freesurfer: Connecting the Freesurfer Software with R

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Abstract We present the package **freesurfer**, a set of R functions that interface with Freesurfer, a commonly-used open-source software package for processing and analyzing structural neuroimaging data. The **freesurfer** package performs operations on nifti image objects in R using command-line functions from Freesurfer, and returns R objects back to the user. **freesurfer** allows users to process neuroanatomical images and provides functionality to convert and read the output of the Freesurfer pipelines. We present an example of the analysis of structural magnetic resonance images, which demonstrates how R users can interface with Freesurfer and analyze the results of the complete Freesurfer analysis pipeline.

Abstract

```
#' if (have_fs()) {
      df = aparcs_to_bg(subjects = "bert", measure = "thickness")
      print(head(df))
#' if (have_fs()){
      img = oro.nifti::nifti(array(rnorm(5*5*5), dim = c(5,5,5)))
      mri_info(img)
#'}
# \dontrun{
#' if (have_fs()){
       mri_watershed("/path/to/T1.nii.gz")
#'}
#'}
#' @examples \dontrun{
#' if (have_fs()){
#'
       mri_normalize("/path/to/T1.nii.gz")
#' }
#'
  #' if (have_fs()) {
      img = oro.nifti::nifti(array(rnorm(5*5*5), dim = c(5,5,5)))
#'
      res = mri_convert(img, outfile = tempfile(fileext = ".mgz"))
#'}
  #' if (have_fs()) {
      img = oro.nifti::nifti(array(rnorm(5*5*5), dim = c(5,5,5)))
      mnc = nii2mnc(img)
#'
      img_file = mnc2nii(mnc)
#'}
#' if (have_fs()) {
#'
      img = oro.nifti::nifti(array(rnorm(5*5*5), dim = c(5,5,5)))
#'
      mnc = nii2mnc(img)
#'
      img_file = mnc2nii(mnc)
#'}
  #' if (have_fs()) {
      img = oro.nifti::nifti(array(rnorm(5*5*5), dim = c(5,5,5)))
#'
      mask = img > 1
#'
      res = mri_mask(img, mask)
#'}
#' @examples \dontrun{
#' if (have_fs()){
```

```
#' nu_correct("/path/to/T1.nii.gz")
#' }
#' }
```

Introduction

Freesurfer is a commonly-used software for processing and analyzing anatomical neuroimaging data (Fischl, 2012), developed by the Laboratory for Computational Neuroimaging at the Athinoula A. Martinos Center for Biomedical Imaging. This software provides open-source, command-line tools for image processing tasks such as brain extraction/skull-stripping (Ségonne et al., 2004), bias-field correction (Sled et al., 1998), segmentation of structures within the brain (Fischl et al., 2002, 2004), and image registration (Fischl et al., 1999; Reuter et al., 2010). Many of these functions are used extensively in medical imaging pipelines and Freesurfer has a complete pipeline packaged with its software as well

We have previously published a similar adaptation of the FSL imaging software (Jenkinson et al., 2012) to R, called fslr (Muschelli et al., 2015). Again, we note that there exist a number of R packages for reading and manipulating image data, including AnalyzeFMRI (Bordier et al., 2011), RNiftyReg (Clayden, 2015), and fmri (Tabelow and Polzehl, 2011) (see the Medical Imaging CRAN task view http://cran.r-project.org/web/views/MedicalImaging.html for more information). Although these packages are useful for performing image analysis, much of the fundamental functionality of image preprocessing and processing that Freesurfer provides are not currently implemented in R. The ANTsR package (https://github.com/stnava/ANTsR) has much of this functionality has been implemented, but this package has not been released onto CRAN. Moreover, having multiple options for image processing through R allows for users to compare methods and the flexibility of using multiple packages to achieve a working data processing pipeline.

In particular, we provide an interface to users to the state-of-the-art anatomical processing implemented in Freesurfer, as well as a suite of tools that simply the analysis of the output of Freesurfer. This will allow R users to implement complete imaging analysis without necessarily learning Freesurfer-specific syntax.

R imaging objects

The freesurfer package relies on the oro.nifti (Whitcher et al., 2011) package implementation of images (referred to as nifti objects) that are in the Neuroimaging Informatics Technology Initiative (NIfTI) format, as well as other common image formats such as ANALYZE. Some Freesurfer functions require other formats, such as MINC (http://www.bic.mni.mcgill.ca/ServicesSoftware/MINC). The Freesurfer installation provides functions to convert from MINC to NIfTI formats and there are implemented in functions such as nii2mnc and mnc2nii in R. Moreover, the mri_convert Freesurfer function has been interfaced in the freesurfer package (same function name), which allows for a more general conversion tool of imaging types for R users than currently implemented in native R.

Reconstruction pipeline in Freesurfer

The Freesurfer pipeline and analysis workflow for neuroanatomical images is based on a structural magnetic resonance image (MRI) of the brain. The specific type of image commonly used in this software is a T1-weighted image, a specific MRI sequence commonly taken. The full pipeline is implemented in the Freesurfer recon-all function, where the "recon" stands for reconstruction (https://surfer.nmr.mgh.harvard.edu/fswiki/recon-all). Using the -all flag in the the recon-all function performs over 30 different steps and takes 20-40 hours to fully process a subject when performing all the steps. This process is the common way of fully processing an T1-weighted image in Freesurfer.

If there are problems with the result of this processing, there are multiple steps users can edit certain parts of the processing, such as skull-stripping, where non-brain tissues are removed from the image. The remainder of the pipeline can be run after these steps. The full pipeline is broken down into 3 separate steps, referred to as autorecon1, autorecon2, and autorecon3, which correspond to the flags in recon-all used to initiate these steps. We have written wrapper functions recon_con1, recon_con2, and recon_con3, respectively, for simplicity to the R user.

R function setup

To use **freesurfer**, a working installation of Freesurfer is required. The following code was run using Freesurfer version freesurfer-Darwin-lion-stable-pub-v5.3.0. The Freesurfer version can be accessed

using the fs_version function. **freesurfer** must also have the path of Freesurfer specified. If using R from a shell environment, and the FREESURFER_HOME environment variable is set (which can be done when installing Freesurfer), **freesurfer** will use this as the path to Freesurfer. If using R through a graphical user interface (GUI) such as RStudio (RStudio, Boston, MA), environmental variables and paths are not explicitly exported. Therefore, FREESURFER_HOME is not set, freesurfer will try the default directories of Mac OSX and Linux. If the user did not perform an standard installation of Freesurfer, the path to Freesurfer can be specified using options (freesurfer .path="/path/to/freesurfer").

We will discuss the setup functions for the **freesurfer** package and how they can be used in analysis and example code. For testing, whether a user has a Freesurfer installation, the have_fs function provides a logical indicator as a result. The fs_dir function will return the directory of the Freesurfer installation.

Structure of Freesurfer analyses

During the installation of Freesurfer, a series of variables are set up in the user's shell environment. One of these variables is SUBJECTS_DIR, which refers to a directory of the output of analysis from all subjects. This setup allows users to simply specify a subject identifier to analyze, rather than a specific path or multiple intermediate files.

This setup may not be desirable if the user prefers to structure the data from multiple studies into different folders. For example, the asegstats2table function takes the anatomical segmentation statistics and convert it to a table. The default argument for asegstats2table is to pass in a subject name rather than a file. The freesurfer asegstats2table function allows the R user to specify a different subject directory to read in the file, while not overridding the default set by SUBJECTS_DIR. This functionality allows users to have separate folders with subjects and read in the data by simply switching the subj_dir argument in the R function.

Similarly to the fs_dir function, the fs_subj_dir function will return the path to the Freesurfer subjects directory if it is set.

Some Freesurfer functions require an image as an input. For those functions, the R freesurfer functions that call those Freesurfer functions will take in a filename or a nifti object. The R code will convert the nifti to the corresponding input required for Freesurfer. From the user's perspective, the input/output process is all within R. The advantage of this approach is that the user can read in an image, do manipulations of the nifti object using standard syntax for arrays, and pass this object into the freesurfer R function. Thus, users can create complete pipelines for the analysis of imaging data by accessing Freesurfer through freesurfer.

Example analyses and use of functions

In the default subjects directory in the Freesurfer installation, there is a subject named "bert", where recon-all was run. In the sub-directory for subject bert, there are 3 folders which we will explore the results: "mri", which contain imaging data, "stats", whic containing statistics based on structures of the brain, and "surf", which contain the surface and curvature output from the Freesurfer processing.

Reading in anatomical statistics

The "aseg.stats" in the "stats" folder of subject bert corresponds to measures and statistics from the anatomical segmentation. The read_aseg_stats function reads this corresponding file and creates a list of 2 different data. frames:

```
file = file.path(fs_subj_dir(), "bert", "stats", "aseg.stats")
out = read_aseg_stats(file)
names(out)
```

The measures element corresponds to global measurements of the brain (e.g.~volume of the brain) as well as measures of gross anatomical structures (e.g.~gray matter).

```
head(out$measures[, c("meaning", "value", "units")])
```

```
meaning value
brain segmentation volume 1193318.000000
brain segmentation volume without ventricles 1174082.000000
brain segmentation volume without ventricles from surf 1173867.217735
left hemisphere cortical gray matter volume 237947.199463
right hemisphere cortical gray matter volume 238312.856735
```

```
6 total cortical gray matter volume 476260.056198 units
1 mm^3
2 mm^3
3 mm^3
4 mm^3
5 mm^3
6 mm^3
```

The structures element corresponds to a set of measures and statistics for a set of fixed anatomical structures.

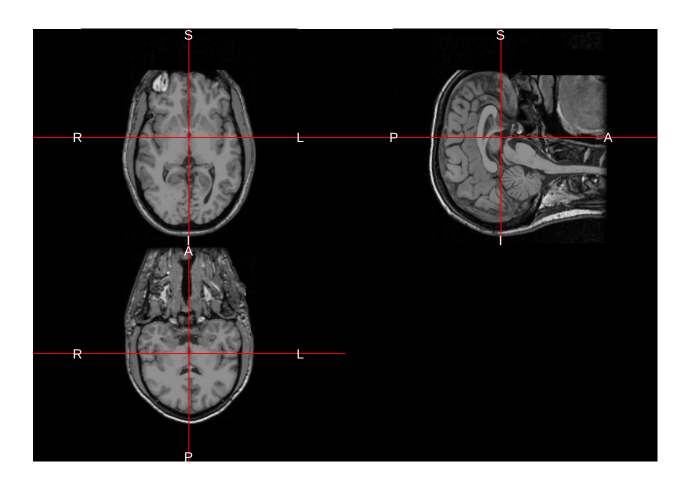
head(out\$structures)

	Index	SegId	NVoxels	Volume_mm3		StructName	normMean
1	1	4	6563	6562.6		Left-Lateral-Ventricle	36.0959
2	2	5	228	228.3		Left-Inf-Lat-Vent	54.8842
3	3	7	15708	15708.2	Left-0	Cerebellum-White-Matter	92.7562
4	4	8	58536	58535.7		Left-Cerebellum-Cortex	77.2709
5	5	10	8150	8150.4		Left-Thalamus-Proper	92.8386
6	6	11	3214	3213.7		Left-Caudate	80.9591
	normSt	tdDev ı	normMin r	normMax norm	nRange		
1	12.	. 2771	16	91	75		
2	10	. 7839	22	87	65		
3	5	.5123	40	107	67		
4	9	. 9521	17	142	125		
5	7.	.0182	49	109	60		
6	8.	. 2079	49	105	56		

MRI conversion

The typical output format from Freesurfer is MGH/MGZ format, which is explained here: https://surfer.nmr.mgh.harvard.edu/fswiki/FsTutorial/MghFormat. As NIfTI formats are one of the most common formats and has been the common format for analysis in the **oro.nifti** and **fslr** packages, it is useful to convert these files to a NIfTI format to use in R. The mri_convert Freesurfer function will be used for that. Here we will use the T1-weighted image from the "bert" subject, convert it to NIfTI, read it into R, and then plot the image.

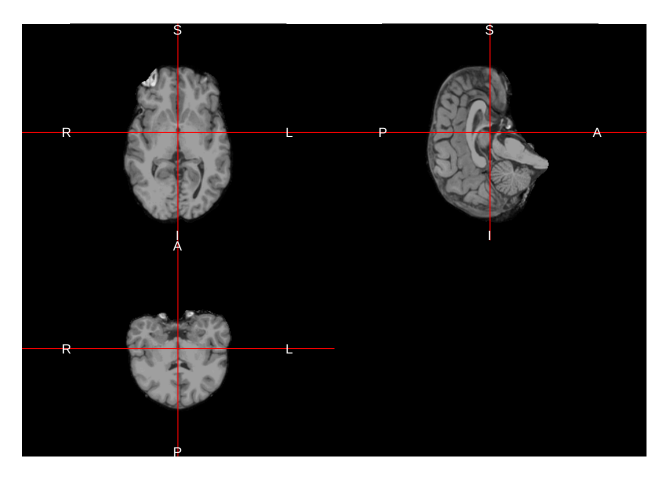
```
library(freesurfer)
t1_mgz = file.path(fs_subj_dir(), "bert", "mri", "T1.mgz")
t1_nii_fname = tempfile(fileext = ".nii.gz")
freesurfer::mri_convert(t1_mgz, t1_nii_fname)
library(fslr)
img = fslr::readnii(t1_nii_fname)
fslr::ortho2(img)
```



Brain extraction

The $mri_watershed$ function will segment the brain from the rest of the image. We cap ass in the nifti object and the output is a brain-extracted nifti object.

ss = mri_watershed(img)
ortho2(ss)



As the result in a nifti object, we can create a mask by standard logical operations:

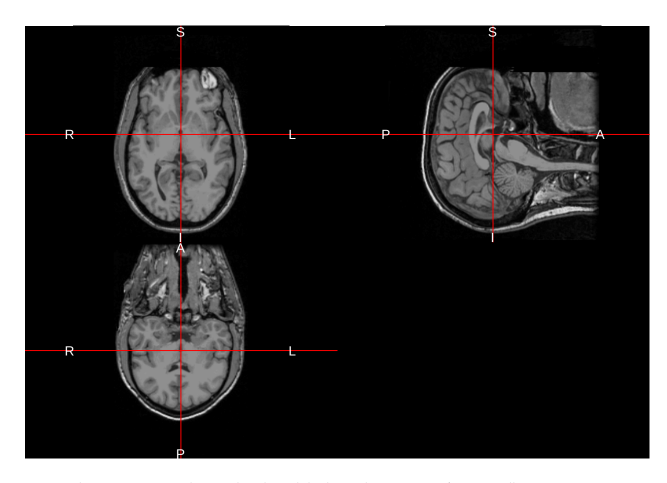
mask = ss > 0

Bias-field correction

MRI images typically exhibit good contrast between soft tissue classes, but intensity inhomogeneities in the radio frequency field can cause differences in the ranges of tissue types at different spatial locations (e.g.~top versus bottom of the brain). These inhomogeneities/non-uniformities can cause problems with algorithms based on histograms, quantiles, or raw intensities (Zhang et al., 2001). Therefore, correction for image inhomogeneities is a crucial step in many analyses. The Freesurfer function nu_correct performs the non-uniformity correction by Sled et al. (1998) and the freesurfer function of the same name will run the correction and return an image.

The Freesurfer nu_correct function requires a MNC format (http://www.bic.mni.mcgill.ca/ServicesSoftware/MINC). For this to work, you can convert the nifti object to a MNC file using nii2mnc and pass that file into nu_correct. The **freesurfer** nu_correct function will run the correction and then convert the output MNC to a NIfTI object.

```
mnc = nii2mnc(img)
print(mnc)
nu_from_mnc = nu_correct(file = mnc)
class(nu_from_mnc)
ortho2(nu_from_mnc)
```



You can also pass in a nifti object in directly, and the **freesurfer** nu_correct function will automatically convert any NIfTI input files, and then run the correction. We can also pass in a mask (generated from above) to run the correction only the areas of the brain.

```
nu_nifti = nu_correct(file = img, mask = mask)
class(nu_from_mnc)
```

Image preprocessing with fslr

We present a complete analysis of structural magnetic resonance imaging (MRI) data performed using fslr and R. Images were obtained from a patient with multiple sclerosis (MS) at 2 different visits (Sweeney et al., 2013), located at bit.ly/FSL_Data. At each visit, the image modalities obtained were T1-weighted (T1), T2-weighted (T2), fluid-attenuated inversion recovery (FLAIR), and proton density (PD). In this example we will perform a MRI bias-field correction using FAST (FMRIB's Automated Segmentation Tool) (Zhang et al., 2001), co-register scans within visits to the T1 image of that visit, and register T1 images between visits. Once these operations have been performed, one can take within-modality difference images to see the changes between visits. We will also register all images to a common stereotaxic template, as this is common in population-based analyses.

Section title in sentence case

This section may contain a figure such as Figure 1.



Figure 1: The logo of R.

Another section

There will likely be several sections, perhaps including code snippets, such as:

```
x <- 1:10
x
```

Conclusion

The neuroimaging community has developed a large collection of tools for image processing and analysis. R has a number of packages to perform operations on images; **EBImage** is one good example (Pau et al., 2010). Much of the fundamental functionality of neuroimage processing is not currently available in R, such as brain extraction and tissue segmentation. We present **fslr** to provide R users functions for image processing and analysis that are based on FSL, an established image processing and analysis software suite. Interfacing R with existing, powerful software provides users with thoroughly-tested software and an additional community of users, which would not be available if the functions were rewritten in R. **fslr** should be easy to use for any standard R user; the workflow allows R users to manipulate array-like nifti objects, pass them to **fslr** functions, which return nifti objects. Moreover, as FSL and R are open source and free, this software is readily available to all users.

There has been an increasing popularity of similar interfacing of tools within the Python community such as Nipype (Gorgolewski et al., 2011) (https://qa.debian.org/popcon.php?package=nipype). As many users of R may not have experience with Python or bash scripting, we believe fslr provides a lower threshold for use in the R community. Other packages provide R users additional neuroimaging processing functionality such as AnalyzeFMRI (Bordier et al., 2011), RNiftyReg (Clayden, 2015), and fmri (Tabelow and Polzehl, 2011).

For example, other inhomogeneity correction methods exist, such as the popular N3 (Sled et al., 1998) and N4 (Tustison et al., 2010), methods which are not implemented in fslr. ANTsR (http://stnava.github.io/ANTsR/index.html) is a currently unpublished R package that interfaces with the ANTs (advanced normalization tools) software suite (Avants et al., 2011). ANTs has implementations of these correction methods, an increased set of registration techniques, and other methods for image processing. Other packages such as this, along with fslr, can create a diverse set of tools for neuroimaging within R, building on preexisting and widely-accepted software.

Most importantly, as **fslr** is based on the R framework, all the benefits of using R are available, such as dynamic documents, reproducible reports, customized figures, and state-of-the-art statistical methods. These benefits provide unique functionality compared to other software packages for neuroimaging.

All data and code processed here is located at https://github.com/muschellij2/FSLR_Data.

Summary

This file is only a basic article template. For full details of *The R Journal* style and information on how to prepare your article for submission, see the Instructions for Authors.

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