## **Tutorial of NeuroRA Version 1.0.8.1**

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This Tutorial of NeuroRA provides information on how to use the NeuroRA including its easy-to-use functions.

Before you read it, you only need to spend a little time learning the basic Python syntax and this toolkit is easy to understand. In addition, it would be better if you are familiar with Python, especially the matrix operations based on NumPy.

If there is anything wrong, difficult to understand or having any useful advice during reading it, you can contact me (zitonglu1996@gmail.com) and I will be happy and thankful to know about it.

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This tutorial consists of these parts:

- Introduction & Installation
- Data Conversion
- Calculate the neural pattern similarity (NPS)
- Calculate the RDM (Representational Dissimilarity Matrices)
- Calculate the correlation coefficient between RDMs
- Visualization for results
- Save as a NIfTI file (for fMRI)
- Others
- Demo

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### **Part 1: Introduction**

**NueorRA** is a Python toolbox for multimode neural data representational analysis.



### **Overview**

Representational Similarity Analysis (RSA) has become a popular and effective method to measure the representation of multivariable neural activity in different modes.

NeuroRA is a novel and easy-to-use toolbox based on Python, which can do some works about RSA among nearly all kinds of neural data, including behavioral, EEG, MEG, fNIRS, ECoG, electrophysiological and fMRI data.

### **Installation**

pip install NeuroRA

## **Required Dependencies**

Numpy: a fundamental package for scientific computing

Matplotlib: a Python 2D plotting library

NiBabel: a package prividing read +/- write access to some common medical

and neuroimaging file formats

*Nilearn*: a Python module for fast and easy statistical learning on NeuroImaging data.

*MNE-Python*: a Python software for exploring, visualizing, and analyzing human neurophysiological data.

## **Paper**

Lu, Z., & Ku, Y. *NeuroRA: A Python toolbox of representational analysis from multi-modal neural data.* (bioRxiv: <a href="https://doi.org/10.1101/2020.03.25.008086">https://doi.org/10.1101/2020.03.25.008086</a>)

# **Part 2: Data Conversion**

transformation

Type of Neural Bata	Data Conversion Scheme	
	Use Nibabel (https://nipy.org/nibabel/) to load fMRI data.	
	import nibabel as nib	
<i>e</i> MDI		
fMRI	fmrifilename = "demo.nii" # the fmri data file name with full address	
	data = nib.load(fmrifilename).get_fdata() # load fMRI data as ndarray	
	Use MATLAB EEGLab (http://sccn.ucsd.edu/eeglab/) to do preprocessing and	
EEG/MEG	obtain .mat files, and use <i>Scipy</i> (https://www.scipy.org) to load EEG data (.mat).	
	import scipy.io as sio	
	filename = "demo.mat" # the EEG/MEG data file name with full address	
	data = sio.loadmat(filename)["data"] # load EEG/MEG data as ndarray	
	Or use MNE (https://mne-tools.github.io) to do preprocessing and return ndarray-	
	type data.	
fNIRS	For raw data from device, use Numpy (http://www.numpy.org) to load fNIRS data	
	(.txt or .csv).	
	import numpy as np	
	txtfilename = "demo.txt" # the fNIRS data file name with full address	
	csvfilename = "demo.csv"	
	data = np.loadtxt(txtfilename) # load fNIRS data as ndarray	
	data = np.loadtxt(csvfilename, delimiter, usecols, unpack)	
ECoG/sEEG	Use Brainstorm (https://neuroimage.usc.edu/brainstorm/) to do preprocessing and	
	obtain .mat files, and use Scipy to load ECoG data (.mat).	
	Use <i>pyABF</i> (https://github.com/swharden/pyABF) to load electrophysiology data	
Electrophysiology	(.abf).	
	import pyabf	
	import pyddi	
	abf = pyabf.ABF("demo.abf") # the electrophysiology data file name with full	
Licetrophysiology	address	
	abf.setSweep(sweepNumber, channel) # access sweep data	
	data = abf.sweepY # get sweep data with sweepY	
Two functions. NumPy	//reshape() & NumPy.transpose(), are recommended for further data	

## Part 3: Calculate the Neural Pattern Similarity

Module: nps\_cal.py

• nps(data, time\_win=5, sub\_opt=0)

A function for calculating the neural pattern similarity

#### Parameters:

data: array

The neural data. The shape of bhv\_data must be [2, n\_subs, n\_trials, n\_chlas, n\_ts]. 2 represent 2 different conditions. n\_subs, n\_trials, n\_chls, n\_ts represent the number of subjects, the number of trials, the number of channels & the number of time-points.

time\_win: int

(Only when time\_opt = 1, time\_win works) The time-window

for each calculation.

sub\_opt: int(0/1)

Calculate the NPS for each subject or not. 1 or 0.

Returns:

nps: array

If sub\_opt=0, return nps (shape: [n\_chls, n\_ts/time\_win]). If

sub\_opt=1, return nps (shape: [n\_subs, n\_chls,

n\_ts/time\_win]).

nps\_fmri(fmri\_data, time\_win=5, sub\_opt=0)

A function for calculating the neural pattern similarity of fMRI data

#### Parameters:

fmri\_data: array

The fmri data. The shape of fmri\_data must be [n\_cons, n\_subs, nx, ny, nz]. n\_cons, n\_subs, nx, ny, nz represent the number of conditions, the number of subjects, the size of the

fMRI data.

ksize: array / list

ksize=[kx, ky, kz] represents that the calculation unit

contains k1\*k2\*k3 voxels.

strides: array / list

strides=[sx, sy, sz] represents the moving steps along the x, y,

z.

Returns:

nps: array

NPS. The shape of nps is  $[n_x, n_y, n_z, 2]$ .

### nps\_fmri\_roi(fmri\_data, mask\_data)

A function for calculating the neural pattern similarity of fMRI data for ROI

#### Parameters:

fmri\_data: array

The fmri data. The shape of fmri\_data must be [n\_cons, n\_subs, nx, ny, nz]. n\_cons, n\_subs, nx, ny, nz represent the number of conditions, the number of subjects, the size of the

fMRI data.

mask\_data: array

The fmri data. The shape of mask\_data must be [nx, ny, nz].

nx, ny, nz represent the size of the fMRI data.

Returns:

nps: array

NPS. The shape of nps is [2].

### Part 4: Calculate the RDM

### Module: *rdm\_cal.py*

bhvRDM(bhv data, sub opt=0, data opt=1)

A function for calculating the RDM based on behavioral data

#### Parameters:

bhv\_data: array

> The behavioral data. If data\_opt=0, the shape of bhv\_data must be [n\_cons, n\_subs]. n\_cons, n\_subs represent the number of conditions & the number of subjects.

sub\_opt: int(0/1)

Calculate the RDM for each subject or not. 1 or 0.

int (0 / 1) data opt:

> If data\_opt=1, each subject's each trial has a value of data. If data\_opt=0, each subject has a value of data, here ignore the

effect of trials.

#### Returns:

rdm/rdms: array

If sub\_opt=0, return only one rdm (shape: [n\_cons, n\_cons]). If sub\_opt=1, return rdms (shape: [n\_subs, n\_cons, n\_cons]).

eegRDM(EEG\_data, time\_win=5, sub\_opt=0, chl\_opt=0, time\_opt=0)

A function for calculating the RDM based on EEG/MEG/fNIRS data

#### Parameters:

EEG\_data: array

> The EEG/MEG/fNIRS data. The shape of EEG\_data must be [n\_cons, n\_subs, n\_trials, n\_chls, n\_ts]. n\_cons, n\_subs, n\_trials, n\_chls, n\_ts represent the number of conditions, subjects, trials, channels, frequencies and time-points.

time\_win: int

(Only when time\_opt = 1, time\_win works) The time-window

for each calculation.

sub\_opt: int(0/1)

Calculate the RDM for each subject or not. 1 or 0.

chl\_opt: int(0/1)

Calculate the RDM for each channel or not. 1 or 0.

time\_opt: int(0/1)

Calculate the RDM for each time-point or not. 1 or 0.

#### Returns:

rdm/rdms: array

If sub\_opt=0 and chl\_opt=0 and time\_opt=0, return only one rdm (shape: [n\_cons, n\_cons]).

If sub\_opt=0 and chl\_opt=0 and time\_opt=1, return rdms (shape: [n\_ts/time\_win, n\_cons, n\_cons]).

If sub\_opt=0 and chl\_opt=1 and time\_opt=0, return rdms (shape: [n\_chls, n\_cons, n\_cons]).

If sub\_opt=1 and chl\_opt=0 and time\_opt=0, return rdms (shape: [n\_subs, n\_cons, n\_cons]).

If sub\_opt=0 and chl\_opt=1 and time\_opt=1, return rdms (shape: [n\_chls, n\_ts/time\_win, n\_cons, n\_cons]).

If sub\_opt=1 and chl\_opt=0 and time\_opt=1, return rdms (shape: [n\_subs, n\_ts/time\_win, n\_cons, n\_cons]).

If sub\_opt=1 and chl\_opt=1 and time\_opt=0, return rdms (shape: [n\_subs, n\_chls, n\_cons, n\_cons]).

If sub\_opt=1 and chl\_opt=1 and time\_opt=1, return rdms (shape: [n\_subs, n\_chls, n\_ts/time\_win, n\_cons, n\_cons]).

## ecogRDM(ele\_data, time\_win=5, opt="all")

A function for calculating the RDM based on ECoG/electrophysiological data

### Parameters:

ele\_data: array

The ECoG/electrophysiological data. The shape of ele\_data must be [n\_cons, n\_trials, n\_chls, n\_ts]. n\_cons, n\_trials, n\_chls, n\_ts represent the number of conditions, trials,

channels, frequencies and time-points.

time\_win: int

The time-window for each calculation.

opt: string ("channel" or "time" or "all")

Calculate the RDM for each channel or for each time-point or

not. "channel" or "time" or "all".

#### Returns:

rdm/rdms: array

If opt="channel", return rdms (shape: [n\_chls, n\_cons,

n\_cons]).

If opt="time", return rdms (shape: [n\_ts/time\_win, n\_cons,

n\_cons])

### fmriRDM(fmri\_data, ksize=[3, 3, 3], strides=[1, 1, 1])

A function for calculating the RDM based on fMRI data

#### Parameters:

fmri\_data: array

The fmri data. The shape of fmri\_data must be [n\_cons, n\_subs, nx, ny, nz]. n\_cons, n\_subs, nx, ny, nz represent the number of conditions, the number of subjects, the size of the

fMRI data.

ksize: array / list

ksize=[kx, ky, kz] represents that the calculation unit

contains k1\*k2\*k3 voxels.

strides: array / list

strides=[sx, sy, sz] represents the moving steps along the x, y,

z.

#### Returns:

rdms: array

Return rdms for each calculation unit. The shape of rdms is [n\_x, n\_y, n\_z, n\_cons, n\_cons]. Here, n\_x, n\_y, n\_z represent the number of calculation units along the x, y, z.

### fmriRDM\_roi(fmri\_data, mask\_data)

A function for calculating the RDM based on fMRI data of a ROI

#### Parameters:

fmri\_data: array

The fmri data. The shape of fmri\_data must be [n\_cons, n\_subs, nx, ny, nz]. n\_cons, n\_subs, nx, ny, nz represent the number of conditions, the number of subjects, the size of the

fMRI data.

mask\_data: array

The mask data. The shape of mask\_data must be [nx, ny, nz].

Returns:

rdm: array

Return the rdm for the ROI. The shape of rdm is [n\_cons,

n\_cons]

### Part 5: Calculate the Correlation Coefficient

## Module: *rdm\_corr.py*

### rdm\_correlation\_spearman(RDM1, RDM2, rescale=False)

A function for calculating the Spearman correlation coefficient between two RDMs

#### Parameters:

RDM1: array

The shape of RDM1 must be [n\_cons, n\_cons].

RDM2: array

The shape of RDM2 must be [n\_cons, n\_cons].

rescale: Boolean (True or False)

Rescale the values in RDM or not. True or False.

### Returns:

corr: array

The corr contains two values: correlation coefficient and p-

value. The shape of corr is [2,].

### rdm\_correlation\_pearson(RDM1, RDM2, rescale=False)

A function for calculating the Pearson correlation coefficient between two RDMs

#### Parameters:

RDM1: array

The shape of RDM1 must be [n\_cons, n\_cons].

RDM2: array

The shape of RDM2 must be [n\_cons, n\_cons].

rescale: Boolean (True or False)

Rescale the values in RDM or not. True or False.

Returns:

corr: array

The corr contains two values: correlation coefficient and p-

value. The shape of corr is [2,].

### rdm\_correlation\_kendall(RDM1, RDM2, rescale=False)

A function for calculating the Kendalls tau correlation coefficient between two RDMs

#### Parameters:

RDM1: array

The shape of RDM1 must be [n\_cons, n\_cons].

RDM2: array

The shape of RDM2 must be [n\_cons, n\_cons].

rescale: Boolean (True or False)

Rescale the values in RDM or not. True or False.

Returns:

corr: array

The corr contains two values: correlation coefficient and p-

value. The shape of corr is [2,].

#### rdm\_similarity(RDM1, RDM2, rescale=False)

A function for calculating the Cosine Similarity between two RDMs

#### Parameters:

RDM1: array

The shape of RDM1 must be [n\_cons, n\_cons].

RDM2: array

The shape of RDM2 must be [n\_cons, n\_cons].

#### Returns:

similarity: float

The Cosine Similarity.

### rdm\_distance(RDM1, RDM2, rescale=False)

A function for calculating the Euclidean Distances between two RDMs

#### Parameters:

RDM1: array

The shape of RDM1 must be [n\_cons, n\_cons].

RDM2: array

The shape of RDM2 must be [n\_cons, n\_cons].

rescale: Boolean (True or False)

Rescale the values in RDM or not. True or False.

Returns:

dist: float

The Euclidean Distance.

### rdm\_permutation (RDM1, RDM2, iter=1000)

A function for permutation test between two RDMs

#### Parameters:

RDM1: array

The shape of RDM1 must be [n\_cons, n\_cons].

RDM2: array

The shape of RDM2 must be [n\_cons, n\_cons].

iter: int

The number of iterations.

#### Returns:

p: float

The p-value.

### Module: corr\_cal.py

bhvANDeeg\_corr(bhv\_data, eeg\_data, sub\_opt=0, bhv\_data\_opt=1, chl\_opt=0, time\_opt=0, time\_win=5, method="spearman", rescale=False)

A function for calculating the Similarity/Correlation Coefficient between behavioral data and EEG/MEG/fNIRS data

#### Parameters:

bhv\_data: array

The behavioral data. If data\_opt=0, the shape of bhv\_data must be [n\_cons, n\_subs]. n\_cons, n\_subs represent the number of conditions & the number of subjects.

eeg\_data: array

The EEG/MEG/fNIRS data. The shape of eeg\_data must be [n\_cons, n\_subs, n\_trials, n\_chls, n\_ts]. n\_cons, n\_subs, n\_trials, n\_chls, n\_ts represent the number of conditions, subjects, trials, channels, frequencies and time-points.

sub\_opt: int(0/1)

Calculate the RDM for each subject or not. 1 or 0.

bhv\_data\_opt: int(0/1)

If bhv\_data\_opt=1, each subject's each trial has a value of data. If bhv\_data\_opt=0, each subject has a value of data, here ignore the effect of trials.

time\_win: int

The time-window for each calculation.

chl\_opt: int(0/1)

Calculate the RDM for each channel or not. 1 or 0.

time\_opt: int(0/1)

Calculate the RDM for each time-point or not. 1 or 0.

method: string ("spearman" or "pearson" or "kendall" or "similarity"

or "distance")

They correspond to different methods of calculation. "spearman" or "pearson" or "kendall" or "similarity" or

"distance".

rescale: Boolean (True or False)

Rescale the values in RDM or not. True or False.

#### Returns:

corr/corrs: array

The correlation coefficients corresponding to the RDMs.

bhvANDecog\_corr(bhv\_data, ele\_data, time\_win=5, ecog\_opt="allin", method="spearman", rescale=False)

A function for calculating the Similarity/Correlation Coefficient between behavioral data and ECoG/electricophysiological data

#### Parameters:

bhv\_data: array

The behavioral data. If data\_opt=0, the shape of bhv\_data must be [n\_cons, n\_subs]. n\_cons, n\_subs represent the number of conditions & the number of subjects.

ele\_data: array

The ECoG/electrophysiological data. The shape of ele\_data must be [n\_cons, n\_trials, n\_chls, n\_ts]. n\_cons, n\_trials, n\_chls, n\_ts represent the number of conditions, trials,

channels, frequencies and time-points.

time\_win: int

The time-window for each calculation.

opt: string ("channels" or "time" or "all")

Calculate the RDM for each channel or for each time-point or

not. "channels" or "time" or "all".

method: string ("spearman" or "pearson" or "kendall" or "similarity"

or "distance")

They correspond to different methods of calculation. "spearman" or "pearson" or "kendall" or "similarity" or

#### "distance".

rescale: Boolean (True or False)

Rescale the values in RDM or not. True or False.

#### Returns:

corr/corrs: array

The correlation coefficients corresponding to the RDMs.

bhvANDfmri\_corr(bhv\_data, fmri\_data, bhv\_data\_opt=1, ksize=[3, 3, 3], strides=[1, 1, 1], method="spearman", rescale=False)

A function for calculating the Similarity/Correlation Coefficient between behavioral data and fMRI data

#### Parameters:

bhv\_data: array

The behavioral data. If data\_opt=0, the shape of bhv\_data must be [n\_cons, n\_subs]. n\_cons, n\_subs represent the number of conditions & the number of subjects.

fmri\_data: array

The fmri data. The shape of fmri\_data must be [n\_cons, n\_subs, nx, ny, nz]. n\_cons, n\_subs, nx, ny, nz represent the number of conditions, the number of subjects, the size of the fMRI data.

bhv\_data\_opt: int(0/1)

If bhv\_data\_opt=1, each subject's each trial has a value of data. If bhv\_data\_opt=0, each subject has a value of data, here ignore the effect of trials.

ksize: array / list

ksize=[kx, ky, kz] represents that the calculation unit contains k1\*k2\*k3 voxels.

strides: array / list

strides=[sx, sy, sz] represents the moving steps along the x, y, z.

method: string ("spearman" or "pearson" or "kendall" or "similarity" or "distance")

They correspond to different methods of calculation.

"spearman" or "pearson" or "kendall" or "similarity" or

"distance".

rescale: Boolean (True or False)

Rescale the values in RDM or not. True or False.

Returns:

corrs: array

The correlation coefficients corresponding to the RDMs. The

shape of corrs is  $[n_x, n_y, n_z, 2]$ .

eegANDfmri\_corr(eeg\_data, fmri\_data, chl\_opt=0, ksize=[3, 3, 3], strides=[1,
1, 1], method="spearman", rescale=False)

A function for calculating the Similarity/Correlation Coefficient between EEG/MEG/fNIRS data and fMRI data

#### Parameters:

eeg\_data: array

The EEG/MEG/fNIRS data. The shape of eeg\_data must be [n\_cons, n\_subs, n\_trials, n\_chls, n\_ts]. n\_cons, n\_subs, n\_trials, n\_chls, n\_ts represent the number of conditions, subjects, trials, channels, frequencies and time-points.

fmri\_data: array

The fmri data. The shape of fmri\_data must be [n\_cons, n\_subs, nx, ny, nz]. n\_cons, n\_subs, nx, ny, nz represent the number of conditions, the number of subjects, the size of the

fMRI data.

chl\_opt: int(0/1)

Calculate the RDM for each channel or not, 1 or 0.

ksize: array / list

ksize=[kx, ky, kz] represents that the calculation unit

contains k1\*k2\*k3 voxels.

strides: array / list

strides=[sx, sy, sz] represents the moving steps along the x, y,

z.

method: string ("spearman" or "pearson" or "kendall" or "similarity"

or "distance")

They correspond to different methods of calculation. "spearman" or "pearson" or "kendall" or "similarity" or

"distance".

rescale: Boolean (True or False)

Rescale the values in RDM or not. True or False.

**Returns:** 

corrs: array

The correlation coefficients corresponding to the RDMs.

Module: corr\_cal\_by\_rdm.py

eegrdms\_corr(demo\_rdm, EEG\_rdms, method="spearman", rescale=False)

A function for calculating the Similarity/Correlation Coefficient between RDMs based on EEG/MEG/fNIRS data and a demo RDM

#### Parameters:

demo\_rdm: array

The shape must be [n\_cons, n\_cons].

EEG\_rdm: array

The shape must be [n\_ts, n\_cons, n\_cons] or [n\_chls, n\_cons,

n\_cons].

method: string ("spearman" or "pearson" or "kendall" or "similarity"

or "distance")

They correspond to different methods of calculation.

"spearman" or "pearson" or "kendall" or "similarity" or

"distance".

rescale: Boolean (True or False)

Rescale the values in RDM or not. True or False.

Returns:

corrs: array

The correlation coefficients corresponding to the RDMs.

### • fmrirdms\_corr(demo\_rdm, fMRI\_rdms, method="spearman", rescale=False)

A function for calculating the Similarity/Correlation Coefficient between RDMs based on fMRI data and a demo RDM

### Parameters:

demo\_rdm: array

The shape must be [n\_cons, n\_cons].

fmri\_rdm: array

The shape must be [n\_x, n\_y, n\_z, n\_cons, n\_cons].

method: string ("spearman" or "pearson" or "kendall" or "similarity"

or "distance")

They correspond to different methods of calculation. "spearman" or "pearson" or "kendall" or "similarity" or

"distance".

rescale: Boolean (True or False)

Rescale the values in RDM or not. True or False.

Returns:

corrs: array

The correlation coefficients corresponding to the RDMs. The

shape of corrs is [n\_x, n\_y, n\_z, 2].

## Part 6: Save as a NIfTI file (for fMRI)

Module: corr\_to\_nii.py

◆ corr\_save\_nii(corrs, filename, corr\_mask=None, affine, size=[60, 60, 60], ksize=[3, 3, 3], strides=[1, 1, 1], p=1, r=0, similarity=0, distance=0, correct\_method=None, correct\_n=27, plotrlt=True, img\_background=None)

A function for saving the correlation coefficients as a .nii file

#### Parameters:

corrs: array

corrs represent the correlation coefficients. Its shape must

be [n\_x, n\_y, n\_z, 2].

filename: string

The filename of the NIfTI file. Don't need a suffix.

corr\_mask: Niimg-like object or the filename

A file for correcting the RSA result. It can just be one of your

fMRI data file in your experiment.

affine: array / list

An affine array that tells you the position of the image array

data in a reference space.

size: array / list

size=[x, y, z] represents that the size of the original data.

ksize: array / list

ksize=[kx, ky, kz] represents that the calculation unit

contains k1\*k2\*k3 voxels.

strides: array / list

strides=[sx, sy, sz] represents the moving steps along the x, y,

z.

p, r, similarity, distance: *float* 

They represent the threshold value for calculation.

correct\_method: None / 'FWE' / 'FDR'

The method for correction.

correct\_n: int

The number of voxels used in correction.

plotrlt: Boolean (True or False)

Plot the RSA result or not.

img\_background: Niimg-like object or the filename

The background image that the ROI/mask will be plotted on top of. If there is no special background requirement, set it as None.

#### Returns:

img\_nii: array

The matrix form of the NIfTI file.

## **Part 7: Visualization for Results**

Module: *rsa\_plot.py* 

plot\_rdm(rdm, rescale=False, conditions=None, con\_fontsize=12, cmap=None)

A function for plotting the RDM

#### Parameters:

rdm: array / list

A representational dissimilarity matrix.

rescale: Boolean (True or False)

Rescale the values in RDM or not. True or False.

conditions: None / string list / string array

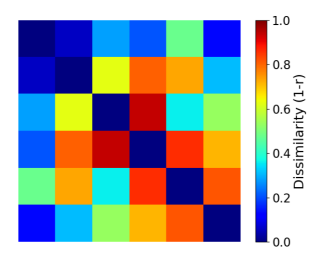
The labels of the conditions.

con\_fontsize: int

Font size of the condition-labels.

cmap: *Matplotlib colormap / string* 

The colormap of the figure.



plot\_rdm\_withvalue(rdm, fontsize=10, conditions=None, con\_fontsize=12, cmap=None)

### A function for plotting the RDM with visible values

#### Parameters:

rdm: array / list

A representational dissimilarity matrix.

fontsize: int / float

Font size of the visible values

conditions: None / string list / string array

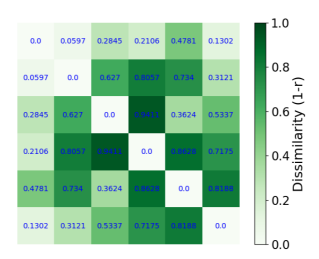
The labels of the conditions.

con\_fontsize: int

Font size of the condition-labels.

cmap: *Matplotlib colormap / string* 

The colormap of the figure.



### plot\_corrs\_by\_time(corrs, labels=None, time\_unit=[0, 1])

A function for plotting the correlation coefficients by time sequence

#### Parameters:

corrs: array

corrs represent the correlation coefficients point-by-point.

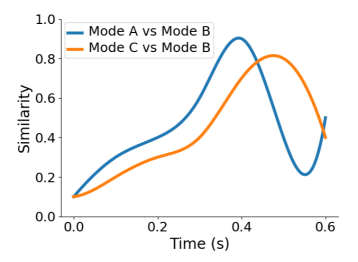
Its shape must be [n\_cons, ts, 2] or [n\_cons, ts].

labels: None / string array / string list

labels represent the names of conditions of RSA results.

time\_unit: array / list

time\_unit=[start\_t, t\_step]. Here, start\_t represents the start time and t\_step represents the time between two adjacent time-points.



plot\_corrs\_hotmap(eegcorrs, chllabels=None, time\_unit=[0, 0.1], lim=[0, 1], smooth=False, figsize=None, cmap=None)

*A function for plotting the correlation coefficients by time sequence* 

#### Parameters:

eegcorrs: array

eegcorrs represent each channels' correlation coefficients point-by-point. Its shape must be [n\_chls, ts, 2] or [n\_chls, ts].

chllabels: None / string array / string list

labels represent the names of channels.

time\_unit: array / list

time\_unit=[start\_t, t\_step]. Here, start\_t represents the start time and t\_step represents the time between two adjacent time-points.

lim: array / list

lim=[lower, upper]. Here, min\_r and max\_r represent the upper limit and lower limit for plotting of the r-values.

smooth: Boolean (True or False)

True or False represents smoothing the results or not.

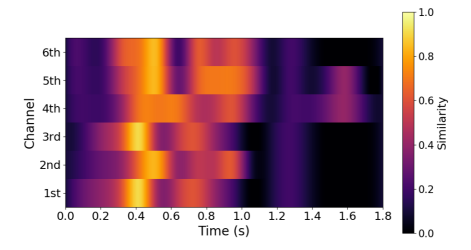
figsize: None / array / list

The size of the Matplotlib figure. Such as figsize=[6.4, 4.8], if

figsize=None, the size will be automatical.

cmap: Matplotlib colormap / string

The colormap of the figure.



plot\_nps\_hotmap(similarities, chllabels=None, time\_unit=[0, 0.1], lim=[0, 1], abs=False, smooth=False, figsize=None, cmap=None)

A function for plotting the correlation coefficients by time sequence

#### Parameters:

similarities: array

similarities represent correlation coefficients point-by-

point. Its shape must be [n\_chls, ts].

chllabels: None / string array / string list

labels represent the names of channels.

time\_unit: array / list

time\_unit=[start\_t, t\_step]. Here, start\_t represents the start time and t\_step represents the time between two adjacent

time-points.

lim: array / list

lim=[lower, upper]. Here, min\_r and max\_r represent the

upper limit and lower limit for plotting of the r-values.

abs: Boolean (True or False)

True or False represents changing the similarities into

absolute values or not.

smooth: Boolean (True or False)

True or False represents smoothing the results or not.

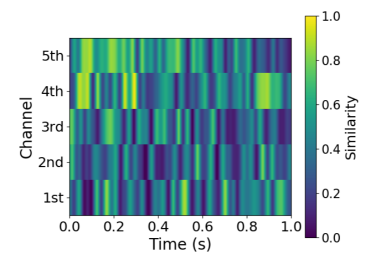
figsize: None / array / list

The size of the Matplotlib figure. Such as figsize=[6.4, 4.8], if

figsize=None, the size will be automatical.

cmap: *Matplotlib colormap / string* 

The colormap of the figure.



### plot\_brainrsa\_region(img, threshold=None, background=get\_bg\_ch2())

A function for plotting the RSA-result regions by 3 cuts (frontal, axial, and lateral)

#### Parameters:

img: string

The file path of the 3-D image of the RSA result.

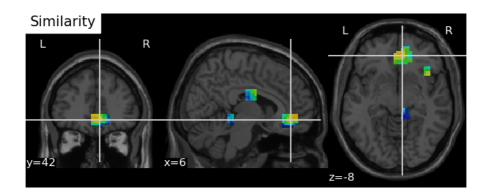
threshold: None or int

If it is an int, threshold is the number of voxels used in correction. Only the similarity clusters consisting more than threshold voxels will be visualized. If it is None, the

threshold-correction won't work.

### background: Niimg-like object or the filename

The background image that the ROI/mask will be plotted on top of.



plot\_brainrsa\_montage (img, threshold=None, slice=[6, 6, 6], background=get\_bg\_ch2bet())

A function for plotting the RSA-result by different cuts

#### Parameters:

img: string

The file path of the 3-D image of the RSA result.

threshold: None or int.

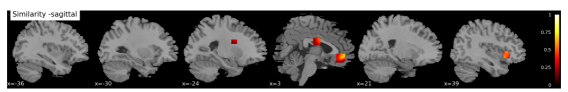
If it is an int, threshold is the number of voxels used in correction. Only the similarity clusters consisting more than threshold voxels will be visualized. If it is None, the threshold-correction won't work.

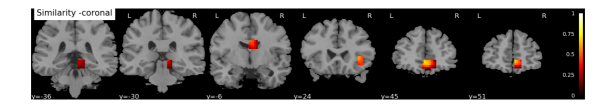
slice: array

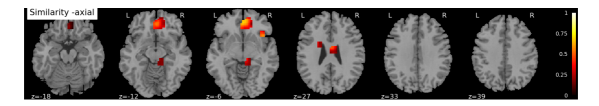
slice=[nx, ny, nz]. The nx, ny, nz is the number of cuts in the x, y, z directions.

### background:string

The file path of the background image that the ROI/mask will be plotted on top of.







### plot\_brainrsa\_glass (img, threshold=None)

A function for plotting the 2-D projection of the RSA-result

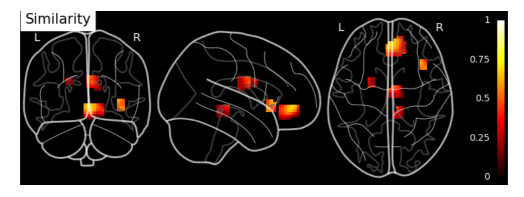
#### Parameters:

img: string

The file path of the 3-D image of the RSA result.

threshold: None or int

If it is an int, threshold is the number of voxels used in correction. Only the similarity clusters consisting more than threshold voxels will be visualized. If it is None, the threshold-correction won't work.



### plot\_brainrsa\_surface (img, threshold=None)

A function for plotting the RSA-result into a brain surface

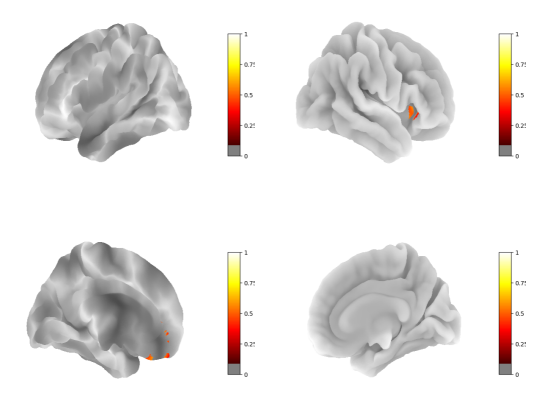
#### Parameters:

img: string

The file path of the 3-D image of the RSA result.

#### threshold: None or int

If it is an int, threshold is the number of voxels used in correction. Only the similarity clusters consisting more than threshold voxels will be visualized. If it is None, the threshold-correction won't work.



### • plot\_brainrsa\_rlts (img, threshold=None, slice=[6, 6, 6], background=None)

A function for plotting the RSA-result by a set of images

#### Parameters:

img: string

The file path of the 3-D image of the RSA result.

threshold: None or int

If it is an int, threshold is the number of voxels used in correction. Only the similarity clusters consisting more than threshold voxels will be visualized. If it is None, the threshold-correction won't work.

slice: array

slice=[nx, ny, nz]. The nx, ny, nz is the number of cuts in the

x, y, z directions.

## background:string

The file path of the background image that the ROI/mask will be plotted on top of. If there is no special background requirement, set it as None.

### Part 8: Others

## Module: *stuff.py*

### ♦ limtozero(x)

A function for zeroing the value close to zero.

Parameters:

x: float

A value.

Returns:

0

### get\_affine(file\_name)

A function for getting the affine.

Parameters:

file\_name: string

The file\_name of a fMRI file.

Returns:

affine: array

An affine array that tells you the position of the image array

data in a reference space.

### • fwe\_correct(p, size=[60, 60, 60], n=64)

A function for FWE correction.

Parameters:

p: array

A 3-D array of p-values, the number of p-value is the same as

the number of RSA calculation units in fMRI. Users can get p by corrs (Code: p = corrs[:, :, :, 1]).

size: array

size=[x, y, z] represents that the size of the original data.

n: array

The number of voxels used in correction.

#### Returns:

correctp: array

The FWE corrected p-values. A 3-D array of p-values, the number of p-value is the same as the number of RSA calculation units in fMRI.

### ♦ fdr\_correct(p, size=[60, 60, 60], n=64)

A function for FDR correction.

#### Parameters:

p: array

A 3-D array of p-values, the number of p-value is the same as the number of RSA calculation units in fMRI. Users can get  $\boldsymbol{p}$ 

by corrs (Code: *p* = *corrs[:, :, :, 1]*).

size: array / list

size=[x, y, z] represents that the size of the original data.

n: int

The number of voxels used in correction.

#### Returns:

correctp: array

The FDR corrected p-values. A 3-D array of p-values, the number of p-value is the same as the number of RSA calculation units in fMRI.

### correct\_by\_threshold(img, threshold)

A function for fMRI correction by threshold (the number of voxels).

#### Parameters:

img: array

A 3-D array of the RSA result.

threshold: n

The number of voxels used in correction. Only the similarity clusters consisting more than threshold voxels will be

visualized.

Returns:

img: array

A 3-D array of the threshold-corrected RSA result.

### get\_bg\_ch2()

A function for getting the path of 'ch2.nii.gz'.

Returns:

path: string

The path of the file "ch2.nii.gz".

### get\_bg\_ch2bet()

A function for getting the path of 'ch2bet.nii.gz'.

Returns:

path: string

The path of the file "ch2bet.nii.gz".

### datamask(fmri\_data, mask\_data)

A function for filtering the data by a ROI mask.

Parameters:

fmri\_data: array

The fmri data. The shape of  $fmri\_data$  must be [nx, ny, nz].

nx, ny, nz represent the size of the fMRI data.

mask\_data: array

The fmri data. The shape of mask\_data must be [nx, ny, nz].

nx, ny, nz represent the size of the fMRI data.

#### Parameters:

newfmri\_data: array

The new fmri data. The shape of fmri\_data must be [nx, ny, nz]. nx, ny, nz represent the size of the fMRI data.

### Part 9: Demo

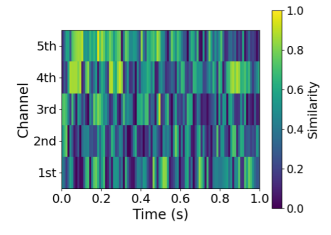
### The EEG/MEG Demo

Here is a demo based on the publicly available visual-92-categories-task MEG datasets. (*Reference:* Cichy, R. M., Pantazis, D., & Oliva, A. "Resolving human object recognition in space and time." Nature neuroscience (2014): 17(3), 455-462.) MNE-Python has been used to load this dataset.

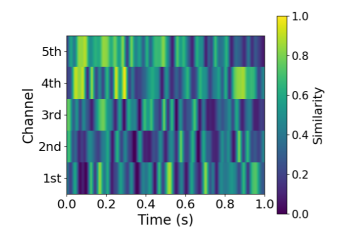
```
# -*- coding: utf-8 -*-
' a demo based on visual-92-categories-task MEG data '
# Users can learn how to use Neurora to do research based on EEG/MEG etc
data.
__author__ = 'Zitong Lu'
import numpy as np
import os.path as op
from pandas import read_csv
import mne
from mne.io import read_raw_fif
from mne.datasets import visual_92_categories
from neurora.nps_cal import nps
from neurora.rdm_cal import eegRDM
from neurora.rdm_corr import rdm correlation spearman
from neurora.corr_cal_by_rdm import rdms_corr
from neurora.rsa_plot import plot_rdm, plot_corrs_by_time,
plot_nps_hotmap, plot_corrs_hotmap
                  Section 1: loading example data
""" Here, we use MNE-Python toolbox for loading data and processing """
""" you can learn this process from MNE-Python (https://mne-
tools.github.io/stable/index.html) """
data_path = visual_92_categories.data_path()
fname = op.join(data_path, 'visual_stimuli.csv')
conds = read_csv(fname)
conditions = []
```

```
for c in conds.values:
   cond tags = list(c[:2])
   cond_tags += [('not-' if i == 0 else '') + conds.columns[k]
                 for k, i in enumerate(c[2:], 2)]
   conditions.append('/'.join(map(str, cond_tags)))
event_id = dict(zip(conditions, conds.trigger + 1))
print(event id)
sub_id = [0, 1, 2]
megdata = np.zeros([3, 92, 306, 1101], dtype=np.float32)
subindex = 0
for id in sub_id:
   fname = op.join(data_path, 'sample_subject_'+str(id)+'_tsss_mc.fif')
   raw = read_raw_fif(fname)
   events = mne.find_events(raw, min_duration=.002)
   events = events[events[:, 2] <= 92]</pre>
   subdata = np.zeros([92, 306, 1101], dtype=np.float32)
   for i in range(92):
       epochs = mne.Epochs(raw, events=events, event id=i + 1,
baseline=None,
                          tmin=-0.1, tmax=1, preload=True)
       data = epochs.average().data
       subdata[i] = data
   megdata[subindex] = subdata
   subindex = subindex + 1
# the shape of MEG data: megdata is [3, 92, 306, 1101]
# n_subs = 3, n_conditions = 92, n_channels = 306, n_timepoints = 1101
(-100ms to 1000ms)
"""********
                                                        ******
                      Section 2: Preprocessing
# shape of megdata: [n_subs, n_cons, n_chls, n_ts] -> [n_cons, n_subs,
n chls, n ts]
megdata = np.transpose(megdata, (1, 0, 2, 3))
# shape of megdata: [n_cons, n_subs, n_chls, n_ts] -> [n_cons, n_subs,
n_trials, n_chls, n_ts]
# here data is averaged, so set n_trials = 1
megdata = np.reshape(megdata, [92, 3, 1, 306, 1101])
"""**** Section 3: Calculating the neural pattern similarity ****""
```

```
# Get data under different condition
# Here we calculate the neural pattern similarity (NPS) between two
stimulus
# Seeing Humanface vs. Seeing Non-Humanface
# get data under "humanface" condtion
megdata humanface = megdata[12:24]
# get data under "nonhumanface" condition
megdata_nonhumanface = megdata[36:48]
# Average the data
avg_megdata_humanface = np.average(megdata_humanface, axis=0)
avg_megdata_nonhumanface = np.average(megdata_nonhumanface, axis=0)
# Create NPS input data
# Here we extract the data from first 5 channels between 0ms and 1000ms
nps_data = np.zeros([2, 3, 1, 5, 1000]) # n_cons=2, n_subs=3, n_chls=5,
n ts=1000
nps_data[0] = avg_megdata_humanface[:, :, :5, 100:1100] # the start time
of the data is -100ms
nps_data[1] = avg_megdata_nonhumanface[:, :, :5, 100:1100] # so 100:1200
corresponds 0ms-1000ms
# Calculate the NPS with a 10ms time-window
# (raw sampling requency is 1000Hz, so here
time_win=10ms/(1s/1000Hz)/1000=10)
nps = nps(nps_data, time_win=10)
# Plot the NPS results
plot_nps_hotmap(nps, time_unit=[0, 0.01], abs=True)
```



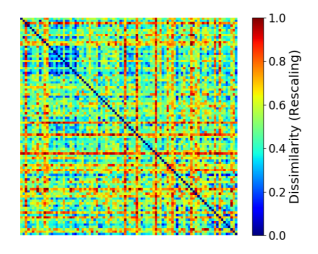
```
# Smooth the results and plot
plot_nps_hotmap(nps, time_unit=[0, 0.01], abs=True, smooth=True)
```



```
"""***** Section 4: Calculating single RDM and Plotting

# Calculate the RDM based on the data during 190ms-210ms
rdm = eegRDM(megdata[:,:,:,:,290:310])

# Plot this RDM
plot_rdm(rdm, rescale=True)
```



```
# Calculate the RDMs by a 10ms time-window
# (raw sampling requency is 1000Hz, so here
time_win=10ms/(1s/1000Hz)/1000=10)
rdms = eegRDM(megdata, time_win=10, time_opt=1)
```

```
# Plot the RDM of 0ms, 50ms, 100ms, 150ms, 200ms
times = [0, 10, 20, 30, 40, 50]
for t in times:
    plot_rdm(rdms[t], rescale=True)
```

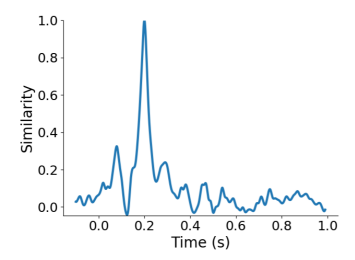
```
# RDM of 200ms
rdm_sample1 = rdms[30]
# RDM of 800ms
rdm_sample2 = rdms[90]

# calculate the correlation coefficient between these two RDMs
corr = rdm_correlation_spearman(rdm_sample1, rdm_sample2, rescale=True)
print(corr)
```

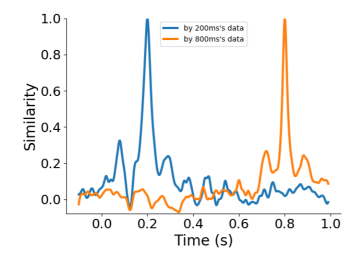
SpearmanrResult(correlation=0.02665680483550596, pvalue=0.08462337954774739)

```
# Calculate the representational similarity between 200ms and all the
time points
corrs1 = rdms_corr(rdm_sample1, rdms)

# Plot the corrs1
corrs1 = np.reshape(corrs1, [1, 110, 2])
plot_corrs_by_time(corrs1, time_unit=[-0.1, 0.01])
```



```
# Calculate and Plot multi-corrs
corrs2 = rdms_corr(rdm_sample2, rdms)
corrs = np.zeros([2, 110, 2])
corrs[0] = corrs1
corrs[1] = corrs2
labels = ["by 200ms's data", "by 800ms's data"]
plot_corrs_by_time(corrs, labels=labels, time_unit=[-0.1, 0.01])
```



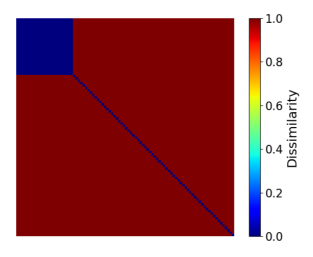
```
"""***** Section 8: Calculating the RDMs for each channels *****""

# Calculate the RDMs for the first six channels by a 10ms time-window
between 0ms and 1000ms
rdms_chls = eegRDM(megdata[:,:,:,:6, 100:1100], chl_opt=1,
time_opt=1, time_win=10)

# Create a 'human-related' coding model RDM
```

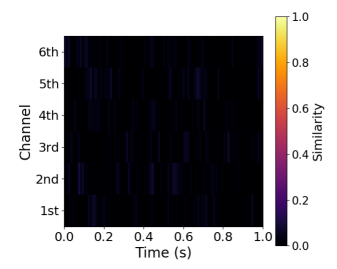
```
model_rdm = np.ones([92, 92])
for i in range(92):
    for j in range(92):
        if (i < 24) and (j < 24):
            model_rdm[i, j] = 0
        model_rdm[i, i] = 0

# Plot this coding model RDM
plot_rdm(model_rdm)</pre>
```

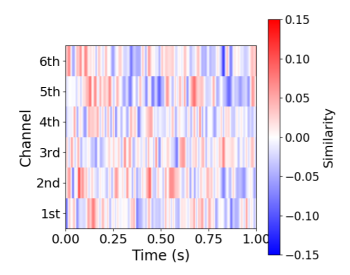


```
# Calculate the representational similarity between the neural
activities and the coding model for each channel
corrs_chls = rdms_corr(model_rdm, rdms_chls)

# Plot the representational similarity results
plot_corrs_hotmap(corrs_chls, time_unit=[0, 0.01])
```



```
# Set more parameters and re-plot
plot_corrs_hotmap(corrs_chls, time_unit=[0, 0.01], lim=[-0.15, 0.15],
smooth=True, cmap='bwr')
```



## The fMRI Demo

Here is a demo based on the publicly available Haxby fMRI datasets. (*Reference*: Haxby, J. V. (2001). Distributed and Overlapping

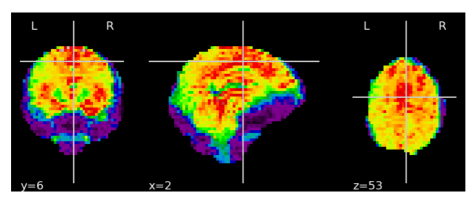
Representations of Faces and Objects in Ventral Temporal Cortex.

Science, 293(5539), 2425–2430.) Nilearn has been used to load this dataset and plot some results in this demo.

```
from neurora.rdm_cal import fmriRDM_roi, fmriRDM
from neurora.corr_cal_by_rdm import fmrirdms corr
from neurora.corr_to_nii import corr_save_nii
                                                         *****
                  Section 1: Loading example data
""" Here, we use Nilearn toolbox for loading data and processing """
""" you can learn this process from Nilearn
(http://nilearn.github.io/index.html) """
# load Haxby dataset (here, we only use subject2's data for this
example)
haxby_dataset = datasets.fetch_haxby()
# load the fMRI data filename & mask data filename
func filename = haxby dataset.func[0]
mask_filename = haxby_dataset.mask
# read label information of the experiment
labelinfo = pd.read_csv(haxby_dataset.session_target[0], sep='_')
labels = labelinfo['labels']
"""*******
                                                        *******
                       Section 2: Preprocessing
# get mask data NumPy array
maskdata = nib.load(mask_filename).get_data()
# get the size of the data
nx, ny, nz = maskdata.shape
# labels of seven ategories
categories = ["face", "cat", "house", "chair", "shoe", "bottle",
"scissors"]
# numbe of conidtions: 7
ncon = len(categories)
# get fmri data under 7 conditions
# here we average the data under different conditions
fmri_data = np.full([ncon, nx, ny, nz], np.nan)
for i in range(ncon):
```

```
img = mean_img(index_img(func_filename,
labels.isin([categories[i]])))
    fmri_data[i] = datamask(img.get_data(), maskdata)
    np.savetxt("demo02/data"+str(i+1)+".txt", np.reshape(fmri_data[i],
    [nx*ny*nz]))

# get fmri data under 'face'-condition
face_img = nib.Nifti1Image(fmri_data[0], affine=img.affine)
# have a look
plotting.plot_epi(face_img)
plotting.show()
```



```
# reshaoe the data: [ncon, nx, ny, nz] -> [ncon, nsubs, nx, ny, nz]
# here just one subject's data
fmri_data = np.reshape(fmri_data, [ncon, 1, nx, ny, nz])

"""**Section 3: Calculating the neural pattern similarity (for ROI)**"""

# get mask of 'mask_face' in the dataset
mask_face_filename = haxby_dataset.mask_face[0]
mask_face_data = nib.load(mask_face_filename).get_data()

# get input data under two condition
# here, "face"-condition vs. "cat"-condition
nps_fmri_data = fmri_data[[0, 6]]

# calculate the neural pattern similarity (NPS) for ROI between two stimulus
nps_roi = nps_fmri_roi(nps_fmri_data, mask_face_data)

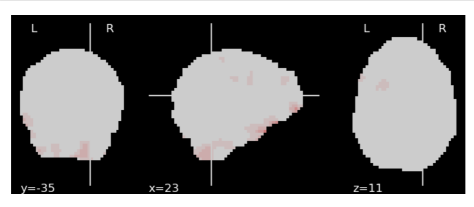
# print the NPS result
print(nps_roi)
```

```
"""Section 4: Calculating the neural pattern similarity (Searchlight)"""

# calculate the neural pattern similarity (NPS) between two stimulus
nps = nps_fmri(nps_fmri_data)

# convert the NPS results into a .nii file
savefilename = "nps_img"
affine = get_affine(mask_filename)
corr_save_nii(nps, filename=savefilename, affine=affine, size=[nx, ny, nz], plotrlt=False)

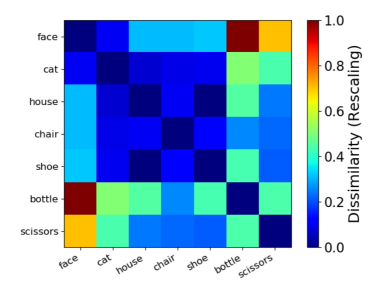
# have a look
plotting.plot_epi(savefilename+".nii")
plotting.show()
```

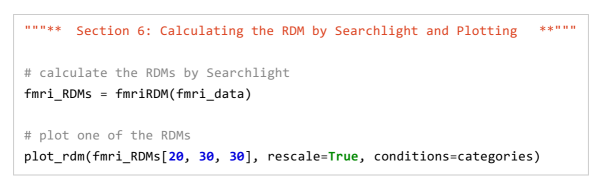


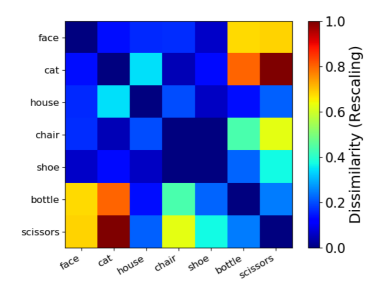
```
# get mask of "mask_vt" in the dataset
mask_vt_filename = haxby_dataset.mask_face[0]
mask_vt_data = nib.load(mask_vt_filename).get_data()

# calculate the RDM for ROI
rdm_roi = fmriRDM_roi(fmri_data, mask_vt_data)

# plot the RDM
plot_rdm(rdm_roi, rescale=True, conditions=categories)
```







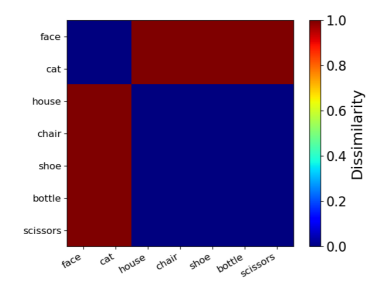
```
"""*** Section 7: Calculating the representational similarities ***"""

"""*** between a coding model and neural activities ***"""

# Create a RDM for "animate-inanimate" coding model

# which means the representations of animate matters are highly similar

# and the representations of inanimate matters are highly similar
```



```
# calculate the similarities between model RDM and searchlight RDMs
corrs = fmrirdms_corr(model_RDM, fmri_RDMs)

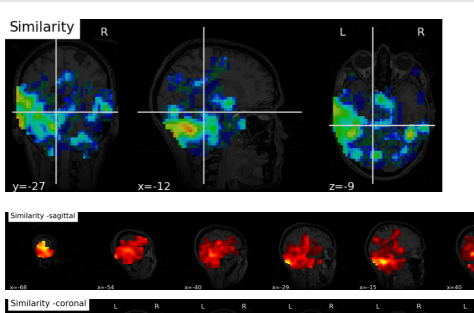
"""***** Section 8: Saving the RSA result and Plotting *****""

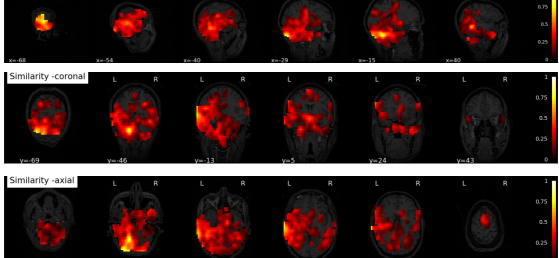
# load the filename of anatomical image as the background for plotting ant_filename = haxby_dataset.anat[0]

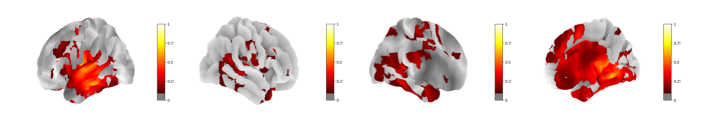
# get the affine info affine = get_affine(mask_filename)

# save the RSA result as a .nii file # and visualize the result automatically # p < 0.05, FDR-correct rsarltfilename = "demo2_rsarlt_img"</pre>
```

img = corr\_save\_nii(corrs, filename=rsarltfilename, affine=affine,
corr\_mask=mask\_filename, size=[40, 64, 64], p=0.05, plotrlt=True,
img\_background=ant\_filename, correct\_method="FDR")







- # Users can plot the RSA results independently by functions below
- # >> from neurora.rsa\_plot import plot\_brainrsa\_regions
- # >> from neurora.rsa\_plot import plot\_brainrsa\_montage
- # >> from neurora.rsa\_plot import plot\_brainrsa\_glass
- # >> from neurora.rsa\_plot import plot\_brainrsa\_surface