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Progress Report

Seed-based Functional Connectivity Analysis of Hippocampal Network of Patients Suffering from Major Depressive Disorder

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Preface

The basis for this project stemmed from the fact that not many researches have been conducted in Nepal regarding the diagnosis of mental disorders. Nonetheless, in other countries, many researches and studies have been conducted regarding the functional connectivity of different brain regions in depression and other mental disorders. However, till date, there is no solid evidence that could be used for the clinical diagnosis of mental disorders. Our project intends to review past researches and keep up with the studies related to Major Depressive Disorder and brain functional connectivity. In addition to that, we have selected hippocampal circuity as the region of interest for our purposes and the overall project is going to revolve around how functional connectivity of hippocampal network in MDD patients differ from that of healthy people.

- Authors

Abstract

The following document is the progress report of the task that have been accomplished so far. The data selection and group matching based on age and sex is attained from statistical analysis. Preprocessing of fMRI images is necessary to remove unwanted artifacts and transform the data into a standard format. The neuroimaging data processing tools that are used to carry out preprocessing are SPM, AFNI. Preprocessing steps: Removing first few TR scans, despising, skull stripping, Slice timing correction, motion correction has been completed. After that segmentation of images using segmentation feature in SPM12 has been achieved. The normalization and smoothing preprocessing steps are in progress. Additional works to achieve these results include conversion of fMR images to compatible format using MRIcron, documentation of various AFNI commands that are used in preprocessing steps. Before, calculating functional connectivity, covariates are needed to be removed. Future works include: removal of covariates, region of interest drawing and functional connectivity analysis and finally the statistical analysis to compare the Healthy controls and MDD patients.

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AFN	II Analysis of Functional Neuroimages
BDI	Beck Depression Index
BOL	LD Blood Oxygen Level Dependent
CSF	Cerebrospinal Fluid
fMR	I Functional Magnetic Resonance Imaging
GM	Gray Matter
HCs	Healthy Controls
MDI	D Major Depressive Disorder
MR	Magnetic Resonance
rs R	esting State
rsFC	C Resting-state Functional Connectivity
SCA	Seed-based Correlation Analysis
sMR	I Structural Magnetic Resonance Imaging
SPM	I Statistical Parametric Mapping
WM	White Matter

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1 Introduction

The following is the progress report of our project. We have briefly discussed individual tasks performed since the commence of out project till the reporting period. Our project aims to perform analysis of the functional connectivity of the hippocampus in patients suffering from Major Depressive Disorder and acquire a comprehensive idea about how the functional connectivity in the hippocampus region changes in people who are suffering from major depressive disorder compared to healthy controls.

Functional connectivity is the communication between distinct brain regions that are not anatomically connected but remain simultaneously active while performing cognitive tasks. In this project we will assess the resting state functional connectivity, which essentially is the functional connectivity of various brain regions in resting state. Functional Magnetic Resonance Imaging (fMRI) is a well established imaging technique for the detection and delineation of the brain, based on the measure of neural activation in the brain. The fMR imaging modality is sensitive to fluctuations in the BOLD signal which reflects neuronal activation, or neuronal activity. This sensitivity allows fMRI technique to localize the exact location of activity in the brain which is caused due to sensory stimulation or cognitive function.

1.1 Background

The specific objective of this project is to perform seed-based functional connectivity analysis to study how the normal function of a particular area of a healthy brain gets disrupted in diseased conditions. Major Depressive Disorder being one of the major mental health illnesses in our country, this project aims to analyze the functional changes in the brain of MDD patients compared to healthy controls. Furthermore, in our literature review, it was found that the hippocampus of the brain is one of the major regions affected by a variety of neurodegenerative and mental disorders. For this reason, we aim to assess the functional connectivity of the hippocampal region of the brain.

1.2 Rationale

The absence of biological markers makes it exceptionally difficult for neurologists to diagnose a person with a psychiatric disorder. The diagnostic procedures that are the gold standard for the diagnosis of neurodegenerative and psychiatric disorders, in the present day, are wholly based on behavioral observations and patient reported symptoms, both of which do not have a molecular or radiological basis. Although there have been countless studies conceptualizing the possibility of implementation of various functional imaging modalities for deciphering the etiology and the functional effects of various mental disorders, the findings from these studies do not appear amongst the diagnostic criteria. A critical barrier to the clinical translation of many such findings is the reverse inference fallacy.

Since neurological disorders are multifaceted and are influenced by more than one factors

and neuroimaging results can be heavily influenced by external factors such as patient movement and instrumental artifacts. The information from structural radiology modalities describe the shape, size and integrity of brain structure, but they do not provide any information about the brain function. Nonetheless, as we will discuss in the coming sections of this progress report, combining structural MRI and functional MRI can be a promising technique to characterize normal and abnormal brain function, which can act as an auspicious biomarker for neurodegenerative or psychiatric disorders to determine the risk, progression and therapeutic effectiveness. This project can lay the foundations for further research and development in this particular field.

2 Methodology and Review on Individual Tasks

2.1 System Setup

For the following project, the primary software tool that we will use for functional connectivity analysis is AFNI. AFNI (Analysis of Functional NeuroImages) is an open-source software, distributed freely under the GNU General Public License. AFNI is used for processing, analyzing and displaying several MRI modalities such as anatomical MRI, functional MRI (FMRI) and diffusion weighted (DW) data. AFNI runs virtually on any UNIX based system such as macOS and GNU Linux. For financial reasons, we opted to install GNU Linux. Along with AFNI, we will also use SPM (Statistical Parametric Mapping) which is an image processing package in GNU Octave. GNU Octave, and all of its associated programs, is freely distributed under the terms of the GNU General Public License.

GNU Linux is an open-source Unix-like operating system. The Linux kernel is licensed under the terms of GNU General Public License version 2 (GPL-2.0). Popular Linux distributions include Ubuntu, Linux Mint, Fedora, and Arch. After we collected the SSDs that we requested for, we installed various linux distributions, namely Ubuntu, Linux Mint and Arch Linux. After installation of Linux, we installed AFNI and other related software on our Linux systems.

2.1.1 Runtime Environment

Here is a list of the major softwares used, along with their version information, during the runtime for the codes and results included in the following report:

- Operating System: Arch Linux x86_64, kernel version 5.16.8-arch1-1
- AFNI, version AFNI_22.0.03 'Hadrian'
- GNU Octave, version 6.4.0

2.2 Data Acquisition

The functional and anatomical MR images for the progression of this study were acquired from the SRPBS Multidisorder MRI Dataset. The datasets included were obtained from the Dec-Nef Project Brain Data Repository, which is a repository of neurological images gathered by a consortium as a part of the Japanese Strategic Research Program for the Promotion of Brain Science supported by the Japanese Advanced Research and Development Programs for Medical Innovation (AMED). Furthermore, the MR datasets included in this repository were diagnosed by trained and experienced neurologists, which assures that our study will be tilted more towards accuracy and efficiency **dataset**. We acquired functional MR as well as T1 weighted images of subjects from two distinct categories. The first set of subjects are categorized as healthy controls, and the second set of subjects are categorized as those who are suffering from major depression. The subjects from the first category, i.e. healthy controls will be referred to

as "HC" and the subjects from the second category, i.e. depressed subjects will be referred to as "MDD patients".

2.3 Data Selection

The SRBPS dataset originally had the MR images of more than 1400 volunteers. All of these volunteers had undergone a standardized clinical evaluation protocol, which involved a general and neurological evaluation. This opts for the accuracy of our study.

The following is a brief summary of the tasks performed during this step:

Selection of Data					
Status during the reporting period Completed					
Actions Out of the data of 1400 volunteers from the SRBPS datase healthy controls (diagnosis 0) and 15 depressed patients (diagno were manually selected by just eyeballing the data sheet include the data acquired from the SRBPS public repository.					
	• The datasheet contained data of volunteers from various different site. Subjects of either sex, male or female, were selected from only one specific site, HUH in particular. The reason behind this was to make sure that the images that we will be working on were acquired from the same MRI scanner. This ensure consistency of data.				
Decisions	 Majorly depressed patients were labeled with a BDI (Beck Depression Index) greater than 30. So subjects having BDI > 30 were only selected. 				
	• Trying to acquire data of people from a certain age group was a failure because there was not just enough data of the volunteers with BDI > 30. So we had to settle with subjects of the age group 20 to 50.				
	Once we had the subjects decided, two statistical tests were performed:				
Testing & Verification	 Chi-square test t-test 				
	Chi-square test was performed to check the goodness of fit between healthy controls and depressed subjects based on sex and the t-test was performed to check the goodness of fit based on age.				

The results of both t-test and the chi-square tests were positive (Appendix A), therefore we concluded that the subjects we selected for both categories, HC and MDD were well matched.

2.4 Data Preparation

After we finalized the subjects on which we will be working on, the image data that we acquired were converted into appropriate formats that could be used for further processing. The original image data that we acquired were in DICOM image format.

The following is a brief summary of the tasks performed during this step:

Conversion of image data from DICOM format to NIfTI format					
Status during the reporting period	Completed				
Actions	Created a BASH script to add the NIfTI extension i.e. ".nii" in order to convert the image data of 15 HC as well as 15 MDD patients from DICOM format to NIfTI format. (Appendix B)				
Decisions	=				
Testing & Verifica-	The images after conversion to NIfTI were opened in AFNI to verify				
tion	that it could understand and render the image data in its interface.				

Furthermore, once the image data were converted into the NIfTI format, which could be understood by a variety of software tools that we use, the multiple 3D rsfMRI image data was converted into a single 4D BOLD data for each subject from either category. This step was necessary because, unlike an anatomical MR image which is a single volume, a functional MR image is acquired in blocks, where each block represents the functional MR signal acquired at a given time. Therefore is essential to merge these multiple 3D functional MR image volumes into a single 4D functional MR image, where time is the fourth dimension. To achieve this, we use a graphical file conversion tool called dcm2nii. The graphical interface was used to select all the 3D rsfMRI volumes to create a compressed file format containing the 4D BOLD data.

After these two steps, our image data for both HCs as well as MDD patients were prepared for further processing.

2.5 Skull Stripping

Since high resolution structural images contain considerable amounts of non-brain tissue such as eyeballs, bone, skin, amongst other tissues. Skull stripping improves the quality and accuracy of the normalization and templates that will be created for skull-stripped images. The skull stripping was achieved in AFNI. The skull stripping process completes in the following steps:

Extraction of the brain tissue							
Status during the reporting period	Completed						
Actions	AFNI provides a skull stripping tool called 3dSkullStrip. A shell script was created to implement this 3dSkullStrip program to extract						
Decisions	 After data preparation, our next step would be segmentation of various brain structures from the brain. But before doing just that, the brain tissues must be extracted by removing the surrounding skull in order to isolate the brain tissue from non-brain tissue from an MRI image of a brain. We decided that it would be better to strip the skull from T1 images in order to exclude gross spatial image non-uniformity artifacts and to reposition the brain in a reasonable manner. 						
Testing & Verification	The generated files were visualized in the AFNI interface. The following figures represent a basic visualization of the results for an axial image with z at 0.404 mm. Figure 1: Axial Brain Image with Skull and without skull						

Once the brain tissue is extracted from the T1 image, we proceed to image segmentation.

2.6 Image Segmentation

Image segmentation is the process of partitioning a digital image into multiple image segments, or image regions which essentially are a similar set of pixels. The primary goal of image segmentation is feature extraction, to simplify and change the representation of an image into something that is more meaningful and easier to analyze. Most, if not all, of the analysis of medical images requires some form of segmentation or feature extraction. Segmentation makes it easier to analyze any given image, as it distinguishes structures, regions or tissue classes of interest from other details in the image. Segmentation depends on a variety of features that are contained in the image. The features can be either color or texture or something else. The primary reason for image segmentation for our purposes is to reduce the information for easy analysis.

For our purposes, the image segmentation part involves segmentation of the gray-matter, white-matter and the cerebrospinal fluid in the T1 weighted image. The segmentation is carried out using SPM12 in GNU Octave. SPM (Statistical Parametric Mapping) is an image processing package of octave which requires the images to be in the NIfTI file format. SPM12 employs an algorithm that performs segmentation by characterizing intensity distributions of different tissue classes. Gray-matter contains high densities of unmyelinated (lacking a myelin sheath) neurons, white-matter contains high densities of myelinated neurons and CSF contains an ultrafiltrate of plasma and protein. This renders the intensities of these tissues differently in the image. SPM uses region based segmentation. Region based segmentation essentially extracts different regions of the brain into separate files. Region based segmentation further includes:

• Threshold Segmentation:

In threshold segmentation, the image grayscale information processing is directly divided based on the gray values of various targets. Segmentation effect can be obtained if the target and background have high contrast. Threshold detection can be employed either locally or globally on the entire image. Local thresholding involves selecting variable segmentation threshold for different regions of the image based on the target regions and backgrounds. Global thresholding on the other hand only uses a single threshold for the entire image.

• Regional Growth segmentation:

In regional growth segmentation, a particular seed pixel is selected and an intensity uniformity constraint is set. Then all voxels around the seed are examined to see if their intensities are sufficiently similar to those already in the region. Those pixels that satisfy the uniformity constraint are added or merged around the seed pixel.

In an ideal case, the files generated at the end of the segmentation are meaningful and contain a distinct set of pixels that represent a distinct brain region. In MR imaging system,

the inherent magnetic inhomogeneities may cause a variation in intensity of a particular tissue across the field of view. In addition to that, the intensity of a single voxel may be composed of signal from more than one tissue type. This is called partial volume effect. This partial volume effect has consequences in classification of the brain-tissue into gray-matter, white-matter and CSF in T1-weighted images. Partial volume effects between white-matter (bright) and CSF (dark) result in voxels with an inbetween intensity which can be misclassified as gray matter. Here is a brief overview of the tasks performed and their description during image segmentation:

Segmentation of Gray matter, White matter and CSF							
Status during the	Status during the Completed. (The results of segmentation are improper for certain						
reporting period subjects so this needs to be redone)							
Actions	Actions Used the interactive GUI of SPM12 to automatically identify as segment different tissue types within the images. A batch of T1 scar to be segmented were specified, along with a list of tissues that we to be identified.						
Decisions	 Segmentation was only performed for T1-weighted images of healthy controls. 						
Testing & Verification	SPM produces several files with the segmented data. Files beginning with "c1" is what the algorithm identifies as the gray matter; files beginning with "c2" is what the algorithm identifies as white matter and files beginning with "c3" is what the algorithm identifies as the CSF. Each of these files were visualized for all subjects in AFNI to verify that segmentation was performed correctly.						

The output of the segmentation will be used for achieving a more accurate inter-subject alignment using DARTEL. The segmentation image data will also be used to generate a common mask.

2.6.1 Template Creation using DARTEL

DARTEL (Diffeomorphic Anatomical Registration Through Exponentiated Lie Algebra) is a toolbox available in SPM. DARTEL can be used to create templates that have a more accurate inter-subject alignment of various tissues. This is achieved by modeling the shape of each brain using millions of parameters. DARTEL achieves accuracy of such proportions by generating its own increasingly crisp average template data, to which the data are iteratively aligned. Each iteration makes individual images fit each other more precisely.

We need DARTEL because it allows an accurate inter-subject registration of brain images which is necessary for both proper separation of different segment, different tissue classes and after it gets specially normalized towards common global MNI we will be able to do group wise statistics and to extrapolate those findings to result of other studies.

The following is a brief summary of the tasks performed during this step:

Creating DARTEL Templates				
Status during the reporting period	Completed			
Actions	DARTEL was used to achieve a more accurate alignment of tissues classes of various all the subjects. DARTEL can be accessed from the interactive GUI of SPM12 and multiple files can be specified for batch processing. DARTEL iterates over the specified files and automatically generates the templates.			
Decisions	 Segmentation was only performed for T1-weighted images of healthy controls. The version of GNU Octave that we used for segmentation had a slight bug which needed manual intervention before the images could be segmented. The source code for the segmentation script had to be modified. 			
	• Some members of the team opted for an older version of Octave.			
Testing & Verification	SPM produces several files with the segmented data. Files beginning with "c1" is what the algorithm identifies as the gray matter; files beginning with "c2" is what the algorithm identifies as white matter and files beginning with "c3" is what the algorithm identifies as the CSF. Each of these files were visualized for all subjects in AFNI to verify that segmentation was performed correctly.			

2.7 Image Preprocessing

Image pre-processing is the most important aspect of our project. Pre-processing is performed to improve the signal quality of the raw data obtained from the MRI scanner. Image pre-processing involves steps like artifact detection, baseline correction, realignment, movement correction, co-registration, normalization, and smoothing. The image preprocessing is entirely done with AFNI.

The following image preprocessing steps were accomplished:

2.7.1 Preprocessing of BOLD fMR Images

The preprocessing of an fMRI image is basically done to improve the quality of the image so as to analyse it in a better way. Preprocessed images can suppress undesired distortions and enhance some features which are necessary for the particular purpose we are working towards. An fMRI volume contains not only the signal that we are interested in, which are the changes in oxygen levels of blood flowing to a certain brain region, but also fluctuations in the signal that can be caused by a variety of reasons such as involuntary or voluntary movement of subjects during the scan, random drifts, breathing artifacts, and heartbeats. These fluctuation in the signal need to completely reduced if possible otherwise they need to be reduced as much as possible.

The image resolution of a raw fMRI image is extremely low. The fMR signals are incredibly faint and are very brief, due to which they need to be acquired extremely fast before the signal disappears. This is the primary cause of low resolution of fMR images. Preprocessing of BOLD fMR images include reconstruction of the image, along with improving the quality of the image and head motion corrections. Head movement artifacts can lead to misinterpretation of the image data during analysis. Each 3D acquisition in a scan is collected in a small unit of 3D grid, which is referred to as a voxel. A single voxel represents a single image intensity value and ideally, voxels will always represent the same part of the brain in each acquisition, rather than vary from one 3D image to the next. To correct small head motion artifacts, AFNI's motion correction tool employs a linear least squares algorithm that attempts to align each 3D image acquired to the first image acquired in the scan. In the preprocessing of fMRI BOLD data, we will also perform masking. A mask in image processing is analogous to filter in signal processing. The general purpose of filtering and applying masks is to remove unwanted signals, or rather unwanted pixels from an image. In our case, masking is necessary to remove the BOLD data outside of the brain region. Masking was also accomplished using AFNI.

AFNI offers a set of programs that can be used for batch processing using a shell script. We implemented an array of AFNI programs through a shell script to perform preprocessing of BOLD fMRI image data. AFNI further divides the NIfTI files into two separate file formats: 'BRIK' and 'HEAD'. The image binary i.e. the actual 3D volumes are stored in a '.BRIK' and the header information originally contained in the DICOM files is stored in a '.HEAD' file. AFNI uses these files as its inputs.

The following is a brief summary of the tasks performed during this step:

Preprocessing of BOLD FMRI Data					
Status during the reporting period Completed					
Actions	In the preprocessing of BOLD FMRI data we implemented a shell script to adjust the slice timing along with head motion corrections (Appendix D). The slice timing adjustment involves reconstruction of the fMRI image and head motion correction involves re-registrations of the voxels with respect to a base. Furthermore, the functional MR signals, or BOLD data outside the brain region was also masked out using a auto mask feature in AFNI.				
Decisions	 Each 3D brain image is composed of multiple 2D slices and although the slices are acquired at essentially the same time, the duration of the scan separates the first scan from the last. So the 2D slices need to be aligned to the same point in time using interpolation. After this step the 2D image slices are reconstructed into an 3D image. In an MRI imaging system, few MR signals generated at the very beginning of the scan are usually omitted. For this reason, the we excluded the first 5 TRs from our image data. Generally, there lies some outliers in almost every statistical data. For head motion correction, we will use the TR with the least number of outliers in the BOLD EPI as the base. 				
Testing & Verification The file generated at each step was visualized in AFNI. The visuation comparisons and results are discussed in a bit more detail in settion 3.					

2.7.2 Preprocessing: Alignment of BOLD EPI to T1 Image

The second preprocessing step is to align the fMRI image, which was processed in the previous step, on top of the anatomical T1 weighted image. Structural MR image only provides information about brain anatomy. Nonetheless, the structural MRI provides enough information about the structure of the brain that complements functional MRI in a number of ways. Since, the brain function ultimately depends on the integrity of the brain structure, the underlying tissue integrity allows one to examine the functional signals. Essentially, structural MRI provides an anatomical reference for visualization of activation patterns and regions of interest to extract functional connectivity information. This step will allow us to map the BOLD signals in the EPI fMRI image to specific brain regions in the T1-weighted image.

Alignment of BOLD EPI to T1 Image				
Status during the reporting period	Completed			
Actions	In this step, a shell script was run to align the BOLD EPI and the T1 image and to spatially normalize the aligned data to a standard template (Appendix E).			
Decisions	• It was decided that we will use the standard TT_icmb452 brain atlas in talairach space as the template for normalizing our image data.			
Testing & Verification	The file generated at each step was visualized in AFNI. The visual comparisons and results are discussed in a bit more detail in section 3.			

2.8 Creating a Common GM Mask

Once the BOLD EPI image and the T1-weighted are aligned and spatially normalized, we will apply a GM mask to extract the functional connectivity information of only the GM region of the brain. For achieving this, we first need to create a common GM mask for all the subjects.

The following is a brief summary of the tasks performed during this step:

Creating a Common GM Mask				
Status during the reporting period	Started			
Actions	This is step a step in progress in our project. Here we will use AFNI to create a common grey matter mask by using the segmented data generated in the previous step.			
Decisions	• The GM mask created initially created using the segmented image data was improper. Most of the pixels that were supposed to be there were missing. In the next attempt, we will generate another GM mask using the data that was aligned in the previous step.			
Testing & Verification	The file generated after the first attempt were visualized in AFNI. After visualizations, we found most of the image pixels were missing form the GM mask.			

3 Results and Discussions

The following section discusses on the results achieved after the above steps:

3.1 Data Selection

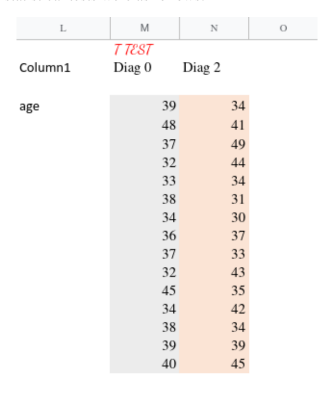
The following image represents an excel table for the selected participants.

	A	В	С	D	E	F	G
1	participant_id	site	diagnosis	age	sex	hand	BDI-II
2	sub-0236	HUH	0	39	1	1	0
3	sub-0240	HUH	0	48	1	1	4
4	sub-0244	HUH	0	37	2	1	3
5	sub-0292	HUH	0	32	2	1	2
- 6	sub-0294	HUH	0	33	1	1	4
7	sub-0310	HUH	0	38	1	1	0
8	sub-0312	HUH	0	34	2	1	6
9	sub-0320	HUH	0	36	2	1	4
10	sub-0322	HUH	0	37	2	1	5
11	sub-0326	HUH	0	32	1	1	6
12	sub-0338	HUH	0	45	2	1	1
13	sub-0346	HUH	0	34	1	1	0
14	sub-0353	HUH	0	38	1	1	0
15	sub-0355	HUH	0	39	2	1	5
16	sub-0359	HUH	0	40	2	1	5
17	sub-0249	HUH	2	34	2	1	39
18	sub-0259	HUH	2	41	1	1	33
19	sub-0265	HUH	2	49	1	1	33
20	sub-0267	HUH	2	44	2	1	33
21	sub-0273	HUH	2	34	2	1	37
22	sub-0277	HUH	2	31	2	1	47
23	sub-0285	HUH	2	30	1	1	40
24	sub-0287	HUH	2	37	2	1	39
25	sub-0289	HUH	2	33	2	1	53
26	sub-0299	HUH	2	43	1	1	36
27	sub-0301	HUH	2	35	1	1	34
28	sub-0319	HUH	2	42	2	1	46
29	sub-0323	HUH	2	34	2	1	42
30	sub-0331	HUH	2	39	1	1	43
31	sub-0349	HUH	2	45	2	1	32

Figure 2: Excel table containing selected subjects

Once the participants were selected, statistical tests were performed to test the goodness of fit based on age and sex.

The results of the statistical tests were as follows:



Ttest P value 0.75152766

Figure 3: Pivot table for T-Test

For the t-test, our null hypothesis is that the true age difference between the subjects from the two categories are zero. Since, the p-value is greater than 0.05, we accept our hypothesis and conclude that the selected subjects are well-matched based on their age.

T	U	U		W
	CHI TEST			
Count of sex	diagnosis			
sex		0	2	Grand Total
1		7	6	13
2		8	9	17
Grand Total		15	15	30
Count of sex	Column Labels			
Row Labels		0	2	Grand Total
F		8.5	8.5	17
M		6.5	6.5	13
Grand Total		15	15	30

CHI TEST P value 0.128735049

Figure 4: Pivot table for Chi-square Test

For, the chi-square test, out null hypothesis is that there is no significant difference between observed and expected frequency of males and females. Since, the p-value is greater than 0.05, it means the selection of males and females have equal probability.

3.2 Preprocessing Results

By running the command of 3dskullstrip, the scans of skull were removed from the structural images i.e, T1 images of all the subjects. The skull stripped structural images of all subjects were viewed in MRIcron so as to conform the error free command execution.

(skull stripped images)

3.4 nikin Image Segmentation Running the HC subjects for segmentation process, results the formation of rc1 images for gray matter, rc2 images for white matter and rc3 images for CSF that will be used for the common masking. (segmented images)

3.5 nikin Scripts Preprocessing

4 Conclusion and Further Work

As a closing note for the following progress report, the extracted dataset are statistically analysed so as to obtain the accuracy which would remove the differences that are created from manual analysis. All the MDD and HC subjects are also converted to NIfTI format to access in the AFNI formats from DICOM.Also, we successfully segmented the structural images of HC that resulted in isolation of gray matter, white matter and CSF which will further be used in common masking. Also, the images are skull stripped using AFNI commands to each subjects and are further used to preprocessed so that the images would be free from noises and any distortions by running bash scripts to each subjects. Thus these are the tasks that are performed so far to meet our objectives and successfully achieved our hypothetical results. Mentioning about our further works, in a preprocessed images, statistical tests will be implemented to for a thorough analysis of the functional connectivity of the seed. Specifically, we plan to assess functional connectivity between various regions of the brain and hippocampal area. To assess functional connectivity in the brain region, Resting-state analyses, that is, time series correlations in BOLD fMRI data acquired in a task-free state will be used. A statistical approach to image analysis makes it possible to discover spatial and temporal patterns that correspond to the performance of specific tasks and specific diagnoses. Such statistical methods have only begun to be applied to clinical disorders but show promise for increasing the "specificity" of brain imaging markers for mental illness.

References

Appendix

A Data Selection

B Data Prep

```
#! /bin/bash
function appendDotnii () {
  for file in $(ls); do
    if [[ -f $file ]]
    then
     mv "\{file\}" "\{basename \{file\}\}.nii"
 done
for category in $(ls -d */); do
 pushd $category
  for subject in (ls -d */); do
    pushd $subject
   pushd rsfmri/
   appendDotnii
   popd
   popd
 done
 popd
done
```

C Skull Stripping

```
#! /bin/bash
WORKING DIR=$HOME/Functional-Connectivity/Subjects/
# Create an array with the names of the directory in which the data
# for healthy controls and majorly depressed patient are stored.
CATEGORIES=(HC MDD)
# Change into the working directory if it exists
if [[ -d "${WORKING_DIR}" ]]
then
 pushd ${WORKING_DIR}
if [ ! -d ''Skull_Stripped_Data'' ] ; then
   mkdir -p Skull_Stripped_Data/HC
   mkdir -p Skull_Stripped_Data/MDD
for CATEGORY in "${CATEGORIES[@]}"
do
 pushd ${CATEGORY}
  for SUBJECT in $(ls)
   pushd ${SUBJECT}/t1/
    SUBJECT ID=$(echo "${SUBJECT}" | cut -d '-' -f 2)
    3dSkullStrip -input defaced_mprage.nii \
```

D Preprocessing: BOLD FMRI Data

```
#! /bin/bash
# In this script you will preprocess BOLD data (4D fMRI data) #
# including slicing timing and head motion correction
WORKING_DIR=''$HOME/Functional-Connectivity/Processed_Data/''
"BOLD_fMRI_Data"
# Create an array with the names of the directory in which the data
# for healthy controls and majorly depressed patient are stored.
CATEGORIES= (HC MDD)
# Move to the working directory
if [ -d "${WORKING_DIR}" ]
then
 pushd ${WORKING_DIR}
for CATEGORY in "${CATEGORIES[@]}"
 pushd ${CATEGORY}
 for SUBJECT in $(ls *.nii)
   SUBJECT_ID=$(basename -s .nii ${SUBJECT} | cut -d '_' -f 2)
   # Convert a dataset from NIfTI to .BRIK and .HEAD
   # -verbose because I want to be able to see what's going on #
   3dcalc -prefix ${SUBJECT_ID}-BOLD-EC-tmp
-a ${SUBJECT} -expr 'a' -verbose
   # Exclude the first 5 TRs
   # In the previous step, ${SUBJECT_ID}-BOLD-EC-tmp+orig.HEAD and
# ${SUBJECT_ID}-BOLD-EC-tmp+orig.BRIK files were created.
   3dcalc -prefix ${SUBJECT_ID}-BOLD-EC1
     -a ${SUBJECT ID}-BOLD-EC-tmp+orig[5..$] -expr 'a'
   ## Despiking:
   3dDespike -prefix ${SUBJECT_ID}-BOLD-EC
${SUBJECT_ID}-BOLD-EC1+orig
   ## Count the outliers in each TR
   3dToutcount -automask -range ${SUBJECT_ID}-BOLD-EC+orig
     > outliers-BOLD-EC.1D
```

```
## Find the TR with least outliers:
    ## This is a perl script.
    base=$(cat outliers-BOLD-EC.1D |
      perl -0777an -F"\n" -e
       $i=0;
       $small=999999;
       map {
         /\s*(\d+)/;
         if ($small > $1) {
           \$small = \$1;
           $ind=$i;
         } ;
       $i++;
       } @F;
       print $ind')
    # Using the TR with least outliers as base for head motion
    # correction and spatial normalization
    3dcalc -prefix ${SUBJECT_ID}-BOLD-EC-base
      -a "${SUBJECT_ID}-BOLD-EC+orig[${base}]" -expr 'a'
    # Slice timing and head motion correction
    # head motion parameters are stored in
    # ${SUBJECT_ID}-BOLD-EC-motion.1D
    \# This is the same with running 3dTshift first and then use
    # 3dvolreg itself. However, due to the lack of slice scan order,
    # we have to ignore this slice time step.
    3dvolreg -verbose -tshift 0
      -base ${base}
      -1Dfile ${subject}-BOLD-EC-motion.1D
      -prefix ${SUBJECT_ID}-BOLD-EC-volreg
      ${SUBJECT_ID}-BOLD-EC+orig
    rm -f ${SUBJECT_ID}-BOLD-EC+orig*
    ## Brain mask of the subject
    3dAutomask -prefix ${SUBJECT_ID}-BOLD-EC-mask
      -dilate 1 ${SUBJECT_ID}-BOLD-EC-volreg+orig
    # Maskout the functional BOLD outside of the brain
    3dcalc -prefix ${SUBJECT_ID}-BOLD-EC-volreg-mask
      -a ${SUBJECT_ID}-BOLD-EC-volreg+orig
-b ${SUBJECT_ID}-BOLD-EC-mask+orig -expr 'a*b'
    gzip *.BRIK
  done
  popd
done
```

E Preprocessing: Alignment of BOLD EPI to T1 Image

```
#! /bin/bash
# In this script we will spatially normalize the anatomical dataset
# into TT space, we will also use the same transformation to normalize
# the BOLD dataset into the MNI space.

WORKING_DIR="${HOME}/Functional-Connectivity/Processed_Data/"\
"Spatially_Normalized_Data"
```

```
# Create an array with the names of the directory in which the data
\# for healthy controls and majorly depressed patient are stored.
CATEGORIES= (HC MDD)
# Change into the working directory if it exists
if [ -d "${WORKING_DIR}" ]
then
   pushd ${WORKING_DIR}
    echo "${WORKING_DIR}: doesn't exist"
   exit 111;
fi
# Use a nested loop to get into the directory for each subject of each
# category
for CATEGORY in "${CATEGORIES[@]}"
 pushd ${CATEGORY}
  for SUBJECT in $(ls)
    # Grab the SUBJECT ID
    # Subjects are stored in NORM-SUBJECT_ID
    # Split NORM-SUBJECT ID at the delimiter and return the
    # specified field
    SUBJECT_ID=$(echo "${SUBJECT}" | cut --delimiter '-' --fields 2)
    pushd ${SUBJECT}
    # Spatially normalize the anatomical dataset to MNI space #
    # Uniformly distribute the white matter in the brain tissue.
    3dUnifize -prefix BOLD-${SUBJECT_ID}-EC-T1
    -input ANAT-${SUBJECT_ID}.nii
                                                                 \
    # Remove the skull and extract the brain tissue from T1-weighted
    # MR image.
    3dSkullStrip -prefix ${SUBJECT ID}-T1-NoSkull
        -input BOLD-${SUBJECT_ID}-EC-T1+tlrc.
    # Shift the center of DSET to the center of BASE. '_shft' will be
    # appended at the end of DSET. Use the center of mass of the
    # volume as the center (By default, center is the center of
    # volume's grid).
    @Align_Centers -cm
        -base ${SUBJECT_ID}-BOLD-EC-base+orig.
        -dset ${SUBJECT_ID}-T1-NoSkull+tlrc.
    # Linearly align the anatomical dataset with the EPI dataset. The
    # EPI dataset cab be the functional-MR image. The -epi_base option
    # specifies the starting sub-brick for the alignment.
   align_epi_anat.py -anat ${SUBJECT_ID}-T1-NoSkull_shft+tlrc.
    -epi ${SUBJECT_ID}-BOLD-EC-base+orig.
        -epi_base 0
        -suffix _alBOLDEC
        -anat_has_skull no
        -epi_strip 3dAutomask
        -volreg off
       -tshift off
        -resample off
```

```
# Create a symbolic link for the atlas you want to use. Atlases
# are downloaded during AFNI installation. To find the location of
# the atlases run 'afni_system_check.py -check_all | grep atlas'.
\# If the atlases were not downloaded; download them from
# "https://bit.ly/3BpKN2K". Once the tarball has been extracted
# and placed into whichever location the afni binaries are stored
# in, they are immediately available for use.
# We will use TT_icbm452+tlrc atals
ln -sf /opt/afni/TT_icbm452+tlrc.* ./
# Shift (roughly) the center of the aligned anatomical dataset
\# with the standard template. The transformation information will
# be stored in the base image of BOLD data as well. Again,
# "_shft" will be appended to the DSET and CHILD.
@Align_Centers -base ./TT_icbm452+tlrc.
    -dset ${SUBJECT_ID}-T1-NoSkull_shft_alBOLDEC+tlrc.
    -child ${SUBJECT_ID}-BOLD-EC-base+orig.
# Align SOURCE to BASE and save the transformation matrix for each
# sub-brick into a 1D file. The cost function that defines the
# matching between SOURCE and BASE is the lpa (Local Pearson
# Correlation Abs). Also comput a weight function using 3dAutomask
# alogrithm plus some blurring of the base image.
3dAllineate -prefix ${SUBJECT_ID}-T1_to_T1_Allineate
    -base ./TT icbm452+tlrc.
    -source ${SUBJECT_ID}-T1-NoSkull_shft_alBOLDEC_shft+tlrc.
    -1Dmatrix_save T1_to_T1_Allineate.aff12.1D
    -source_automask
    -cost lpa
    -autoweight
    -cmass
# Compute a non-linearly warped version of SOURCE dataset to match
# the BASE dataset. Gaussian blur the SOURCE and BASE before doing
\sharp the alignmnet. By default, the blur values is 2.345 (for no good
# reason). Set the blur values to 0 if you do not want to blur the
# inputs.
3dQwarp -prefix T1-NoSkull_shft_alBOLDEC_shft
    -blur 0 0
    -base ./TT_icbm452+tlrc.
    -source ${SUBJECT_ID}-T1_to_T1_Allineate+tlrc
\# So, the 3dcopy command seems to overwrite one of the files \# created above. The old file needs to be deleted for 3dcopy to
# execute. ${SUBJECT_ID}-T1-NoSkull_shft_alBOLDEC_shft+tlrc.* will
# be recreated by 3dcopy.
rm --force ${SUBJECT_ID}-T1-NoSkull_shft_alBOLDEC_shft+tlrc.*
# Copy one dataset to the other. 3dCopy foo bar copies foo+orig.
# to bar+orig. and foo+tlrc. to bar+tlrc. This program copies
# entire datasets and not jsut sub-bricks.
3dcopy T1-NoSkull_shft_alBOLDEC_shft
    ${SUBJECT_ID}-T1-NoSkull_shft_alBOLDEC_shft
# Spatially normalize the BOLD dataset to MNI space #
# Create a copy of the original volume registartion mask.
3dcopy ${SUBJECT_ID}-BOLD-EC-volreg-mask+orig.
    ${SUBJECT_ID}-BOLD-EC-volreg-mask_temp
```

```
# Shift (roughly) the center of the 4D BOLD fMRI data to the space
    # using the same parameters as the base image. The base image has
    # similar alignmnet as the anatomical image. (As explained
    # earlier).
   @Align_Centers -cm
        -base ${SUBJECT_ID}-BOLD-EC-base_shft+orig
       -dset ${SUBJECT ID}-BOLD-EC-volreg-mask temp+orig.
    ## Apply a nonlinear transformation and resample to 3x3x3 mm
    3dNwarpApply -prefix ${SUBJECT_ID}-BOLD-EC-volreg-Nwarp
       -source ${SUBJECT_ID}-BOLD-EC-volreg-mask_temp_shft+orig.
        -nwarp 'T1-NoSkull_shft_alBOLDEC_shft_WARP+tlrc. '\
       'T1_to_T1_Allineate.aff12.1D'
    3dcopy ${SUBJECT_ID}-BOLD-EC-volreg-Nwarp+orig
        ${SUBJECT_ID}-BOLD-EC-volreg-Nwarp_Backup
    3drefit -view tlrc ${SUBJECT_ID}-BOLD-EC-volreg-Nwarp+orig
    ## get tissuse-based signal before smooth
    3dDeconvolve -float -polort A
       -errts ${SUBJECT_ID}-BeforeSmooth-lp
       -bucket BeforeSmooth-bucket
       -input ${SUBJECT_ID}-BOLD-EC-volreg-Nwarp+tlrc.
    3dBandpass -band 0.01 0.08
        -prefix ${SUBJECT_ID}-BeforeSmooth-lp-bp
       -input ${SUBJECT_ID}-BeforeSmooth-lp+tlrc
    # Spatial smoothing with 6 mm FWHM. The value for FWHM is
    # adjustable
    3dmerge -doall
       -1blur_fwhm 6
       -prefix ${SUBJECT_ID}-BOLD-EC-volreg-Nwarp-blur6
       ${SUBJECT_ID}-BOLD-EC-volreg-Nwarp+tlrc
    # Linear detrending
    3dDeconvolve -float -polort A
       -bucket bucket
       -errts ${SUBJECT_ID}-BOLD-EC-volreg-Nwarp-blur6-lp
       -input ${SUBJECT ID}-BOLD-EC-volreg-Nwarp-blur6+tlrc.
    # Temporal filtering band pass 0.01-0.08 Hz
    3dBandpass -band 0.01 0.08
       -prefix ${SUBJECT_ID}-BOLD-EC-volreg-Nwarp-blur6-lp-bp
       -input ${SUBJECT_ID}-BOLD-EC-volreg-Nwarp-blur6-lp+tlrc
    3dresample -master ./TT_icbm452+tlrc.
      -input _{s}
     -prefix ${SUBJECT_ID}-BOLD-EC-volreg-Nwarp-blur6-lp-bp-resampled
   gzip *.BRIK
   popd
 done
 popd
done
popd
```

F Gantt Chart (Unchanged)

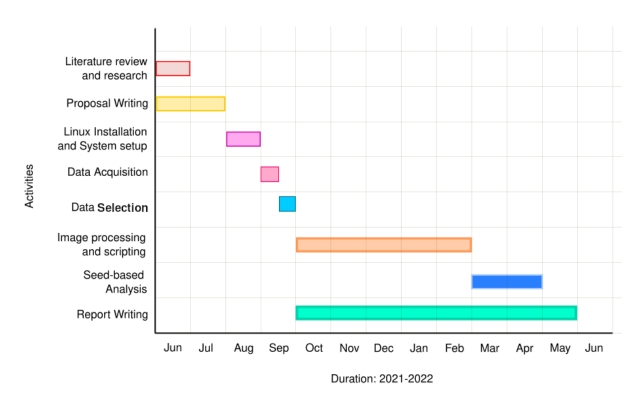


Figure 5: Proposed Workflow (Unchanged)