Cell-type-specific-GSA on test dataset

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Here we will show a simple workflow for carrying out cell type-specific gene set analysis (CT-GSA). First let's load the CAR-seq package which is designed for carrying out cell type-specific differential expression analysis.

#installation of mremaR

```
#library("devtools");
#install_github("osedo/mremaR")
```

load the library

```
library(mremaR)
```

#installation of CARseq

```
#devtools::install_github("chongjin/CARseq")
library(CARseq)
```

##Simulation here is shown as we are running the test dataset. No need to do simulations on real data.

Now we will simulate gene expression for samples which are a mixture of six different cell types. This defaults to 1,000 genes in 100 cases and 100 controls. In each cell type 5% of genes are assigned a log-fold-change drawn from a normal distribution with a mean of plus/minus 4 and a standard deviation of 0.2.

```
sims <- CTexpSimulation(g = 1000, n = 100, p = 1, m_lfc = 0, sd_lfc = 1) simsbulk.expression[c(1:5), c(1:5)] # bulk expression counts
```

```
##
           sample.1 sample.2 sample.3 sample.4 sample.5
                                                7
## gene.1
                  9
                           11
                                      7
                                                          7
                  6
                                                7
                                                          2
## gene.2
                            3
                                      3
## gene.3
                  7
                           11
                                      6
                                               11
                                                         10
## gene.4
                183
                          177
                                    137
                                              156
                                                        178
## gene.5
                 69
                           78
                                     50
                                               79
                                                         54
```

```
sims$ols.mixture.estimate[c(1:5),] # cell type proportion estimates
```

```
## sample.1 cellType.2 cellType.3 cellType.4 cellType.5 cellType.6 ## sample.1 0.04885907 0.13231213 0.06446669 0.18004738 0.2467514 0.3275633 ## sample.2 0.06670770 0.05758956 0.04622072 0.09654129 0.3207856 0.4121551 ## sample.3 0.06885271 0.03949473 0.01930733 0.07285703 0.1528079 0.6466803 ## sample.4 0.08728981 0.05957621 0.05453721 0.10650392 0.2766101 0.4154828 ## sample.5 0.04492365 0.03171203 0.07128882 0.12597111 0.1707001 0.5554043
```

```
table(sims$trait) # sample traits
```

```
## 0 1
## 50 50
```

These outputs can then be passed to the CAR-seq package. The important results for us are the shrunken_lfc and shrunken_lfcSE matrices

```
# this will take a minute or two to run
res <- run_CARseq(
   count_matrix = sims$bulk.expression,
   cellular_proportions = sims$ols.mixture.estimate,
   groups = sims$trait,
   cores = 10  ##number of cores can be changed
)</pre>
```

As we are running on a test data, we are simulating a few gene sets. For original data, use open available datasets or the link provided below:

```
#kegg gene sets data

#kegg.gs <- "https://raw.githubusercontent.com/osedo/GSA-MREMA/main/real_data/kegg.RData"

#load(url(kegg.gs))
```

Since the gene set mentioned above contains genes in ENTREZ IDs, I recommend converting the gene names from Ensembl IDs to ENTREZ IDs. Alternatively, you can load a gene set dataset that uses Ensembl IDs or another dataset with your preferred ID format using ensembldb or gProfiler R packages.

Let's simulate a few gene sets (no need for the real dataset, just load kegg.gs in place of gs)

```
# a few power gene sets, gs1 enriched for DE genes in cellType1, same for gs2 and so on
# we will give sets of 100 approx 20% DE gene
g <- 1000
gs.power <- lapply(c(1:6), function(x){
  ct_de <- which(abs(sims\simulated.lfc[,x]) > log2(2.5))
  ct_nonde <- c(1:g)[!c(1:g) %in% ct_de]
  ct_de <- paste0("gene.", ct_de)</pre>
  ct_nonde <- paste0("gene.", ct_nonde)</pre>
  genes per set <- 100
  p \leftarrow mean(abs(sims\$simulated.lfc[,x]) > log2(2.5))
  enrichment <- 3
  c(sample(ct_de, genes_per_set*enrichment*p),
    sample(ct_nonde, (genes_per_set-genes_per_set*enrichment*p)))
})
# random gene sets
gs.null <- lapply(c(7:20), function(x){
  paste0("gene.", sample.int(1000, 100))
gs <- c(gs.power, gs.null)
names(gs) <- paste0("gs", 1:20)
```

Let's run mrema() on all cell-types

##parametric tests

```
mrema_res <- mrema(postdata = postdata, raw.gs = gs, threshold = 2, DF = 4, ncores = 1)
mrema_res
})
names(mrema.res) <- colnames(res$p)
saveRDS(mrema.res, file = "samplepara.RDS")</pre>
```

##non-parametric tests – doesnt make any assumptions about the LFC distribution of a group of genes

We can now launch the shiny app to investigate our results

```
mremApp() #parametric tests
REshine() #non-parametric tests
```

If there are any issues related to mremaR package, please contact - d.oshea20@universityofgalway.ie (Dónal O'Shea - Wrote the package)

Queries can also be sent to m.chopra1@universityofgalway.ie / mehak.chopra@yale.edu