P vs NP, differential measurement $_{HBG}$ $_{11/5/2018}$

This script use the differential measure (z-score) between tumor/non-tumor cells

```
### this block perform z-score calculations
library("microbenchmark")
library("matrixStats")
## the GO terms for biological processes (BP)
GOcategory.file <- read.csv("../Data/human.bp.term.csv",header=TRUE, stringsAsFactors=F)
bp.go.cat <- GOcategory.file$GO.term</pre>
bp.dim <- length(bp.go.cat)</pre>
conn.dim <- 20
hspin <- matrix(as.numeric(unlist(read.table("human.rnf43.PvNP.bp.txt", header=F, sep=","))), nrow=bp.d
obs <- c(hspin)
perm <- c()
for (i in 1:100) {
    name <- paste("ms02.human", "/", "rnf43.PvNP", "/", "ms02.", i, ".bp.matrix.csv", sep="")</pre>
    mat <- matrix(as.numeric(unlist(read.table(name, header=F, sep=","))), nrow=bp.dim, ncol=conn.dim)</pre>
    perm <- rbind(perm, c(mat))</pre>
mean <- colMeans(perm)</pre>
std <- colSds(perm)</pre>
zscore <- round((obs - mean)/std, 3)</pre>
z.pvnp <- matrix(zscore, nrow=bp.dim, ncol=conn.dim)</pre>
\#write.table(z, file="human.rnf43.PvNP.bp.z.csv", sep=",", row.names=F, col.names=F, quote=F)
library('gplots')
## Attaching package: 'gplots'
## The following object is masked from 'package:stats':
##
##
       lowess
z.pvnp <- t(z.pvnp)
rownames(z.pvnp) <- c("5%","10%","15%","20%","25%","30%","35%","40%","45%","50%",
                  "55%", "60%", "65%", "70%", "75%", "80%", "85%", "90%", "95%", "100%")
```

```
colnames(z.pvnp) <- bp.go.cat</pre>
colors = c(seq(min(z.pvnp), -10.1, length=100), seq(-9.9, 9.9, length=100), seq(10.1, max(z.pvnp), length=100))
my palette <- colorRampPalette(c("blue2", "white", "red2"))(n = 299)
png(filename = "17.PvsNP.z.hm.png", width=6, height=5.5, res=1200, unit="in")
heatmap.2(z.pvnp, col=my_palette, breaks=colors, dendrogram='none', Rowv=F, Colv=F,
          trace='none', offsetRow = 0, offsetCol = 0,
          xlab="Biological Process Terms", ylab="PDAC vs Normal Cells, Quantiles",
          margins = c(2,3.5), key.title = "Color Bar", key.xlab="Z-score", key.ylab=NA,
          revC = T,
          labCol = NA, #labRow =,
          #srtCol=45, adjCol=c(1,0),
          \#lmat=rbind(c(0,3,4), c(2,1,0)), lwid=c(1.5,4,2),
          scale="none", symbreaks=T, symm=F, symkey = F)
dev.off()
## pdf
##
#correlations to quantile 20 (95%-100%) genes
y.val <- c()
for (i in 1:20) {
 y.val2 \leftarrow c()
 for (j in 1:20) {
     cor.val <- cor.test(z.pvnp[j,], z.pvnp[i,])</pre>
     y.val2 <- rbind(y.val2, cor.val$estimate)</pre>
  }
 y.val <- cbind(y.val, y.val2)</pre>
rownames(y.val) <- c("5%","10%","15%","20%","25%","30%","35%","40%","45%","50%",
                    "55%", "60%", "65%", "70%", "75%", "80%", "85%", "90%", "95%", "100%")
colnames(y.val) <- c("5%","10%","15%","20%","25%","30%","35%","40%","45%","50%",
                    "55%", "60%", "65%", "70%", "75%", "80%", "85%", "90%", "95%", "100%")
my_palette2 <- colorRampPalette(c("blue2", "white", "red2"))(9)</pre>
png(filename = "17.PvsNP.z.hm.cor.matrix.png", width=6, height=5.5, res=1200, unit="in")
heatmap.2(y.val, col=my_palette2, dendrogram='none', Rowv=F, Colv=F,
          trace='none', offsetRow = 0, offsetCol = 0,
          xlab="Quantiles of Bayes Factors", ylab="Quantiles of Bayes Factors",
          margins = c(3.5,3.5), key.title = "Color Bar", key.xlab="Pearson Correlation", key.ylab=NA,
          revC = T,
          #labCol = NA, #labRow =,
          #srtCol=45, adjCol=c(1,0),
          \#lmat=rbind(c(0,3,4), c(2,1,0)), lwid=c(1.5,4,2),
          scale="none", symbreaks=T, symm=F, symkey = F)
dev.off()
## pdf
##
   2
library(GO.db)
```

Loading required package: AnnotationDbi

```
## Loading required package: stats4
## 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, sd, var, xtabs
## The following objects are masked from 'package:base':
##
##
       anyDuplicated, append, as.data.frame, basename, cbind,
##
       colMeans, colnames, colSums, dirname, do.call, duplicated,
##
       eval, evalq, Filter, Find, get, grep, grepl, intersect,
##
       is.unsorted, lapply, lengths, Map, mapply, match, mget, order,
       paste, pmax, pmax.int, pmin, pmin.int, Position, rank, rbind,
##
       Reduce, rowMeans, rownames, rowSums, sapply, setdiff, sort,
##
       table, tapply, union, unique, unsplit, which, which.max,
##
       which.min
## Loading required package: Biobase
## Welcome to Bioconductor
##
##
       Vignettes contain introductory material; view with
##
       'browseVignettes()'. To cite Bioconductor, see
       'citation("Biobase")', and for packages 'citation("pkgname")'.
##
## Attaching package: 'Biobase'
## The following objects are masked from 'package:matrixStats':
##
##
       anyMissing, rowMedians
## Loading required package: IRanges
## Loading required package: S4Vectors
## Attaching package: 'S4Vectors'
## The following object is masked from 'package:gplots':
##
##
       space
## The following object is masked from 'package:base':
##
##
       expand.grid
##
```

```
hspin <- matrix(as.numeric(unlist(read.table("human.rnf43.all.bp.txt", header=F, sep=","))), nrow=bp.di
obs <- c(hspin)
perm <- c()
for (i in 1:100) {
        name <- paste("ms02.human", "/", "rnf43.heatmap", "/", "ms02.", i, ".bp.matrix.csv", sep="")</pre>
        mat <- matrix(as.numeric(unlist(read.table(name, header=F, sep=","))), nrow=bp.dim, ncol=conn.dim)</pre>
        perm <- rbind(perm, c(mat))</pre>
}
mean <- colMeans(perm)</pre>
std <- colSds(perm)</pre>
zscore <- round((obs - mean)/std, 3)</pre>
z.p <- matrix(zscore, nrow=bp.dim, ncol=conn.dim)</pre>
z.p \leftarrow t(z.p)
rownames(z.p) \leftarrow c("5\%","10\%","15\%","20\%","25\%","30\%","35\%","40\%","45\%","50\%","50\%","50\%","50\%","50\%","50\%","50\%","50\%","50\%","50\%","50\%","50\%","50\%","50\%","50\%","50\%","50\%","50\%","50\%","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\","50\"
                                      "55%", "60%", "65%", "70%", "75%", "80%", "85%", "90%", "95%", "100%")
colnames(z.p) <- bp.go.cat</pre>
hspin <- matrix(as.numeric(unlist(read.table("human.rnf43.NP.bp.txt", header=F, sep=","))), nrow=bp.dim
obs <- c(hspin)
perm <- c()
for (i in 1:100) {
        name <- paste("ms02.human", "/", "rnf43.NP", "/", "ms02.", i, ".bp.matrix.csv", sep="")</pre>
        mat <- matrix(as.numeric(unlist(read.table(name, header=F, sep=","))), nrow=bp.dim, ncol=conn.dim)</pre>
        perm <- rbind(perm, c(mat))</pre>
}
mean <- colMeans(perm)</pre>
std <- colSds(perm)</pre>
zscore <- round((obs - mean)/std, 3)</pre>
z.np <- matrix(zscore, nrow=bp.dim, ncol=conn.dim)</pre>
z.np \leftarrow t(z.np)
rownames(z.np) <- c("5%","10%","15%","20%","25%","30%","35%","40%","45%","50%",
                                      "55%", "60%", "65%", "70%", "75%", "80%", "85%", "90%", "95%", "100%")
colnames(z.np) <- bp.go.cat</pre>
enrich \leftarrow names(z.pvnp[20, which(z.pvnp[20,] > 8)])
suppress <- names(z.pvnp[20, which(z.pvnp[20,] < -8)])</pre>
enriched.terms <- c("GO.ID", "GO.Term", "Z.Tumor", "Z.Normal", "Z.Differential")
for (i in 1:length(enrich)) {
    id <- as.character(enrich[i])</pre>
    term <- Term(GOID(id))</pre>
    seri <- which(bp.go.cat %in% id)</pre>
    z.tumor <- z.p[20,seri]</pre>
    z.normal <- z.np[20,seri]</pre>
    z.diff <- z.pvnp[20,seri]</pre>
    enriched.terms <- rbind(enriched.terms, c(id, term, z.tumor, z.normal, z.diff))</pre>
```

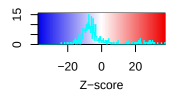
```
enriched.terms
##
##
  enriched.terms "GO.ID"
                   "GD:0006886"
##
##
                   "GD:0006888"
##
                   "GD:0006890"
                   "GD:0008286"
##
##
                   "GO:0015031"
                   "GD:0015721"
##
                   "GD:0015991"
##
                   "GO:0016241"
##
##
                   "GD:0019216"
##
                   "GO:0034220"
                   "GO:0043401"
##
                   "GD:0045165"
##
                   "GO:0045893"
##
##
                   "GD:0060070"
                   "GO:0061024"
##
                   "GO:1904837"
##
                   GD:0006886
##
   enriched.terms "GO.Term"
##
##
                   "intracellular protein transport"
##
                   "ER to Golgi vesicle-mediated transport"
##
                   "retrograde vesicle-mediated transport, Golgi to ER"
##
                   "insulin receptor signaling pathway"
                   "protein transport"
##
                   "bile acid and bile salt transport"
##
##
                   "ATP hydrolysis coupled proton transport"
##
                   "regulation of macroautophagy"
                   "regulation of lipid metabolic process"
##
                   "ion transmembrane transport"
##
                   "steroid hormone mediated signaling pathway"
##
##
                   "cell fate commitment"
                   "positive regulation of transcription, DNA-templated"
##
##
                   "canonical Wnt signaling pathway"
##
                   "membrane organization"
##
                   "beta-catenin-TCF complex assembly"
##
##
  enriched.terms "Z.Tumor" "Z.Normal" "Z.Differential"
                   "-5.797"
                             "-12.979"
                                        "9.586"
##
##
                   "0.447"
                             "-5.762"
                                        "10.926"
##
                   "5.891"
                             "1.758"
                                        "8.685"
                   "-4"
                             "-9.397"
                                         "11.969"
##
##
                   "-4.142"
                             "-8.97"
                                         "10.599"
                   "-0.11"
                             "-0.828"
                                        "8.931"
##
##
                   "6.554"
                             "2.113"
                                         "8.124"
                   "3.269"
                             "-1.876"
                                         "8.35"
##
                   "-8.752" "-11.681"
                                        "8.712"
##
                   "-4.452" "-9.392"
                                         "9.001"
##
##
                   "-5.947" "-7.538"
                                         "8.016"
                   "-7.727" "-9.86"
                                         "9.338"
##
                   "-11.599" "-11.275" "9.389"
##
```

```
##
                   "-8.112" "-11.894"
                                         "11.8"
##
                   "-6.643" "-9.363"
                                         "8.481"
                                         "8.535"
##
                   "-0.481" "-1.023"
write.table(enriched.terms, file="17.PvNP.bp.enriched.csv", sep="\t", col.names = F, row.names = F, quo
suppressed.terms <- c("GO.ID", "GO.Term", "Z.Tumor", "Z.Normal", "Z.Differential")
for (i in 1:length(suppress)) {
  id <- as.character(suppress[i])</pre>
  term <- Term(GOID(id))</pre>
  seri <- which(bp.go.cat %in% id)</pre>
  z.tumor <- z.p[20,seri]</pre>
  z.normal <- z.np[20,seri]</pre>
  z.diff <- z.pvnp[20,seri]</pre>
  suppressed.terms <- rbind(suppressed.terms, c(id, term, z.tumor, z.normal, z.diff))</pre>
suppressed.terms
##
## suppressed.terms "GO.ID"
##
                     "GD:0000184"
                     "GO:0000245"
##
##
                     "GD:0000375"
##
                     "GO:0000381"
##
                     "GD:0000398"
                     "GO:0006351"
##
##
                     "GD:0006357"
##
                     "GD:0006364"
                     "GD:0006368"
##
##
                     "GD:0006369"
##
                     "GD:0006370"
##
                     "GD:0006405"
                     "GO:0006406"
##
                     "GD:0006412"
##
##
                     "GD:0006413"
                     "GD:0006614"
##
                     "GO:0007186"
##
                     "GO:0008380"
##
                     "GD:0016070"
##
                     "GD:0031124"
##
##
                     GD:0000184
##
  suppressed.terms "GO.Term"
                     "nuclear-transcribed mRNA catabolic process, nonsense-mediated decay"
##
##
                     "spliceosomal complex assembly"
##
                     "RNA splicing, via transesterification reactions"
                     "regulation of alternative mRNA splicing, via spliceosome"
##
##
                     "mRNA splicing, via spliceosome"
##
                     "transcription, DNA-templated"
##
                     "regulation of transcription by RNA polymerase II"
##
                     "rRNA processing"
##
                     "transcription elongation from RNA polymerase II promoter"
##
                     "termination of RNA polymerase II transcription"
                     "7-methylguanosine mRNA capping"
##
                     "RNA export from nucleus"
##
##
                     "mRNA export from nucleus"
```

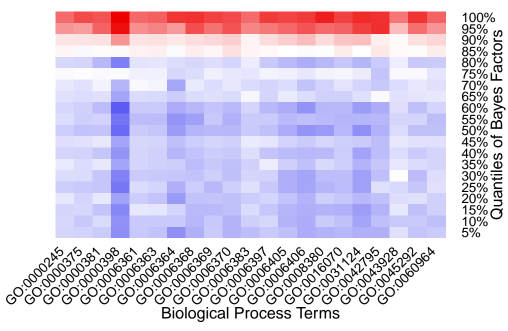
```
##
                    "translational initiation"
##
                    "SRP-dependent cotranslational protein targeting to membrane"
                    "G protein-coupled receptor signaling pathway"
##
##
                    "RNA splicing"
##
                    "RNA metabolic process"
##
                    "mRNA 3'-end processing"
##
  suppressed.terms "Z.Tumor" "Z.Normal" "Z.Differential"
                    "14.405"
                              "16.471"
                                          "-11.518"
##
##
                    "21.839"
                              "26.653"
                                          "-9.467"
                    "26.615"
                              "33.108"
                                          "-8.647"
##
                    "25.057"
                              "30.565"
                                          "-8.374"
##
                                          "-25.573"
                    "37.487" "45.921"
##
                    "-37.573" "-35.083"
##
                                         "-11.756"
                    "-30.787" "-29.925"
##
                                         "-10.889"
##
                    "29.684"
                              "33.13"
                                          "-9.211"
                    "30.011" "36.357"
                                          "-9.318"
##
##
                    "28.085"
                              "36.305"
                                          "-9.622"
                    "26.45"
                              "33.574"
                                          "-8.672"
##
##
                    "27.319" "34.43"
                                         "-11.203"
##
                    "28.886" "35.153"
                                         "-13.816"
                    "14.239"
                              "15.477"
                                          "-8.634"
##
                    "13.262" "14.146"
                                          "-8.171"
##
                    "11.217" "13.523"
                                          "-10.395"
##
##
                    "-48.543" "-51.947"
                                         "-9.709"
##
                    "33.127"
                              "40.361"
                                          "-11.357"
##
                    "27.901"
                              "33.063"
                                          "-10.41"
                                         "-15.223"
##
                    "31.361" "40.541"
write.table(suppressed.terms, file="17.PvNP.bp.suppressed.csv", sep="\t", col.names = F, row.names = F,
# PDAC cells
p.pos <-z.p[, which(z.p[20,] > 20)]
heatmap.2(p.pos, col=my_palette, dendrogram='none',
          trace='none', Rowv=F, Colv=F, revC=T, #labCol=NA,
          xlab="Biological Process Terms", ylab="Quantiles of Bayes Factors",
          margins = c(5.5,4), key.title = NA, key.xlab="Z-score", key.ylab=NA,
          scale="none", symbreaks=T, symm=F, symkey = F,
          srtCol=45, adjCol=c(1,0),
          main="Tumor cells, top 5% enriched")
```

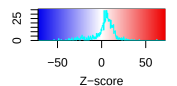
"translation"

##

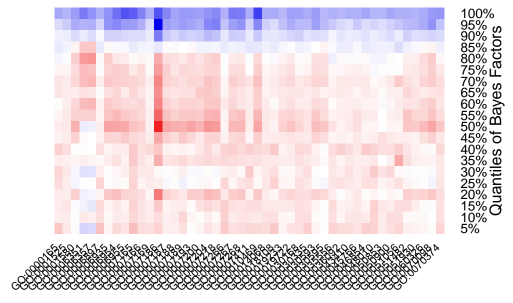


Tumor cells, top 5% enriched

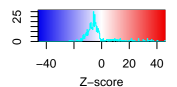




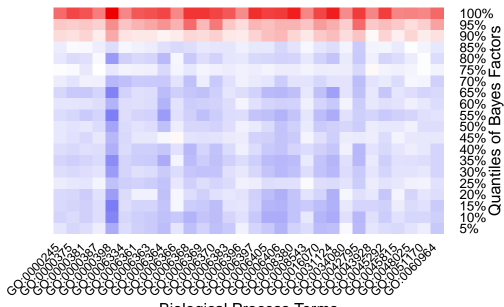
Tumor cells, top 5% suppressed



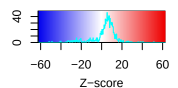
Biological Process Terms



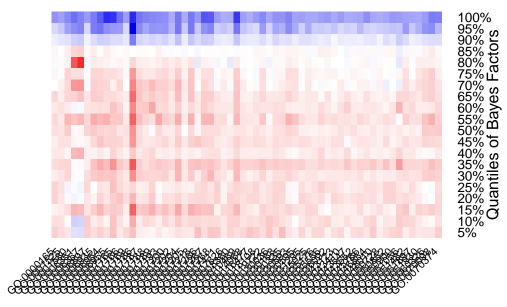
Normal cells, top 5% enriched



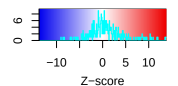
Biological Process Terms



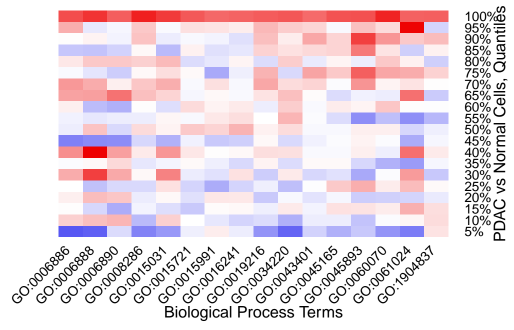
Normal cells, top 5% suppressed

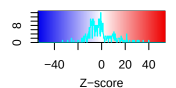


Biological Process Terms



Tumor vs normal cells, Enriched





Fumor vs normal cells, suppressed

