# Olivia R package

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### Introduction

Here we demonstrate how to perform association analyses of continuous phenotypes using the Olivia package with RNA-seq data based on the pipeline proposed in the manuscript "Benchmarking Association Analyses of Continuous Exposures with RNA-seq in Observational Studies" https://www.biorxiv.org/content/10.1101/2021.02.12.430989v1.abstract.

# Installation and require packages

To install, open R and type:

```
library("devtools")
install_github("nkurniansyah/Olivia")
library(Olivia)
```

Olivia require external packages from CRAN (dplyr) and Bioconductor(qvalue)

```
install.packages("dplyr")

BiocManager::install("qvalue")

Load packages
library(dplyr)
library(qvalue)
```

### Load example data

#### Load raw gene counts matrix

First we load the transcripts, where were obtained from https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE151243.

Note: we reformatted the transcripts matrix into desired form and embedded them into Olivia package.

```
data(rnaseq_count_matrix)
rnaseq_count_matrix[1:5,1:5]
```

```
##
                   10L_S26_L006_R1_001 10N_S15_L003_R1_001 11L_S35_L007_R1_001
## ENSG0000000003
                                                                              525
## ENSG00000000005
                                      5
                                                                                2
                                                          19
## ENSG0000000419
                                    883
                                                        1058
                                                                              609
## ENSG0000000457
                                    790
                                                        1009
                                                                              619
## ENSG0000000460
                                    206
                                                         289
                                                                              272
##
                   11N_S25_L005_R1_001 12L_S14_L003_R1_001
                                    701
## ENSG00000000003
## ENSG0000000005
                                     64
                                                          25
## ENSG0000000419
                                    547
                                                         576
                                                         650
## ENSG0000000457
                                    887
## ENSG0000000460
                                    214
                                                         334
```

#### Load simulated phenotypes

We simulated in advance a data.frame of phenotypes.

```
data(phenotype)
head(phenotype)
```

```
##
                       Age Sex Trait.1 Trait.2
## 10L_S26_L006_R1_001
                            0 16.06608 15.58321
## 10N S15 L003 R1 001
                       19
                            0 21.20045 20.61345
## 11L_S35_L007_R1_001
                       20
                            0 14.44867 13.88567
                       21
## 11N_S25_L005_R1_001
                            0 35.89606 34.84859
## 12L_S14_L003_R1_001
                       22
                            0 24.09078 24.42536
## 12N_S27_L006_R1_001
                            0 29.61045 26.62854
                       21
```

We define the trait of interest to study as an exposure associated with genes. The trait/phenotype has to correspond to a column name in the phenotype data frame.

```
trait <- "Trait.1"
```

We will adjust our analysis to the simulated covariates Age and Sex. The covariates have to correspond to column names in the phenotype data frame. In the analysis, we will use a string defining the regression model (just the covariates part of it), so we define it here:

```
covariates_string <- "Age + as.factor(Sex)"</pre>
```

Note that we can also define the string to be "Age + as.factor(Sex)", or use interaction terms, like one would use in regression functions in R.

Match the (simulated) individuals between the phenotype and the RNA-seq count matrix. Make sure the there are matching IDs.

```
IDs_both <- intersect(rownames(phenotype), colnames(rnaseq_count_matrix))
rnaseq_matrix <- rnaseq_count_matrix[, IDs_both]
phenotypes <- phenotype[match(IDs_both,rownames(phenotype)),]</pre>
```

# Normalize the RNA-seq dataset

We use median normalization in Olivia to reduce package dependencies. However, users can use different normalization method using different packages, for example: estimateSizeFator(DESeq2) or TMM(edgeR). Here we show how each of these is used. There are no downstream differences in how the methods are applied once the data is normalized.

#### Median normalization

```
median_norm<- median_normalization(rnaseq_matrix)</pre>
```

#### estimateSizeFactor

This method implemented in DESeq2.

#### TMM (Trimmed Mean of M-values)

This method implemented in edgeR

```
BiocManager::install("edgeR")
library(edgeR)
counts <- DGEList(rnaseq_matrix)

dgList<- calcNormFactors(counts, method = "TMM")</pre>
```

```
TMM_norm<- cpm(dgList)</pre>
```

### Filtering transcripts

Remove lowly express gene counts

```
clean_count_matrix <- apply_filters(count_matrix = median_norm,</pre>
                                    median_min = 1,
                                    expression sum min = 10,
                                    \max \min = 10,
                                    range_min = 5,
                                    prop_zero_max = 0.5)
## applying filters on a transcript count matrix of 58051 transcripts, across 40 individuals
## Computing transtripts characteristics...
## Appying filters...
## There are 23987 transcripts with median
##
                     value lower than 1
## There are 14190 transcripts with expression sum
##
                     value lower than 10
## There are 22297 transcripts with maximum expression
                     value lower than 10
##
## There are 17188 transcripts with maximum
##
                     expression range value lower than 5
## There are 21923 transcripts with propotion
##
                     of zero counts higher than 0.5
## Removing 24834 unique transcripts not passing requested filters
```

After filtering gene counts, there are 33217 remaining for differential expression analysis.

# Perform differential expression analysis

## Computing empirical p-values

We show how we perform differential expression analysis on all transcripts using empirical p-value(quantile empirical p-values and Storey empirical p-values, as these are reffered to in the manuscript). In order to generate p-values under the null, we create a "residual permuted" trait 100 times and perform differential expression analysis, and use the resulting p-values/z-score as our null p-values/z-score. (See manuscript).

```
set.seed(12)
quantile_emp<-lm_count_mat_emp_pval(clean_count_matrix, pheno=phenotypes, trait, covariates_string,
                                  n_permute=100, gene_IDs=NULL,log_transform = "log_replace_half_min",
                                  stat_type="p_value", empirical_type = "quantile",
                                  t_df = NULL, outcome_type ="continous")
## Performing residual permutation to generate permuted trait...
## performing differential expression analysis on 100 permuted traits
```

```
## Run quantile empirical p-values using p_value
head(quantile_emp)
##
              geneID
                            beta
                                                t_stat t_stat_df
                                                                       p_value
## 1 ENSG0000000000 0.02140173 0.01055381
                                             2.0278677
                                                               36 5.002424e-02
## 2 ENSG00000000005
                      0.16670092 0.02639112
                                             6.3165534
                                                               36 2.629257e-07
## 3 ENSG0000000419 0.00588087 0.01466879
                                             0.4009102
                                                               36 6.908559e-01
## 4 ENSG0000000457 0.00800729 0.01044979 0.7662634
                                                               36 4.485152e-01
## 5 ENSG0000000460 -0.01315765 0.01333808 -0.9864726
                                                               36 3.304853e-01
## 6 ENSG00000000938 -0.06909230 0.02051793 -3.3674103
                                                               36 1.817907e-03
##
          fdr bh
                    emp_pvals bh_emp_pvals
## 1 0.296579900 5.330042e-02
                                 0.3159968
## 2 0.004366802 1.505253e-07
                                 0.0025000
## 3 0.904505585 6.962342e-01
                                 0.9115463
## 4 0.788532027 4.553238e-01
                                 0.8004866
## 5 0.702422623 3.371201e-01
                                 0.7165153
## 6 0.068018905 2.079658e-03
                                 0.0778805
tophits<-quantile_emp[which(quantile_emp$bh_emp_pvals< 0.05),]</pre>
head(tophits)
                                                   t_stat t_stat_df
##
                geneID
                              beta
                                                                         p_value
       ENSG0000000005
                        0.16670092 0.026391120
                                                                 36 2.629257e-07
## 2
                                                6.316553
## 231 ENSG0000009709
                        0.12205262 0.028668757
                                                4.257339
                                                                 36 1.413253e-04
## 248 ENSG0000010278
                        0.03556122 0.007342435
                                                4.843246
                                                                 36 2.423317e-05
## 328 ENSG00000013503 0.02283551 0.006028366
                                                3.788009
                                                                 36 5.569283e-04
## 337 ENSG00000013810 -0.05450117 0.013567542 -4.017026
                                                                 36 2.868601e-04
## 369 ENSG00000018280 -0.11359856 0.029790212 -3.813285
                                                                 36 5.179180e-04
##
            fdr bh
                      emp pvals bh emp pvals
       0.004366802 1.505253e-07
                                  0.00250000
## 231 0.036960240 1.207213e-04
                                  0.03351724
## 248 0.032198126 1.746094e-05
                                  0.02320000
## 328 0.047191580 5.870488e-04
                                  0.04974490
## 337 0.039758853 2.832887e-04
                                  0.03933884
## 369 0.046122473 5.403859e-04
                                  0.04812332
```

# Perform differential expression analysis using multiple exposure

We show how we perform differential expression analysis on all transcripts using emprical p-value(quantile empirical p-values and Storey empirical p-values) using multiple exposure.

## Computing empirical p-values

```
## Run quantile empirical p-values using p_value
head(quantile_emp_multi)
##
              geneID beta_Trait.1 beta_Trait.2 chisq_stat chisq_stat_df
                                    0.11755276 6.4796146
## 1 ENSG0000000003 -0.095383807
                                                                       2
## 2 ENSG0000000000 0.381867680
                                   -0.21658030 41.2674590
                                                                       2
## 3 ENSG00000000419 -0.145930094
                                    0.15280829 2.0975976
                                                                       2
                                                                       2
## 4 ENSG00000000457 -0.039683211
                                    0.04800380 0.9365473
## 5 ENSG0000000460 -0.115913899
                                                                       2
                                    0.10343131 2.0195879
                                                                       2
## 6 ENSG00000000938 -0.005696896
                                   -0.06381189 11.2401449
##
                        fdr_bh
                                  emp_pvals bh_emp_pvals
          p_value
## 1 3.917144e-02 2.651878e-01 5.391155e-02
                                               0.3649745
## 2 1.093667e-09 3.632834e-05 1.204203e-06
                                               0.0200000
## 3 3.503583e-01 6.809041e-01 3.728952e-01
                                               0.7246800
## 4 6.260822e-01 8.451406e-01 6.418391e-01
                                               0.8663758
## 5 3.642940e-01 6.923092e-01 3.866234e-01
                                               0.7347363
## 6 3.624378e-03 9.232925e-02 7.779149e-03
                                               0.1980963
quantile_emp_multi<-quantile_emp_multi[quantile_emp_multi$bh_emp_pvals< 0.05,]
quantile_emp_multi
##
                 geneID beta_Trait.1 beta_Trait.2 chisq_stat chisq_stat_df
## 2
        ENSG00000000005
                           0.3818677
                                      -0.21658030
                                                     41.26746
## 8164 ENSG00000144821
                                                                          2
                           0.1339460
                                      -0.03822377
                                                    39.15829
                                     emp_pvals bh_emp_pvals
##
                           fdr_bh
             p_value
## 2
        1.093667e-09 3.632834e-05 1.204203e-06
                                                        0.02
## 8164 3.139679e-09 5.214536e-05 1.204203e-06
                                                        0.02
```

# Perform differential expression analysis using permutation

When testing only a handful of genes, we may not want to perform transfcriptome-wide association analysis, and therefore empirical p-values using the quantile or Storey's approach cannot be computed (not enough tests to generate the null distribution). Therefore, in this case one needs to permute the specific gene many times.

Here we show how to perform differential expression analysis on selected transcripts when computing a permutation p-value for each gene based on permutations for this gene only. We suggest running 100000 permutations.

#### perm\_res

```
beta
                                               t_stat t_stat_df
             geneID
                                        se
                                                                p_value
## 3 ENSG00000211888 -0.100704705 0.03780596 -2.6637255
                                                            36 0.01148709
## 2 ENSG00000100416 -0.011504192 0.01349350 -0.8525725
                                                            36 0.39953313
## 1 ENSG00000039650 -0.031589560 0.01114510 -2.8343890
                                                            36 0.00748205
## 4 ENSG00000249700 -0.007725497 0.01928230 -0.4006523
                                                            36 0.69104411
## 5 ENSG00000264932 0.021374024 0.03900637 0.5479623
                                                            36 0.58709966
##
    perm_pval
## 3 0.01164
## 2 0.40085
## 1 0.00621
## 4 0.69280
## 5 0.58565
```