

MRFscRNAseq Vignette

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Contents

Installing MRFscRNAseq from GitHub

```
if (!requireNamespace("devtools", quietly = TRUE))  
  install.packages("devtools")  
library(devtools)  
install_github("eddiehli/MRFscRNAseq", quiet = TRUE)  
library(MRFscRNAseq)
```

Main Function

The main function is

```
?get_DE_MRF()
```

It requires three input values: (1) `data`: a summary statistics matrix where rows are genes and columns are cell types; (2) `g_g`: the gene to gene binary network matrix; and (3) `c_c`: the cell type dependency binary network matrix. The usage of this function will be illustrated on a simulated data set.

Example

The `data_example` is a list object that contains the expression array, the gene to gene binary network matrix, and the cell type dependency binary network matrix. The expression array profiles 200 genes across 18 cell types in 2 groups. Each cell type has 200 cells.

```
data = data_example  
gene_gene = data$gene_gene  
gene_gene[1:10, 1:10]  
#>      G1 G2 G3 G4 G5 G6 G7 G8 G9 G10  
#> G1   0  0  0  1  0  1  1  0  1   1  
#> G2   0  0  0  0  0  0  0  0  0   0  
#> G3   0  0  0  0  0  1  0  0  0   0  
#> G4   1  0  0  0  0  1  0  0  0   0  
#> G5   0  0  0  0  0  0  0  0  0   0  
#> G6   1  0  1  1  0  0  1  0  1   0  
#> G7   1  0  0  0  0  1  0  0  0   0
```

```
#> G8 0 0 0 0 0 0 0 0 0 0 0
#> G9 1 0 0 0 0 0 1 0 0 0 0
#> G10 1 0 0 0 0 0 0 0 0 0 0
cell_cell = data$cell_cell
cell_cell[1:10, 1:10]
#>      C1 C2 C3 C4 C5 C6 C7 C8 C9 C10
#> C1 0 1 0 0 0 0 0 0 1 1 0
#> C2 1 0 0 0 0 0 0 0 1 1 0
#> C3 0 0 0 1 1 1 1 0 0 0 1
#> C4 0 0 1 0 1 1 1 0 0 0 1
#> C5 0 0 1 1 0 1 1 0 0 0 1
#> C6 0 0 1 1 1 0 1 0 0 0 1
#> C7 0 0 1 1 1 1 0 0 0 0 1
#> C8 1 1 0 0 0 0 0 0 0 1 0
#> C9 1 1 0 0 0 0 0 0 1 0 0
#> C10 0 0 1 1 1 1 1 0 0 0 0
```

Then we use our proposed MRF models to obtain DE results, we first convert the raw expression values to z scores using two-sample t-tests,

```
zz = get_z_scores(data)
```

Then we ran the MRF model. Note that here we set `iterEM = 20`, `iterGibbsPost = 500`, `brPost = 200` for illustrative purposes only. The default values are `iterEM = 200`, `iterGibbsPost = 20000`, `brPost = 10000`.

```
MRF_Results = get_DE_MRF(zz, gene_gene, cell_cell,
                          iterEM = 1, iterGibbsPost = 100, brPost = 50)
#>
#> Estimating model parameters, 100 % completed
#> Posterior sampling, 1 % completed Posterior sampling, 2 % completed Posterior sampling, 3 % complet
```

The MRF parameters are

```
MRF_Results$paraMRF
#>      Gamma  Beta_Gene  Beta_Cell
#> -0.64773991 0.02400703 0.03791838
```

and the DE states for each gene in each cell type are

```
results = get_DE_results(MRF_Results, alpha = 0.05)
head(results)
#>      C1 C2 C3 C4 C5 C6 C7 C8 C9 C10 C11 C12 C13 C14 C15 C16 C17 C18
#> G1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 0 0 0
#> G2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
#> G3 0 0 0 0 0 0 0 1 0 0 0 0 0 0 1 0 0 0
#> G4 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 0
#> G5 0 0 0 0 0 0 0 0 0 0 1 0 0 0 1 0 0 0
#> G6 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 0 0
```