

Ensemble of Gene Set Enrichment Analyses

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May 15, 2016

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1 Introduction

The *EGSEA* package implements the Ensemble of Gene Set Enrichment Analysis (EGSEA) algorithm that utilizes the analysis results of eleven prominent GSE algorithms in the literature to calculate collective significance scores for each gene set. These methods include: *ora* [1], *globaltest* [2], *plage* [3], *safe* [4], *zscore* [5], *gage* [6], *ssgsea* [7], *roast* [8], *PADOG* [9], *camera* [10] and *GSVA* [11]. The *ora*, *gage*, *camera* and *gsva* methods depend on a competitive null hypothesis while the remaining seven methods are based on a self-contained hypothesis. Conveniently, *EGSEA* is not limited to these eleven GSE methods and new GSE tests can be easily integrated into the framework. The *plage*, *zscore* and *ssgsea* algorithms are implemented in the *GSVA* package and *camera* and *roast* are implemented in the *limma* package. *EGSEA* was implemented with parallel computation enabled using the *parallel* package. There are two levels of parallelism in EGSEA: (i) parallelism at the method-level and (ii) parallelism at the experimental contrast level. A wrapper function was written for each individual GSE method to utilize existing R packages and create a universal interface for all methods. The *ora* method was implemented using the *phyper* function from the *stats* package, which estimates the hypergeometric distribution for a 2×2 contingency table.

RNA-seq reads are first aligned to the reference genome and mapped reads are assigned to annotated genomic features to obtain a summarized *count matrix*. The *EGSEA* package was developed so that it can accept a count matrix or a *voom* object. Most of the GSE methods were intrinsically designed to work with microarray expression values and not with RNA-seq counts, hence the *voom* transformation is applied to the count matrix to generate an expression matrix applicable for use with these methods [12]. Since gene set tests are most commonly applied when two experimental conditions are compared, a design matrix and a contrast matrix are used to construct the experimental comparisons of interest. The target collection of gene sets is indexed so that the gene identifiers can be substituted with the indices of genes in the rows of the count matrix. The GSE analysis is then carried out by each of the selected methods independently and an FDR value is assigned to each gene set. Lastly, the ensemble functions are invoked to calculate collective significance scores for each gene set.

The *EGSEA* package also allows for performing over-representation analysis on the EGSEA gene set collections that were adopted from MSigDB, KEGG and GeneSetDB databases.

2 Citation

- Monther Alhamdoosh, Milica Ng, Nicholas J. Wilson, Julie M. Sheridan, Huy Huynh, Michael J. Wilson and Matthew E. Ritchie. Combining multiple tools outperforms individual methods in gene set enrichment analyses.

3 Installation instructions

The *EGSEA* package was developed so that it harmonizes with the existing R packages in the CRAN repository or the Bioconductor project.

3.1 System prerequisites

EGSEA does not require any software package or library to be installed before it can be installed regardless of the operating system.

3.2 R package dependencies

The *EGSEA* package depends on several R packages that are not in the Bioconductor project. These packages are listed below:

- *HTMLUtils* facilitates automated HTML report creation, in particular framed HTML pages and dynamically sortable tables. It is used in *EGSEA* to generate the stats tables. To install it, type in the R console
`install.packages("HTMLUtils")`
- *hwriter* has easy-to-use and versatile functions to output R objects in HTML format. It is used in this package to create the HTML pages of the EGSEA report. To install it,
`install.packages("hwriter")`
- *ggplot2* is an implementation of the grammar of graphics in R. It is used in this package to create the summary plots. To install it, type
`install.packages("ggplot2")`
- *gplots* has various R programming tools for plotting data. It is used in *EGSEA* to create heatmaps. To install it, run
`install.packages("gplots")`
- *stringi* allows for fast, correct, consistent, portable, as well as convenient character string/text processing in every locale and any native encoding. It is used in generating the HTML pages. To install this package, type
`install.packages("stringi")`
- *parallel* handles running much larger chunks of computations in parallel. It is used to carry out gene set tests on parallel. It is usually installed with R.

3.2.1 Bioconductor packages

The Bioconductor packages that need to be installed in order for *EGSEA* to function properly are: *PADOG*, *GSVA*, *AnnotationDbi*, *topGO*, *pathview*, *gage*, *globaltest*, *limma*, *edgeR*, *safe*, *org.Hs.eg.db*, *org.Mm.eg.db*, *org.Rn.eg.db*. These packages can be installed from Bioconductor using the following commands in R console

```
source("http://www.bioconductor.org/biocLite.R")
biocLite(c("PADOG", "GSVA", "AnnotationDbi", "topGO", "pathview",
          "gage", "globaltest", "limma", "edgeR", "safe", "org.Hs.eg.db",
          "org.Mm.eg.db", "org.Rn.eg.db"))
```

3.2.2 Essential data package

The gene set collections that are used by *EGSEA* were preprocessed and converted into R data objects to be used by the EGSEA functions. The data objects are stored in an R package, named *EGSEAdata*. It contains

the gene set collections that are used by *EGSEA* to perform gene set testing. *EGSEAdata* can be installed from Bitbucket.

To install packages from Bitbucket, *devtools* should be installed. *devtools* Package devtools is available at CRAN. For Windows this seems to depend on having Rtools for Windows installed. You can download and install this from: <http://cran.r-project.org/bin/windows/Rtools/>

To install *devtools*, run

```
install.packages("devtools")
```

To install *EGSEAdata*, run in R console the following commands

```
library(devtools)
install_bitbucket("malhamdoosh/egseadata", ref = "Stable_Release")
```

3.3 Installation

3.3.1 Bioconductor

In R console, type

```
source("http://bioconductor.org/biocLite.R")
biocLite("EGSEA")
```

3.3.2 Bitbucket

To install EGSEA from bitbucket, type in the R console

```
library(devtools)
install_bitbucket("malhamdoosh/egsea", ref = "Stable_Release")
```

4 Quick start

4.1 EGSEA gene set collections

The Molecular Signatures Database (MSigDB) [13] v5.0 was downloaded from <http://www.broadinstitute.org/gsea/msigdb> (05 July 2015, date last accessed) and the human gene sets were extracted for each collection (h, c1, c2, c3, c4, c5, c6, c7). Mouse orthologous gene sets of these MSigDB collections were adopted from <http://bioinf.wehi.edu.au/software/MSigDB/index.html> [10]. EGSEA uses Entrez Gene identifiers [14] and alternate gene identifiers must be first converted into Entrez IDs. KEGG pathways [15] for mouse and human were downloaded using the *gage* package. To extend the capabilities of EGSEA, a third database of gene sets was downloaded from the GeneSetDB [16] <http://genesetdb.auckland.ac.nz/sourcedb.html> project. In total, more than 20,000 gene sets have been collated along with annotation information for each set (where available).

The *EGSEA* package has four main functions that utilizes the gene set collections of *EGSEAdata*. They map the dataset Entrez gene IDs into the available gene sets of each collection and create indexes for each gene set

collection. They also compile annotation information for each gene set to be integrated with the final EGSEA report. These functions are:

- `buildKEGGIdxEZID` indexes the KEGG pathway gene sets and loads gene set annotation. Type `?buildKEGGIdxEZID` in the console to see how to use this function.
- `buildMSigDBIdxEZID` indexes the MSigDB gene sets and loads gene set annotation. Type `?buildMSigDBIdxEZID` in the console to see how to use this function.
- `buildGeneSetDBIdxEZID` indexes the GeneSetDB gene sets and loads gene set annotation. Type `?buildGeneSetDBIdxEZID` in the console to see how to use this function.
- `buildIdxEZID` indexes the MSigDB and KEGG gene sets and loads gene set annotation. Type `?buildIdxEZID` in the console to see how to use this function.

The above functions require a vector of Entrez Gene IDs and the species name. To use the output of these functions with the [EGSEA](#) functions, the order of gene ids in the `entrezIDs` parameter should match that of the row names of the count matrix or the voom object.

4.2 A simple example

The [EGSEA](#) package basically performs gene set enrichment analysis on a voom object generated by the `voom` function from the [limma](#) package. It was primarily developed to extend the limma-voom RNA-seq analysis pipeline.

To quickly start with [EGSEA](#) analysis, an example on analyzing a human IL-13 dataset is presented here. This experiment aims to identify the biological pathways and diseases associated with the cytokine Interleukin 13 (IL-13) using gene expression measured in peripheral blood mononuclear cells (PBMCs) obtained from 3 healthy donors. The expression profiles of *in vitro* IL-13 stimulation were generated using RNA-seq technology for 3 PBMC samples at 24 hours. The transcriptional profiles of PBMCs without IL-13 stimulation were also generated to be used as controls. Finally, an IL-13R α 1 antagonist was introduced into IL-13 stimulated PBMCs and the gene expression levels after 24h were profiled to examine the neutralization of IL-13 signaling by the antagonist. Only two samples were available for the last condition. Single-end 100bp reads were obtained via RNA-seq from total RNA using a HiSeq 2000 Illumina sequencer. TopHat was used to map the reads to the human reference genome (GRCh37.p10). HTSeq was then used to summarize reads into a gene-level count matrix. The TMM method from the [edgeR](#) package was used to normalize the RNA-seq counts.

To perform EGSEA analysis on this dataset, the [EGSEA](#) package is first loaded:

```
library(EGSEA)

## Loading required package: Biobase
## Loading required package: BiocGenerics
## Loading required package: parallel
##
## Attaching package: 'BiocGenerics'
## The following objects are masked from 'package:parallel':
##
##   clusterApply, clusterApplyLB, clusterCall, clusterEvalQ, clusterExport,
##   clusterMap, parApply, parCapply, parLapply, parLapplyLB, parRapply,
##   parSapply, parSapplyLB
```

```
## The following objects are masked from 'package:stats':  
##  
##   IQR, mad, xtabs  
  
## The following objects are masked from 'package:base':  
##  
##   Filter, Find, Map, Position, Reduce, anyDuplicated, append, as.data.frame,  
##   cbind, colnames, do.call, duplicated, eval, evalq, get, grep, grepl,  
##   intersect, is.unsorted, lapply, lengths, mapply, match, mget, order, paste,  
##   pmax, pmax.int, pmin, pmin.int, rank, rbind, rownames, sapply, setdiff,  
##   sort, table, tapply, union, unique, unsplit  
  
## Welcome to Bioconductor  
##  
##   Vignettes contain introductory material; view with 'browseVignettes()'. To  
##   cite Bioconductor, see 'citation("Biobase")', and for packages  
##   'citation("pkgname")'.  
  
## Loading required package:  gage  
## Loading required package:  AnnotationDbi  
## Loading required package:  stats4  
## Loading required package:  IRanges  
## Loading required package:  S4Vectors  
  
##  
## Attaching package:  'S4Vectors'  
  
## The following objects are masked from 'package:base':  
##  
##   colMeans, colSums, expand.grid, rowMeans, rowSums  
  
## Loading required package:  topGO  
## Loading required package:  graph  
## Loading required package:  GO.db  
  
##  
## Loading required package:  SparseM  
  
##  
## Attaching package:  'SparseM'  
  
## The following object is masked from 'package:base':  
##  
##   backsolve  
  
##  
## groupGOTerms:  GOBPterm, GOMFterm, GOCCterm environments built.
```

```
##
## Attaching package: 'topGO'

## The following object is masked from 'package:IRanges':
##
##   members

## The following object is masked from 'package:gage':
##
##   geneData

## Loading required package: pathview

## Loading required package: org.Hs.eg.db

##

## #####
## Pathview is an open source software package distributed under GNU General
## Public License version 3 (GPLv3). Details of GPLv3 is available at
## http://www.gnu.org/licenses/gpl-3.0.html. Particullary, users are required to
## formally cite the original Pathview paper (not just mention it) in publications
## or products. For details, do citation("pathview") within R.
##

## The pathview downloads and uses KEGG data. Non-academic uses may require a KEGG
## license agreement (details at http://www.kegg.jp/kegg/legal.html).
## #####

##

## KEGG.db contains mappings based on older data because the original resource was
## removed from the the public domain before the most recent update was produced.
## This package should now be considered deprecated and future versions of
## Bioconductor may not have it available. Users who want more current data are
## encouraged to look at the KEGGREST or reactome.db packages

##

##

##

##
```

Then, the voom data object of this experiment from *EGSEAdata* is loaded to perform the EGSEA analysis:

```
library(EGSEAdata)
data(il13.data)
v = il13.data$voom
names(v)

## [1] "genes" "targets" "E" "weights" "design"

v$design

## X24 X24IL13 X24IL13Ant X40513 X40913
```

```
## 1 0 1 0 0 0
## 2 0 0 1 0 0
## 3 1 0 0 1 0
## 4 0 1 0 1 0
## 5 1 0 0 0 1
## 6 0 1 0 0 1
## 7 0 0 1 0 1
## 8 1 0 0 0 0
## attr("assign")
## [1] 1 1 1 2 2
## attr("contrasts")
## attr("contrasts")$`d1$samples$group`
## [1] "contr.treatment"
##
## attr("contrasts")$`d1$samples$Date`
## [1] "contr.treatment"

contrasts = il13.data$contra
contrasts

##           Contrasts
## Levels      X24IL13 - X24 X24IL13Ant - X24IL13
##   X24                -1                0
##   X24IL13              1               -1
##   X24IL13Ant            0                1
##   X40513                0                0
##   X40913                0                0
```

Before the EGSEA function is called gene set collection(s) needs to be preprocessed and indexed using EGSEA indexing functions that were presented earlier. For example, to use the KEGG pathway collections without the Metabolism pathways, type

```
# prepare gene set collections
gs.annots = buildIdxEZID(entrezIDs = rownames(v$E), species = "human",
  msgdb.gsets = "c5", kegg.exclude = c("Metabolism"))

## [1] "Reading Broad Gene Sets ... "
## [1] "Created the gs.annot$original for c5 \n..."
## [1] "Created the gs.annot$idx for c5 ..."
## [1] "Created the gs.annot$anno for c5 ..."
## [1] "Building KEGG pathways annotation object ... "

names(gs.annots)

## [1] "c5" "kegg"
```

Finally, the EGSEA analysis can be invoked using the `egsea` function as follows

```
# perform the EGSEA analysis set display.top = 20 to display
# more gene sets. It takes longer time to run.
gsa = egsea(voom.results = v, contrasts = contrasts, gs.annots = gs.annots,
  symbolsMap = v$genes, baseGSEAs = egsea.base()[-2], display.top = 3,
```



```

## INFLAMMATORY_RESPONSE          M10617 0.000000e+00
## DEFENSE_RESPONSE              M3458 0.000000e+00
## FATTY_ACID_BIOSYNTHETIC_PROCESS M11481 8.080358e-07
## [1] "GO graphs are being generated for top-ranked GO terms based on \np-values ... "

##
## Building most specific GOs .....
## ( 9828 GO terms found. )

##
## Build GO DAG topology .....
## ( 13656 GO terms and 32783 relations. )

##
## Annotating nodes .....
## ( 11749 genes annotated to the GO terms. )

##
## Building most specific GOs .....
## ( 3395 GO terms found. )

##
## Build GO DAG topology .....
## ( 3908 GO terms and 4929 relations. )

##
## Annotating nodes .....
## ( 11797 genes annotated to the GO terms. )

##
## Building most specific GOs .....
## ( 1403 GO terms found. )

##
## Build GO DAG topology .....
## ( 1664 GO terms and 3283 relations. )

##
## Annotating nodes .....
## ( 12389 genes annotated to the GO terms. )
## [1] "X24IL13 - X24"

## Loading required package: Rgraphviz
## Loading required package: grid

##
## Attaching package: 'grid'

```

```

## The following object is masked from 'package:topGO':
##
##   depth
##
## Attaching package: 'Rgraphviz'
##
## The following objects are masked from 'package:IRanges':
##
##   from, to
##
## The following objects are masked from 'package:S4Vectors':
##
##   from, to
##
## [1] "X24IL13Ant - X24IL13"
## [1] "Heat maps are being generated for top-ranked gene sets based on \nlogFC ... "
## [1] "Summary plots are being generated ... "
## E:\biocbld\bbs-3.3-bioc\tmpdir\RtmpaGW3rg\Rbuild10589227c88\EGSEA\vignettes\il13-egsea-report/
## [1] "EGSEA is running on the provided data and kegg gene sets"
## [1] "   Running CAMERA for X24IL13 - X24"
## [1] "   Running CAMERA for X24IL13Ant - X24IL13"
## [1] "Running CAMERA on all \ncontrasts ... COMPLETED "
## 203 categories formed
## [1] "   Running SAFE for X24IL13 - X24"
## Warning: only 20 unique resamples exist
##       switching to exhaustive permutation
## [1] "   Running SAFE for X24IL13Ant - X24IL13"
## Warning: only 20 unique resamples exist
##       switching to exhaustive permutation
## [1] "Running SAFE on all \ncontrasts ... COMPLETED "
## [1] "   Running GAGE for X24IL13 - X24"
## [1] "   Running GAGE for X24IL13Ant - X24IL13"
## [1] "Running GAGE on all \ncontrasts ... COMPLETED "
## [1] "   Running PADOG for X24IL13 - X24"
## [1] "   Running PADOG for X24IL13Ant - X24IL13"
## [1] "Running PADOG on all \ncontrasts ... COMPLETED "
## [1] "   Running PLAGE for X24IL13 - X24"
## [1] "   Running PLAGE for X24IL13Ant - X24IL13"
## [1] "Running PLAGE on all \ncontrasts ... COMPLETED "
## [1] "   Running ZSCORE for X24IL13 - X24"
## [1] "   Running ZSCORE for X24IL13Ant - X24IL13"
## [1] "Running ZSCORE on all \ncontrasts ... COMPLETED "
## [1] "   Running GSVA for X24IL13 - X24"
## [1] "   Running GSVA for X24IL13Ant - X24IL13"
## [1] "Running GSVA on all \ncontrasts ... COMPLETED "
## [1] "   Running SSGSEA for X24IL13 - X24"
## [1] "   Running SSGSEA for X24IL13Ant - X24IL13"
## [1] "Running SSGSEA on all \ncontrasts ... COMPLETED "

```

```

## [1] "    Running GLOBALTEST for X24IL13 - X24"
## [1] "    Running GLOBALTEST for X24IL13Ant - X24IL13"
## [1] "Running GLOBALTEST on all \ncontrasts ... COMPLETED "
## [1] "    Running ORA for X24IL13 - X24"
## [1] "    Running ORA for X24IL13Ant - X24IL13"
## [1] "Running ORA on all \ncontrasts ... COMPLETED "
## [1] "Writing out the top-ranked gene sets for each contrast .. \nKEGG gene sets"
## [1] "The top gene sets for contrast X24IL13 - X24 are:"
##                                     Type           p.adj
## Asthma                             Disease 2.816639e-09
## Intestinal immune network for IgA production Signaling 7.624705e-08
## Amoebiasis                         Disease 1.110637e-07
## [1] "The top gene sets for contrast X24IL13Ant - X24IL13 are:"
##                                     Type           p.adj
## NOD-like receptor signaling pathway Signaling 3.635776e-06
## Asthma                             Disease 7.619107e-07
## Toll-like receptor signaling pathway Signaling 3.155081e-08
## Pathway maps are being generated for top-ranked
## pathways based
## on logFC ...
## [1] "    X24IL13 - X24"
## [1] "    X24IL13Ant - X24IL13"
## [1] "Heat maps are being generated for top-ranked gene sets based on \nlogFC ... "
## [1] "Summary plots are being generated ... "
## [1] "Comparison summary plots are being generated ... "
## Pathway maps are being generated for top-ranked
## comparative
## pathways based on logFC ...

topSets(gsa, contrast = 1, gs.label = "kegg", number = 10)

## [1] "Asthma"
## [2] "Intestinal immune network for IgA production"
## [3] "Amoebiasis"
## [4] "Viral myocarditis"
## [5] "Endocrine and other factor-regulated calcium reabsorption"
## [6] "Legionellosis"
## [7] "HTLV-I infection"
## [8] "Prion diseases"
## [9] "Toxoplasmosis"
## [10] "Proteoglycans in cancer"

topSets(gsa, contrast = 1, gs.label = "kegg", sort.by = "ora",
        number = 10, names.only = FALSE)

##                                     p.value           p.adj vote.rank avg.rank
## Cytokine-cytokine receptor interaction 0.000000e+00 0.000000e+00      5      56.0
## Staphylococcus aureus infection        9.992007e-15 6.761258e-13     10      69.2
## Hematopoietic cell lineage             0.000000e+00 0.000000e+00      5      50.9

```

## Phagosome	2.950185e-10	1.197775e-08	130	68.0
## Tuberculosis	1.567893e-08	2.652353e-07	5	59.2
## Leishmaniasis	7.178571e-09	1.619166e-07	20	80.0
## Cell adhesion molecules (CAMs)	4.376893e-09	1.110637e-07	10	79.0
## Rheumatoid arthritis	5.168753e-08	7.494692e-07	10	96.0
## Asthma	5.550027e-11	2.816639e-09	10	26.0
## Type I diabetes mellitus	9.536817e-06	8.417278e-05	100	98.2
##	med.rank	min.pvalue	min.rank	avg.logFC
## Cytokine-cytokine receptor interaction	47.0	2.687045e-16	1	0.7563135
## Staphylococcus aureus infection	74.5	1.836804e-15	2	0.6701979
## Hematopoietic cell lineage	14.0	8.441405e-12	2	0.7557242
## Phagosome	64.0	1.685628e-11	4	0.5389219
## Tuberculosis	58.0	1.103390e-09	4	0.4968563
## Leishmaniasis	40.0	1.915394e-09	6	0.5805793
## Cell adhesion molecules (CAMs)	86.0	1.007610e-08	7	0.5597125
## Rheumatoid arthritis	90.0	1.760956e-08	8	0.5947603
## Asthma	13.0	1.490995e-07	9	0.7024946
## Type I diabetes mellitus	110.0	2.629903e-07	10	0.6005818
##	Significance	camera	safe	gage
## Cytokine-cytokine receptor interaction	100.00000	30	76	1
## Staphylococcus aureus infection	81.88529	87	114	9
## Hematopoietic cell lineage	99.92208	12	16	2
## Phagosome	42.86011	129	87	26
## Tuberculosis	32.80426	111	83	33
## Leishmaniasis	39.58128	181	26	18
## Cell adhesion molecules (CAMs)	39.07866	14	98	10
## Rheumatoid arthritis	36.57453	202	14	8
## Asthma	60.30259	10	70	11
## Type I diabetes mellitus	24.56941	128	121	23
##	padog	plage	zscore	
## Cytokine-cytokine receptor interaction	1	170	64	
## Staphylococcus aureus infection	62	6	93	
## Hematopoietic cell lineage	2	128	203	
## Phagosome	13	41	96	
## Tuberculosis	12	42	74	
## Leishmaniasis	18	31	146	
## Cell adhesion molecules (CAMs)	81	152	113	
## Rheumatoid arthritis	71	159	161	
## Asthma	39	9	21	
## Type I diabetes mellitus	99	51	143	
##	gsva	ssgsea	globaltest	ora
## Cytokine-cytokine receptor interaction	78	118	21	1
## Staphylococcus aureus infection	134	182	3	2
## Hematopoietic cell lineage	103	6	34	3
## Phagosome	124	137	23	4
## Tuberculosis	119	109	4	5
## Leishmaniasis	180	145	49	6
## Cell adhesion molecules (CAMs)	193	31	91	7
## Rheumatoid arthritis	98	157	82	8
## Asthma	15	11	65	9
## Type I diabetes mellitus	146	165	96	10

```
topSets(gsa, contrast = "comparison", gs.label = "kegg", number = 10)
```

```
## [1] "Asthma"
## [2] "Viral myocarditis"
## [3] "Amoebiasis"
## [4] "HTLV-I infection"
## [5] "Legionellosis"
```

```
## [6] "NOD-like receptor signaling pathway"
## [7] "Intestinal immune network for IgA production"
## [8] "Cytokine-cytokine receptor interaction"
## [9] "Malaria"
## [10] "Endocrine and other factor-regulated calcium reabsorption"
```

The `egsea` returns a list of elements, one for each gene set collection and one for the comparative analysis. Each element is also a list of two elements: the `top.gene.sets`, which stores the top *display.top* gene sets for each contrast and the `test.results`, which stores the EGSEA test results for all gene sets along with the ensemble and individual rankings.

The EGSEA report of this experiment can be launched from [./il13-egsea-report/index.html](#).

Finally, the EGSEA analysis can be run with all the gene set collections that are available in the *EGSEAdata* package as follows

```
gs.annots = buildIdxEZID(entrezIDs = rownames(v$E), species = "human")

## [1] "Reading Broad Gene Sets ... "
## [1] "Created the gs.annot$original for h \n..."
## [1] "Created the gs.annot$idx for h ..."
## [1] "Created the gs.annot$anno for h ..."
## [1] "Created the gs.annot$original for c1 \n..."
## [1] "Created the gs.annot$idx for c1 ..."
## [1] "Created the gs.annot$anno for c1 ..."
## [1] "Created the gs.annot$original for c2 \n..."
## [1] "Created the gs.annot$idx for c2 ..."
## [1] "Created the gs.annot$anno for c2 ..."
## [1] "Created the gs.annot$original for c3 \n..."
## [1] "Created the gs.annot$idx for c3 ..."
## [1] "Created the gs.annot$anno for c3 ..."
## [1] "Created the gs.annot$original for c4 \n..."
## [1] "Created the gs.annot$idx for c4 ..."
## [1] "Created the gs.annot$anno for c4 ..."
## [1] "Created the gs.annot$original for c5 \n..."
## [1] "Created the gs.annot$idx for c5 ..."
## [1] "Created the gs.annot$anno for c5 ..."
## [1] "Created the gs.annot$original for c6 \n..."
## [1] "Created the gs.annot$idx for c6 ..."
## [1] "Created the gs.annot$anno for c6 ..."
## [1] "Created the gs.annot$original for c7 \n..."
## [1] "Created the gs.annot$idx for c7 ..."
## [1] "Created the gs.annot$anno for c7 ..."
## [1] "Building KEGG pathways annotation object ... "

gsetdb.annots = buildGeneSetDBIdxEZID(entrezIDs = rownames(v$E),
  species = "human")

## [1] "Building custom pathways annotation object ... "

gs.annots = c(gs.annots, gsetdb.annots)
```

```
names(gs.annots)
```

```
## [1] "h"      "c1"      "c2"      "c3"      "c4"      "c5"      "c6"
## [8] "c7"      "kegg"     "gsdbdrug" "gsdbdis" "gsdbgo"   "gsdbpath" "gsdbreg"
```

5 Ensemble of Gene Set Enrichment Analysis

Given an RNA-seq dataset D of samples from N experimental conditions, K annotated genes $g_k (k = 1, \dots, K)$, L experimental comparisons of interest $C_l (l = 1, \dots, L)$, a collection of gene sets Γ and M methods for gene set enrichment analysis, the objective of a GSE analysis is to find the most relevant gene sets in Γ which explain the biological processes and/or pathways that are perturbed in expression in individual comparisons and/or across multiple contrasts simultaneously. Numerous statistical gene set enrichment analysis methods have been proposed in the literature over the past decade. Each method has its own characteristics and assumptions on the analyzed dataset and gene sets tested. In principle, gene set tests calculate a statistic for each gene individually $f(g_k)$ and then integrate these significance scores in a framework to estimate a set significance score $h(\gamma_i)$.

We propose seven statistics to combine the individual gene set statistics across multiple methods, and to rank and hence identify biologically relevant gene sets. Assume a collection of gene sets Γ , a given gene set $\gamma_i \in \Gamma$, and that the GSE analysis results of M methods on γ_i for a specific comparison (represented by ranks R_i^m and statistical significance scores p_i^m , where $m = 1, \dots, M$ and $i = 1, \dots, |\Gamma|$) are given. The EGSEA scores can then be devised, for each experimental comparison, as follows:

- The p -value score is the average p -value assigned to γ_i
- The minimum p -value score is the smallest p -value calculated for γ_i
- The minimum rank score of γ_i is the smallest rank assigned to γ_i
- The average ranking score is the mean rank across the M ranks
- The median ranking score is the median rank across the M ranks
- The majority voting score is the most commonly assigned bin ranking
- The significance score assigns high scores to the gene sets with strong fold changes and high statistical significance

It is worth noting that the p -value score can only be calculated under the independence assumption of individual gene set tests, and thus it is not an accurate estimate of the ensemble gene set significance, but can still be useful for ranking results. The significance score is scaled into $[0, 100]$ range for each gene set collection. To learn more about the calculation of each EGSEA score, the original paper of this work is available at Section 2.

6 EGSEA report

Since the number of annotated gene set collections in public databases continuously increases and there is a growing trend towards generating dynamic analytical tools, our software tool was developed to enable users to interactively navigate through the analysis results by generating an HTML *EGSEA Report*. The report presents the results in different ways. For example, the *Stats table* displays the top n gene sets (where n is selected by the user) for each experimental comparison and includes all calculated statistics. Hyperlinks are enabled wherever possible, to access additional information on the gene sets such as annotation information. The gene

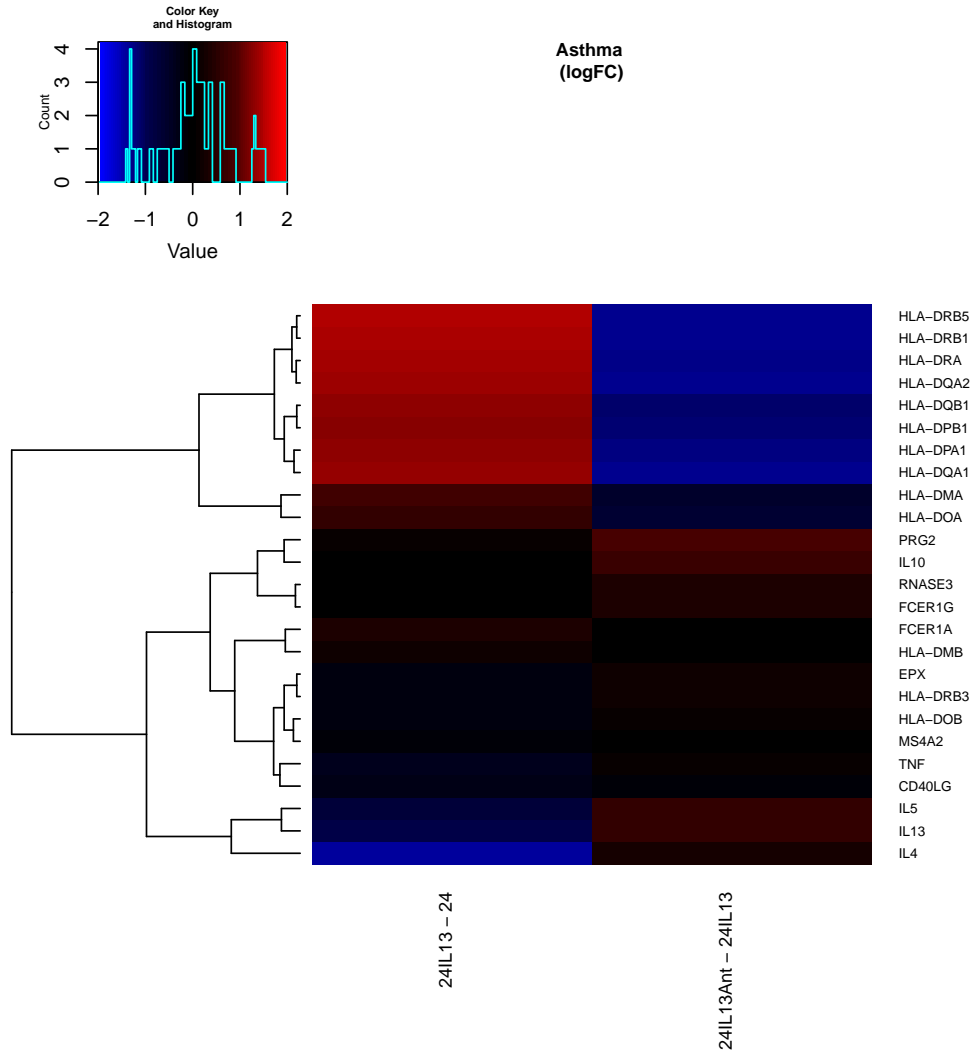


Figure 1: Asthma heatmap for the comparative analysis

expression fold changes can be visualized using heat maps for individual gene sets (Fig. 1) or projected onto pathway maps where available (e.g. KEGG gene sets) (Fig. 2). The most significant Gene Ontology (GO) terms for each comparison can be viewed in a GO graph that shows their relationships (Fig. 3).

Additionally, EGSEA creates summary plots for each gene set collection to visualize the overall statistical significance of gene sets (Fig. 4). Two types of summary plots are generated: (i) a plot that emphasizes the gene regulation direction and the significance score of a gene set and (ii) a plot that emphasizes the set cardinality and its rank. EGSEA also generates a multidimensional scaling (MDS) plot that shows how various GSE methods rank a collection of gene sets (Fig. 5). This plot gives insights into the similarity of different methods on a given dataset. Finally, the reporting capabilities of EGSEA can be used to extend any existing or newly developed GSE method by simply using only that method.

Similar reporting capabilities are also provided for the comparative analysis results of EGSEA (Fig. 6 and Fig. 7).

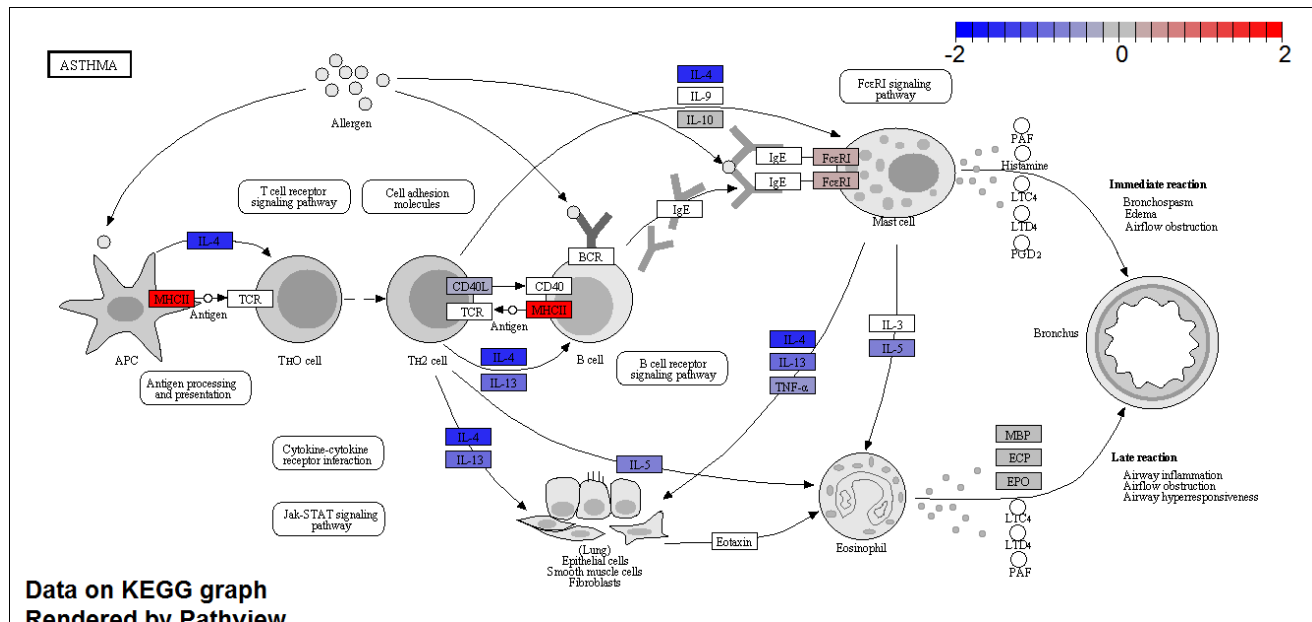


Figure 2: Asthma pathway map for the contrast X24IL13-X24

6.1 Comparative analysis

Unlike most GSE methods that calculate a gene set enrichment score for a given gene set under a single experimental contrast (e.g. disease vs. control), the comparative analysis proposed here allows researchers to estimate the significance of a gene set across multiple experimental contrasts. This analysis helps in the identification of biological processes that are perturbed by multiple experimental conditions simultaneously. Comparative significance scores are calculated for a gene set.

An interesting application of the comparative analysis would be finding pathways or biological processes that are activated by a stimulation with a particular cytokine yet are completely inhibited when the cytokine's receptor is blocked by an antagonist, revealing the functions uniquely associated with the signaling of that particular receptor as in the experiment below.

7 EGSEA on a non-human dataset

Epithelial cells from the mammary glands of female virgin 8-10 week-old mice were sorted into three populations of basal, luminal progenitor (LP) and mature luminal (ML) cells. Three independent samples from each population were profiled via RNA-seq on total RNA using an Illumina HiSeq 2000 to generate 100bp single-end read libraries. The *Rsubread* aligner was used to align these reads to the mouse reference genome (*mm10*) and mapped reads were summarized into gene-level counts using *featureCounts* with default settings. The raw counts are also normalized using the TMM method. Data are available from the GEO database as series GSE63310.

To perform EGSEA analysis on this dataset, the following commands are invoked in the R console

```
# load the mammary dataset
library(EGSEA)
```

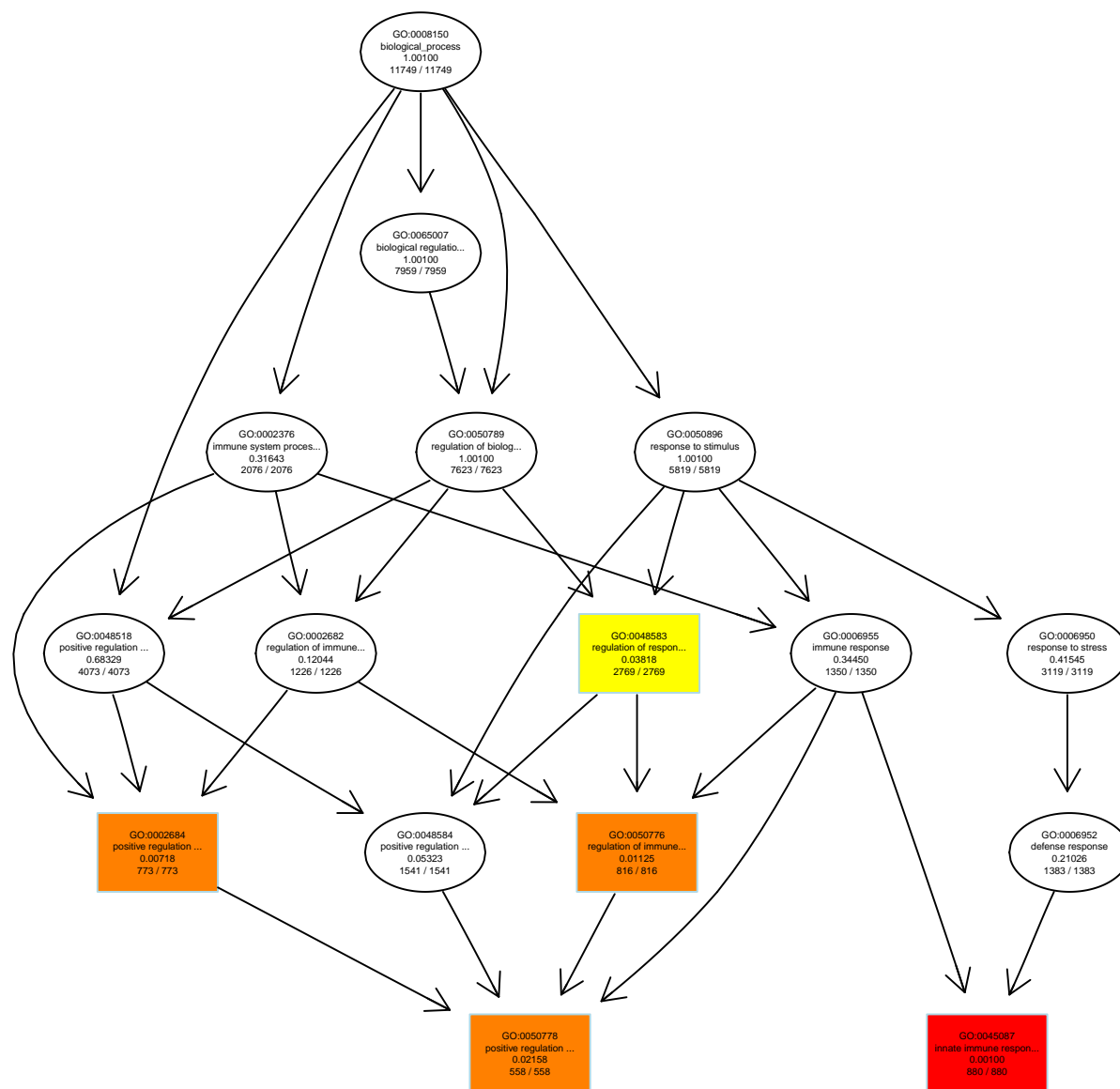


Figure 3: The top significant Biological Processes (BP) from GO terms.

```

library(EGSEAdata)
data(mam.data)
v = mam.data$voom
names(v)
v$design
contrasts = mam.data$contra
contrasts
# build the gene set collections

```

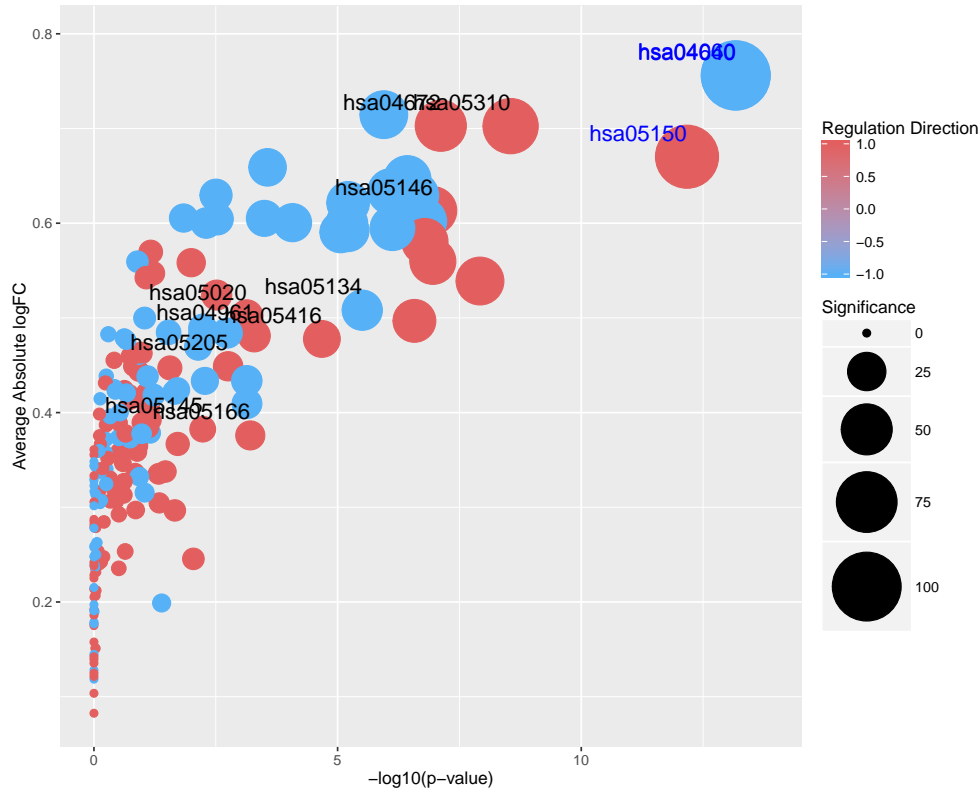


Figure 4: Summary plot for the contrast X24IL13-X24

```
gs.annots = buildIdxEZID(entrezIDs = rownames(v$E), species = "mouse",
  msigdb.gsets = "c2", kegg.exclude = "all")
names(gs.annots)
# create Entrez IDs - Symbols map
symbolsMap = v$genes[, c(1, 3)]
colnames(symbolsMap) = c("FeatureID", "Symbols")
symbolsMap[, "Symbols"] = as.character(symbolsMap[, "Symbols"])
# replace NA Symbols with IDs
na.sym = is.na(symbolsMap[, "Symbols"])
symbolsMap[na.sym, "Symbols"] = symbolsMap[na.sym, "FeatureID"]
# perform the EGSEA analysis set report = TRUE to generate
# the EGSEA interactive report
gsa = egsea(voom.results = v, contrasts = contrasts, gs.annots = gs.annots,
  symbolsMap = symbolsMap, baseGSEAs = egsea.base()[-c(2, 5,
    6, 9)], display.top = 20, sort.by = "med.rank", egsea.dir = "./mam-egsea-report",
  num.threads = 4, report = FALSE)
# show top 20 comparative gene sets in C2 collection
topSets(gsa, contrast = "comparison", gs.label = "c2", number = 20)
```

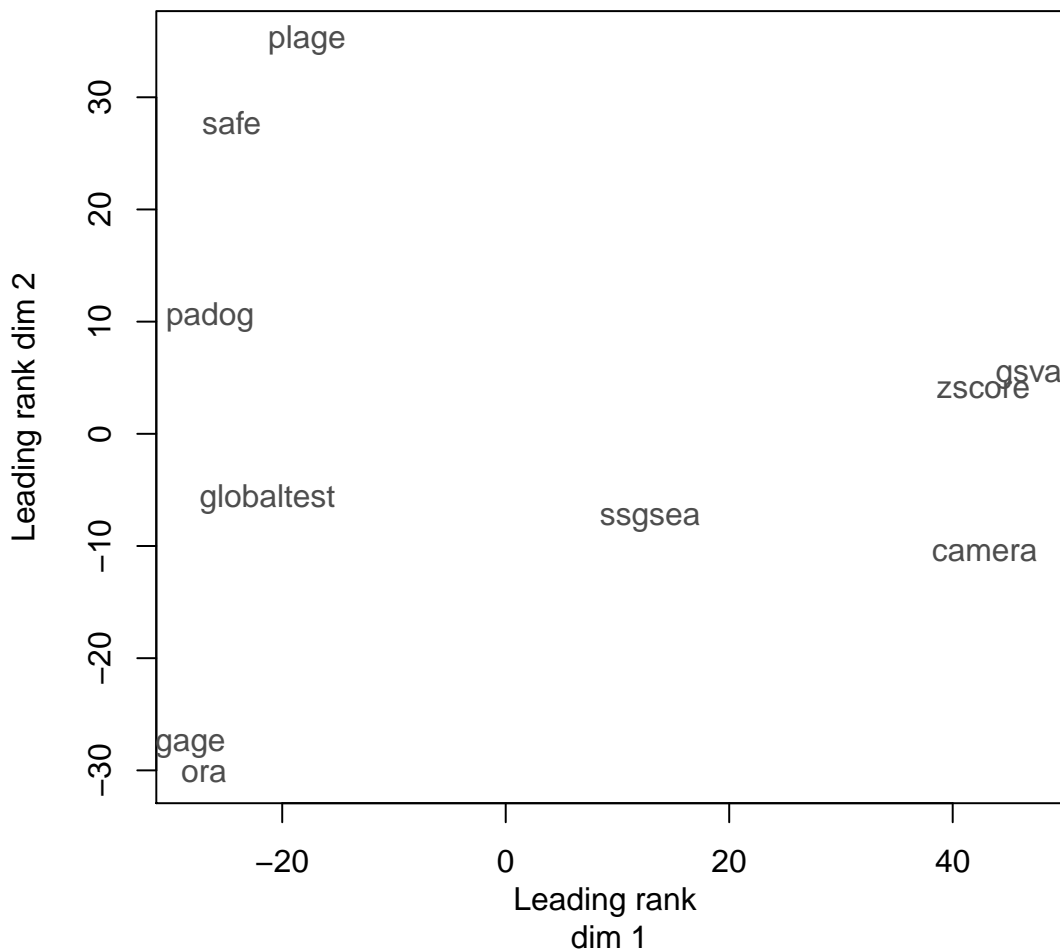


Figure 5: The performance of multiple GSE methods on the contrast X24IL13-X24.

8 EGSEA on a count matrix

The EGSEA analysis can be also performed on the count matrix directly without the need of having a `voom` object in advance. The `egsea.cnt` can be invoked on a count matrix given the group of each sample is provided with design and contrast matrices as it is illustrated in this example. This function uses the `voom` function from the [limma](#) package to convert the RNA-seq counts into expression values.

Here, the IL-13 human dataset is reanalyzed using the count matrix.

```
# load the count matrix and other relevant data
library(EGSEAdata)
data(il13.data.cnt)
cnt = il13.data.cnt$counts
```

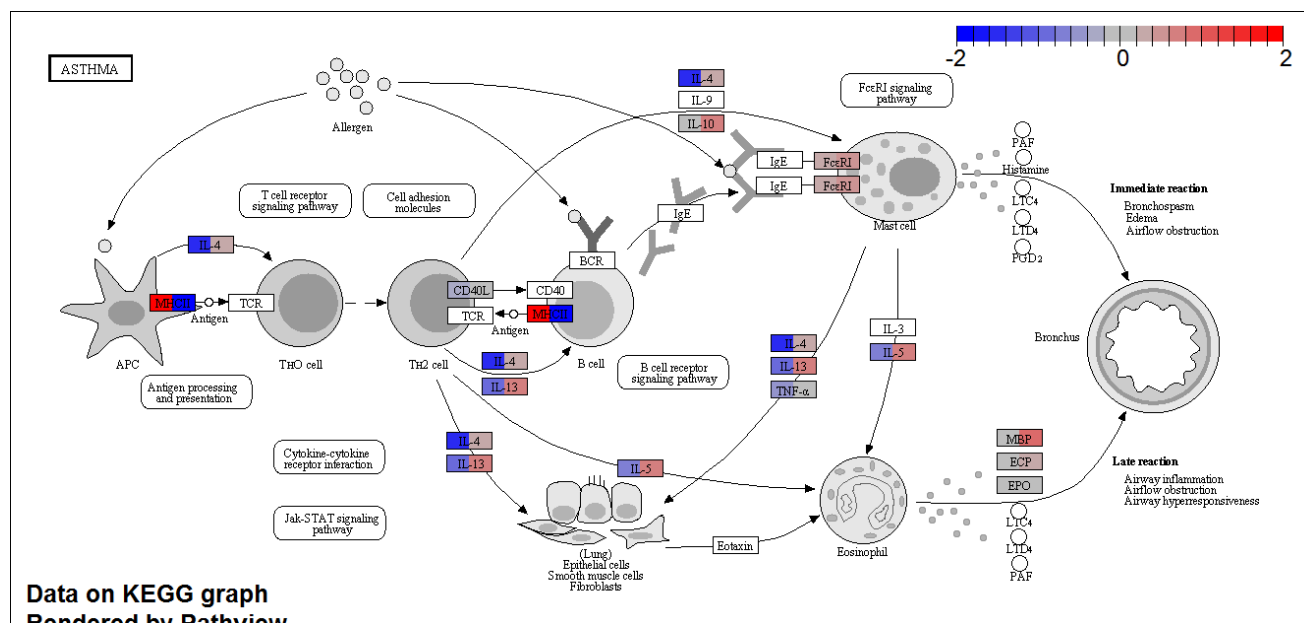


Figure 6: Asthma pathway map for the comparative analysis

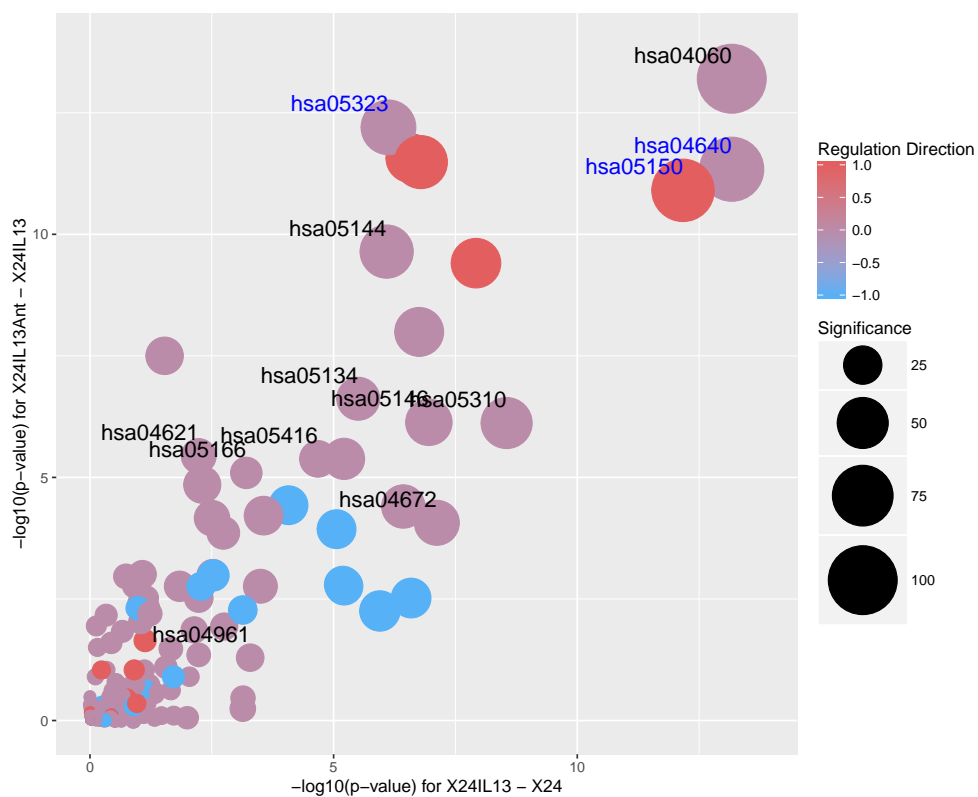


Figure 7: Summary plot for the comparative analysis

```
group = il13.data.cnt$group
group
```

```

design = il13.data.cnt$design
contrasts = il13.data.cnt$contra
genes = il13.data.cnt$genes
# build the gene set collections
gs.annots = buildIdxEZID(entrezIDs = rownames(cnt), species = "human",
  msigdb.gsets = "none", kegg.exclude = c("Metabolism"))
# perform the EGSEA analysis set report = TRUE to generate
# the EGSEA interactive report
gsa = egsea.cnt(counts = cnt, group = group, design = design,
  contrasts = contrasts, gs.annots = gs.annots, symbolsMap = genes,
  baseGSEAs = egsea.base()[-2], display.top = 5, sort.by = "avg.rank",
  egsea.dir = "./il13-egsea-cnt-report", num.threads = 4, report = FALSE)

```

9 EGSEA on a list of genes

Since performing simple over-representation analysis on large collections of gene sets is not readily available in Bioconductor, an ORA analysis was augmented to the [EGSEA](#) package so that all the reporting capabilities of EGSEA are enabled.

To perform ORA using the DE genes of the *X24/IL13-X24* contrast from the IL-13 dataset, cut-off thresholds of $p\text{-value}=0.05$ and $\log FC = 1$ are used to select a subset of DE genes. Then, the `egsea.ora` function is invoked as it is illustrated in the following example

```

# load IL-13 dataset
library(EGSEAdata)
data(il13.data)
vroom.results = il13.data$vroom
contrast = il13.data$contra
# find Differentially Expressed genes
library(limma)

##
## Attaching package: 'limma'

## The following object is masked from 'package:BiocGenerics':
##
## plotMA

vfit = lmFit(vroom.results, vroom.results$design)
vfit = contrasts.fit(vfit, contrast)
vfit = eBayes(vfit)
# select DE genes (Entrez IDs and logFC) at p-value <= 0.05
# and |logFC| >= 1
top.Table = topTable(vfit, coef = 1, number = Inf, p.value = 0.05,
  lfc = 1)
deGenes = as.character(top.Table$FeatureID)
logFC = top.Table$logFC
names(logFC) = deGenes

```

```
# build the gene set collection index
gs.annots = buildIdxEZID(entrezIDs = deGenes, species = "human",
  msigdb.gsets = "none", kegg.exclude = c("Metabolism"))

## [1] "Building KEGG pathways annotation object ... "

# perform the ORA analysis set report = TRUE to generate the
# EGSEA interactive report
gsa = egsea.ora(entrezIDs = deGenes, universe = as.character(voom.results$genes[,
  1]), logFC = logFC, title = "X24IL13-X24", gs.annots = gs.annots,
  symbolsMap = top.Table[, c(1, 2)], display.top = 5, egsea.dir = "./il13-egsea-ora-report",
  num.threads = 4, report = FALSE)

## [1] "EGSEA is running on the provided data and kegg gene sets"
## [1] "    Running ORA for X24IL13-X24"
## [1] "Running ORA on all \ncontrasts ... COMPLETED "
## [1] "Writing out the top-ranked gene sets for each contrast .. \nKEGG gene sets"
## [1] "The top gene sets for contrast X24IL13-X24 are:"
##
##                                     Type          p.adj
## Cytokine-cytokine receptor interaction Signaling 1.830767e-13
## Staphylococcus aureus infection          Disease 1.830767e-13
## Hematopoietic cell lineage              Signaling 3.714218e-10
## Phagosome                              Signaling 5.562572e-10
## Tuberculosis                           Disease 2.912950e-08
```

10 Non-standard gene set collections

Scientists usually have their own lists of gene sets and are interested in finding which sets are significant in the investigated dataset. Additional collections of gene sets can be easily added and tested using the EGSEA algorithm. The `buildCustomIdxEZID` function indexes newly created gene sets and attach gene set annotation if provided. To illustrate the use of this function, assume a list of gene sets is available where each gene set is represented by a character vector of Entrez Gene IDs. In this example, 50 gene sets were selected from the KEGG collection and then they were used to build a custom gene set collection index.

```
library(EGSEAdata)
data(il13.data)
v = il13.data$voom
# load KEGG pathways
data(kegg.pathways)
# select 50 pathways
gsets = kegg.pathways$human$kg.sets[1:50]
gsets[1]

## $`hsa00010 Glycolysis / Gluconeogenesis`
## [1] "10327" "124" "125" "126" "127" "128" "130" "130589" "131"
## [10] "160287" "1737" "1738" "2023" "2026" "2027" "217" "218" "219"
## [19] "220" "2203" "221" "222" "223" "224" "226" "229" "230"
## [28] "2538" "2597" "26330" "2645" "2821" "3098" "3099" "3101" "3939"
```

```
## [37] "3945"      "3948"      "441531"    "501"       "5105"      "5106"      "5160"      "5161"      "5162"
## [46] "5211"      "5213"      "5214"      "5223"      "5224"      "5230"      "5232"      "5236"      "5313"
## [55] "5315"      "55276"     "55902"     "57818"     "669"       "7167"      "80201"     "83440"     "84532"
## [64] "8789"      "92483"     "92579"     "9562"

# build custom gene set collection using these 50 pathways
gs.annots = buildCustomIdxEZID(entrezIDs = rownames(v$E), gsets = gsets,
  species = "human")

## [1] "Building custom pathways annotation object ... "

names(gs.annots)

## [1] "original"    "idx"         "anno"         "label"         "featureIDs" "species"
## [7] "name"

colnames(gs.annots$anno)

## [1] "ID"          "GeneSet"     "NumGenes"
```

The `buildCustomIdxEZID` creates an annotation data frame for the gene set collection if the *anno* parameter is not provided. Once the gene set collection is indexed, it can be used with any of the [EGSEA](#) functions: `egsea`, `egsea.cnt` or `egsea.ora`.

11 Add new GSE method

If you have an interesting gene set test method that you would like to add to the EGSEA framework, please contact us and we will be happy to add your method to the next release of [EGSEA](#).

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