from an input image or nifti to have .mnc extension and be converted to MNC when necessary

Usage

```
checkmnc(file, ...)  
  
## S4 method for signature 'nifti'  
checkmnc(file, ...)  
  
## S4 method for signature 'character'  
checkmnc(file, ...)  
  
## S4 method for signature 'list'  
checkmnc(file, ...)  
  
ensure_mnc(file, ...)
```

Arguments

file	character or nifti object
...	options passed to checking

Value

Character filename of mnc image

Author(s)

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check_fs_result	<i>Check Freesurfer Result</i>
-----------------	--------------------------------

Description

Checks the Freesurfer system command result and will stop or warning based on whether output files exist.

Usage

```
check_fs_result(res, fe_before, fe_after)
```

Arguments

res	(numeric) Result from system command
fe_before	(logical) did the output file exist before the command ran
fe_after	(logical) did the output file exist after the command ran

Value

No return value, called for side effects

construct_subj_dir *Construct Subject Directory*

Description

This function copies files specified by the types of data, determined by the folder Freesurfer put them in, into a temporary directory for easier separation of data and different structuring of data.

Usage

```
construct_subj_dir(
  label = NULL,
  mri = NULL,
  stats = NULL,
  surf = NULL,
  touch = NULL,
  subj = NULL,
  subj_root_dir = tempdir()
)
```

Arguments

label	Files to copy to subj_root_dir/subj/label folder
mri	Files to copy to subj_root_dir/subj/mri folder
stats	Files to copy to subj_root_dir/subj/stats folder
surf	Files to copy to subj_root_dir/subj/surf folder
touch	Files to copy to subj_root_dir/subj/touch folder
subj	Name of subject to make folder for to use for Freesurfer functions. If NULL, a temporary id will be generated
subj_root_dir	Directory to put folder with contents of subj

Value

List with the subject name, the SUBJECTS_DIR to use (the directory that contains the subject name), and the types of objects copied

Examples

```
## Not run:
library(freesurfer)
label = "/Applications/freesurfer/subjects/bert/label/aparc.annot.a2009s.ctab"
mri = c(
  "/Applications/freesurfer/subjects/bert/mri/aparc.a2009s+aseg.mgz",
  "/Applications/freesurfer/subjects/bert/mri/aseg.auto.mgz")
stats = c("/Applications/freesurfer/subjects/bert/stats/lh.aparc.stats",
  "/Applications/freesurfer/subjects/bert/stats/aseg.stats")
surf = "/Applications/freesurfer/subjects/bert/surf/lh.thickness"
touch = NULL

## End(Not run)
```

convert_surface	<i>Convert Freesurfer Surface</i>
-----------------	-----------------------------------

Description

Reads in a surface file from Freesurfer and separates into vertices and faces

Usage

```
convert_surface(infile, ...)
```

Arguments

infile	Input surface file
...	additional arguments to pass to <code>mris_convert</code>

Value

List of 3 elements: a header indicating the number of vertices and faces, the vertices, and the faces

Note

This was adapted from the gist: <https://gist.github.com/mm--/4a4fc7badacfad874102>

Examples

```
if (have_fs()) {
  infile = file.path(fs_subj_dir(),
    "bert", "surf", "rh.pial")
  res = convert_surface(infile = infile)
}
```

freesurferdir	<i>Get Freesurfer's Directory</i>
---------------	-----------------------------------

Description

Finds the FREESURFER_HOME from system environment or getopt("freesurfer.path") for location of Freesurfer functions and returns it

Usage

freesurferdir()

freesurfer_dir()

fs_dir()

Value

Character path

Examples

```
if (have_fs()) {  
  freesurferdir()  
  freesurfer_dir()  
  fs_dir()  
}
```

freesurfer_read3	<i>Freesurfer Read 3 records</i>
------------------	----------------------------------

Description

Reads first 3 records of file and returns the rotated value, for checking for other functions.

Usage

freesurfer_read3(file)

Arguments

file thickness file or anything in surf/ directory from Freesurfer subject

Value

Numeric

Examples

```
if (have_fs()) {  
  bert_dir = file.path(fs_subj_dir(), "bert", "surf")  
  file = file.path(bert_dir, "lh.thickness")  
  out = freesurfer_read3(file)  
}
```

freesurfer_read3_con *Freesurfer Read 3 records*

Description

Reads first 3 records from a connection and returns the rotated value, for checking for other functions.

Usage

```
freesurfer_read3_con(fid)
```

Arguments

fid	connection to a thickness file or anything in surf/ directory from Freesurfer subject
-----	---

Value

Numeric

Examples

```
if (have_fs()) {  
  bert_dir = file.path(fs_subj_dir(), "bert", "surf")  
  file = file.path(bert_dir, "lh.thickness")  
  fid = file(file, open = "rb")  
  out = freesurfer_read3_con(fid)  
}
```

freesurfer_read_curv *Read Freesurfer Curv file*

Description

Reads a Freesurfer curvature file according to the FREESURFER_HOME/matlab/read_curv.m file.

Usage

```
freesurfer_read_curv(file)
```

Arguments

file file name of a curvature file

Value

Numeric vector

Examples

```
if (have_fs()) {  
    bert_dir = file.path(fs_subj_dir(), "bert", "surf")  
    file = file.path(bert_dir, "lh.thickness")  
    fid = file(file, open = "rb")  
    out = freesurfer_read_curv(file)  
}
```

freesurfer_read_surf *Read Freesurfer Surface file*

Description

Reads a Freesurfer Surface file from the surf/ directory from recon-all

Usage

```
freesurfer_read_surf(file)
```

Arguments

file surface file (e.g. lh.inflated)

Value

List of length 2: vertices and faces are the elements

Examples

```

if (have_fs()) {
  fname = file.path(fs_subj_dir(), "bert", "surf", "lh.inflated")
  out = freesurfer_read_surf(fname)
}

```

fs_cmd

*FS Command Wrapper***Description**

This function calls Freesurfer command passed to func

Usage

```

fs_cmd(
  func,
  file,
  outfile = NULL,
  retimg = TRUE,
  reorient = FALSE,
  intern = FALSE,
  opts = "",
  verbose = TRUE,
  samefile = FALSE,
  opts_after_outfile = FALSE,
  frontopts = "",
  add_ext = TRUE,
  bin_app = "bin",
  ...
)

```

Arguments

func	(character) Freesurfer function
file	(character) image to be manipulated
outfile	(character) resultant image name (optional)
retimg	(logical) return image of class nifti
reorient	(logical) If retimg, should file be reoriented when read in? Passed to readnii .
intern	(logical) to be passed to system
opts	(character) operations to be passed to func
verbose	(logical) print out command before running
samefile	(logical) is the output the same file?
opts_after_outfile	(logical) should opts come after the outfile in the Freesurfer command?

frontopts	(character) options/character to put in before filename
add_ext	(logical) should the extension be added to the out file
bin_app	(character) appendix to add to get_fs
...	additional arguments passed to system .

Value

If ret ing then object of class nifti. Otherwise, Result from system command, depends if intern is TRUE or FALSE.

fs_help	<i>Wrapper for getting Freesurfer help</i>
---------	--

Description

This function takes in the function and returns the help from Freesurfer for that function

Usage

```
fs_help(func_name, help.arg = "--help", extra.args = "", ...)
```

Arguments

func_name	Freesurfer function name
help.arg	Argument to print help, usually "--help"
extra.args	Extra arguments to be passed other than --help
...	additional arguments to get_fs

Value

Prints help output and returns output as character vector

Examples

```
if (have_fs()) {
  fs_help(func_name = "mri_watershed")
}
```

`fs_imgext`*Determine extension of image based on FSLOUTPUTTYPE*

Description

Runs `get_fs_output()` to extract `FSLOUTPUTTYPE` and then gets corresponding extension (such as `.nii.gz`)

Usage

```
fs_imgext()
```

Value

Extension for output type

Examples

```
fs_imgext()
```

`fs_lut`*Freesurfer look up table (LUT)*

Description

A `data.frame` with the index, label, and RGBA (red, blue, green, alpha) specification for the segmentations

Usage

```
fs_lut
```

Format

An object of class `data.frame` with 1266 rows and 6 columns.

fs_subj_dir	<i>Determine Freesurfer Subjects Directory</i>
-------------	--

Description

Finds the SUBJECTS_DIR from system environment or `getOption("fs.subj_dir")` for subjects dir

Usage

```
fs_subj_dir()
```

Value

SUBJECTS_DIR, such as `${FREESURFER_HOME}/subjects`

Examples

```
if (have_fs()) {  
  fs_subj_dir()  
}
```

fs_version	<i>Find Freesurfer Version</i>
------------	--------------------------------

Description

Finds the Freesurfer version from `FREESURFER_HOME/build-stamp.txt`

Usage

```
fs_version()
```

Value

If the version file does not exist, it will throw a warning, but it will return an empty string. Otherwise it will be a string of the version.

Note

This will use `fs_dir()` to get the directory of `FREESURFER`

Examples

```
if (have_fs()) {  
  fs_version()  
}
```

get_fs	<i>Create command declaring FREESURFER_HOME</i>
--------	---

Description

Finds the Freesurfer from system environment or `getOption("freesurfer.path")` for location of Freesurfer functions

Usage

```
get_fs(bin_app = c("bin", "mni/bin", ""))
```

Arguments

bin_app	Should bin be added to the freesurfer path? All executables are assumed to be in FREESURFER_HOME/bin/. If not, and bin_app = "", they will be assumed to be in FREESURFER_HOME/.
---------	--

Value

NULL if Freesurfer in path, or bash code for setting up Freesurfer DIR

Note

This will use `Sys.getenv("FREESURFER_HOME")` before `getOption("freesurfer.path")`. If the directory is not found for Freesurfer in `Sys.getenv("FreesurferDIR")` and `getOption("freesurfer.path")`, it will try the default directory `/usr/local/freesurfer`.

Examples

```
if (have_fs()) {
  get_fs()
}
```

get_fs_output	<i>Determine Freesurfer output type</i>
---------------	---

Description

Finds the FSF_OUTPUT_FORMAT from system environment or `getOption("fs.outputtype")` for output type (nii.gz, nii, ANALYZE,etc)

Usage

```
get_fs_output()
```

Value

FSF_OUTPUT_FORMAT, such as nii.gz If none found, uses nii.gz as default

Examples

```
get_fs_output()
```

have_fs	<i>Logical check if Freesurfer is accessible</i>
---------	--

Description

Uses `get_fs` to check if `FreesurferDIR` is accessible or the option `freesurfer.path` is set and returns logical

Usage

```
have_fs(..., check_license = FALSE)
```

Arguments

... options to pass to `get_fs`
 check_license Should a license file be checked to exist?

Value

Logical TRUE is Freesurfer is accessible, FALSE if not

Examples

```
have_fs()
```

mnc2nii	<i>Convert MNC to NIFTI</i>
---------	-----------------------------

Description

This function calls `mnc2nii` to convert MNC files to NIFTI

Usage

```
mnc2nii(file, outfile = NULL)
```

Arguments

file (character) input filename
 outfile (character) output filename

Value

Character filename of output

Examples

```
if (have_fs() && requireNamespace("oro.nifti", quietly = TRUE)) {
  img = oro.nifti::nifti(array(rnorm(5*5*5), dim = c(5,5,5)))
  mnc = nii2mnc(img)
  img_file = mnc2nii(mnc, outfile = tempfile(fileext = ".nii"))
  neurobase::readnii(img_file, verbose = TRUE)
}
```

mnc2nii.help

MNC to NIFTI Help

Description

This calls Freesurfer's mnc2nii help

Usage

```
mnc2nii.help()
```

Value

Result of fs_help

mris_convert

Use Freesurfer's MRIs Converter

Description

This function call mris_convert, a general conversion program for converting between cortical surface file formats

Usage

```
mris_convert(infile, outfile = NULL, ext = ".asc", opts = "", verbose = TRUE)
```

Arguments

infile	(character) file path for input file
outfile	(character) output file path
ext	(character) output file extension, default is set to .asc
opts	(character) additional options to add to front of command
verbose	(logical) print diagnostic messages

Value

Name of output file

Examples

```
if (have_fs()) {  
  bert_surf_dir = file.path(fs_subj_dir(), "bert", "surf")  
  asc_file = mris_convert(  
    infile = file.path(bert_surf_dir, "lh.white")  
  )  
}
```

mris_convert.help

Help file for Freesurfer MRIs Converter

Description

This calls Freesurfer's `mris_convert help`

Usage

```
mris_convert.help()
```

Value

Result of `fs_help`

mris_convert_annot *Convert Annotation file*

Description

This function call `mris_convert`, using the `--annot` option

Usage

```
mris_convert_annot(annot, opts = "", ...)
```

Arguments

annot	(character) annotation or gifti label data
opts	(character) additional options to <code>mris_convert</code>
...	additional arguments to <code>mris_convert</code>

Value

Result of `mris_convert`

Examples

```
if (have_fs()) {  
  bert_dir = file.path(fs_subj_dir(), "bert")  
  gii_file = mris_convert_annot(  
    infile = file.path(bert_dir, "surf", "lh.white"),  
    annot = file.path(bert_dir, "label", "lh.aparc.annot"),  
    ext = ".gii"  
  )  
  gii = mris_convert_annot(  
    infile = file.path(bert_dir, "surf", "lh.white"),  
    annot = gii_file,  
    ext = ".gii"  
  )  
}
```

mris_convert_curv *Convert Curvature file*

Description

This function call `mris_convert`, using the `-c` option

Usage

```
mris_convert_curv(curv, opts = "", ...)
```

Arguments

curv (character) scalar curv overlay file
 opts (character) additional options to `mris_convert`
 ... additional arguments to `mris_convert`

Value

Result of `mris_convert`

Note

The filename of the output may change due to how Freesurfer does curvature conversions and you may need to paste the prefix to get the correct filename, as seen in the example.

Examples

```

if (have_fs()) {
  bert_surf_dir = file.path(fs_subj_dir(), "bert", "surf")
  asc_file = mris_convert_curv(
    infile = file.path(bert_surf_dir, "lh.white"),
    curv = file.path(bert_surf_dir, "lh.thickness")
  )
  if (!file.exists(asc_file)) {
    asc_file = file.path(dirname(asc_file), paste0("lh.",
      basename(asc_file)))
  }
  res = read_fs_table(asc_file, header = FALSE)
  colnames(res) = c("index", "coord_1", "coord_2", "coord_3", "value")
  head(res)
}

```

`mris_convert_normals` *Convert Surface to Surface normals*

Description

This function call `mris_convert`, using the `-n` option

Usage

```
mris_convert_normals(opts = "", ...)
```

Arguments

opts (character) additional options to `mris_convert`
 ... additional arguments to `mris_convert`

Value

Result of [mris_convert](#)

Examples

```
if (have_fs()) {
  bert_dir = file.path(fs_subj_dir(), "bert")
  asc_file = mris_convert_normals(
    infile = file.path(bert_dir, "surf", "lh.white")
  )
  readLines(asc_file, n = 6)
}
```

mris_convert_vertex *Convert Surface to vertex file*

Description

This function call [mris_convert](#), using the `-v` option

Usage

```
mris_convert_vertex(opts = "", ...)
```

Arguments

opts (character) additional options to [mris_convert](#)
... additional arguments to [mris_convert](#)

Value

Result of [mris_convert](#)

Examples

```
if (have_fs()) {
  bert_surf_dir = file.path(fs_subj_dir(), "bert", "surf")
  asc_file = mris_convert_vertex(
    infile = file.path(bert_surf_dir, "lh.white")
  )
  readLines(asc_file, n = 6)
}
```

mris_euler_number *MRI Euler Number*

Description

This function calls mris_euler_number to calculate the Euler Number

Usage

```
mris_euler_number(file, outfile = NULL, opts = "")
```

Arguments

file	(character) input filename
outfile	(character) output filename
opts	(character) additional options to mris_euler_number

Value

Result of system command

Examples

```
## Not run:
if (have_fs() && requireNamespace("oro.nifti", quietly = TRUE)) {
  img = oro.nifti::nifti(array(rnorm(5*5*5), dim = c(5,5,5)))
  res = mris_euler_number(img, outfile = tempfile(fileext = ".mgz"))
}

## End(Not run)
```

mris_euler_number.help

MRI Euler Number Help

Description

This calls Freesurfer's mris_euler_number help

Usage

```
mris_euler_number.help()
```

Value

Result of fs_help

`mri_convert`*Use Freesurfer's MRI Converter*

Description

This function calls `mri_convert` to convert an image

Usage

```
mri_convert(file, outfile, opts = "")
```

Arguments

<code>file</code>	(character) input filename
<code>outfile</code>	(character) output filename
<code>opts</code>	(character) additional options to <code>mri_convert</code>

Value

Result of system command

Examples

```
if (have_fs() && requireNamespace("oro.nifti", quietly = TRUE)) {  
  img = oro.nifti::nifti(array(rnorm(5*5*5), dim = c(5,5,5)))  
  res = mri_convert(img, outfile = tempfile(fileext = ".mgz"))  
}
```

`mri_convert.help`*MRI Normalize Help*

Description

This calls Freesurfer's `mri_convert help`

Usage

```
mri_convert.help()
```

Value

Result of `fs_help`

mri_deface

MRI Deface

Description

This calls Freesurfer's `mri_deface`

Usage

```
mri_deface(file, brain_template = NULL, face_template = NULL, ...)
```

Arguments

<code>file</code>	File to pass to <code>mri_deface</code>
<code>brain_template</code>	gca brain template file to pass to <code>mri_deface</code>
<code>face_template</code>	gca face template file to pass to <code>mri_deface</code>
<code>...</code>	Additional arguments to pass to <code>fs_cmd</code>

Value

Result of `fs_cmd`, which type depends on arguments to `...`

Note

If `brain_template` or `face_template` is `NULL`, they will be downloaded.

Examples

```
if (have_fs()){
  base_url = "https://surfer.nmr.mgh.harvard.edu/pub/dist/mri_deface"
  url = file.path(base_url, "sample_T1_input.mgz")
  x = tempfile(fileext = ".mgz")
  out = try({
    utils::download.file(url, destfile = x)
  })
  if (!inherits(out, "try-error")) {
    noface = mri_deface(x)
  } else {
    url = paste0(
      "https://raw.githubusercontent.com/muschellij2/kirby21.t1/master/",
      "inst/visit_1/113/113-01-T1.nii.gz")
    x = tempfile(fileext = ".nii.gz")
    out = try({
      utils::download.file(url, destfile = x)
    })
    noface = mri_deface(x)
  }
}
```

mri_info	<i>MRI information</i>
----------	------------------------

Description

This calls Freesurfer's `mri_info`

Usage

```
mri_info(file, ...)
```

Arguments

file	File to pass to <code>mri_info</code>
...	Additional arguments to pass to <code>fs_cmd</code>

Value

Result of `fs_cmd`, which type depends on arguments to ...

Examples

```
if (have_fs() && requireNamespace("oro.nifti", quietly = TRUE)){
  img = oro.nifti::nifti(array(rnorm(5*5*5), dim = c(5,5,5)))
  mri_info(img)
}
```

mri_info.help	<i>MRI information Help</i>
---------------	-----------------------------

Description

This calls Freesurfer's `mri_info help`

Usage

```
mri_info.help()
```

Value

Result of `fs_help`

mri_mask	<i>Use Freesurfer's MRI Mask</i>
----------	----------------------------------

Description

This function calls `mri_mask` to mask an image

Usage

```
mri_mask(file, mask, outfile = NULL, retimg = TRUE, opts = "", ...)
```

Arguments

<code>file</code>	(character) input filename
<code>mask</code>	(character) mask filename
<code>outfile</code>	(character) output filename
<code>retimg</code>	(logical) return image of class nifti
<code>opts</code>	(character) additional options to <code>mri_mask</code>
<code>...</code>	additional arguments passed to <code>fs_cmd</code> .

Value

Character or nifti depending on `retimg`

Examples

```
if (have_fs() && requireNamespace("oro.nifti", quietly = TRUE)) {
  img = oro.nifti::nifti(array(rnorm(5*5*5), dim = c(5,5,5)))
  mask = img > 1
  res = mri_mask(img, mask)
}
```

mri_mask.help	<i>MRI Normalize Help</i>
---------------	---------------------------

Description

This calls Freesurfer's `mri_mask` help

Usage

```
mri_mask.help()
```

Value

Result of `fs_help`

mri_normalize	<i>Use Freesurfer's MRI Normalize Algorithm</i>
---------------	---

Description

This function calls `mri_normalize` to normalize the values of the image, with white matter voxels around 110.

Usage

```
mri_normalize(file, outfile = NULL, retimg = TRUE, opts = "", ...)
```

Arguments

<code>file</code>	(character) input filename
<code>outfile</code>	(character) output filename
<code>retimg</code>	(logical) return image of class nifti
<code>opts</code>	(character) additional options to <code>mri_normalize</code>
<code>...</code>	additional arguments passed to <code>fs_cmd</code> .

Value

Character or nifti depending on `retimg`

Examples

```
## Not run:
if (have_fs()){
  mri_normalize("/path/to/T1.nii.gz")
}

## End(Not run)
```

mri_normalize.help	<i>MRI Normalize Help</i>
--------------------	---------------------------

Description

This calls Freesurfer's `mri_normalize help`

Usage

```
mri_normalize.help()
```

Value

Result of `fs_help`

mri_segment

Use Freesurfer's MRI Segmentation Algorithm

Description

This function calls `mri_segment` to segment tissues from an image

Usage

```
mri_segment(file, outfile = NULL, retimg = TRUE, opts = "", ...)
```

Arguments

<code>file</code>	(character) input filename
<code>outfile</code>	(character) output filename
<code>retimg</code>	(logical) return image of class nifti
<code>opts</code>	(character) additional options to <code>mri_segment</code>
<code>...</code>	additional arguments passed to <code>fs_cmd</code> .

Value

Character or nifti depending on `retimg`

Note

NOT COMPLETE

mri_segment.help

MRI Segment Help

Description

This calls Freesurfer's `mri_segment help`

Usage

```
mri_segment.help()
```

Value

Result of `fs_help`

mri_surf2surf	<i>Use Freesurfer's mri_surf2surf function to resamples one cortical surface onto another</i>
---------------	---

Description

This function calls Freesurfer `mri_surf2surf` to resample one cortical surface onto another

Usage

```
mri_surf2surf(
  subject = NULL,
  target_subject = NULL,
  trg_type = c("curv", "w", "mgh", "nii"),
  src_type = c("curv", "w"),
  outfile = NULL,
  hemi = c("lh", "rh"),
  sval = c("thickness"),
  subj_dir = NULL,
  opts = "",
  verbose = TRUE
)
```

Arguments

<code>subject</code>	(character) vector of subject name
<code>target_subject</code>	(character) vector of target subject name
<code>trg_type</code>	(character) target file type, can be curv, paint (w), mgh, or nii
<code>src_type</code>	(character) source file type, can be curv or paint (w)
<code>outfile</code>	(character) output filename
<code>hemi</code>	(character) hemisphere to run statistics
<code>sval</code>	(character) source file
<code>subj_dir</code>	(character path) if a different subjects directory is to be used other than SUBJECTS_DIR from shell, it can be specified here. Use with care as if the command fail, it may not reset the SUBJECTS_DIR back correctly after the error
<code>opts</code>	(character) additional options to <code>mri_surf2surf</code>
<code>verbose</code>	(logical) print diagnostic messages

Value

Name of output file

Examples

```

if (have_fs()) {
  out = mri_surf2surf(
    subject = 'bert',
    target_subject = 'fsaverage',
    trg_type = 'curv',
    src_type = 'curv',
    hemi = "rh",
    sval = "thickness")
}

```

mri_surf2surf.help *Freesurfer's mri_surf2surf Help*

Description

This calls Freesurfer's `mri_surf2surf help`

Usage

```
mri_surf2surf.help()
```

Value

Result of `fs_help`

mri_watershed *Use Freesurfer's MRI Watershed Algorithm*

Description

This function calls `mri_watershed` to extract a brain from an image, usually for skull stripping.

Usage

```
mri_watershed(file, outfile = NULL, retimg = TRUE, opts = "", ...)
```

Arguments

<code>file</code>	(character) input filename
<code>outfile</code>	(character) output filename
<code>retimg</code>	(logical) return image of class <code>nifti</code>
<code>opts</code>	(character) additional options to <code>mri_watershed</code>
<code>...</code>	additional arguments passed to <code>fs_cmd</code> .

Value

Character or nifti depending on retimg

Examples

```
## Not run:
if (have_fs()){
  mri_watershed("/path/to/T1.nii.gz")
}

## End(Not run)
```

mri_watershed.help *MRI Watershed Help*

Description

This calls Freesurfer's mri_watershed help

Usage

```
mri_watershed.help()
```

Value

Result of fs_help

nii2mnc *Convert NIfTI to MNC*

Description

This function calls nii2mnc to convert NIfTI to MNC files

Usage

```
nii2mnc(file, outfile = NULL)
```

Arguments

file (character) input filename
 outfile (character) output filename

Value

Character filename of output

Examples

```

if (have_fs() && requireNamespace("oro.nifti", quietly = TRUE)) {
  img = oro.nifti::nifti(array(rnorm(5*5*5), dim = c(5,5,5)))
  mnc = nii2mnc(img)
  img_file = mnc2nii(mnc)
}

```

nii2mnc.help

Convert NIFTI to MNC Help

Description

This calls Freesurfer's `mnc2nii help`

Usage

```
nii2mnc.help()
```

Value

Result of `fs_help`

nu_correct

Use Freesurfer's Non-Uniformity Correction

Description

This function calls `nu_correct` to correct for non-uniformity

Usage

```
nu_correct(file, mask = NULL, opts = "", verbose = TRUE, ...)
```

Arguments

<code>file</code>	(character) input filename
<code>mask</code>	(character or nifti) Mask to use for correction.
<code>opts</code>	(character) additional options to <code>mri_segment</code>
<code>verbose</code>	print diagnostic messages
<code>...</code>	additional arguments passed to <code>fs_cmd</code> .

Value

Object of class `nifti` depending on `retimg`

Examples

```
## Not run:
if (have_fs()){
  nu_correct("/path/to/T1.nii.gz")
}

## End(Not run)
```

nu_correct.help

Non-Uniformity Correction Help

Description

This calls Freesurfer's nu_correct help

Usage

```
nu_correct.help()
```

Value

Result of fs_help

readmgz

Read MGH or MGZ File

Description

This function calls `mri_convert` to convert MGH/MGZ files to NIfTI, then reads it in using [readnii](#)

Usage

```
readmgz(file)
```

```
readmgh(file)
```

Arguments

file (character) input filename

Value

Object of class `nifti`

readmnc	<i>Read MNC File</i>
---------	----------------------

Description

This function calls `mnc2nii` to convert MNC files to NIFTI, then reads it in using `readnii`

Usage

```
readmnc(file)
```

Arguments

file (character) input filename

Value

Object of class `nifti`

read_annotation	<i>Read Freesurfer annotation file</i>
-----------------	--

Description

Reads Freesurfer binary annotation files that contain information on vertex labels and colours for use in analyses and brain area lookups.

Usage

```
read_annotation(path, verbose = TRUE)
```

Arguments

path path to annotation file, usually with extension `annot`
 verbose logical.

Details

This function is heavily based on Freesurfer's `read_annotation.m` Original Author: Bruce Fischl
 CVS Revision Info: \$Author: greve \$ \$Date: 2014/02/25 19:54:10 \$ \$Revision: 1.10 \$

Value

list of 3 with vertices, labels, and colortable

Examples

```

if (have_fs()) {
  bert_dir = file.path(fs_subj_dir(), "bert")
  annot_file = file.path(bert_dir, "label", "lh.aparc.annot")
  res = read_annotation(annot_file)
}

```

read_aseg_stats	<i>Read Anatomical Segmentation Statistics</i>
-----------------	--

Description

Reads an aseg.stats file from an individual subject

Usage

```
read_aseg_stats(file)
```

Arguments

file aseg.stats file from Freesurfer

Value

List of 2 data.frames, one with the global measures and one with the structure-specific measures.

Examples

```

if (have_fs()) {
  file = file.path(fs_subj_dir(), "bert", "stats", "aseg.stats")
  out = read_aseg_stats(file)
}

```

read_fs_label	<i>Read Label File</i>
---------------	------------------------

Description

Reads an label file from an individual subject

Usage

```
read_fs_label(file)
```

Arguments

file label file from Freesurfer

Value

data.frame with 5 columns:

vertex_num: Vertex Number
 r_coord: Coordinate in RL direction
 a_coord: Coordinate in AP direction
 s_coord: Coordinate in SI direction
 value: Value of label (depends on file)

Examples

```
if (have_fs()) {
  file = file.path(fs_subj_dir(), "bert", "label", "lh.BA1.label")
  if (!file.exists(file)) {
    file = file.path(fs_subj_dir(), "bert", "label", "lh.BA1_exvivo.label")
  }
  out = read_fs_label(file)
}
```

read_fs_table

Read Freesurfer Table Output

Description

This function reads output from a Freesurfer table command, e.g. `aparcstats2table`, `asegstats2table`

Usage

```
read_fs_table(file, sep = NULL, stringsAsFactors = FALSE, header = TRUE, ...)
```

Arguments

file (character path) filename of text file
 sep separator to override attribute of file, to pass to [read.table](#).
 stringsAsFactors (logical) passed to [read.table](#)
 header Is there a header in the data
 ... additional arguments to [read.table](#)

Value

data.frame from the file

Examples

```

if (have_fs()) {
  outfile = aparstats2table(subjects = "bert",
                           hemi = "lh",
                           meas = "thickness")
  df = read_fs_table(outfile)
  seg_outfile = asegstats2table(subjects = "bert", meas = "mean")
  df_seg = read_fs_table(seg_outfile)
}
## Not run:
### using the pipe
if (requireNamespace("magrittr", quietly = TRUE)) {
  df_seg = asegstats2table(subjects = "bert", meas = "mean") %>%
    read_fs_table
}

## End(Not run)

```

recon

Reconstruction from Freesurfer

Description

Reconstruction from Freesurfer with most of the options implemented.

Usage

```

recon(
  infile,
  outdir = NULL,
  subjid,
  motioncor = TRUE,
  nuintensitycor = TRUE,
  talairach = TRUE,
  normalization = TRUE,
  skullstrip = TRUE,
  gcareg = TRUE,
  canorm = TRUE,
  careg = TRUE,
  rmneck = TRUE,
  skull_lta = TRUE,
  calabel = TRUE,
  normalization2 = TRUE,
  segmentation = TRUE,
  fill = TRUE,
  tessellate = TRUE,
  smooth1 = TRUE,
  inflate1 = TRUE,

```

```

qsphere = TRUE,
fix = TRUE,
finalsurfs = TRUE,
smooth2 = TRUE,
inflate2 = TRUE,
cortribbon = TRUE,
sphere = TRUE,
surfreg = TRUE,
contrasurfreg = TRUE,
avgcurv = TRUE,
cortparc = TRUE,
parcstats = TRUE,
cortparc2 = TRUE,
parcstats2 = TRUE,
aparc2aseg = TRUE,
verbose = TRUE,
opts = ""
)

```

Arguments

<code>infile</code>	Input filename (dcm or nii)
<code>outdir</code>	Output directory
<code>subjid</code>	subject id
<code>motioncor</code>	When there are multiple source volumes, this step will correct for small motions between them and then average them together. The input are the volumes found in file(s) <code>mri/orig/XXX.mgz</code> . The output will be the volume <code>mri/orig.mgz</code> . If no runs are found, then it looks for a volume in <code>mri/orig</code> (or <code>mri/orig.mgz</code>). If that volume is there, then it is used in subsequent processes as if it was the motion corrected volume. If no volume is found, then the process exits with errors.
<code>nuintensitycor</code>	Non-parametric Non-uniform intensity Normalization (N3), corrects for intensity non-uniformity in MR data, making relatively few assumptions about the data. This runs the MINC tool <code>'nu_correct'</code> . By default, four iterations of <code>nu_correct</code> are run. The flag <code>'-nuiterations'</code> specification of some other number of iterations.
<code>talairach</code>	computes the affine transform from the orig volume to the MNI305 atlas using the MINC program <code>mritotal</code> . Creates the files <code>mri/transform/talairach.auto.xfm</code> and <code>talairach.xfm</code> .
<code>normalization</code>	Performs intensity normalization of the orig volume and places the result in <code>mri/T1.mgz</code>
<code>skullstrip</code>	Removes the skull from <code>mri/T1.mgz</code> and stores the result in <code>mri/brainmask.auto.mgz</code> and <code>mri/brainmask.mgz</code> . Runs the <code>mri_watershed</code> program.
<code>gcareg</code>	Computes transform to align the <code>mri/nu.mgz</code> volume to the default GCA atlas found in <code>FREESURFER_HOME/average</code> . Creates the file <code>mri/transforms/talairach.lta</code> .
<code>canorm</code>	Further normalization, based on GCA model. Creates <code>mri/norm.mgz</code> .
<code>careg</code>	Computes a nonlinear transform to align with GCA atlas. Creates the file <code>mri/transform/talairach.m3z</code> .

rmneck	The neck region is removed from the NU-corrected volume <code>mri/nu.mgz</code> . Makes use of transform computed from prior CA Register stage. Creates the file <code>mri/nu_noneck.mgz</code> .
skull_lta	Computes transform to align volume <code>mri/nu_noneck.mgz</code> with GCA volume possessing the skull. Creates the file <code>mri/transforms/talairach_with_skull.lta</code> .
calabel	Labels subcortical structures, based in GCA model. Creates the files <code>mri/aseg.auto.mgz</code> and <code>mri/aseg.mgz</code> .
normalization2	Performs a second (major) intensity correction using only the brain volume as the input (so that it has to be done after the skull strip). Intensity normalization works better when the skull has been removed. Creates a new <code>brain.mgz</code> volume. If <code>-noaseg</code> flag is used, then <code>aseg.mgz</code> is not used by <code>mri_normalize</code> .
segmentation	Attempts to separate white matter from everything else. The input is <code>mri/brain.mgz</code> , and the output is <code>mri/wm.mgz</code> . Uses intensity, neighborhood, and smoothness constraints. This is the volume that is edited when manually fixing defects. Calls <code>mri_segment</code> , <code>mri_edit_wm_with_aseg</code> , and <code>mri_preless</code> . To keep previous edits, run with <code>-keepwmedits</code> . If <code>-noaseg</code> is used, then <code>mri_edit_wm_aseg</code> is skipped.
fill	This creates the subcortical mass from which the orig surface is created. The mid brain is cut from the cerebrum, and the hemispheres are cut from each other. The left hemisphere is binarized to 255. The right hemisphere is binarized to 127. The input is <code>mri/wm.mgz</code> and the output is <code>mri/filled.mgz</code> . Calls <code>mri_fill</code> . If the cut fails, then seed points can be supplied (see <code>-cc-crs</code> , <code>-pons-crs</code> , <code>-lh-crs</code> , <code>-rh-crs</code>). The actual points used for the cutting planes in the corpus callosum and pons can be found in <code>scripts/ponscut.cut.log</code> . This is the last stage of volumetric processing. If <code>-noaseg</code> is used, then <code>aseg.mgz</code> is not used by <code>mri_fill</code> .
tessellate	This is the step where the orig surface (ie, <code>surf/?h.orig.nofix</code>) is created. The surface is created by covering the filled hemisphere with triangles. Runs <code>mri_tessellate</code> . The places where the points of the triangles meet are called vertices. Creates the file <code>surf/?h.orig.nofix</code> Note: the topology fixer will create the surface <code>?h.orig</code> .
smooth1	Calls <code>mris_smooth</code> . Smooth1 is the step just after tessellation
inflate1	Inflation of the <code>surf/?h.smoothwm(.nofix)</code> surface to create <code>surf/?h.inflated</code> .
qsphere	automatic topology fixing. It is a quasi-homeomorphic spherical transformation of the inflated surface designed to localize topological defects for the subsequent automatic topology fixer.
fix	Finds topological defects (ie, holes in a filled hemisphere) using <code>surf/?h.qsphere.nofix</code> , and changes the orig surface (<code>surf/?h.orig.nofix</code>) to remove the defects. Changes the number of vertices. All the defects will be removed, but the user should check the orig surface in the volume to make sure that it looks appropriate. Calls <code>mris_fix_topology</code> .
finalsurfs	Creates the <code>?h.white</code> and <code>?h.pial</code> surfaces as well as the thickness file (<code>?h.thickness</code>) and curvature file (<code>?h.curv</code>). The white surface is created by "nudging" the orig surface so that it closely follows the white-gray intensity gradient as found in the T1 volume. The pial surface is created by expanding the white surface so that it closely follows the gray-CSF intensity gradient as found in the T1 volume. Calls <code>mris_make_surfaces</code> .
smooth2	the step just after topology fixing.

inflate2	inflate2 is the step just after topology fixing
cortribbon	Creates binary volume masks of the cortical ribbon, ie, each voxel is either a 1 or 0 depending upon whether it falls in the ribbon or not. Saved as ?h.ribbon.mgz. Uses mgz regardless of whether the -mgz option is used.
sphere	Inflates the orig surface into a sphere while minimizing metric distortion. This step is necessary in order to register the surface to the spherical atlas. (also known as the spherical morph). Calls mris_sphere. Creates surf/?h.sphere.
surfreg	Registers the orig surface to the spherical atlas through surf/?h.sphere. The surfaces are first coarsely registered by aligning the large scale folding patterns found in ?h.sulc and then fine tuned using the small-scale patterns as in ?h.curv. Calls mris_register. Creates surf/?h.sphere.reg.
contrasurfreg	Same as ipsilateral but registers to the contralateral atlas. Creates lh.rh.sphere.reg and rh.lh.sphere.reg.
avgcurv	Resamples the average curvature from the atlas to that of the subject. Allows the user to display activity on the surface of an individual with the folding pattern (ie, anatomy) of a group. Calls mris_paint. Creates surf/?h.avg_curv.
cortparc	Assigns a neuroanatomical label to each location on the cortical surface. Incorporates both geometric information derived from the cortical model (sulcus and curvature), and neuroanatomical convention. Calls mris_ca_label. -cortparc creates label/?h.aparc.annot, and -cortparc2 creates /label/?h.aparc.a2005s.annot.
parcstats	Runs mris_anatomical_stats to create a summary table of cortical parcellation statistics for each structure, including 1. structure name 2. number of vertices 3. total surface area (mm ²) 4. total gray matter volume (mm ³) 5. average cortical thickness (mm) 6. standard error of cortical thickness (mm) 7. integrated rectified mean curvature 8. integrated rectified Gaussian curvature 9. folding index 10. intrinsic curvature index. For -parcstats, the file is saved in stats/?h.aparc.stats. For -parcstats2, the file is saved in stats/?h.aparc.a2005s.stats.
cortparc2	see cortparc argument
parcstats2	see cortparc2 argument
aparc2aseg	Maps the cortical labels from the automatic cortical parcellation (aparc) to the automatic segmentation volume (aseg). The result can be used as the aseg would.
verbose	print diagnostic messages
opts	Additional options

Value

Result of `system`

reconner *Reconstruction Helper for recon from Freesurfer*

Description

Wrapper for the recon-all function in Freesurfer

Usage

```
reconner(  
  infile = NULL,  
  outdir = NULL,  
  subjid = NULL,  
  verbose = TRUE,  
  opts = "-all",  
  force = FALSE  
)
```

Arguments

infile	Input filename (dcm or nii)
outdir	Output directory
subjid	subject id
verbose	print diagnostic messages
opts	Additional options
force	Force running of the reconstruction

Value

Result of `system`

Note

If you set `infile = NULL`, then you can omit the `-i` flag in `recon-all`

recon_all	<i>Reconstruction from Freesurfer for All Steps</i>
-----------	---

Description

Reconstruction from Freesurfer for All Steps

Usage

```
recon_all(  
  infile = NULL,  
  outdir = NULL,  
  subjid = NULL,  
  verbose = TRUE,  
  opts = "-all",  
  ...  
)
```

Arguments

infile	Input filename (dcm or nii)
outdir	Output directory
subjid	subject id
verbose	print diagnostic messages
opts	Additional options
...	arguments passed to reconner

Value

Result of [system](#)

Note

If you would like to restart a recon-all run, change opts so that opts = "-make all"

recon_con1	<i>Reconstruction from Motion Correction to Skull Strip</i>
------------	---

Description

Reconstruction from Freesurfer for Step 1-5 (Motion Correction to Skull Strip), which calls -autorecon1 in recon-all

Usage

```
recon_con1(infile, outdir = NULL, subjid, verbose = TRUE)
autorecon1(infile, outdir = NULL, subjid, verbose = TRUE)
recon_con2(infile, outdir = NULL, subjid, verbose = TRUE)
autorecon2(infile, outdir = NULL, subjid, verbose = TRUE)
recon_con3(infile, outdir = NULL, subjid, verbose = TRUE)
autorecon3(infile, outdir = NULL, subjid, verbose = TRUE)
```

Arguments

infile	Input filename (dcm or nii)
outdir	Output directory
subjid	subject id
verbose	print diagnostic messages

Value

Result of `system`

Note

See <https://surfer.nmr.mgh.harvard.edu/fswiki/recon-all> for the steps of each autorecon1-3. If you set `infile = NULL`, then you can omit the `-i` flag in recon-all.

run_check_fs_cmd *Run and Check a Freesurfer Command*

Description

Checks whether an output filename exists before a command has run, prints and runs the command, and then checks the output from the result.

Usage

```
run_check_fs_cmd(cmd, outfile, verbose = TRUE)
```

Arguments

cmd	Command to be run
outfile	Output file to be produced
verbose	print diagnostic messages

Value

Invisible NULL

See Also

[check_fs_result](#)

set_fs_subj_dir *Set Freesurfer Subjects Directory*

Description

Sets the SUBJECTS_DIR variable in the system environment or options("fs.subj_dir" = x)

Usage

```
set_fs_subj_dir(x = file.path(fs_dir(), "subjects"))
```

Arguments

x	path to SUBJECTS_DIR defaults to file.path(fs_dir(), "subjects")
---	--

Value

No return value, called for side effects ('SUBJECTS_DIR' environment variable set, and 'fs.subj_dir' option set)

surface_to_obj	<i>Convert Freesurfer Surface to Wavefront OBJ</i>
----------------	--

Description

Reads in a surface file from Freesurfer and converts it to a Wavefront OBJ file

Usage

```
surface_to_obj(infile, outfile = NULL, ...)
```

Arguments

infile	Input surface file
outfile	output Wavefront OBJ file. If NULL, a temporary file will be created
...	additional arguments to pass to convert_surface

Value

Character filename of output file

Examples

```
if (have_fs()) {  
  infile = file.path(fs_subj_dir(),  
                    "bert", "surf", "rh.pial")  
  res = surface_to_obj(infile = infile)  
}
```

surface_to_triangles	<i>Convert Freesurfer Surface to Triangles</i>
----------------------	--

Description

Reads in a surface file from Freesurfer and converts it into triangles

Usage

```
surface_to_triangles(infile, ...)
```

Arguments

infile	Input surface file
...	additional arguments to pass to convert_surface

Value

Matrix of triangles with the number of rows equal to the number of faces (not the triplets - total faces)

Examples

```

if (have_fs()) {
infile = file.path(fs_subj_dir(),
                  "bert", "surf", "rh.pial")
right_triangles = surface_to_triangles(infile = infile)
infile = file.path(fs_subj_dir(),
                  "bert", "surf", "lh.pial")
left_triangles = surface_to_triangles(infile = infile)
if (requireNamespace("rgl", quietly = TRUE)) {
  rgl::open3d()
  rgl::triangles3d(right_triangles,
                  color = rainbow(nrow(right_triangles)))
  rgl::triangles3d(left_triangles,
                  color = rainbow(nrow(left_triangles)))
}
infile = file.path(fs_subj_dir(),
                  "bert", "surf", "rh.inflated")
right_triangles = surface_to_triangles(infile = infile)
infile = file.path(fs_subj_dir(),
                  "bert", "surf", "lh.inflated")
left_triangles = surface_to_triangles(infile = infile)
if (requireNamespace("rgl", quietly = TRUE)) {
  rgl::open3d()
  rgl::triangles3d(left_triangles,
                  color = rainbow(nrow(left_triangles)))
  rgl::triangles3d(right_triangles,
                  color = rainbow(nrow(right_triangles)))
}
}

```

surf_convert

Convert Surface Data to ASCII

Description

This function calls `mri_convert` to convert a measure from surfaces to an ASCII file and reads it in.

Usage

```
surf_convert(file, outfile = NULL)
```


Arguments

file (character) input filename of curvature measure
outfile (character) output filename (if wanted to be saved)

Value

data.frame

Examples

```
if (have_fs()) {  
  fname = file.path(fs_subj_dir(), "bert", "surf", "lh.thickness")  
  out = surf_convert(fname)  
}
```

tracker

Tract Reconstruction Helper for trac-all from Freesurfer

Description

Wrapper for the trac-all function in Freesurfer

Usage

```
tracker(infile, outdir = NULL, subjid, verbose = TRUE, opts = "")
```

Arguments

infile Input filename (dcm or nii)
outdir Output directory
subjid subject id, if NULL, the basename of the infile will be used
verbose print diagnostic messages
opts Additional options

Value

Result of `system`

trac_all	<i>Tract Reconstruction Helper for trac-all from Freesurfer for All Steps</i>
----------	---

Description

Wrapper for the trac-all function in Freesurfer for All Steps

Usage

```
trac_all(infile, outdir = NULL, subjid, verbose = TRUE, opts = "")
```

Arguments

infile	Input filename (dcm or nii)
outdir	Output directory
subjid	subject id
verbose	print diagnostic messages
opts	Additional options

Value

Result of `system`

trac_prep	<i>Tract Reconstruction for Each Step</i>
-----------	---

Description

Reconstruction from Freesurfer for Preprocessing, Bedpost, and Path reconstruction

Usage

```
trac_prep(infile, outdir = NULL, subjid, verbose = TRUE)
```

```
trac_bedpost(infile, outdir = NULL, subjid, verbose = TRUE)
```

```
trac_path(infile, outdir = NULL, subjid, verbose = TRUE)
```

Arguments

infile	Input filename (dcm or nii)
outdir	Output directory
subjid	subject id
verbose	print diagnostic messages

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Value

Result of [system](#)

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