# Brief Introduction to R package gact

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## Contents

The practical is based on the R package gact (Rohde et al.2022)). This package provides an infrastructure for working with large-scale genomic association data linked to different types of genomic features.

The most recent version of gact can be obtained from github:

```
library(devtools)
devtools::install_github("psoerensen/gact")
```

## Load packages used

```
library(gact)
library(qgg)

library(org.Hs.eg.db)
library(reactome.db)

library(corrplot)
```

### Download and install GDT database

The function gact() dowload and install the GDT database:

```
# Set working for database
setwd("C://Users//au223366//Dropbox//Projects//balder")

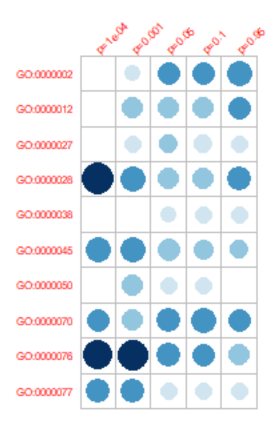
# Download data bases from repository
GACTdb <- gact(version = "t2dm-gact-0.0.1", task = "download")</pre>
```

#### Extract data from GDT database

The function getStat() extract data from the database:

```
# Extract data from T2D for genomic feature Genes
stat <- getStat(GACTdb = GACTdb, trait = "t2d", feature = "Genes")
head(stat)</pre>
```

```
##
                   Ensembl Gene ID
                                     Symbol m
                                                    stat
## ENSG00000121410 ENSG00000121410
                                      A1BG 58 0.00000 1.0000
## ENSG00000175899 ENSG00000175899
                                       A2M 153 20.78100 0.1477
## ENSG00000256069 ENSG00000256069
                                     A2MP1 64 0.00000 1.0000
## ENSG00000171428 ENSG00000171428
                                       NAT1 135 0.00000 1.0000
## ENSG00000156006 ENSG00000156006
                                       NAT2 102 20.90052 0.0951
## ENSG00000196136 ENSG00000196136 SERPINA3 64 0.00000 1.0000
# Extract data from T2D for genomic feature Gene Ontology (GD) where
# output format is a data frame
stat <- getStat(GACTdb = GACTdb, trait = "t2d", feature = "GO")</pre>
head(stat)
##
                   GO ID
                           m
                                   stat
## GD:0000002 GD:0000002 2165 143.36901 0.3208
## GD:0000012 GD:0000012 1885 107.38110 0.3577
## GD:0000027 GD:0000027 1867 72.77531 0.5031
## GD:0000028 GD:0000028 861 71.43454 0.2242
## GD:0000038 GD:0000038 2397 63.58818 0.6599
## GD:0000045 GD:0000045 8223 749.36205 0.2368
# Extract data from T2D for genomic feature Gene Ontology (GO) where
# output format is list
stat <- getStat(GACTdb = GACTdb, trait = "t2d", feature = "GO", format = "list",</pre>
   cls = c("p=1e.04", "p=0.001", "p=0.05", "p=0.1", "p=0.95"))
str(stat)
## List of 3
## $ m : Named int [1:4547] 2165 1885 1867 861 2397 8223 1050 3519 1077 5192 ...
   ..- attr(*, "names")= chr [1:4547] "G0:0000002" "G0:0000012" "G0:0000027" "G0:0000028" ...
## $ stat: num [1:4547, 1:5] 0 0 0 17 0 ...
   ..- attr(*, "dimnames")=List of 2
##
    ....$ : chr [1:4547] "GD:0000002" "GD:0000012" "GD:0000027" "GD:0000028" ...
##
    ....$ : chr [1:5] "p=1e.04" "p=0.001" "p=0.05" "p=0.1" ...
## $ p : num [1:4547, 1:5] 1 1 1 0.0781 1 ...
    ..- attr(*, "dimnames")=List of 2
##
##
    ....$ : chr [1:4547] "GD:0000002" "GD:0000012" "GD:0000027" "GD:0000028" ...
     ....$ : chr [1:5] "p=1e.04" "p=0.001" "p=0.05" "p=0.1" ...
# Plot results for genomic feature Gene Ontology (GO)
colbar <- colorRampPalette(c("#FFFFFF", "#D1E5FO", "#92C5DE", "#4393C3",</pre>
    "#2166AC", "#053061"))
corrplot(-log10(stat$p[1:10, ]), is.corr = FALSE, tl.cex = 0.7, tl.srt = 45,
   col = colbar(6), cl.pos = "n", mar = c(1, 1, 1, 1))
```



### Extract and write data from GDT database

The function writeStat() extract and write data from the database:

```
writeStat(GACTdb = GACTdb, feature = "GO", trait = "t2d", file.csv = "go_t2dm_gcta.csv")
writeStat(GACTdb = GACTdb, feature = "Pathways", trait = "t2d", file.csv = "pathways_t2dm_gcta.csv")
writeStat(GACTdb = GACTdb, feature = "ProteinComplexes", trait = "t2d",
    file.csv = "proteincomplexes_t2dm_gcta.csv")
writeStat(GACTdb = GACTdb, feature = "ChemicalComplexes", trait = "t2d",
    file.csv = "chemicalcomplexes_t2dm_gcta.csv")
writeStat(GACTdb = GACTdb, feature = "Genes", trait = "t2d", file.csv = "genes_t2dm_gcta.csv")
```