A Quick Guide for the MixfMRI Package

Wei-Chen Chen¹ and Ranjan Maitra²

¹pbdR Core Team Silver Spring, MD, USA

²Department of Statistics Iowa State University Ames, IA, USA

Contents

1.	Introduction	1
	1.1. Dependent Packages	1
	1.2. The Main Function	
	1.3. Datasets	2
	1.4. Examples	3
	1.5. Workflows	3
2.	Demonstrations	3
	2.1. 2D Phantoms	3
	2.2. 2D Simulations	
	2.3. 2D Clustering	5

@ 2018 Wei-Chen Chen and Ranjan Maitra.

Permission is granted to make and distribute verbatim copies of this vignette and its source provided the copyright notice and this permission notice are preserved on all copies. This publication was typeset using IAT_EX .

Warning: The findings and conclusions in this article have not been formally disseminated by the U.S. Food and Drug Administration and should not be construed to represent any determination or policy of any University, Institution, Agency, Administration and National Laboratory.

Ranjan Maitra and this research were supported in part by the National Institute of Biomedical Imaging and Bioengineering (NIBIB) of the National Institutes of Health (NIH) under its Award No. R21EB016212. The content of this paper however is solely the responsibility of the authors and does not represent the official views of either the NIBIB or the NIH.

This document is written to explain the main function of **MixfMRI** (?), version 0.1-0. Every effort will be made to ensure future versions are consistent with these instructions, but features in later versions may not be explained in this document.

1. Introduction

The main purpose of this vignette is to demonstrate basic usage of **MixfMRI** which is prepared to implement developed methodology, simulation studies, data analyses in "A Practical Model-based Segmentation Approach for Accurate Activation Detection in Single-Subject functional Magnetic Resonance Imaging Studies" (?). The methodology mainly utilizes model-based clustering (unsupervised) of functional Magnetic Resonance (fMRI) data that identifies regions of brain activation associated with the performance of a task or the application of a stimulus. The implemented methods include 2D and 3D unsupervised segmentation analyses for fMRI signals. For simplification, only 2D clustering is demonstrated in this vignette. In this package, the data on fMRI signals are on the form of the p-values at each voxel of a Statistical Parametric Map (SPM).

The clustering and segmentation analyses identify activated voxels/signals (in terms of small p-values) from normal brain behaviors within a single subject but also use spatial context.

Note that the *p*-values may be derived from statistical models where typically a proper experiment design is required. These *p*-values are our data, and are used for analysis and no statements about significance level of *p*-values associated with activated voxels/signals are needed. Our analysis approach allows for the prespecification of *a priori* expected upper bounds for the proportion of activated voxels/signals that can guide in determining activated voxels.

For large datasets, the methods and analyses are also implemented in a distributed manner especially using SPMD programming framework. The package also includes workflows which utilize SPMD techniques. The workflows serve as examples of data analyses and large scale simulation studies. Several workflows are also built in to automatically process clusterings, hypotheses, merging clusters, and visualizations. See Section 1.5 and files in MixfMRI/inst/workflow/ for more information.

1.1. Dependent Packages

The **MixfMRI** package depends on other R packages to be functional even though they are not always required. This is because some examples, functions and workflows of the **MixfMRI** may need utilities of those dependent packages. For instance,

Imports: MASS, Matrix, RColorBrewer, fftw, MixSim, EMCluster.

Enhances: pbdMPI, oro.nifti.

1.2. The Main Function

The main function, fclust(), implements model-based clustering using the EM algorithm (?) for fMRI signal data and provides unsupervised clustering results that identify activated regions in the brain. The fclust() function contains an initialization method and EM algorithms for clustering fMRI signal data which have two parts:

- PV.gbd for *p*-value of signals associated with voxels, and
- X.gbd for voxel information/locations in either 2D or 3D,

where PV.gbd is of length N (number of voxels) and X.gbd is of dimension $N \times 2$ or $N \times 3$ (for 2D or 3D). Each signal (per voxel) is assumed to follow a mixture distribution of K components with mixing proportion ETA. Each component has two independent coordinates (one for each part) with density functions: Beta and multivariate Normal distributions, for each part of fMRI signal data.

Beta Density:

The first component (k = 1) is restricted by min.1st.prop and Beta(1, 1) (equivalently, the standard uniform) distribution. The rest $k = 2, 3, \ldots, K - 1$ components have different Beta(alpha, beta) distributions with alpha < 1 < beta for all k > 1 components. This coordinate mainly represents the results of test statistics for determining activation of voxels (those that have smaller *p*-values). Note that the test statistics may be developed/smoothed/-computed from a time course model associated with voxel behaviors. See the main paper ? for information.

Multivariate Normal Density:

The logarithm of the multivariate normal density is used as a penalty to provide regularization of the estimated parameters and in the estimated activation. model.X = "I" is for identity covariance matrix of this multivariate Normal distribution, and "V" for unstructured covariance matrix. ignore.X = TRUE is to ignore X.gbd and normal density, *i.e.* there is no regularization and only the Beta density is used. Note that this coordinate (for each axis) is recommended to be normalized in the (0, 1) scale which is on the same scale of Beta density. From a modeling perspective, rescaling X.gbd does not have an effect.

In this package, the two parts PV.gbd and X.gbd are assumed to be independent because the latter comes through the addition of a penalty term to the log likelihood of the voxel-wise data on *p*-values. The goal of the main function is to provide spatial clusters (in addition to the PV.gbd) indicating spatial correlations.

Currently, APECMa (?) and EM algorithms are implemented with EGM algorithm (?) to speed up convergence when MPI and **pbdMPI** (?) are available. RndEM initialization (?) with a specific way of choosing initial seeds is implemented for obtaining good initial values that has the potential to increase the chances of convergence.

1.3. Datasets

The package has been built with several datasets including

- three 2D phantoms, shepp0fMRI, shepp1fMRI, and shepp2fMRI,
- one 3D dataset, pstats, with *p*-values obtained from the SPM obtained after running the Analysis of Functional Neuroimaging (AFNI) software (???) on the imagination dataset of ?.
- two small 2D voxels datasets, pval.2d.complex and pval.2d.mag, in p-values
- two toy examples, toy1 and toy2.

1.4. Examples

The scripts in MixfMRI/demo/ have several examples that demonstrate the main function, the example datasets and other utilities in this package. For a quick start,

- the scripts MixfMRI/demo/fclust2d.r and MixfMRI/demo/fclust3d.r show the basic usage of the main function fclust() using the two toy datasets,
- the scripts MixfMRI/demo/maitra_2d.r and MixfMRI/demo/shepp.r show and visualize examples on how to generate simulated datasets with given overlap levels, and
- the scripts MixfMRI/demo/alter_*.r show alternative methods.

1.5. Workflows

The package also has several workflows established for simulation studies. The main examples are located in MixfMRI/inst/workflow/simulation/. See the file create_simu.txt that generates scripts for simulations.

The files under MixfMRI/inst/workflow/spmd/ have the main scripts for the workflows. Note that MPI and **pbdMPI** are required for workflows because these simulations require potentially long computing times.

2. Demonstration

The examples presented below are simulated and are not necessarily meant to represent meaningful activation study on the brain. Their purpose is to demonstrate our segmentation methodology in activation detection.

2.1. 2D Phantoms

Three 2D phantoms built in the $\mathbf{MixfMRI}$ package can be displayed in R from the demo as simple as

Maitra's Phantoms

```
R> demo(maitra_phantom, package = "MixfMRI", ask = F)
```

which performs the code in MixfMRI/demo/maitra_phantom.r. The R command should give a plot similar to Figure 1 containing three different simulated 2D slices of a hypothesized brain. Each phantom may have different amounts of activated voxels (in terms of smaller *p*-values). Colors represent different activation intensities. The total proportions of truly active voxels are listed in the title of each phantom.



The examples used below mimic some active regions (in 2D) depending on different types of stimuli that may trigger responses in the brain. Hypothetically, the voxels may be active by regions, but each region may not be active in the same way (or magnitude) even though they may need to collectively respond to the stimuli (for example, due to time delay, response order, or sensitivity of study design).

As an example, only 3.968% of voxels in the shepp2fMRI phantom are active and indicated by two different colors (blue and brown) for different activation types where *p*-values may be smaller than 0.05 and may follow two Beta distributions (with different configurations) for the truly active voxels and one uniform distribution (i.e. Beta(1,1)) for the truly inactive voxels.

The following code provides some counts for each groups of active and inactive voxels in the shepp2fMRI phantom.

Summary of shepp2fMRI Phantoms									
> table	<pre>> table(shepp2fMRI, useNA="always")</pre>								
shepp2fl	shepp2fMRI								
0	1	2	<na></na>						
13408	472	82	51574						

The summary says that this phantom has three kinds of activations with group ids: 0, 1, and 2. There are 13,408 voxels belonging to cluster 0 (inactive), followed by 472 voxels belonging to cluster 1 (active & highlighted in blue in Figure 1), and 82 voxels belonging to cluster 2 (active & high lighted in brown in Figure 1). There are 51,574 pixels (NA) of this imaging dataset which are not within the brain (contour by the black line in Figure 1). See Section 2.2 for information of generating p-values from a mixture of three Beta distributions.

2.2. 2D Simulations

The MixfMRI provides a function gendataset(phantom, overlap) to generate *p*-values of activations. The function needs two arguments: phantom and overlap. The phantom is a map containing voxel group id's where *p*-values will be simulated from a mixture Beta distribution with certain mixture level specified by the overlap argument. The example can be found in MixfMRI/demo/maitra_2d.r and can be done in R as simple as

	Simulations of Active Voxels					
(R>	<pre>demo(maitra_2d,</pre>	package =	"MixfMRI",	ask = F)		

Note that the overlap represents similarity of activation signals. The higher the overlap, the more difficult it becomes to distinguish between activation and inactivation and also the kinds of activation.

The command above should give a plot similar to Figure 2 containing group id's on the left and their associated *p*-values for stimulus responses on the right. The top row displays examples for phantom shepp1fMRI, and the bottom row displays examples for phantom shepp2fMRI.

- Inside the brain, the group id 2's are indicated by white (active is highly associated with stimula due to experiment design), 1's are indicated by light gray (slightly active), and 0's are indicated by dark gray (inactive). Note that the region with white color was the region colored blue and the light gray region corresponds to the region colored brown in Figure 1
- The simulated *p*-values are colored by a map using a red-orange-yellow palette from 0 to 1. Note that small *p*-values (redder voxels) may also occur at truly inactive voxels.

See Figure 3 for the distribution of simulated *p*-values for the phantom shepp2fMRI.

The methodology and analyses implemented in this package aim to identify those active voxels in spatial clusters. For example, regions of active voxels associated with imagining the playing of certain sports. When an experiment was conducted/designed to detect brain behaviors, the statistical model and the p-values of the treatment effect should be able to reflect the voxel activations. Typically, the statistical tests are done independently voxel-by-voxel due to complexity of computation and modeling. This package provides post hoc clustering that adds spatial contents to p-values and helps to isolate meaningful regions clouded with many small p-values. See ? for information of clustering performance and comprehensive assessments for this post hoc approach.

2.3. 2D Clustering

The example can be found in MixfMRI/demo/maitra_2d_fclust.r as simple as

	Clustering of Active Voxels								
(R>	<pre>demo(maitra_2d_fclust, package = "MixfMRI", ask = F)</pre>								

This demo (explained below) is to cluster the simulated p-values (see Section 2.2) using the developed method.

Code of maitra_2d_fclust.r

library(MixfMRI, quietly = TRUE)



Figure 2: Activated regions of voxels and simulated *p*-values.

In the code above, the histogram of simulated *p*-values is plotted in Figure 3. Then, the fclust(X.gbd, PV.gbd, K = 3) groups voxels in three clusters. At the end, ret saves the clustering results. The print(ret) from the above code will show the results below in detail:

• N is the total number of voxels to be clustered/segmented

- K is the total number of segments.
- n.class is the number of voxels in each segment
- ETA is the mixing proportion of each segment
- BETA is the set of parameters of the Beta distributions (by column)
- MU is the centers of the segments (the spatial location inside the brain)
- SIGMA is the dispersion of the segments

Figure 3: Activation (*p*-values) Distribution. The x-axis is for the *p*-values.



The numbers of voxels for each segment, in this example, are 13,394, 184, and 384 associated with new cluster ids: 0, 1, and 2, respectively. Comparing with the true classifications (see the table in Section 2.1), the adjusted Rand index gives 0.9749 indicating good agreement between the truly active and the activated (as determined by our segmentation methodology) results.

Outputs of Clustering

```
R> print(ret)
Algorithm: apecma
                    Model.X: I
                                Ignore.X: FALSE
 Convergence: 1
                   iter:
                        16
                             abs.err: 0.02091979
                                                    rel.err: 7.375343e-07
- N:
     13962
                     К: З
                           logL: 28364.52
                 2
            p.X:
                       -56557.25 ICL-BIC: -55712.18
                   BIC:
 AIC: -56693.04
_
 n.class: 13394 184 384
```

```
- init.class.method:
- ETA: (min.1st.prop: 0.8 max.PV: 0.1)
[1] 0.95266704 0.01307847 0.03425449
- BETA: (2 by K)
    [,1]
                 [,2]
                           [,3]
     1 1.127244e-01 0.04237128
[1,]
[2,]
      1 4.429518e+04 1.00000130
- MU: (p.X by K)
         [,1]
                   [,2]
                         [,3]
[1,] 0.5013538 0.3105460 0.5917377
[2,] 0.5076080 0.3718145 0.3749842
- SIGMA: (d.normal by K)
          [,1]
                                  [,3]
                      [,2]
[1,] 0.01271198 0.0001859628 0.009462210
[2,] 0.02186662 0.0011582052 0.004284016
R> RRand(ret$class, shepp2fMRI[id] + 1)
  Rand adjRand Eindex
 0.9964 0.9749 1.7012
```