# FreeSurfer Multiprocessing Pipeline

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# Module freesurfer\_wrapper

# freesurfer\_wrapper

**freesurfer\_wrapper** aims to facilitate the creation of a multiprocessing pipeline using FreeSurfer. It is a Python wrapper to execute parallel runs of recon-all and some pial edits algorithms.

# Requirements

- Docker<sup>1</sup>
- FreeSurfer license key<sup>2</sup>

# **Preparation**

- 1) Place the license in a license.txt file in the same folder as the Dockerfile.
- 2) Place your dataset folder in the same folder as the Dockerfile.
- 3) Build the docker image:

```
docker build -t fs_wrapper .
```

#### Overview run

The main script has 4 commands, each of them is explained below.

```
python run.py -h
usage: run.py [-h] {recon,segment_hip_amg,edit,recon_edit} ...
```

Command-line wrapper tool to execute parallel runs of FreeSurfer recon-all and some pial edits algorithms.

```
positional arguments:
```

```
{recon,segment_hip_amg,edit,recon_edit}
```

```
recon Run FreeSurfer recon-all.
```

segment\_hip\_amg Run segmentation of hippocampal subfields and nuclei of the amygdala.

edit Run mri\_gcut and mri\_binarize for pial edits.

recon\_edit Re-run recon-all for pial edits.

#### optional arguments:

-h, --help show this help message and exit

#### recon

#### Run FreeSurfer recon-all.

<sup>&</sup>lt;sup>1</sup>https://www.docker.com/

<sup>&</sup>lt;sup>2</sup>https://surfer.nmr.mgh.harvard.edu/registration.html

#### segment\_hip\_amg

Run segmentation of hippocampal subfields and nuclei of the amygdala. Original script by Juan Eugenio Iglesias. For more information, please consult FS official documentation<sup>3</sup>.

#### edit

```
Run mri_gcut and mri_binarize for pial edits.
```

## recon\_edit

```
Re-run recon-all for pial edits.
```

<sup>&</sup>lt;sup>3</sup>https://surfer.nmr.mgh.harvard.edu/fswiki/HippocampalSubfieldsAndNucleiOfAmygdala

#### How to run recon

#### Create the input file

A tab-separated file is needed as input with the following characteristics:

- Each line must represent a single scan.
- First column: unique ID.
- · Second column: path to dcm/nii file.

For the ADNI dataset, you can create this file using the create\_recon\_input.py script. This will create a recon\_input.txt file. The script will combine the subject ID and the session ID to create a unique ID.

```
docker run --rm -it -v $(pwd):/root/freesurfer_wrapper fs_wrapper \
python scripts/create_recon_input.py
```

As an example, a recon\_input.txt file for some ADNI records will look like this:

```
137_S_1414_S46193 ADNI/137_S_1414/MP-RAGE/2008-02-26_11_57_53.0/S46193/ADNI_137_S_1414_MR_MP-RAGE_
137_S_1414_S72806 ADNI/137_S_1414_MP-RAGE/2009-08-26_11_06_33.0/S72806/ADNI_137_S_1414_MR_MP-RAGE_
```

For other datasets you can try to edit the PATH PATTERN variable in scripts/create\_recon\_input.py.

#### Run recon

Now you can run the recon command using Docker. This can take several hours.

```
docker run --rm -it -v $(pwd):/root/freesurfer_wrapper fs_wrapper \
python run.py recon -i recon_input.txt
```

This command will use the maximum number of CPUs. You can append the -p <INT> flag where <INT> is the number of parallel runs you want.

#### How to run segment hip amg

This step requires that a whole brain T1 scan of the subject has been analyzed with the main FreeSurfer stream ("recon-all"). After running recon, you can use the same input table to run segment\_hip\_amg.

```
docker run --rm -it -v $(pwd):/root/freesurfer_wrapper fs_wrapper \
python run.py segment_hip_amg -i recon_input.txt
```

#### How to run edit

## Create the input file

After running recon you can check your results using freeview. Please refer to Manual quality analysis section to use a custom script.

If any skull edits are necessary, you need to create an input table to run edit with the following characteristics:

- Each line must represent a single scan.
- First column: unique ID.
- · Second column: path to dcm/nii file.
- Third column: the tissue ratio for WM edits.

Tissue ratio is the threshold to value (%) of WM intensity. The value should be >0 and <1; larger values would correspond to cleaner skull-strip but higher chance of brain erosion.

You can copy the recon input table, keep only the lines for the scans that need pial edits, and add the column with the tissue ratio values.

As an example, a edit input.txt file for some ADNI records will look like this:

```
137_S_1414_S46193 ADNI/137_S_1414/MP-RAGE/2008-02-26_11_57_53.0/S46193/ADNI_137_S_1414_MR_MP-RAGE_
137_S_1414_S72806 ADNI/137_S_1414_MP-RAGE/2009-08-26_11_06_33.0/S72806/ADNI_137_S_1414_MR_MP-RAGE_
```

#### Run edit

Now you can run the edit command using Docker.

```
docker run --rm -it -v $(pwd):/root/freesurfer_wrapper fs_wrapper \
python run.py edit -i edit_input.txt
```

This command will use the maximum number of CPUs. You can append the -p <INT> flag where <INT> is the number of parallel runs you want.

You can check the resulting edited mask using freeview:

```
SUBJECTS_DIR=$(pwd)/FS_OUTPUTS
freeview -recon <UNIQUE_ID> -v brainmask.gcutsT$<TISSUE_RATIO>.mgz:colormap=heat:opacity=0.5
```

If still not good, change the tissue ratio value in the input file and run edit again. When all masks are OK, proceed to recon\_edit command.

#### How to run recon\_edit

#### Create the input file

The input file is the table used for edit with the final values for tissue ratio.

#### Run recon edit

recon\_edit will re-run parts of FS recon-all using the edited masks. This can take several hours.

```
docker run --rm -it -v $(pwd):/root/freesurfer_wrapper fs_wrapper python run.py recon_edit -i edit_i
```

This command will use the maximum number of CPUs. You can append the -p <INT> flag where <INT> is the number of parallel runs you want.

#### How to check for completed runs and hard errors

FreeSurfer's recon-all command creates different logs while running. The recon-all.done log is created only for completed runs. The recon-all.error is created for hard failures.

You can check these logs using a custom script. The script was written to work on ADNI folder structure. For other datasets you can try to edit the PATH PATTERN variable in scripts/check logs.py.

#### Done

```
docker run --rm -it -v $(pwd):/root/freesurfer_wrapper fs_wrapper \
python scripts/check_logs.py done
You can also pipe the output to bash word count command to get a quick count:
```

```
docker run --rm -it -v $(pwd):/root/freesurfer_wrapper fs_wrapper \
python scripts/check_logs.py done | wc -l
```

#### Error

```
docker run --rm -it -v $(pwd):/root/freesurfer_wrapper fs_wrapper \
python scripts/check_logs.py error
```

#### How to restart after a computer failure

If the execution of the pipeline is halted by a computer failure or system restart then you have to update the input file of the recon command.

```
bash scripts/update_recon_input.sh
```

This will remove all "done" samples from the original recon\_input. It will also delete the folders from the samples that were running when the failure happened. These samples will run again from scratch.

Now run again the recon command but using the new input file (<YYYY-MM-DD>\_recon\_input.txt).

#### **Quality control**

#### **Automated quality analysis**

The tool is packaged with **qatools-python**<sup>4</sup> version 1.2 for quality control measurements. This script was developed by Reuter DeepMI Lab<sup>5</sup> as a revision, extension, and translation to the Python language of the Freesurfer QA Tools.

```
docker run --rm -it -v $(pwd):/root/freesurfer_wrapper fs_wrapper \
python scripts/qatools-python/qatools.py --subjects_dir FS_OUTPUTS --output_dir QC \
--screenshots --outlier --fornix
```

This will create qatools-results.csv file; screenshots, outliers and fornix folders inside the QC folder. Please consult gatools-python docs<sup>6</sup> for a full explanation of each QC measurement.

#### Manual quality analysis

You can visually inspect each result using freeview. We provide a script to speed up the opening process of each scan. The script also prompts the user about the result of the QC after each window of freeview is closed. The result is saved to manual\_QC.txt

It is not possible to run freeview using Docker, the graphical user interface cannot be displayed. Therefore, you need to have FreeSurfer/freeview installed in your host machine.

#### Step 1

Set SUBJECTS\_DIR environment variable. Here, the results are stored inside the FS\_OUTPUTS directory. export SUBJECTS DIR=\$(pwd)/FS OUTPUTS

#### Step 2

Run the view.py script.

python scripts/view.py

#### **Sub-modules**

- · freesurfer wrapper.run
- freesurfer\_wrapper.scripts

# Module freesurfer\_wrapper.run

Command-line wrapper tool to execute parallel runs of FreeSurfer recon-all and some pial edits algorithms.

This file can also be imported as a module and contains the following functions:

- \* argument\_parser parser for command-line options, arguments and sub-commands.
- \* run\_command -
- \* handle\_workers creates a pool of parallel worker processes running commands.
- \* worker invokes a subprocess running the command.
- \* recon formats recon-all command string.
- \* edit formats mri\_gcut and mri\_binarize command string.
- \* recon edit formats a cp and recon-all command string.
- \* parse\_input\_file parses the input tables.

#### **Functions**

# Function argument\_parser

```
def argument_parser(
    args: list
```

<sup>&</sup>lt;sup>4</sup>https://github.com/Deep-MI/qatools-python

<sup>5</sup>https://deep-mi.org/

 $<sup>^6</sup>$  scripts/qatools-python/README.md#description

```
) -> ArgumentParser.parse_args
```

Parser for command-line options, arguments and sub-commands.

**Parameters** 

args : list Command-line arguments list

Returns

#### Parser

#### Function edit

```
def edit(
    edit_args: list
) -> str
```

Formats mri\_gcut and mri\_binarize command string. mri\_gcut performs skull stripping algorithm based on graph cut. mri\_binarize binarizes the edited mask.

**Parameters** 

```
edit_args: list mri gcut and mri binarize arguments list
```

Returns

```
mri_gcut [args] && mri_binarize [args]
```

#### Function handle\_workers

```
def handle_workers(
   p: int,
   command: function,
   input_file: str
)
```

Creates a pool of parallel worker processes running commands. Workers will be called until all lines from the input file are processed.

**Parameters** 

```
p: int The number of parallel processes.
```

 $\label{lem:command-line} \textbf{command-line string to pass the worker.}$ 

input\_file: str Tab-separated .txt file.

Returns

#### None

#### Function parse\_input\_file

```
def parse_input_file(
        input_file: str
) -> List[List[str]]
```

Parses the input tables.

**Parameters** 

```
input_file: str Tab-separated .txt file.
```

Returns

File lines and columns parsed as a list of lists.

```
Function recon
```

```
def recon(
        recon_args: list
    ) -> str

Formats recon-all command string.

Parameters

recon_args: list recon-all arguments list

Returns

recon-all [args]

Function recon_edit
    def recon_edit(
        recon_edit_args: list
    ) -> str
```

Formats a cp and recon-all command string. cp replaces the original brainmask with the edited brainmask.gcutsT{tissue\_ratio}.mgz. recon-all re-runs -autorecon2-wm -autorecon3 stream with the new mask.

**Parameters** 

```
recon_edit_args: list cp and recon-all arguments list
Returns
cp [args] && recon-all [args]
```

Function run\_command

```
def run_command(
     args
)
```

Pass the appropriate command function to the worker handler.

Parameters

args : list Command-line arguments list

Returns

None

# Function worker

```
def worker(
    cmd: str
) -> <function run at 0x7f1070968670>
```

Invokes a subprocess running the command.

Parameters

cmd: str Command-line string

Returns

subprocess.run()

# Namespace freesurfer\_wrapper.scripts

#### **Sub-modules**

- freesurfer wrapper.scripts.check logs
- freesurfer\_wrapper.scripts.create\_recon\_input
- freesurfer\_wrapper.scripts.view

# Module freesurfer\_wrapper.scripts.check\_logs

Script to check recon-all logs for each run

usage: python check\_logs.py <done|error>

Please edit the PATH PATTERN variable with the appropriate pathname pattern to find each file.

This file can also be imported as a module and contains the following functions:

```
* get_logs - get the path for each log based on pathname pattern.
* print_id_from_logs - prints the IDs from a list of logs.
```

#### **Functions**

```
Function get_logs
```

```
def get_logs(
    path_pattern: str
) -> list
```

Get the path for each log based on pathname pattern.

**Parameters** 

path\_pattern: str Glob pathname pattern to find each log.

Returns

#### List of log paths

```
Function print_id_from_logs
```

```
def print_id_from_logs(
    logs: list
)
```

Prints the IDs from a list of logs.

**Parameters** 

logs: list List of log paths

Returns

None

# Module freesurfer\_wrapper.scripts.create\_recon\_input

Script to create recon input table

This script creates an input table based on the directory organization of the image files.

Please edit the PATH\_PATTERN variable with the appropriate pathname pattern to find each file.

This file can also be imported as a module and contains the following functions:

```
* create_input_file - creates the input table.
```

#### **Functions**

Creates a two column text file to be used as input for the main script recon command. First column: unique ID (combines SUBJECT ID and SESSION ID). Second column: path to DICOM file.

**Parameters** 

path\_pattern: str Glob pathname pattern to find each DICOM file.

Returns

None

# Module freesurfer\_wrapper.scripts.view

Script to open the scans in sequence and register if passed or not in QC analysis

This script identifies the subjects present in the folder set in SUBJECTS\_DIR environment variable. Using this list, it automatically opens freeview and asks the user input for the result of QC analysis. Results are saved in manual\_QC.txt file.

This file can also be imported as a module and contains the following functions:

```
\ast get_subjects - returns a list of subjects inside the folder.
```

\* freeview - returns a freeview command formated as string

#### **Functions**

```
Function freeview
```

```
def freeview(
    subject_id: str
)
```

Returns a freeview command formated as string.

**Parameters** 

```
subject_id: str FreeSurfer subject_id.
```

Returns

str

# $\textbf{Function} \ \texttt{get\_subjects}$

```
def get_subjects(
    subjects_dir='/subjects'
)
```

Returns a list of subjects inside the folder.

**Parameters** 

subjects\_dir: str, default=os.environ['SUBJECTS\_DIR'] FreeSurfer SUBJECTS\_DIR.

Returns

list

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