

metaboGSE: Gene Set Enrichment Analysis via Integration of Metabolic Networks and RNA-Seq Data

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2018-11-22

1 Introduction

The *metaboGSE* package is designed for the integration of transcriptomic data and genome-scale metabolic networks (GSMN) by constructing condition-specific series metabolic sub-networks by means of RNA-seq data and providing a gene set enrichment analysis with the aid of such sub-network series (Tran et al., 2018).

2 Installation

metaboGSE depends on the *sybil* package and was evaluated with GLPK and COIN-OR Clp solvers via the *glpkAPI* and *clpAPI* packages, respectively, available in CRAN¹. The solvers and their R interface API package are not automatically installed along with *metaboGSE*. The *simplex* method from GLPK and *inibARRIER* method from Clp were investigated and yielded identical results. *glpkAPI simplex* was observed to be somehow faster. Other solvers and methods implemented for *sybil* should also work, however have not been tested. Eventually it should be mentioned that *metaboGSE* is very demanding for the solvers and may reveal sporadic computational instability, possibly due to the solver compilation and execution environment.

3 Usage

3.1 Sybil settings

```
library(metaboGSE)
SYBIL_SETTINGS("SOLVER", "glpkAPI")
SYBIL_SETTINGS("METHOD", "simplex")
SYBIL_SETTINGS("OPT_DIRECTION", "max")
```

3.2 Data preparation

3.2.1 Metabolic networks

A metabolic network can be imported from tabular or SBML inputs (see *sybil*'s manual for more details). Two *Yarrowia lipolytica* models in normoxic and hypoxic environments are provided in the *iMK735* dataset (Kavšček et al., 2015) and presented in the *MetaNetX/MNXref* namespace (Moretti et al., 2016). They are identical apart from the bounds of exchange oxygen flux.

```
data(iMK735)
iMK735[1]

## $hypoxia
## $hypoxia$model
## model name:          hypoxia
## number of compartments 8
```

¹<https://cran.r-project.org>

```
##                               Golgi apparatus
##                               cytoplasm
##                               endoplasmic reticulum
##                               extracellular region
##                               mitochondrion
##                               nucleus
##                               peroxisome
##                               vacuole
## number of reactions:         1329
## number of metabolites:      1112
## number of unique genes:     709
## objective function:          +1 R1835AC86
##
## $hypoxia$obj
## [1] 4.113629
##
## $hypoxia$comp
## model name:                  hypoxia_clean
## number of compartments      9
##                               Golgi apparatus
##                               cytoplasm
##                               endoplasmic reticulum
##                               extracellular region
##                               mitochondrion
##                               nucleus
##                               peroxisome
##                               vacuole
##                               RECO_PUSH_MNXC3
## number of reactions:         818
## number of metabolites:      605
## number of unique genes:     469
## objective function:          +1 R1835AC86
```

3.2.2 RNA-seq data

Normalized (or raw) RNA-seq counts should be provided as a matrix with gene per row and library per column. The `exprMaguire` (Maguire et al., 2014) dataset contains two matrices representing log2 voom-normalized count (`expr`) and RPKM (`pkmExpr`) per library.

```
data(exprMaguire)
names(exprMaguire)
```

```
## [1] "expr"      "pkmExpr"
```

```
str(exprMaguire$expr, vec.len=3)
```

```
## num [1:469, 1:22] 0.848 7.836 8.244 8.62 ...
## - attr(*, "dimnames")=List of 2
## ..$ : chr [1:469] "euk:Q6C902_YARLI" "euk:Q6CDH6_YARLI" "euk:Q6CED5_YARLI" ...
## ..$ : chr [1:22] "DN2" "DN3" "SH1" ...
```

3.2.3 Gene set annotation

Here we apply `metaboGSE` for Gene Ontology enrichment analysis. A mapping between genes and GO terms should be provided. The `yarli2GO` dataset contains such a mapping in a `list` format. The pre-built

`topGOdata` object `yarliGOdata` is also provided.

```
data(yarli2GO)
length(yarli2GO)
```

```
## [1] 4747
```

```
str(head(yarli2GO))
```

```
## List of 6
## $ euk:3HAO_YARLI : chr [1:7] "GO:0000334" "GO:0005737" "GO:0006569" "GO:0008198" ...
## $ euk:ACEA_YARLI : chr [1:6] "GO:0004451" "GO:0006097" "GO:0006099" "GO:0009514" ...
## $ euk:ACEB_YARLI : chr [1:4] "GO:0004451" "GO:0005759" "GO:0019629" "GO:0046421" ...
## $ euk:ACH1_YARLI : chr [1:6] "GO:0003986" "GO:0005739" "GO:0005829" "GO:0006083" ...
## $ euk:ACOX1_YARLI: chr [1:5] "GO:0003995" "GO:0003997" "GO:0005777" "GO:0033540" ...
## $ euk:ACOX2_YARLI: chr [1:5] "GO:0003995" "GO:0003997" "GO:0005777" "GO:0033540" ...
```

3.2.4 Pre-built data

The tutorial below shows a simple example to be executed in a reasonable computing time. The datasets of pre-built series of metabolic sub-networks and gene set enrichment (Tran et al., 2018) are also provided.

```
data(yarliSubmnets)
names(yarliSubmnets)
```

```
## [1] "DN" "SH" "SN" "UH" "UN" "WH" "WN"
```

```
data(yarliGSE)
length(yarliGSE)
```

```
## [1] 135
```

3.3 Analysis

3.3.1 Model rescuing

Both models of iMK735 grow, nevertheless we should still apply the rescue procedure to produce the rescue reactions for all metabolites in growth and maintenance (if any) reactions (see (Tran et al., 2018)). For instance, we here perform the rescue process while targeting a growth of 20% of the initial objective value for the hypoxic model.

```
target.ratio <- 0.2
hmodel <- iMK735$hypoxia$model
hobj <- iMK735$hypoxia$obj
hmodel.rescue <- rescue(model = hmodel,
                        target = c(target.ratio*hobj),
                        react = c(which(obj_coef(hmodel) == 1)))
```

```
## SYBIL_SETTINGS(TOLERANCE) has been set to 1e-12
## SYBIL_SETTINGS(OPT_DIRECTION) has been set to min
```

In the `hmodel.rescue` object, the `rescued` field represents the rescued model \mathcal{M}'' , which is the same as the initial model \mathcal{M} since the latter grows, as presented in Fig 1. The `rescue` field presents the rescue model \mathcal{M}' , which contains rescue reactions on every metabolite in growth and maintenance reactions, and the `coef` field indicates the coefficients of those rescue reactions used for optimization in the rescue process. RECO_PUSH_MNXC3 denotes the compartment of artificial metabolites, e.g. X' . The rescue procedure can also be performed on maintenance reactions, which are restricted to non-empty fixed fluxes.

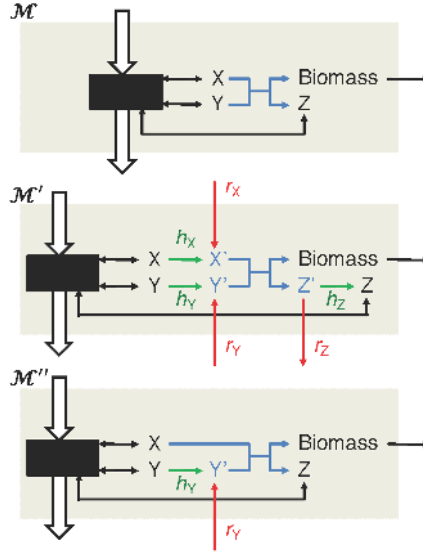


Figure 1: Schema of GSMN rescue process. \mathcal{M} , original GSMN with growth reaction $X + Y \rightarrow Z + \text{Biomass}$. \mathcal{M}' , expanded GSMN with the full set of rescue (r_x) and help (h_x) reactions for every metabolite x in the biomass reaction. \mathcal{M}'' , example of a minimal rescued GSMN in the particular case where only metabolite Y needs to be rescued. (Tran et al., 2018)

```
hmodel.rescue$rescue
```

```
## model name:          hypoxia
## number of compartments 9
##                      Golgi apparatus
##                      cytoplasm
##                      endoplasmic reticulum
##                      extracellular region
##                      mitochondrion
##                      nucleus
##                      peroxisome
##                      vacuole
##                      RECO_PUSH_MNXC3
## number of reactions:  1413
## number of metabolites: 1154
## number of unique genes: 709
## objective function:   +1 R1835AC86
```

```
head(hmodel.rescue$coef)
```

```
##                      R1835AC86
## RECO_PUSH_MNXC32_MNXC3  3.47947112
## RECO_PUSH_MNXC3_MNXC3  0.06531679
## RECO_PUSH_MNXC89557_MNXC3  5.28960592
## RECO_PUSH_MNXC2_MNXC3  0.06531679
## RECO_PUSH_MNXC93970_MNXC3 202.83975659
## RECO_PUSH_MNXC31_MNXC3  35.71428571
```

3.3.2 Model cleaning

We set the TOLERANCE to 1e-8, which indicates that values less than 1e-8 are considered as 0, to deal with numerical imprecision.

```
SYBIL_SETTINGS("TOLERANCE", 1e-08)
```

The blocked reactions from the rescue models are determined by a flux variability analysis via the `fluxVar` function from *sybil*. Those reactions as well as related genes and metabolites are then removed from the models. `mc.cores` can be set appropriately to perform parallel computation of `fluxVar`. A high `mc.cores` is recommended as `fluxVar` is time consuming.

```
## not run
mc.cores <- 10
fva <- multiDel(model=hmodel.rescue$rescue,
               nProc=mc.cores,
               todo="fluxVar",
               fixObjVal=F,
               del1=react_id(hmodel.rescue$rescue))
reacs.blo <- names(which(setNames(unlist(lapply(fva, blReact)),
                                   react_id(hmodel.rescue$rescue))))
hmodel.clean <- rmReact(hmodel.rescue$rescue, reacs.blo, rm_met=T)
##
```

`hmodel.clean`, considered as the comprehensive model for hypoxic conditions, can be loaded from the `iMK735` dataset. It is noted that the TOLERANCE set in `SYBIL_SETTINGS("TOLERANCE")` determined the threshold for a reaction to be blocked. It should be adequately selected so that multiple (if any) input models, for example those in hypoxic and normoxic environment, contain the same gene set after cleaning by removing blocked reaction and propagating.

```
hmodel.clean <- iMK735$hypoxia$comp
hmodel.clean
```

```
## model name:          hypoxia_clean
## number of compartments 9
##                      Golgi apparatus
##                      cytoplasm
##                      endoplasmic reticulum
##                      extracellular region
##                      mitochondrion
##                      nucleus
##                      peroxisome
##                      vacuole
##                      RECO_PUSH_MNXC3
## number of reactions:  818
## number of metabolites: 605
## number of unique genes: 469
## objective function:    +1 R1835AC86
```

Now we convert the growth objective of the comprehensive model to a weighted objective function on rescue reactions with the determined coefficients. Hereafter, the goal is to minimize this function.

```
SYBIL_SETTINGS("OPT_DIRECTION", "min")
hmodel.weight <- changeObjFunc(hmodel.clean, react=rownames(hmodel.rescue$coef),
                              obj_coef=hmodel.rescue$coef)
hmodel.weight
```

```
## model name:          hypoxia_clean
## number of compartments 9
##                      Golgi apparatus
```

```
## cytoplasm
## endoplasmic reticulum
## extracellular region
## mitochondrion
## nucleus
## peroxisome
## vacuole
## RECO_PUSH_MNXC3
## number of reactions: 818
## number of metabolites: 605
## number of unique genes: 469
## objective function: +0.0653167864141084 RECO_PUSH_MNXC3 +0.0653167864141084 RECO_PUSH_MNXC3
```

```
optimizeProb(hmodel.weight)
```

```
## solver: glpkAPI
## method: simplex
## algorithm: fba
## number of variables: 818
## number of constraints: 605
## return value of solver: solution process was successful
## solution status: solution is optimal
## value of objective function (fba): 0.000000
## value of objective function (model): 0.000000
```

The obtained objective of 0 above indicates that there is no need to rescue the `hmodel.clean` since it grows.

3.3.4 Weighting scheme

We now compute weights for rescue reactions to account for the importance and dependency of metabolites to rescue (see Weighting scheme for model fitness in Tran et al. (2018))

```
mc.cores <- 1
rescue.weight <- (weightReacts(hmodel.weight, mc.cores=mc.cores, gene.num=1))$weight
str(rescue.weight, vec.len=2)
```

```
## Named num [1:45] 0.0229 0.2028 ...
## - attr(*, "names")= chr [1:45] "RECO_PUSH_MNXC3" "RECO_PUSH_MNXC1" ...
```

3.3.4 GO annotation

We compute the set of preliminary GO terms in *biological process* category using `topGO` with *fisher* statistic and *weight01* algorithm. The whole GO annotation and gene universe are used. The aim of the following R script is to preliminarily filter the set of GO terms of interest. The resulting 135 GO terms are filtered by p -value < 0.1 and contain at least 3 genes and at most 50 genes from the model.

```
G02geneID <- inverseList(yarli2GO)
length(G02geneID)
```

```
## [1] 4629
```

```
str(head(G02geneID), vec.len=3)
```

```
## List of 6
## $ G0:0000001: chr [1:13] "euk:GEM1_YARLI" "euk:MDM12_YARLI" "euk:Q6C1G8_YARLI" ...
## $ G0:0000002: chr [1:27] "euk:B5FVH9_YARLI" "euk:CCM1_YARLI" "euk:DML1_YARLI" ...
## $ G0:0000007: chr "euk:Q6C0U6_YARLI"
```

```
## $ GO:0000009: chr [1:7] "euk:F2Z612_YARLI" "euk:F2Z6C7_YARLI" "euk:Q6C1C8_YARLI" ...
## $ GO:0000010: chr "euk:CQ1_YARLI"
## $ GO:0000011: chr [1:9] "euk:ACT_YARLI" "euk:Q6C134_YARLI" "euk:Q6C1E5_YARLI" ...
```

```
gene.name <- names(yarli2GO)
gene.list <- factor(as.integer(gene.name %in% sybil::allGenes(hmodel.clean)))
names(gene.list) <- gene.name
GOdata <- new("topGOdata",
              ontology = "BP",
              nodeSize = 5,
              allGenes = gene.list,
              annot     = annFUN.gene2GO,
              gene2GO   = yarli2GO
            )
result <- runTest(GOdata, statistic="fisher", algorithm="weight01")
table <- GenTable(GOdata,
                  weight = result,
                  orderBy = "weight",
                  numChar = 10000,
                  topNodes = result@geneData[4]
                )
table$weight <- as.numeric(sub("<", "", table$weight))
table <- table[!is.na(table$weight), ]
MINSIG <- 3
MAXSIG <- 50
WCUTOFF <- 0.1
GO.interest <- table[table$Significant >= MINSIG & table$Significant <= MAXSIG &
                     table$weight < WCUTOFF, ]$GO.ID
GO2geneID.interest.proteome <- genesInTerm(GOdata, GO.interest)
GO2geneID.interest <- lapply(GO2geneID.interest.proteome, function(git) {
  intersect(sybil::allGenes(hmodel.clean), git)
})
length(GO.interest)
```

```
## [1] 136
```

```
str(head(GO2geneID.interest), vec.len=3)
```

```
## List of 6
## $ GO:0046323: chr [1:23] "euk:F2Z653_YARLI" "euk:Q6C0EO_YARLI" "euk:Q6C0K5_YARLI" ...
## $ GO:0015986: chr [1:17] "euk:ATP6_YARLI" "euk:ATP8_YARLI" "euk:ATP9_YARLI" ...
## $ GO:0015991: chr [1:15] "euk:ATP9_YARLI" "euk:Q6CHD8_YARLI" "euk:Q6CH91_YARLI" ...
## $ GO:0006696: chr [1:19] "euk:Q6C8C2_YARLI" "euk:ERG6_YARLI" "euk:Q6C231_YARLI" ...
## $ GO:0006099: chr [1:16] "euk:PRPC_YARLI" "euk:Q6C450_YARLI" "euk:Q6C823_YARLI" ...
## $ GO:0006096: chr [1:12] "euk:KPYK_YARLI" "euk:PFKA_YARLI" "euk:G3P_YARLI" ...
```

GO.interest contains other GO terms than those in GO2geneID, as topGO allows propagating in the gene ontology.

3.3.5 Expression-based gene removal

`step` indicates the difference of gene numbers to remove between consecutive sub-model constructions, then determines numbers of genes to remove in the simulation. Here we set `step = 50` and `draw.num = 4` to reduce the computing time in this tutorial, i.e. the 0, 50, 100, etc. first genes in certain ranking will be successively removed from the comprehensive model, and 4 random removals will be performed. The series of

metabolic sub-networks is constructed for the hypoxic *upc2Δ* (UH) condition with various gene rankings as below.

```
cond <- "UH"
step <- 50
draw.num <- 4
reps.i <- grep(cond, colnames(exprMaguire$expr), value=T)
ranks <- mclapply(reps.i, mc.cores=mc.cores, function(ri) {
  data.frame(
    # ranks1. voom-normalized expression
    expr = exprMaguire$expr[, ri, drop=T],
    # ranks2. pkm normalized expression
    pkmExpr = exprMaguire$pkmExpr[, ri, drop=T],
    # ranks3. relative expression power 1
    relExpr1 = relativeExpr(exprMaguire$expr, power=1)[, ri, drop=T],
    # ranks4. relative expression power 2
    relExpr2 = relativeExpr(exprMaguire$expr, power=2)[, ri, drop=T],
    # ranks5. relative expression power 3
    relExpr3 = relativeExpr(exprMaguire$expr, power=3)[, ri, drop=T],
    # ranks6. reverse expression (the worst)
    revExpr = 1/(1 + exprMaguire$expr[, ri, drop=T]),
    # ranks7. z-score expression
    zExpr = zscoreExpr(exprMaguire$expr)[, ri, drop=T]
  )
})
names(ranks) <- reps.i
fitnessUH <- fitness(model      = hmodel.weight,
                    ranks      = ranks,
                    rescue.weight = rescue.weight,
                    step       = step,
                    draw.num    = draw.num,
                    mc.cores    = mc.cores)
submnetsUH <- submnet(model      = hmodel.weight,
                    fn          = fitnessUH,
                    rank.best   = "expr",
                    gene.sets   = G02geneID.interest,
                    mc.cores    = mc.cores)
```

It may happen that several genes have identical values for their expression in a sample, which triggers a random ranking among these genes, and thus introduces an unexpected difference among samples. To overcome such a randomness, a negligible value could be added to the expression of each gene in each sample using the overall expression in all the samples.

```
## not run
pseudo.rank <- base::rank(rowSums(exprMaguire$expr),
                        ties.method='first')/nrow(exprMaguire$expr)*1e-6
exprMaguire$expr <- exprMaguire$expr + pseudo.rank
##
```

ranks can be set to the only expression you want to use, e.g. **expr**, and **draw.num** set to 0 to skip the ranking evaluation step. It is observed that the **optimizeProb** function from *sybil* may stay in an infinite loop with some system configuration. A timeout limit should be given in the **fitness** function of *metaboGSE*. It helps to stop the occasional infinite loop in **optimizeProb**. Such a limit is set by default to 12 seconds, which is sufficient for the model iMK735 from *Y. lipolytica* and iMM1415 from mouse. A higher limit may be required for a larger model.


```
submnetsUH$condition
```

```
## [1] "UH"
```

```
knitr::kable(submnetsUH$gene.del)
```

	0	50	100	150	200	250	300	350	400	450
gene.del	0	50	100	150	200	250	300	350	400	450

```
knitr::kable(submnetsUH$fitness.random, digits=3)
```

		0	50	100	150	200	250	300	350	400	450
F.random.1	1	0.614	0.600	0.439	0.429	0.421	0.435	0.419	0.423	0.418	
F.random.2	1	0.552	0.641	0.444	0.443	0.444	0.419	0.437	0.418	0.425	
F.random.3	1	0.683	0.502	0.457	0.436	0.420	0.422	0.419	0.417	0.417	
F.random.4	1	0.509	0.584	0.461	0.426	0.434	0.439	0.418	0.425	0.417	

```
knitr::kable(submnetsUH$fitness.ranks$UH1, digits=3)
```

		0	50	100	150	200	250	300	350	400	450
expr	1	0.958	0.653	0.653	0.575	0.483	0.436	0.418	0.417	0.417	
pkmExpr	1	0.966	0.653	0.652	0.556	0.502	0.418	0.418	0.418	0.417	
relExpr1	1	0.729	0.675	0.637	0.492	0.483	0.436	0.436	0.436	0.418	
relExpr2	1	0.695	0.653	0.577	0.561	0.443	0.443	0.443	0.418	0.417	
relExpr3	1	0.695	0.653	0.577	0.561	0.443	0.438	0.418	0.417	0.417	
revExpr	1	0.509	0.466	0.436	0.427	0.427	0.424	0.423	0.422	0.421	
zExpr	1	0.737	0.652	0.596	0.491	0.469	0.436	0.436	0.418	0.418	

The `yarliSubmnets` dataset contains the series of sub-networks built with `step = 1` and `draw.num = 50`, indicating the gene-by-gene removal.

```
data(yarliSubmnets)
```

```
str(yarliSubmnets$UH$gene.del)
```

```
## num [1, 1:470] 0 1 2 3 4 5 6 7 8 9 ...
## - attr(*, "dimnames")=List of 2
## ..$ : chr "gene.del"
## ..$ : chr [1:470] "0" "1" "2" "3" ...
```

```
dim(yarliSubmnets$UH$fitness.random)
```

```
## [1] 50 470
```

```
str(yarliSubmnets$UH$fitness.ranks)
```

```
## List of 3
## $ UH1: num [1:7, 1:470] 1 1 1 1 1 1 1 1 1 ...
## ..- attr(*, "dimnames")=List of 2
## .. ..$ : chr [1:7] "expr" "pkmExpr" "relExpr1" "relExpr2" ...
## .. ..$ : chr [1:470] "0" "1" "2" "3" ...
## $ UH2: num [1:7, 1:470] 1 1 1 1 1 1 1 1 1 ...
## ..- attr(*, "dimnames")=List of 2
```

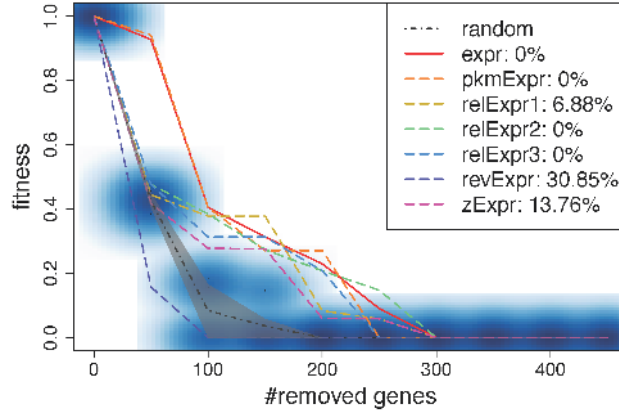


Figure 2: submnetsUH with step = 50 and draw.num = 4

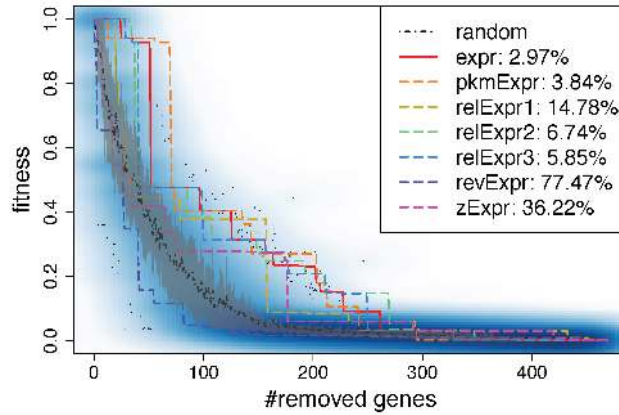


Figure 3: yarliSubmnets with step = 1 and draw.num = 50

```
## .. .$ : chr [1:7] "expr" "pkmExpr" "relExpr1" "relExpr2" ...
## .. .$ : chr [1:470] "0" "1" "2" "3" ...
## $ UH3: num [1:7, 1:470] 1 1 1 1 1 1 1 1 1 1 ...
## ..- attr(*, "dimnames")=List of 2
## .. .$ : chr [1:7] "expr" "pkmExpr" "relExpr1" "relExpr2" ...
## .. .$ : chr [1:470] "0" "1" "2" "3" ...
```

The sub-network construction can be visualized via the `simulateSubmnet` function, which produces a plot for each condition. Figures 2 and 3 show the fitness of submodels obtained by removing genes following different rankings for the hypoxic *upc2Δ* condition.

```
simulateSubmnet(sgd      = submnetsUH,
                mc.cores = mc.cores)
```

3.3.6 GO term enrichment

We evaluate the significance of given gene sets with the `metaboGSE` function and randomization tests. `nrand = 1000` is used in the test for the significance of the gene sets against random sets in each individual condition. `nperm = 1000` is used in the test for the significance of difference between conditions.

```
## not run
GSE <- metaboGSE(yarliSubmnets, method="perm", nperm=1000, nrand=1000,
```

```
mc.cores=mc.cores, prefix="/tmp/summary")
##
```

This step is time consuming. The yarliGSE dataset can be loaded instead. Note that yarliG0data should be also loaded, since the constructed G0data may not match the pre-built G0data due to G0.db and topG0 version difference.

```
data(yarliGSE)
data(yarliG0data)
GSE <- yarliGSE
G0data <- yarliG0data
str(GSE[["G0:0006696"]], vec.len=2)
```

```
## List of 2
## $ res :List of 8
## ..$ GS.ID      : chr "G0:0006696"
## ..$ Description: chr "ergosterol biosynthetic process"
## ..$ statistic  : num 0.184
## ..$ p.Cond     : Named num [1:7] 0.000104 0.002104 ...
## ..- attr(*, "names")= chr [1:7] "DN" "SH" ...
## ..$ p.Val      : num 0.014
## ..$ auc        : Named num [1:22] 0.287 0.267 ...
## ..- attr(*, "names")= chr [1:22] "DN2" "DN3" ...
## ..$ pw.posthoc : 'data.frame': 21 obs. of 4 variables:
## .. ..$ cond1    : chr [1:21] "DN" "DN" ...
## .. ..$ cond2    : chr [1:21] "SH" "SN" ...
## .. ..$ statistic: chr [1:21] "0.17" "0.065" ...
## .. ..$ p.Val    : chr [1:21] "0.174825174825175" "0.0689310689310689" ...
## ..$ contrast   : Named chr [1:3] "((SH,WH),((UH,WN),(DN,(SN,UN))))" "DN,SN,UH,UN,WN" ...
## ..- attr(*, "names")= chr [1:3] "newick" "group1" ...
## $ plot:List of 3
## ..$ gs.fracs      : num [1:470, 1:22] 1 1 1 1 1 ...
## ..- attr(*, "dimnames")=List of 2
## .. ..$ : NULL
## .. ..$ : chr [1:22] "DN2" "DN3" ...
## ..$ xout          : num [1:44] 0 0.334 ...
## ..$ gs.fracs.itp.mean: num [1:44, 1:7] 0 0 0 0 0 ...
## ..- attr(*, "dimnames")=List of 2
## .. ..$ : NULL
## .. ..$ : chr [1:7] "DN" "SH" ...
```

```
GSE[["G0:0006696"]]$res$p.Val
```

```
## [1] 0.01398601
```

```
GS.sig.all <- as.data.frame(t(sapply(GSE, function(gsm) {
  c(GS.ID=gsm$res$GS.ID,
    Description=gsm$res$Description,
    Statistic=gsm$res$Statistic,
    p.Cond=if (is.null(gsm$res$p.Cond)) NA else min(gsm$res$p.Cond),
    p.Val=gsm$res$p.Val,
    Genes=paste(intersect(genesInTerm(G0data, gsm$res$GS.ID)[[1]],
                          sybil::allGenes(hmodel.clean)), collapse=","))
})), stringsAsFactors=F)
GS.sig.all$FDR <- p.adjust(as.numeric(GS.sig.all$p.Val), method="BH")
GS.sig.all <- GS.sig.all[!is.na(GS.sig.all$FDR), ]
```

```
dim(GS.sig.all)
```

```
## [1] 135 7
```

```
GS.sig <- GS.sig.all[as.numeric(GS.sig.all$FDR) < 0.05, , drop=F]
head(GS.sig[order(as.numeric(GS.sig$Statistic), decreasing=T), c(1:5,7)])
```

##	GS.ID	Description
##	G0:0019878 G0:0019878	lysine biosynthetic process via aminoadipic acid
##	G0:0009098 G0:0009098	leucine biosynthetic process
##	G0:0000162 G0:0000162	tryptophan biosynthetic process
##	G0:0006772 G0:0006772	thiamine metabolic process
##	G0:0009085 G0:0009085	lysine biosynthetic process
##	G0:0009082 G0:0009082	branched-chain amino acid biosynthetic process
##	Statistic	p.Val
##	G0:0019878 0.641290467573758	2.08328993145976e-05
##	G0:0009098 0.585836597550887	0.000124997395887586
##	G0:0000162 0.570029210757622	6.24986979437928e-05
##	G0:0006772 0.563256553843708	6.24986979437928e-05
##	G0:0009085 0.493367522482941	0.000229161892460574
##	G0:0009082 0.468685229785908	0.000312493489718964
##	FDR	
##	G0:0019878 0.02334204	
##	G0:0009098 0.01348651	
##	G0:0000162 0.01348651	
##	G0:0006772 0.01556136	
##	G0:0009085 0.02052296	
##	G0:0009082 0.01348651	

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