# HFR\_ai: a toolbox for finding Homologous Functional Regions Across Individuals

Part0: Download HFR\_ai.tar.gz and add it in MATLAB

Part1: Individual parcellation

Part2: HFR\_ai

Part3: Create FC matrix

If you will use this toolbox, please make sure you have installed FREESURFER and add the directory of "matlab", which is in the FREESURFER software, into your Matlab (<a href="https://surfer.nmr.mgh.harvard.edu/fswiki/">https://surfer.nmr.mgh.harvard.edu/fswiki/</a>)

Organized by Meiling Li, 2017/05/31

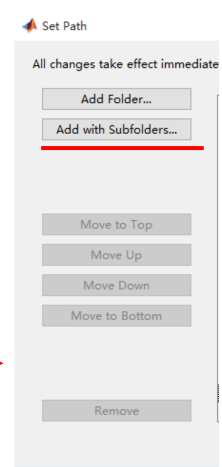
## Part0: Download HFR\_ai.tar.gz and add it in MATLAB

- a. Download this toolbox here: <a href="http://nmr.mgh.harvard.edu/bid/DownLoad.html">http://nmr.mgh.harvard.edu/bid/DownLoad.html</a>
- b. Unpack the HFR\_ai.tar.gz in your directory like tar -zxvf HFR\_ai.tar.gz you will get a directory named HFR\_ai
- c. Open MATLAB in your terminal, and add this toolbox(HFR\_ai) in the toolstrip like:

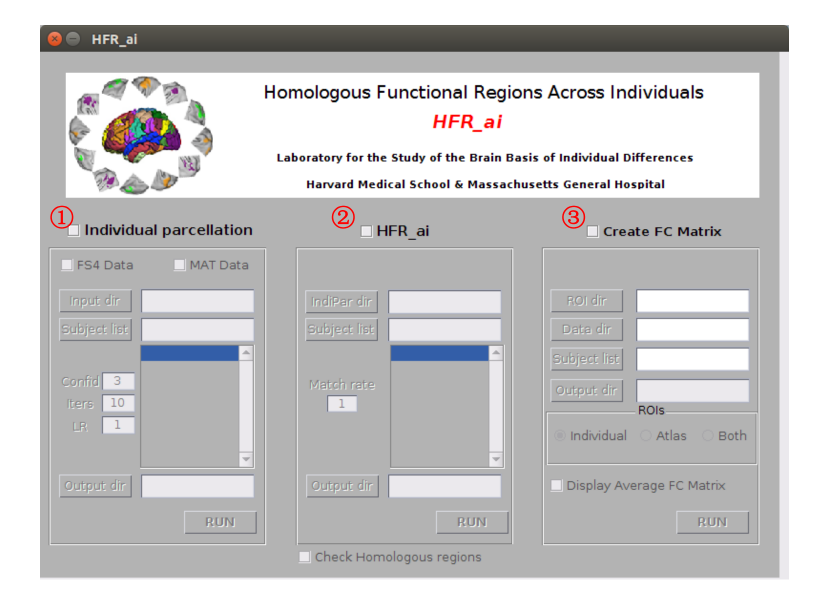
#### **HOME->Set Path -> Add with Subfolders -> Save -> Close**

or use the MATLAB command "addpath(genpath('/Path/HFR\_ai'))" if you use only once.





d. Type "HFR\_ai" and Return in the command window, then you should see the following interface, if you have added this toolbox successfully.



HFR\_ai Pipeline summary:

- Individual parcellation. Parcellating individualized functional networks proposed in Wang et al. 2015, NN.
- HFR\_ai. Finding the homologous functional regions across individuals.
- 3. Create FC Matrix. Calculating the ROI-ROI functional connectivity based on ROIs from HFR\_ai and the corresponding ROIs from atlas.

## Part1: Individual parcellation

### Input dir:

the preprocessed data projected on fsaverage4.

The supported data type is ".nii.gz"(FS4 Data), and ".mat" (MAT Data). The preprocessed data is suggested to name and place like the following structure (next page).

### Subject list:

a TXT file recorded all the subject ID you would like to do

parcellation.

SubList.txt x sub\_001 sub\_002 sub\_003 sub\_004 sub\_005

#### Confid:

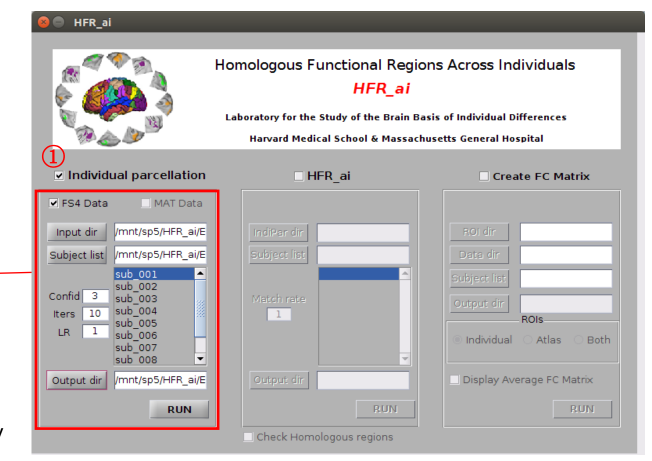
Confidence, the default value is 3, if the input value is extremely large, it will select the vertices with maximum confidence as the seed in the interation.

#### Iters:

the terminating iteration number in the parcellation. The default value is 10.

#### LR:

whether to combine the same network of left hemisphere and right hemisphere, "1" means "Yes", "0" means "No". The default value is 1.



# Data Directory Structure

sub\_003

### FS4 Data

1. Project Name

2. Subject Names

3. surf

4. Data on fsaverage4

Ih. \*\_fsaverage6\_sm6\_fsaverage4.nii.gz

sub\_001

surf

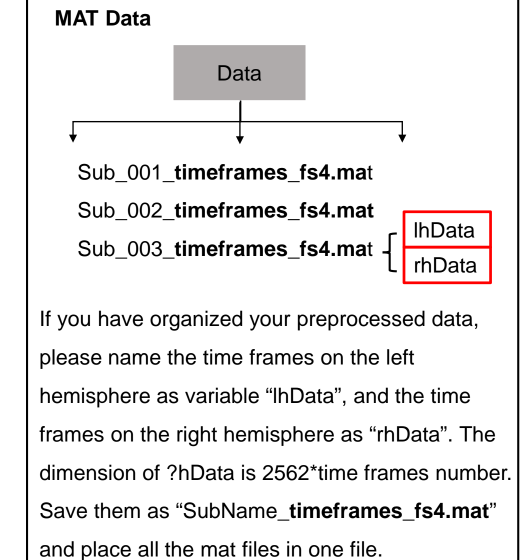
Data

sub\_002

surf

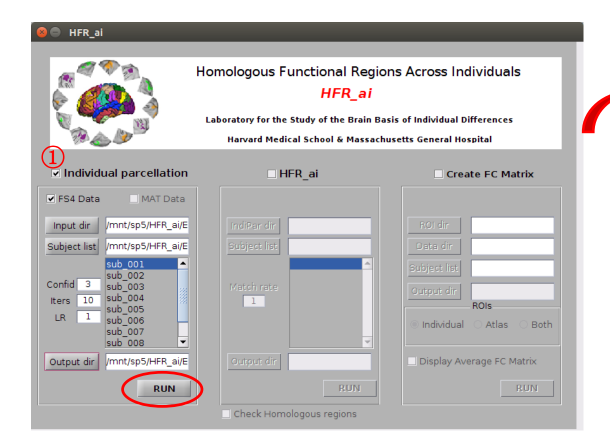
rh.\* \_fsaverage6\_sm6\_fsaverage4.nii.gz

Note: The last dimension of the data is the number of time frames



### FS4 Data example:

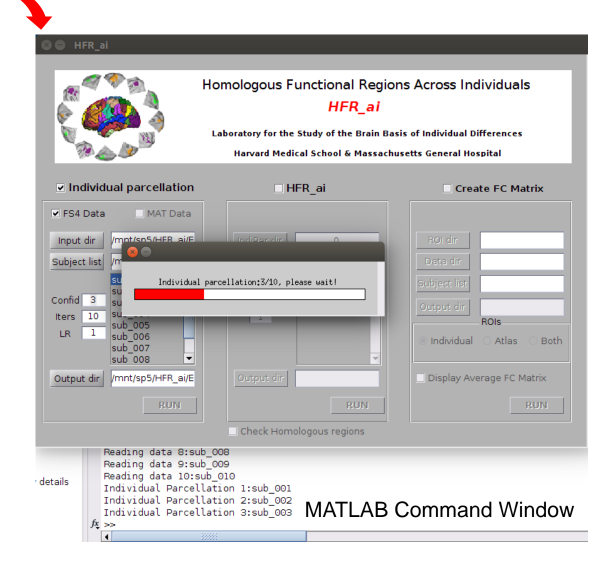
Ih.sub\_001\_bld013\_rest\_reorient\_skip\_faln\_mc\_g1000000000\_bpss\_resid\_fsaverage6\_sm6\_fsaverage4.nii.gz rh.sub\_001\_bld013\_rest\_reorient\_skip\_faln\_mc\_g1000000000\_bpss\_resid\_fsaverage6\_sm6\_fsaverage4.nii.gz



Click the RUN after all be set, then you will see the individual parcellation starts running!

Do not close the wait bar! Just wait!

This Part will generate two files: /Output dir/Organized Data /Output dir/IndiPar



Part2: HFR\_ai

### Indipar dir:

Individual parcellation results obtained from Part1.

eg: /Output dir/IndiPar

### Subject list:

a TXT file recorded all the subject names you would like to find Homologous functional regions across these subjects.

```
SubList.txt x
sub_001
sub_002
sub_003
sub_004
sub_005
```

#### *Iter*:

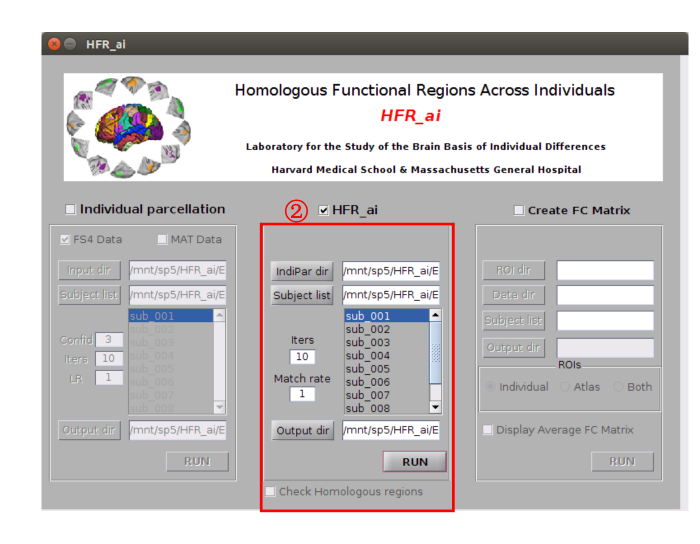
Iter time (parcellation result) you would like to be further used, to get the homologous functional regions.

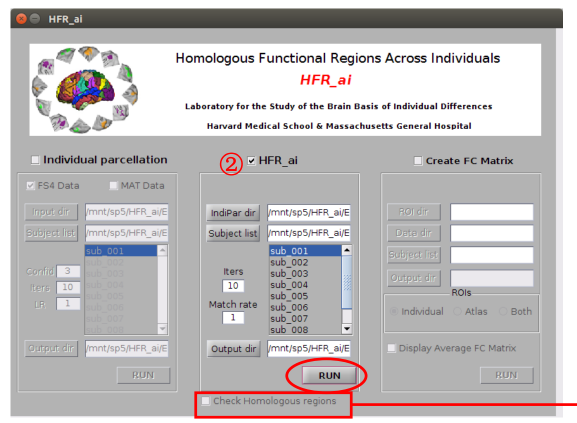
#### Match Rate:

The Match rate (0~1) means the minimum ratio that subjects have a given ROI when matching the ROIs across subjects.

Eg: "1" means all subjects have matched that ROI, the kept ROI is homologous in function across all individuals.

"0.8" means that at least 80% subjects have matched ROI( i.e. we can find this ROI in at least 80% subjects.)





Click the RUN after all be set, then you will see the HFR\_ai starts running!

It may take some time to match the ROIs according to the subjects number. After generated the homologous regions, please check them.



It will display the homologous ROIs on the surface (shown on fsaverage5 using the corresponding ROIs from Atlas). If the number of homologous regions across individuals is extremely small and the homologous ROIs even can not cover the most of the surface, it is not suggested to do further analysis using the individualized ROIs.

Homologous functional regions can be found here:

### /Output dir/MatchRate1/

Which also includes the corresponding ROIs on atlas.

### Part3: Create FC Matrix

#### ROI dir:

Homologous regions obtained from Part2.

eg: /Output dir/MatchRate1/

Do NOT select the "Indi\_Matched\_ROIs" or "GrpTemplate\_Matched\_ROIs".

#### Data dir:

The preprocessed resting state fMRI data(organized data)

Eg: /Output dir/OrganizedData

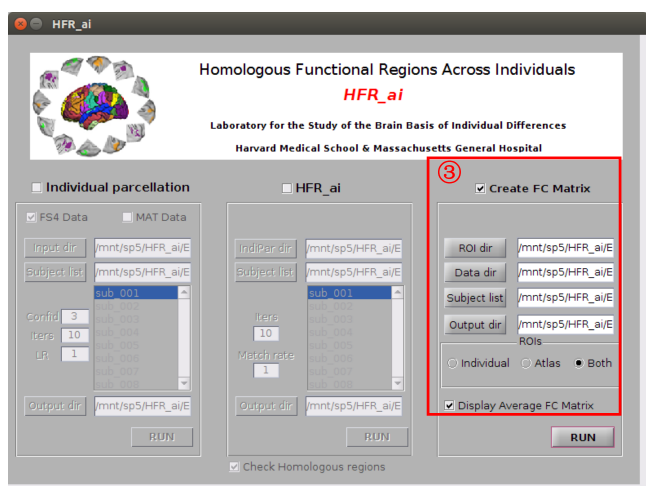
### Subject list:

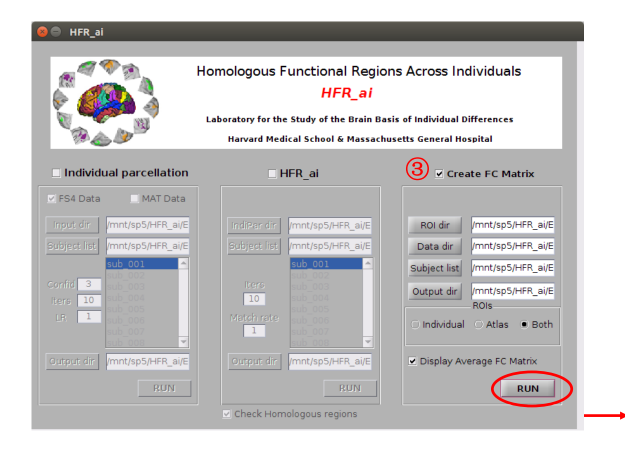
a TXT file recorded all the subject names you would like to calculate the ROI-ROI FC and check the mean FC across these subjects.

Sub\_001
sub\_002
sub\_003
sub\_004
sub\_005

#### ROIs:

ROIs used to calculate the ROI-ROI functional connectivity.





Click the RUN after all be set, then you will see the average FC matrix.

Check the FC Matrix here:

/Output dir/MatchRate1/ROI2ROIFC\_Indi

/Output dir/MatchRate1/ROI2ROIFC\_Atlas

