tmod: Analysis of Transcriptional Modules

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Abstract

The package tmod provides blood transcriptional modules described by Chaussabel et al. (2008) and by Li et al. (2014) as well as metabolic profiling clusters from Weiner et al. (2012). Furthermore, the package includes tools for testing the significance of enrichment of the modules as well as visualisation of the features (genes, metabolites etc.) and modules. This vignette is a tutorial for the package.

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Basic data analysis

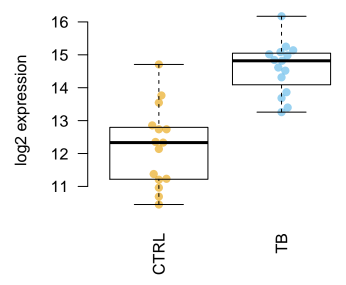
The Gambia data set

In the following, we will use the Egambia data set included in the package. The data set has been generated by Maertzdorf et al. (2011) and has the GEO ID GSE28623. The data is already background corrected and normalized, so we can proceed with a differential gene expression analysis. Note that only a bit over 5000 genes from the original set of over 45000 probes is included.

```
design <- cbind(Intercept=rep(1, 30), TB=rep(c(0,1), each= 15))</pre>
E <- as.matrix(Egambia[,-c(1:3)])</pre>
fit <- eBayes( lmFit(E, design))</pre>
tt <- topTable(fit, coef=2, number=Inf,
  genelist=Egambia[,1:3] )
head(tt, 10)
##
         GENE_SYMBOL
## 4178
              FAM20A
## 20799
              FCGR1B
## 4122
               BATF2
## 23567
             ANKRD22
## 20498
               SEPT4
## 20360
               CD274
## 2513
                 AIM2
## 24032
              GOLSYN
## 1337
                 ETV7
## 467
            SERPING1
##
                                                                GENE NAME
                                                                              F.G
## 4178
                          family with sequence similarity 20, member A"
                                                                           54757
                Fc fragment of IgG, high affinity Ib, receptor (CD64)"
## 20799
                 basic leucine zipper transcription factor, ATF-like 2 116071
## 4122
## 23567
                                                ankyrin repeat domain 22 118932
## 20498
                                                                 septin 4
                                                                            5414
## 20360
                                                           CD274 molecule
                                                                           29126
## 2513
                                                    absent in melanoma 2
                                                                            9447
## 24032
                                                 Golgi-localized protein
                                                                           55638
                                                            ets variant 7
## 1337
                                                                           51513
## 467
         serpin peptidase inhibitor, clade G (C1 inhibitor), member 1"
                                                                             710
##
             logFC
                                              P.Value
                                                        adj.P.Val
                      AveExpr
                                      t
## 4178
          2.955829
                    4.007327
                               6.200637 3.423267e-07 0.001898886 6.457171
## 20799
                               5.946113 7.552423e-07 0.002094665 5.741043
          2.391490 13.401207
                               5.797752 1.198442e-06 0.002215920 5.322491
## 4122
          2.680837 10.398520
## 23567 2.763908
                               5.624092 2.057601e-06 0.002692116 4.832003
                    8.651749
## 20498 3.286528
                    4.223270
                               5.480564 3.215558e-06 0.002692116 4.426508
## 20360
                               5.463149 3.394453e-06 0.002692116 4.377314
          2.377399
                    7.334747
## 2513
          1.966342
                    9.933621
                               5.462879 3.397298e-06 0.002692116 4.376553
## 24032 -2.534812 2.221666 -5.362575 4.639596e-06 0.003018586 4.093323
## 1337
          2.844012 8.075046 5.345142 4.897651e-06 0.003018586 4.044119
          2.639069 7.708228 5.150375 8.958000e-06 0.004969002 3.495088
## 467
OK, we see some of the genes known to be prominent in the human host response to TB. We can display one
of these using tmod's showGene function (it's just a boxplot combined with a beeswarm, nothing special):
group <- rep( c("CTRL", "TB"), each=15)</pre>
showGene(E["20799",], group,
  main=Egambia["20799", "GENE_SYMBOL"])
```

library(limma)
library(tmod)
data(Egambia)

FCGR1B



Fine, but what about the modules?

Transcriptional module analysis

There are two main functions in tmod to understand which modules are significantly enriched¹

The first one, tmodHGtest, is simply a hypergeometric test on two groups of genes: 'foreground' (fg), or the list of differentially expressed genes, and 'background' (bg) – the gene universe, i.e., all genes present in the analysis. The gene identifiers used currently by tmod are HGNC identifiers, and we will use the GENE_SYMBOL field from the Egambia data set.

In this particular example, however, we have almost no genes which are significantly differentially expressed after correction for multiple testing: the power of the test with 10 individuals in each group is too low. For the sake of the example, we will therefore relax our selection. Normally, I'd use a q-value threshold of at least 0.001.

```
fg <- tt$GENE_SYMBOL[tt$adj.P.Val < 0.05 & abs( tt$logFC ) > 1]
res <- tmodHGtest(fg=fg, bg=tt$GENE_SYMBOL)</pre>
res
##
                                                                    Title b B
## LI.M112.0 LI.M112.0
                                               complement activation (I) 4 11
## LI.M11.0
              LI.M11.0
                                              enriched in monocytes (II) 4 20
## LI.M75
                                                 antiviral IFN signature 3 10
                LI.M75
## LI.S4
                 LI.S4
                                              Monocyte surface signature 3 10
## LI.S5
                 LI.S5
                                                     DC surface signature 4 34
## LI.M165
               LI.M165
                              enriched in activated dendritic cells (II) 3 19
## LI.M4.3
               LI.M4.3 myeloid cell enriched receptors and transporters 2
## LI.M16
                LI.M16
                                          TLR and inflammatory signaling 2 5
##
                   N
                             Ε
                                    P. Value
                                               adj.P.Val
## LI.M112.0 47 4826 37.33849 2.480096e-06 0.0008581134
## LI.M11.0
             47 4826 20.53617 3.414323e-05 0.0059067783
## LI.M75
             47 4826 30.80426 9.906126e-05 0.0085687989
```

¹If you work with limma, there are other, more efficient and simpler to use functions. See "Working with limma" below.

The columns in the above table contain the following:

- ID The module ID. IDs starting with "LI" come from Li et al. (S. Li et al. 2014), while IDs starting with "DC" have been defined by Chaussabel et al. (Chaussabel et al. 2008).
- Title The module description
- b Number of genes from the given module in the fg set
- B Number of genes from the module in the bg set
- n Size of the fg set

[1] 25

- N Size of the bg set
- E Enrichment, calcualted as (b/n)/(B/N)
- P.Value P-value from the hypergeometric test
- adj.P.Val P-value adjusted for multiple testing using the Benjamini-Hochberg correction

Well, IFN signature in TB is well known. However, the numbers of genes are not high: n is the size of the foreground, and b the number of genes in fg that belong to the given module. N and B are the respective totals – size of bg+fg and number of genes that belong to the module that are found in this totality of the analysed genes. If we were using the full Gambia data set (with all its genes), we would have a different situation.

Another approach is to sort all the genes (for example, by the respective p-value) and perform a U-test on the ranks of (i) genes belonging to the module and (ii) genes that do not belong to the module. This is a bit slower, but often works even in the case if the power of the statistical test for differential expression is low. That is, even if only a few genes or none at all are significant at acceptable thresholds, sorting them by the p-value or another similar metric can nonetheless allow to get meaningful enrichments².

Moreover, we do not need to set arbitrary thresholds, like p-value or logFC cutoff.

```
<- tt$GENE SYMBOL
res2 <- tmodUtest(1)
head(res2)
                  ID
                                                                U
                                                                   N1
                                                                             AUC
                                                     Title
## LI.M37.0 LI.M37.0 immune activation - generic cluster 352659 100 0.7462103
## LI.M37.1 LI.M37.1
                                                            50280
                              enriched in neutrophils (I)
                                                                    12 0.8703781
## LI.S4
               LI.S4
                               Monocyte surface signature
                                                            43220
                                                                    10 0.8974252
              LI.M75
## LI.M75
                                  antiviral IFN signature
                                                            42996
                                                                   10 0.8927741
## LI.M11.0 LI.M11.0
                               enriched in monocytes (II)
                                                            74652
                                                                   20 0.7766542
## LI.M67
              LI.M67
                                activated dendritic cells
                                                            28095
                                                                    6 0.9714730
                 P.Value
                             adj.P.Val
## LI.M37.0 1.597067e-17 5.525852e-15
## LI.M37.1 4.530577e-06 6.569127e-04
## LI.S4
            6.853638e-06 6.569127e-04
            8.632649e-06 6.569127e-04
## LI.M75
## LI.M11.0 9.492958e-06 6.569127e-04
## LI.M67
            3.200305e-05 1.811391e-03
nrow(res2)
```

²The rationale is that the non-significant p-values are not associated with the test that we are actually performing, but merely used to sort the gene list. Thus, it does not matter whether they are significant or not.

This list makes a lot of sense, and also is more stable than the other one: it does not depend on modules that contain just a few genes. Since the statistics is different, the b, B, n, N and E columns in the output have been replaced by the following:

- U The Mann-Whitney U statistics
- N1 Number of genes in the module
- AUC Area under curve a measure of the effect size

The U test is simple, however has two main issues. Firstly, it detects enrichments as well as depletions – that is, modules which are enriched at the bottom of the list (e.g. modules which are never, ever regulated in a particular comparison) will be detected as well. This is often undesirable. Secondly, large modules will be reported as significant even if the actual effect size (i.e., AUC) is modest or very small, just because of the sheer number of genes in a module. Unfortunately, also the reverse is true: modules with a small number of genes, even if they consist of highly up- or down-regulated genes from the top of the list will not be detected.

The CERNO test, described by Yamaguchi et al. (Yamaguchi et al. 2008), is based on Fisher's method of combining probabilities. In summary, for a given module, the ranks of genes from the module are logarithmized, summed and multiplied by -2:

$$f_{CERNO} = -2 \cdot \sum_{i=1}^{N} \ln \frac{R_i}{N_{tot}}$$

This statitic has the χ^2 distribution with $2 \cdot N$ degrees of freedom, where N is the number of genes in a given module and N_{tot} is the total number of genes (Yamaguchi et al. 2008).

The CERNO test is actually much more practical than the U test for most purposes.

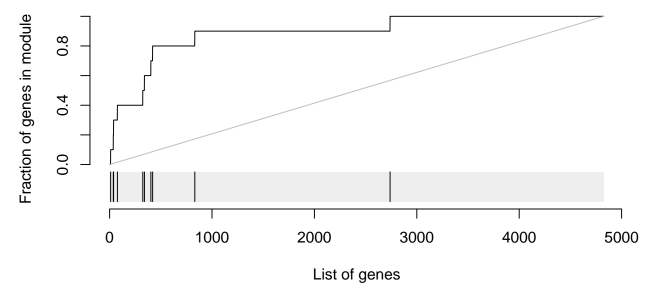
```
1 <- tt$GENE_SYMBOL
res2 <- tmodCERNOtest(1)
head( res2 )</pre>
```

```
##
                    ID
                                                       Title
                                                                        N1
                                                                 cerno
## LI.M37.0
              LI.M37.0 immune activation - generic cluster 426.35781 100
## LI.M11.0
              LI.M11.0
                                 enriched in monocytes (II) 113.80864
## LI.S4
                                 Monocyte surface signature
                 LI.S4
                                                              76.37298
## LI.M112.0 LI.M112.0
                                  complement activation (I)
                                                              73.67987
                                                                        11
                LI.M75
                                                              65.29854
## LI.M75
                                    antiviral IFN signature
## LI.M16
                LI.M16
                                                              46.33475
                             TLR and inflammatory signaling
                                                                         5
                   AUC
##
                             cES
                                      P. Value
                                                 adj.P.Val
             0.7462103 2.131789 1.824844e-18 6.313962e-16
## LI.M37.0
## LI.M11.0
             0.7766542 2.845216 5.255069e-09 9.091270e-07
## LI.S4
             0.8974252 3.818649 1.606057e-08 1.852319e-06
## LI.M112.0 0.8455773 3.349085 1.722322e-07 1.489809e-05
## LI.M75
             0.8927741 3.264927 1.045914e-06 7.192190e-05
## LI.M16
             0.9790500 4.633475 1.247201e-06 7.192190e-05
```

Here, the results are similar, however CERNO test was able to detect another module – "TLR and inflammatory signaling". Although only 5 genes are in this module (which is why U test could not detect it), the genes are all on the top of the list of the differentially regulated genes.

Let us now investigate in more detail the module LI.M75, the antiviral interferon signature. We can use the evidencePlot function to see how the module is enriched in the list 1.

```
evidencePlot(1, "LI.M75")
```



In essence, this is a receiver-operator characteristic (ROC) curve, and the area under the curve (AUC) is related to the U-statistic, from which the P-value in the tmodUtest is calculated, as AUC = $\frac{U}{n_1 \cdot n_2}$. Both the U statistic and the AUC are reported. Moreover, the AUC can be used to calculate effect size according to the Wendt's formula (Wendt 1972) for rank-biserial correlation coefficient:

$$r = 1 - \frac{2 \cdot U}{n_1 \cdot n_2} = 1 - 2 \cdot AUC$$

In the above diagram, we see that nine out of the 10 genes that belong to the LI.M75 module and which are present in the Egambia data set are ranked among the top 1000 genes (as sorted by p-value).

Working with multiple sets of comparisons

Working with limma

Given the popularity of the limma package, tmod includes functions to easily integrate with limma. In fact, if you fit a design / contrast with limma function lmFit and calculate the p-values with eBayes(), you can directly use the resulting object in tmodLimmaTest and tmodLimmaDecideTests³.

```
res.l <- tmodLimmaTest(fit, Egambia$GENE_SYMBOL)
length(res.1)
## [1] 2
names(res.1)
## [1] "Intercept" "TB"
head(res.1$TB)
##
                                                       Title
                                                                 cerno
## LI.M37.0
              LI.M37.0 immune activation - generic cluster 414.27395 100
## LI.M11.0
              LI.M11.0
                                 enriched in monocytes (II) 105.61794
## LI.M112.0 LI.M112.0
                                  complement activation (I)
                                                              75.62229
                                                                        11
## LI.S4
                                 Monocyte surface signature
                                                              69.97439
```

 $^{^3}$ The function tmodLimmaDecideTests is described in the next section

```
## LI.M75
                LI.M75
                                    antiviral IFN signature
                                                              66.10214
## LI.M67
                I.T. M67
                                  activated dendritic cells
                                                              50.35750
##
                   AUC
                             cES
                                      P.Value
                                                 adj.P.Val
             0.7255121 2.071370 4.568772e-17 1.580795e-14
## LI.M37.0
             0.7862464 2.640449 7.921155e-08 9.671792e-06
## LI.M112.0 0.8667988 3.437377 8.385947e-08 9.671792e-06
## LI.S4
             0.8836794 3.498719 1.838992e-07 1.590728e-05
## LI.M75
             0.8645349 3.305107 7.780282e-07 5.383955e-05
## LI.M67
             0.9712310 4.196458 1.208877e-06 6.971189e-05
```

The tmodLimmaTest function uses coefficients and p-values from the limma object to order the genes. By default, the genes are ordered by MSD (Minimum Significant Difference), rather than p-value or log fold change.

The MSD is defined as follows:

$$\label{eq:msd} \text{MSD} = \begin{cases} CI.L & \text{if logFC} > 0 \\ -CI.R & \text{if logFC} < 0 \end{cases}$$

Where logFC is the log fold change, CI.L is the left boundary of the 95% confidence interval of logFC and CI.R is the right boundary. MSD is always greater than zero and is equivalent to the absolute distance between the confidence interval and the x axis. For example, if the logFC is 0.7 with 95% CI = [0.5, 0.9], then MSD=0.5; if logFC is -2.5 with 95% CI = [-3.0, -2.0], then MSD = 2.0.

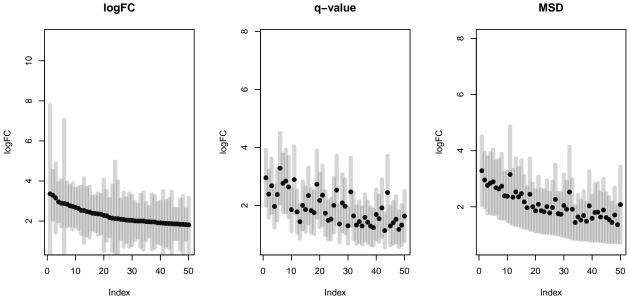
The idea behind MSD is as follows. Ordering genes by decreasing absolute log fold change will include on the top of the list some genes close to background, for which log fold changes are grand, but so are the errors and confidence intervals, just because measuring genes with low expression is loaded with errors. Ordering genes by decreasing absolute log fold change should be avoided.

On the other hand, in a list ordered by p-values, many of the genes on the top of the list will have strong signals and high expression, which results in better statistical power and ultimately with lower p-values – even though the actual fold changes might not be very impressive.

However, by using MSD and using the boundary of the confidence interval to order the genes, the genes on the top of the list are those for which we can *confidently* that the actual log fold change is large. That is because the 95% confidence intervals tells us that in 95% cases, the real log fold change will be anywhere within that interval. Using its bountary closer to the x-axis (zero log fold change), we say that in 95% of the cases the log fold change will have this or larger magnitude (hence, "minimal significant difference").

This can be visualized as follows, using the drop-in replacement for limma's topTable function, tmodLimmaTopTable, which calculates msd as well as confidence intervals. We will consider only genes with positive log fold changes and we will show top 50 genes as ordered by the three different measures:

```
##
         logFC.TB
                          t.TB
                                   msd.TB
                                              SE.TB
                                                            d.TB
                                                                      ciL.TB
       0.02819016
                   0.07556852 -0.7277810 0.3730410
                                                      0.02879949 -0.7277810
## 34
                   3.87981294
                               0.7280616 0.3928453
##
       1.52416640
                                                      1.63977633
                                                                  0.7280616
                   0.17834978 -0.8174289 0.4422934
       0.07888294
                                                      0.09554857 -0.8174289
##
##
       0.15321399
                   0.32385505 -0.8055162 0.4730944
                                                      0.19850746 -0.8055162
  52 -0.23501607 -0.61703279 -0.5368429 0.3808810 -0.24514187 -1.0068750
  62 -0.31952987 -0.55848332 -0.8399144 0.5721386 -0.50066053 -1.4789741
##
         ciR.TB
                   qval.TB
## 34 0.7841613 0.99538447
  36 2.3202712 0.04393162
## 41 0.9751948 0.99504430
## 44 1.1119442 0.99504430
## 52 0.5368429 0.99504430
## 62 0.8399144 0.99504430
x <- x[ x$logFC.TB > 0, ] # only to simplify the output!
x2 <- x[ order(abs(x$logFC.TB), decreasing=T),][1:50,]</pre>
plotCI(x2$logFC.TB, x2$ciL.TB, x2$ciR.TB, "logFC")
x2 <- x[ order(x$qval.TB),][1:50,]</pre>
plotCI(x2$logFC.TB, x2$ciL.TB, x2$ciR.TB, "q-value")
x2 \leftarrow x[ order(x$msd.TB, decreasing=T),][1:50,]
plotCI(x2$logFC.TB, x2$ciL.TB, x2$ciR.TB, "MSD")
```



Black dots are logFCs, and grey bars denote 95% confidence intervals. On the left panel, the top 50 genes ordered by the fold change include several genes with broad confidence intervals, which, despite having a large log fold change, are not significantly up- or down-regulated.

On the middle panel the genes are ordered by p-value. It is clear that the log fold changes of the genes vary considerably, and that the list includes genes which are more and less strongly regulated in TB.

The third panel shows genes ordered by decreasing MSD. There is less variation in the logFC than on the second panel, but at the same time the fallacy of the first panel is avoided. MSD is a compromise between considering the effect size and the statistical significance.

What about enrichments?

```
x <- tmodLimmaTopTable(fit, coef="TB", genelist=Egambia[,1:3])</pre>
x.lfc <- x[ order(abs(x$logFC.TB), decreasing=T),]</pre>
x.qval <- x[ order(x$qval.TB),]</pre>
x.msd <- x[ order(x$msd.TB, decreasing=T),]</pre>
head(tmodCERNOtest(x.lfc$GENE SYMBOL))
##
                    ID
                                                              Title
                                                                        cerno
                              immune activation - generic cluster 381.20058
## LI.M37.0
              LI.M37.0
## LI.M112.0 LI.M112.0
                                         complement activation (I)
                                                                     67.57425
## LI.M75
                LI.M75
                                           antiviral IFN signature
                                                                     59.49036
## LI.S4
                 LT.S4
                                        Monocyte surface signature
                                                                     58.93059
## LI.M67
                LI.M67
                                         activated dendritic cells
                                                                     44.54543
## LI.M165
               LI.M165 enriched in activated dendritic cells (II)
                                                                     84.06285
              N1
                       AUC
                                 cES
                                          P.Value
                                                     adj.P.Val
## LI.M37.0 100 0.7345239 1.906003 1.990299e-13 6.886434e-11
## LI.M112.0 11 0.8340036 3.071557 1.583945e-06 2.740225e-04
## LI.M75
              10 0.8714493 2.974518 8.537502e-06 8.529686e-04
## LI.S4
              10 0.8759759 2.946529 1.041171e-05 8.529686e-04
## LI.M67
               6 0.9406639 3.712119 1.232614e-05 8.529686e-04
## LI.M165
              19 0.7057362 2.212180 2.482939e-05 1.431828e-03
head(tmodCERNOtest(x.qval$GENE_SYMBOL))
                    ID
                                                      Title
                                                                       N1
                                                                 cerno
## LI.M37.0
              LI.M37.0 immune activation - generic cluster 427.03180 100
## LI.M11.0
              LI.M11.0
                                 enriched in monocytes (II) 114.85395
## LI.S4
                 LI.S4
                                Monocyte surface signature
                                                             77.27812
## LI.M112.0 LI.M112.0
                                  complement activation (I)
                                                             74.30410
                                                                        11
## LI.M75
                LI.M75
                                    antiviral IFN signature
                                                             65.41723
## LI.M16
                LI.M16
                            TLR and inflammatory signaling
                                                             46.32281
                   AUC
                                      P.Value
                                                 adj.P.Val
## LI.M37.0 0.7523212 2.135159 1.521279e-18 5.263625e-16
## LI.M11.0 0.7910112 2.871349 3.691442e-09 6.386194e-07
             0.9164037 3.863906 1.131929e-08 1.305491e-06
## LI.S4
## LI.M112.0 0.8595676 3.377459 1.367620e-07 1.182991e-05
             0.8932932 3.270861 1.001270e-06 6.928788e-05
## LI.M75
## LI.M16
             0.9790085 4.632281 1.253436e-06 7.228150e-05
head(tmodCERNOtest(x.msd$GENE SYMBOL))
                                                      Title
##
                    TD
                                                                 cerno
                                                                       N1
## LI.M37.0
              LI.M37.0 immune activation - generic cluster 414.27395 100
## LI.M11.0
              LI.M11.0
                                enriched in monocytes (II) 105.61794
## LI.M112.0 LI.M112.0
                                 complement activation (I)
                                                             75.62229
                                                                        11
                                Monocyte surface signature
## LI.S4
                 LI.S4
                                                             69.97439
## I.T.M75
                                    antiviral IFN signature
                LI.M75
                                                             66.10214
                                                                        10
## LI.M67
                LI.M67
                                  activated dendritic cells
                                                             50.35750
##
                   AUC
                             cES
                                      P. Value
                                                 adj.P.Val
## LI.M37.0 0.7255121 2.071370 4.568772e-17 1.580795e-14
## LI.M11.0 0.7862464 2.640449 7.921155e-08 9.671792e-06
## LI.M112.0 0.8667988 3.437377 8.385947e-08 9.671792e-06
## LI.S4
             0.8836794 3.498719 1.838992e-07 1.590728e-05
## LI.M75
             0.8645349 3.305107 7.780282e-07 5.383955e-05
## LI.M67
             0.9712310 4.196458 1.208877e-06 6.971189e-05
```

In this case, the results of p-value and msd-ordering are very similar.

Comparing tests across experimental conditions

In the above example with the Gambian data set there were only two coefficients calculated in limma, the intercept and the TB. However, often there are several coefficients or contrasts which are analysed simultaneously, for example different experimental conditions or different time points. tmod includes several functions which make it easy to visualize such sets of enrichments.

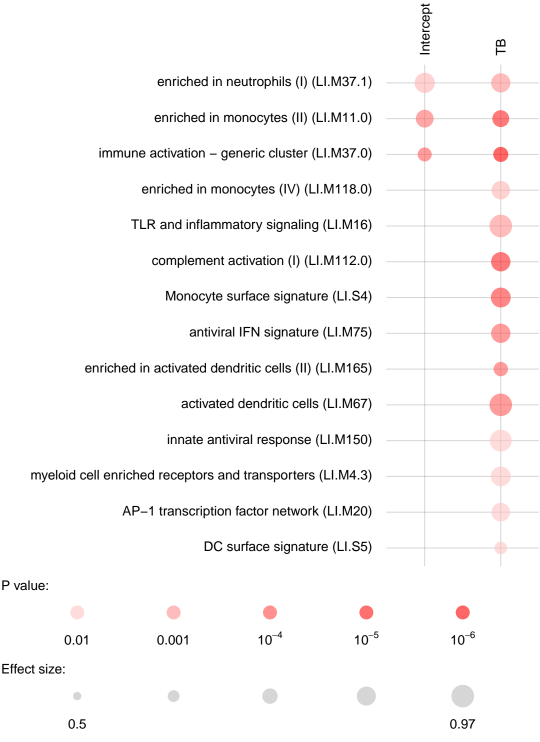
The object res.l created above using the tmod function tmodLimmaTest is a list of tmod results. Any such list can be directly passed on to functions tmodSummary and tmodPanelPlot, as long as each element of the list has been created with tmodCERNOtest or a similar function. tmodSummary creates a table summarizing module information in each of the comparisons made. The values for modules which are not found in a result object (i.e., which were not found to be significantly enriched in a given comparison) are shown as NA's:

head(tmodSummary(res.1), 5)

```
##
                    ID
                                                Title AUC. Intercept
## LI.M11.0
              LI.M11.0
                           enriched in monocytes (II)
                                                           0.8145651
## LI.M112.0 LI.M112.0
                            complement activation (I)
                                                                  NA
## LI.M118.0 LI.M118.0
                           enriched in monocytes (IV)
                                                                  NA
## LI.M124
               LI.M124 enriched in membrane proteins
                                                           0.8807517
## LI.M127
               LI.M127
                           type I interferon response
                                                                  NA
##
              q.Intercept
                             AUC.TB
## LI.M11.0 0.0001137611 0.7862464 9.671792e-06
## LI.M112.0
                       NA 0.8667988 9.671792e-06
## LI.M118.0
                       NA 0.8377967 2.850219e-03
## LI.M124
             0.0114869572
                                  NA
## LI.M127
                       NA 0.9448247 1.043621e-02
```

We can neatly visualize the above information on a heatmap-like representation:

```
tmodPanelPlot(res.1, text.cex=0.8)
```



The sizes of the red blobs on the figure correspond to the effect size, that is, the AUC, while the intensity of the color reflects the q-value from the module enrichment test. We can see that also the intercept term is enriched for genes found in monocytes and neutrophils. Note that by default, todPanelPlot only shows enrichments with p < 0.01, hence a slight difference from the todPanelPlot only shows

The function tmodPanelPlot has many optional arguments for customization, including options for label sizes, p value thresholds and custom functions for plotting the test results instead of just red blobs.

It is often of interest to see which enriched modules go up, and which go down? Specifically, we would like to

see, for each module, how many genes are up-, and how many genes are down-regulated. tmodPanelPlot takes an optional argument, pie, which contains information on significantly regulated genes in modules. We can conveniently generate it from a limma linear fit object with the tmodLimmaDecideTests function:

```
pie <- tmodLimmaDecideTests(fit, genes=Egambia$GENE_SYMBOL)
head(pie$TB[ order( pie$TB[,"Up"], decreasing=T), ])</pre>
```

```
##
              Down Zero Up
## DC.M3.4
                 0
                     11
                          9
## DC.M4.2
                 0
                     16
                          7
## LI.M11.0
                 0
                     16
## LI.M37.0
                 0
                    110
                          4
## LI.M112.0
                      9
## LI.M165
                 0
                     24
data(tmod)
tmod$MODULES["DC.M3.4",]
```

```
## ID Title Category Annotated
## DC.M3.4 DC.M3.4 Interferon DC.M3 Yes
##

## DC.M3.4 http://www.biir.net/public_wikis/module_annotation/V2_Trial_8_Modules_M3.4
## Source SourceID original.ID B
## DC.M3.4 http://www.biir.net/ DC M3.4 53
```

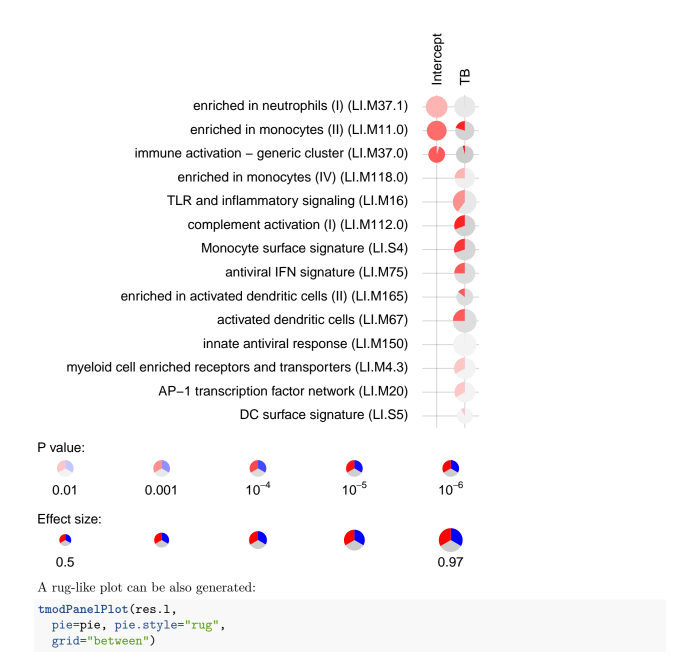
The pie object is a list. Each element of the list corresponds to one coefficient and is a data frame with the columns "Down", "Zero" and "Up" (in that order). Importantly, all names of the "res.l" list must correspond to an item in the pie list.

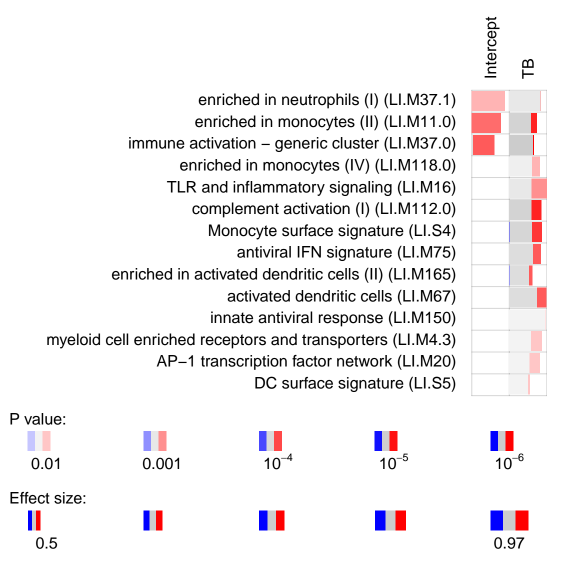
```
all(names(pie) %in% names(res.1))
```

[1] TRUE

We can now use this information in tmodPanelPlot:

```
tmodPanelPlot(res.1, pie=pie, text.cex=0.8)
```





There is also a more general function, tmodDecideTests that also produces a tmodPanelPlot-compatible object, a list of data frames with gene counts. However, instead of taking a limma object, it requires (i) a gene name, (ii) a vector or a matrix of log fold changes, and (iii) a vector or a matrix of p-values. We can replicate the result of tmodLimmaDecideTests above with the following commands:

```
tt.I <-
  topTable(fit, coef="Intercept", number=Inf, sort.by="n")
tt.TB <- topTable(fit, coef="TB", number=Inf, sort.by="n")
pie2 <- tmodDecideTests(Egambia$GENE_SYMBOL,
  lfc=cbind(tt.I$logFC, tt.TB$logFC),
  pval=cbind(tt.I$adj.P.Val, tt.TB$adj.P.Val))
identical(pie[[1]], pie2[[1]])</pre>
```

[1] TRUE

Using other sets of modules

By default, tmod uses the modules published by Li et al. (S. Li et al. 2014) (LI). A second set of modules was published by Chaussabel et al. (Chaussabel et al. 2008) (DC); new module definitions were described by

Banchereau et al. (Banchereau et al. 2012) and can be found on a public website⁴.

Depending on the mset parameter to the test functions, either the LI or DC sets are used, or both, if the mset=all has been specified.

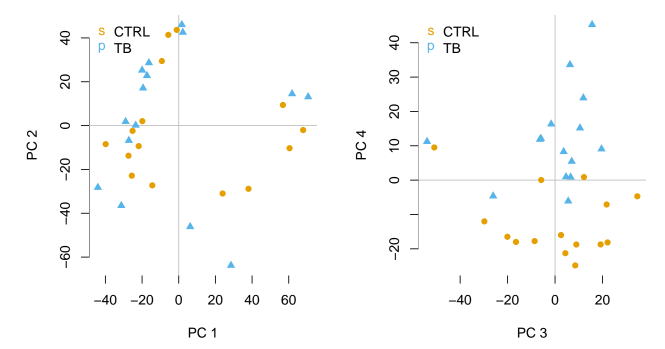
```
<- tt$GENE_SYMBOL
res2 <- tmodUtest(1, mset="all")
head( res2 )
##
                                                              U N1
                                                                          AUC
                  TD
                                                   Title
## LI.M37.0 LI.M37.0 immune activation - generic cluster 352659 100 0.7462103
## DC.M4.2
            DC.M4.2
                                            Inflammation 91352
                                                                 20 0.9503953
## DC.M1.2
            DC.M1.2
                                                          73612
                                                                 17 0.9004196
                                              Interferon
## DC.M3.2
            DC.M3.2
                                            Inflammation
                                                         96366 24 0.8361620
## DC.M5.15 DC.M5.15
                                             Neutrophils
                                                          65289 16 0.8483498
## DC.M7.29 DC.M7.29
                                                          77738 20 0.8087599
                                            Undetermined
                 P.Value
                            adj.P.Val
## LI.M37.0 1.597067e-17 9.678227e-15
## DC.M4.2 1.674762e-12 5.074530e-10
## DC.M1.2 5.703006e-09 9.623646e-07
## DC.M3.2 6.352241e-09 9.623646e-07
## DC.M5.15 7.240084e-07 8.774982e-05
## DC.M7.29 9.084521e-07 9.175366e-05
```

As you can see, the information contained in both module sets is partially redundant.

Functional multivariate analysis

Transcriptional modules can help to understand the biological meaning of the calculated multivariate transformations. For example, consider a principal component analysis (PCA), visualised using the pca3d package (Weiner 2013):

 $^{^4} http://www.biir.net/public_wikis/module_annotation/G2_Trial_8_Modules$



The fourth component looks really interesting. Does it correspond to the modules which we have found before? Each principal component is, after all, a linear combination of gene expression values multiplied by weights (or scores) which are constant for a given component. The i-th principal component for sample j is given by

$$PC_{i,j} = \sum_{k} w_{i,k} \cdot x_{k,j}$$

where k is the index of the variables (genes in our case), $w_{i,k}$ is the weight associated with the i-th component and the k-th variable (gene), and $x_{k,j}$ is the value of the variable k for the sample j; that is, the gene expression of gene k in the sample j. Genes influence the position of a sample along a given component the more the larger their absolute weight for that component.

For example, on the right-hand figure above, we see that samples which were taken from TB patients have a high value of the principal component 4; the opposite is true for the healthy controls. The genes that allow us to differentiate between these two groups will have very large, positive weights for genes highly expressed in TB patients, and very large, negative weights for genes which are highly expressed in NID, but not TB.

We can sort the genes by their weight in the given component, since the weights are stored in the pca object in the "rotation" slot, and use the tmodUtest function to test for enrichment of the modules.

```
o <- order(abs(pca$rotation[,4]), decreasing=TRUE)</pre>
1 <- Egambia$GENE_SYMBOL[o]</pre>
res <- tmodUtest(1)
head(res)
##
                   ID
                                                                   U
                                                                                AUC
                                                       Title
                                                                     N1
## LI.M37.0 LI.M37.0 immune activation - generic cluster 339742 100 0.7188785
## LI.M37.1 LI.M37.1
                               enriched in neutrophils (I)
                                                              50096
                                                                      12 0.8671929
## LI.M75
                                   antiviral IFN signature
                                                              43379
               LI.M75
                                                                      10 0.9007267
```

enriched in monocytes (II)

activated dendritic cells

adj.P.Val

74343

28291

20 0.7734395

34 0.7058762

6 0.9782503

LI.M11.0 LI.M11.0

LI.S5

LI.M67

P. Value

LI.S5

LI.M67

##

DC surface signature 115007

```
## LI.M37.0 3.133111e-14 1.084056e-11

## LI.M37.1 5.405722e-06 6.700097e-04

## LI.M75 5.809333e-06 6.700097e-04

## LI.M11.0 1.185187e-05 1.025187e-03

## LI.S5 1.711493e-05 1.184353e-03

## LI.M67 2.506730e-05 1.445548e-03
```

Perfect, this is what we expected: we see that the neutrophil / interferon signature which is the hallmark of the TB biosignature. What about other components? We can run the enrichment for each component and visualise the results using tmod's functions tmodSummary and tmodPanelPlot. Below, we use the filter.empty option to omit the principal components which show no enrichment at all.

```
# Calculate enrichment for each component
     <- Egambia$GENE_SYMBOL
# function calculating the enrichment of a PC
gn.f <- function(r) {</pre>
    tmodCERNOtest(gs[order(abs(r), decreasing=T)],
                 qval=0.01)
}
x <- apply(pca$rotation, 2, gn.f)
tmodSummary(x, filter.empty=TRUE)[1:5,]
##
                                                                      q.PC3
                     ID
                                              Title
                                                       AUC.PC3
              LI.M11.0 enriched in monocytes (II)
## LI.M11.0
                                                            NA
                                                                         NA
## LI.M112.0 LI.M112.0 complement activation (I)
                                                            NA
                                                                         NA
## LI.M118.0 LI.M118.0 enriched in monocytes (IV)
                                                            NA
                                                                         NA
## LI.M127
               LI.M127 type I interferon response
                                                            NA
                                                                         NA
## LI.M144
               LI.M144
                           cell cycle, ATP binding 0.9894257 0.006051848
               AUC.PC4
                               q.PC4 AUC.PC9 q.PC9 AUC.PC14 q.PC14 AUC.PC30
##
             0.7734395 2.136524e-07
                                                                            NA
## LI.M11.0
                                           NA
                                                 NA
                                                           NA
                                                                  NA
## LI.M112.0 0.7509865 4.910746e-05
                                           NA
                                                 NA
                                                           NA
                                                                  NA
                                                                            NA
## LI.M118.0 0.8528591 5.027869e-05
                                                 NA
                                                                            NA
                                           NA
                                                           NA
                                                                  NA
## LI.M127
             0.9593030 3.706095e-03
                                           NA
                                                 NA
                                                           NA
                                                                  NA
                                                                            NA
## LI.M144
                     NA
                                   NA
                                                           NA
                                                                            NA
                                           NA
                                                 NA
                                                                  NA
##
             q.PC30
## LI.M11.0
                  NA
## LI.M112.0
                  NA
## LI.M118.0
                 NA
```

The following plot shows the same information in a visual form. The size of the blobs corresponds to the effect size (AUC value), and their color – to the q-value.

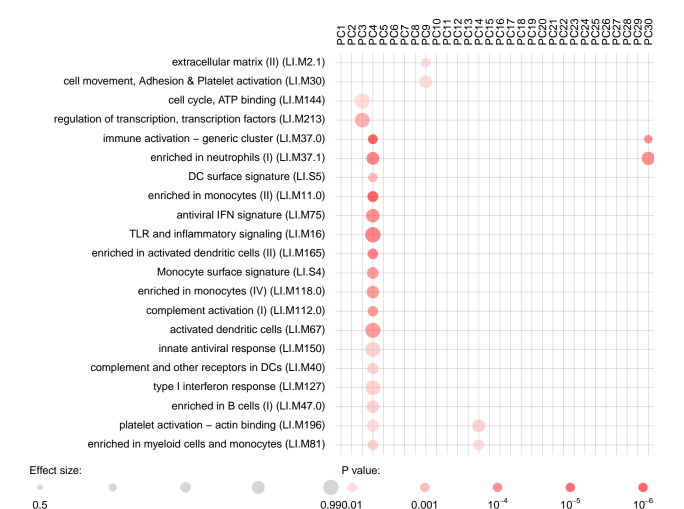
```
tmodPanelPlot(x)
```

NA

NA

LI.M127

LI.M144



However, we might want to ask, for each module, how many of the genes in that module have a negative, and how many have a positive weight? We can use the function tmodDecideTests for that. For each principal component shown, we want to know how many genes have very large (in absolute terms) weights – we can use the "lfc" parameter of tmodDecideTests for that. We define here "large" as being in the top 25% of all weights in the given component. For this, we need first to calculate the 3rd quartile (top 25% threshold). We will show only 10 components:

```
qfnc <- function(r) quantile(r, 0.75)
qqs <- apply(pca$rotation[,1:10], 2, qfnc)
pie <- tmodDecideTests(gs, lfc=pca$rotation[,1:10], lfc.thr=qqs)
tmodPanelPlot(x[1:10], pie=pie,
    pie.style="rug", grid="between")</pre>
```



PCA and tag clouds

For another way of visualizing enrichment, we can use the tagcloud package (Weiner 2014). P-Values will be represented by the size of the tags, while AUC – which is a proxy for the effect size – will be shown by the color of the tag, from grey (AUC=0.5, random) to black (1):

```
library(tagcloud)
```

```
## Loading required package: Rcpp
w <- -log10(res$P.Value)
c <- smoothPalette(res$AUC, min=0.5)
tags <- strmultline(res$Title)
tagcloud(tags, weights=w, col=c)</pre>
```

```
enriched in
               DC surface monocytes (IV)
    enriched in
    B cells (II)
                signature
                              enriched in
                             monocytes (II)
                                                    complement
          platelet activation
                                                    activation (I)
                                myeloid cell enriched
           - actin binding
                             receptors and transporters
                 TLR and inflammatory complement and other
  activated
dendritic cells
                        signaling
                                           receptors in DCs
                                                     Monocyte surface
      immune activation
                                                         signature
                                                  antiviral

    generic cluster

                                               IFN signature
                              enriched in activated
innate antiviral
                enriched in
                                dendritic cells (II)
```

innate antiviral response enriched in dendritic cells (II)

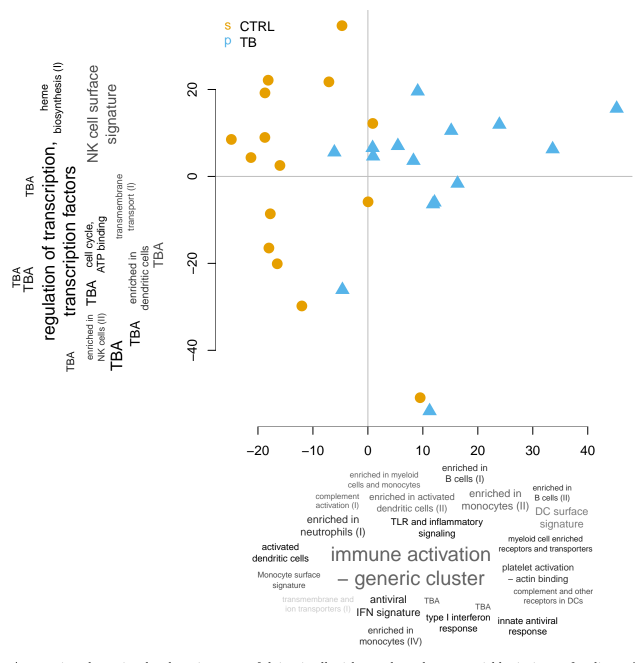
TBA neutrophils (I)

enriched in enriched in myeloid type I interferon

B cells (I) cells and monocytes response

We can now annotate the PCA axes using the tag clouds; however, see below for a shortcut in tmod.

```
par(mar=c(1,1,1,1))
o3 <- order(abs(pca$rotation[,3]), decreasing=TRUE)
13 <- Egambia$GENE_SYMBOL[o3]
res3 <- tmodUtest(13)
layout(matrix(c(3,1,0,2),2,2,byrow=TRUE),
  widths=c(1/3, 2/3), heights=c(2/3, 1/3))
# note -- PC4 is now x axis!!
1<-pca2d(pca, group=group, components=4:3,</pre>
  palette=mypal, radius=1.8)
cols <- as.character(1$colors)</pre>
legend("topleft",
  as.character(1$groups),
  pch=1$shapes,
  col=cols, bty="n")
tagcloud(tags, weights=w, col=c, fvert= 0)
tagcloud(strmultline(res3$Title),
  weights=-log10(res3$P.Value),
  col=smoothPalette(res3$AUC, min=0.5),
 fvert=1)
```



As mentioned previously, there is a way of doing it all with tmod much more quickly, in just a few lines of code:

Note that plot.params are just parameters which will be passed to the pca2d function. However, remember that is must be a list.

To plot the PCA, tmod uses the function pca2d from the pca3d package, but you can actually do it yourself by providing tmodPCA with a suitable function. The only requirement is that the function takes named parameters "pca" and "components":

```
plotf <- function(pca, components) {
  id1 <- components[1]
  id2 <- components[2]
  print(id1)
  print(id2)</pre>
```

```
plot(pca$x[,id1], pca$x[,id2])
}
     <- tmodPCA(pca, genes=Egambia$GENE_SYMBOL,</pre>
   components=3:4, plotfunc=plotf)
## [1] 3
## [1] 4
                                                                                       0
                               4
             type I interferor
                                                                                0
                               30
                                                                                    0
            TLR and inflammatory
                                                                                   0
                                                                                         0
                                                                                 0
                                                                               \infty
                                                                                    0
                                                                        0
                                                          0
                                                                                                    0
                                                                                0
                                                                                           0
                                                        0
                                                                                         00
                                                                               0
                                                                                  0
                                               -40
                                                             -20
                                                                            0
                                                                                         20
                                                  cell cycle,
                                                 ATP binding
                                               regulation of transcription,
                                                 transcription factors
                                                    and signaling
                                             phosphatidylinositol
                                              signaling system
```

Permutation tests

The GSEA approach (Subramanian et al. 2005) is based on similar premises as the other approaches described here. In principle, GSEA is a combination of an arbitrary scoring of a sorted list of genes and a permutation test. Although the GSEA approach has been criticized from statistical standpoint (Damian and Gorfine 2004), it remains one of the most popular tools to analyze gene sets amongst biologists. In the following, it will be shown how to use a permutation-based test with tmod.

A permutation test is based on a simple principle. The labels of observations (that is, their group assignments) are permutated and a statistic s_i is calculated for each *i*-th permutation. Then, the same statistic s_o is calculated for the original data set. The proportion of the permutated sets that yielded a statistic s_i equal to or higher than s_o is the p-value for a statistical hypothesis test.

First, we will set up a function that creates a permutation of the Egambia data set and repeats the limma procedure for this permutation, returning the ordering of the genes.

```
permset <- function(data, design) {
   require(limma)
   data <- data[, sample(1:ncol(data)) ]
   fit <- eBayes(lmFit(data, design))
   tt <- topTable(fit, coef=2, number=Inf, sort.by="n")
   order(tt$P.Value)
}</pre>
```

In the next step, we will generate 100 random permutations. The sapply function will return a matrix with a column for each permutation and a row for each gene. The values indicate the order of the genes in each permutation. We then use the tmod function tmodAUC to calculate the enrichment of each module for each permutation.

[1] 5547 250

We can now calculate the true values of the AUC for each module and compare them to the results of the permutation. The parameters "order.by" and "qval" ensure that we will calculate the values for all the modules (even those without any genes in our gene list!) and in the same order as in the perms variable.

```
fit <- eBayes(lmFit(E, design))
tt <- topTable(fit, coef=2, number=Inf,
   genelist=Egambia[,1:3])
res <- tmodCERNOtest(tt$GENE_SYMBOL, qval=Inf, order.by="n")
all(res$ID == rownames(perms))</pre>
```

```
## [1] TRUE
```

```
##
                                             Title
                                                         AUC
                                                                 adj.P.Val
## LI.M16
            LI.M16 TLR and inflammatory signaling 0.9790500 7.192190e-05
## LI.M59
            LI.M59
                       CCR1, 7 and cell signaling 0.9771973 5.751429e-02
## LI.M67
                        activated dendritic cells 0.9714730 8.363690e-05
            LI.M67
## LI.M150 LI.M150
                         innate antiviral response 0.9498859 9.956972e-03
## LI.M127 LI.M127
                       type I interferon response 0.9455715 1.163487e-02
## LI.S4
             LI.S4
                       Monocyte surface signature 0.8974252 1.852319e-06
##
           perm.P.Val.adj
## LI.M16
## LI.M59
                        Λ
## LI.M67
                        0
## LI.M150
                        0
```

```
## LI.M127 0
## LI.S4 0
```

Although the results are based on a small number of permutations, the results are nonetheless strikingly similar. For more permutations, they improve further. The table below is a result of calculating 100,000 permutations.

ID	Title	AUC	adj.P.Val
LI.M37.0	immune activation - generic cluster	0.7462103	0.00000
LI.M11.0	enriched in monocytes (II)	0.7766542	0.00000
LI.M112.0	complement activation (I)	0.8455773	0.00000
LI.M37.1	enriched in neutrophils (I)	0.8703781	0.00000
LI.M105	TBA	0.8949512	0.00000
LI.S4	Monocyte surface signature	0.8974252	0.00000
LI.M150	innate antiviral response	0.9498859	0.00000
LI.M67	activated dendritic cells	0.9714730	0.00000
LI.M16	TLR and inflammatory signaling	0.9790500	0.00000
LI.M118.0	enriched in monocytes (IV)	0.8774710	0.00295
LI.M75	antiviral IFN signature	0.8927741	0.00295
LI.M127	type I interferon response	0.9455715	0.00295
LI.S5	DC surface signature	0.6833387	0.02336
LI.M188	TBA	0.8684647	0.09894
LI.M165	enriched in activated dendritic cells (II)	0.7197180	0.11600
LI.M240	chromosome Y linked	0.8157171	0.11849
LI.M20	AP-1 transcription factor network	0.8763327	0.12672
LI.M81	enriched in myeloid cells and monocytes	0.7562851	0.13202
LI.M3	regulation of signal transduction	0.7763995	0.14872
LI.M4.3	$\label{eq:myeloid_cell_enriched_receptors} \ \ \text{and} \ \ transporters$	0.8859573	0.15675

Unfortunately, the permutation approach has two main drawbacks. Firstly, it requires a sufficient number of samples – for example, with three samples in each group there are only 6! = 720 possible permutations. Secondly, the computational load is substantial.

Accessing the tmod data

The tmod package stores its data in two data frames and two lists. This object is contained in a list called tmod, which is loaded with data("tmod"). The names mimick the various environments from Annotation.dbi packages, but currently the objects are just two lists and two data frames.

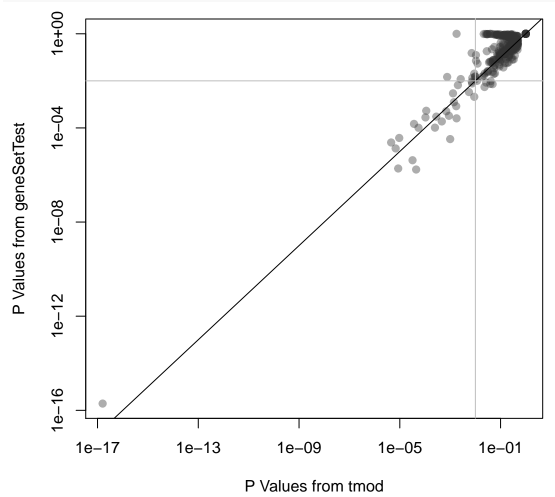
- tmod\$MODULES is a data frame which contains general module information as defined in the supplementary materials for Li et al. (S. Li et al. 2014) and Chaussabel et al. (Chaussabel et al. 2008)
- tmod\$GENES is a data frame which contains general gene information, including columns with HGNC ("primary"), as well as ENTREZ and REFSEQ identifiers.
- tmod\$MODULES2GENES is a list with module IDs (same as in the "ID" column of tmod\$MODULES) as names. Every element of the list is a character vector with IDs ("primary" column of tmod\$GENES) of the genes which are included in this module.
- tmod\$GENES2MODULES is a list with gene IDs (same as in the "primary" column of tmod\$GENES) as names. Every element of the list is a character vector with IDs of the modules in which the gene is found.

Using these variables, one can apply any other tool for the analysis of enriched module sets available, for example, the geneSetTest function from the limma package (Smyth et al. (Smyth 2005)). We will first run tmodUtest setting the qval to Inf to get p-values for all modules. Then, we apply the geneSetTest function to each module:

```
data(tmod)
res <- tmodUtest(tt$GENE_SYMBOL, qval=Inf)
gstest <- function(x) {
   sel <- tt$GENE_SYMBOL %in% tmod$MODULES2GENES[[x]]
   geneSetTest(sel, tt$logFC)
}
gst <- sapply(res$ID, gstest)</pre>
```

Are the results of both statistical approaches similar? tmod uses a very simple statistical test. The approach from geneSetTest is more complex, but similar in principle.

```
plot(res$P.Value, gst,
    log="xy", pch=19,
    col="#33333366",
    xlab="P Values from tmod",
    ylab="P Values from geneSetTest")
abline(0,1)
abline(h=0.01, col="grey")
abline(v=0.01, col="grey")
```



On the plot above, the p-values from tmod are plotted against the p-values from geneSetTest. As you can see, in this particular example, both methods give very similar results.

Using and creating custom sets of modules

It is possible to use any kind of arbitrary or custom gene set definitions. These custom definition of gene sets takes form of a list which is then provided as the mset parameter to the test functions. The list in question must have the following members:

- MODULES A data frame which contains at least the columns "ID" and "Title". The IDs must correspond to the names of MODULES2GENES.
- **GENES** (optional) A data frame which contains at least the column "ID". The gene IDs must correspond to the gene IDs used in MODULES2GENES.
- MODULES2GENES A list. The names of the list are the IDs from the MODULES data frame. The items in the list are character vectors with names of the genes that are associated with each module.
- GENES2MODULES (optional) A list with the reverse mapping from genes to modules. Names on that list must correspond to GENES\$ID, and the character vector members of the list must correspond to MODULES\$ID.

tmod will accept a simple list that contains the above fields. However, the function makeTmod can be used conveniently to create a tmod object.

Here is a minimal definition of such a set:

```
## An object of class "tmod"
## 2 modules, 4 genes
```

Both GENES and GENES2MODULES will be automatically created by makeTmod.

Whether the gene IDs are Entrez, or something else entirely does not matter, as long as they matched the provided input to the test functions.

MSigDB

The MSigDB database from the Broad institute is an interesting collection of gene sets (actually, multiple collections). Unfortunately, MSigDB cannot be distributed or even accessed without a free registration, which imposes a heavy limination on third party tools such as tmod. Use the following guide to download and parse the database such that you can use it with R and tmod.

First, you will need to download the MSigDB in XML format⁵. This file can be accessed at the URL http://www.broadinstitute.org/gsea/msigdb/download_file.jsp?filePath=/resources/msigdb/5.0/msigdb_v5.0.xml – follow the link, register and log in, and save the file on your disk (roughly 65MB).

Importing MSigDB is easy – tmod has a function specifically for that purpose. Once you have downloaded the MSigDB file, you can create the tmod-compatible R object with one command⁶. However, the tmod function tmodImportMsigDB() can also use this format, look up the manual page:

 $^{^{5}}$ Note that even if you register with MSig, it is not possible to download the database directly from R in the XML format.

⁶MSigDB gene sets can be also downloaded as "GMT" files. This format contains less information and is therefore less usable.

```
msig <- tmodImportMSigDB("msigdb_v5.0.xml")</pre>
msig
## An object of class "tmod"
     8430 modules, 32233 genes
That's it – now you can use the full MSigDB for enrichment tests:
res <- tmodCERNOtest(tt$GENE_SYMBOL, mset=msig )</pre>
head(res)
##
            ID
                                                          Title
## M3408 M3408
                         GSE1432 ctrl vs ifng 24h microglia dn
## M3010 M3010
                                          Hecker ifnb1 targets
## M3286 M3286
                  GSE13485 ctrl vs day3 yf17d vaccine pbmc dn
## M3288 M3288
                  GSE13485 ctrl vs day7 yf17d vaccine pbmc dn
## M3311 M3311 GSE13485 pre vs post yf17d vaccination pbmc dn
## M3347 M3347
                               GSE14000 unstim vs 4h lps dc dn
##
                            AUC
                                     cES
                                               P. Value
            cerno N1
## M3408 239.0983 39 0.8014227 3.065363 2.967858e-18
## M3010 244.1219 43 0.8459807 2.838626 4.555892e-17
## M3286 247.0915 45 0.7293732 2.745461 1.408943e-16
## M3288 272.2570 54 0.7222067 2.520898 3.626792e-16
## M3311 229.4948 41 0.7272625 2.798718 6.715323e-16
## M3347 272.0698 55 0.7334883 2.473362 9.792737e-16
##
            adj.P.Val
## M3408 2.501904e-14
## M3010 1.920308e-13
## M3286 3.959129e-13
## M3288 7.643464e-13
## M3311 1.132204e-12
## M3347 1.375880e-12
```

The results are quite typical for MSigDB, which is quite abundant with similar or overlapping gene sets. As the first results, we see, again, interferon response, as well as sets of genes which are significantly upregulated after yellow fever vaccination – and which are also interferon related. We might want to limit our analysis only to the 50 "hallmark" module categories:

```
sel <- msig$MODULES$Category == "H"
tmodCERNOtest(tt$GENE_SYMBOL, mset=msig[sel] )</pre>
```

```
Hallmark interferon gamma response
## M5913 M5913
## M5921 M5921
                                       Hallmark complement
## M5911 M5911
                       Hallmark interferon alpha response
## M5946 M5946
                                      Hallmark coagulation
## M5890 M5890
                         Hallmark tnfa signaling via nfkb
## M5930 M5930 Hallmark epithelial mesenchymal transition
## M5932 M5932
                           Hallmark inflammatory response
## M5953 M5953
                                Hallmark kras signaling up
## M5892 M5892
                         Hallmark cholesterol homeostasis
##
                            AUC
                                      cES
                                               P. Value
             cerno N1
## M5913 221.68317 41 0.7786936 2.703453 8.505170e-15
## M5921 217.81028 56 0.6979148 1.944735 8.607634e-09
## M5911 108.39559 20 0.7563566 2.709890 3.192325e-08
## M5946 179.24580 50 0.6779481 1.792458 1.966824e-06
```

```
## M5890 148.95123 47 0.6484665 1.584588 2.657694e-04
## M5930 212.53461 73 0.6371808 1.455717 2.701053e-04
## M5932 184.53035 62 0.6206393 1.488148 3.457724e-04
## M5953 221.76208 82 0.6046637 1.352208 1.790956e-03
         49.14641 14 0.6138968 1.755229 8.040562e-03
##
            adj.P.Val
## M5913 4.252585e-13
## M5921 2.151909e-07
## M5911 5.320542e-07
## M5946 2.458530e-05
## M5890 2.250878e-03
## M5930 2.250878e-03
## M5932 2.469803e-03
## M5953 1.119347e-02
## M5892 4.466979e-02
```

We see both – the prominent interferon response and the complement activation. Also, in addition, TNF- α signalling via NF- $\kappa\beta$.

Manual creation of tmod module objects: MSigDB

For the purposes of an example, the code below shows how to parse the XML MSigDB file using the R package XML. Essentially, this is the same code that tmodImportMsigDB is using:

```
library(XML)
foo <- xmlParse( "/home/january/Projects/R/pulemodule/vignette/msigdb_v5.0.xml" )
foo2 <- xmlToList(foo)</pre>
```

There are over 10,000 "gene sets" (equivalent to modules in tmod) defined. Each member of foo2 is a named character vector:

```
path1 <- foo2[[1]]
class(path1)</pre>
```

```
## [1] "character"
```

names(path1)

```
[1] "STANDARD_NAME"
                                "SYSTEMATIC_NAME"
    [3] "HISTORICAL_NAMES"
                                "ORGANISM"
##
##
    [5] "PMID"
                                "AUTHORS"
##
   [7] "GEOID"
                                "EXACT_SOURCE"
   [9] "GENESET_LISTING_URL"
                                "EXTERNAL_DETAILS_URL"
                                "CATEGORY_CODE"
## [11] "CHIP"
  [13]
       "SUB_CATEGORY_CODE"
                                "CONTRIBUTOR"
  [15]
       "CONTRIBUTOR_ORG"
                                "DESCRIPTION_BRIEF"
## [17]
       "DESCRIPTION_FULL"
                                "TAGS"
## [19] "MEMBERS"
                                "MEMBERS SYMBOLIZED"
##
  [21] "MEMBERS_EZID"
                                "MEMBERS_MAPPING"
  [23] "FOUNDER NAMES"
                                "REFINEMENT DATASETS"
## [25] "VALIDATION_DATASETS"
```

For our example analysis, we will use only human gene sets. We further need to make sure there are no NULLs in the list.

```
orgs <- sapply(foo2, function(x) x["ORGANISM"])
unique(orgs)</pre>
```

```
foo3 <- foo2[ orgs == "Homo sapiens" ]
foo3 <- foo3[ ! sapply(foo3, is.null) ]</pre>
```

Next, construct the MODULES data frame. We will use four named fields for each vector, which contain the ID (systematic name), description, category and subcategory:

```
modules <- t(sapply(foo3,
  function(x)
    x[ c("SYSTEMATIC_NAME", "STANDARD_NAME", "CATEGORY_CODE", "SUBCATEGORY_CODE") ]))
colnames(modules) <- c( "ID", "Title", "Category", "Subcategory" )
modules <- data.frame(modules, stringsAsFactors=FALSE)</pre>
```

Then, we create the modules to genes mapping and the GENES data frame. For this, we use the MEMBERS_SYMBOLIZED field, which is a comma separated list of gene symbols belonging to a particular module:

```
m2g <- lapply(foo3,
  function(x) strsplit( x["MEMBERS_SYMBOLIZED"], "," )[[1]])
names(m2g) <- modules$ID

msig <- makeTmod(modules=modules, modules2genes=m2g)</pre>
```

From now on, you can use msig with tmod.

Note that it is not necessary to create the members GENES and GENES2MODULES manually. The reverse mapping from genes to modules, GENES2MODULES, will be automatically inferred from MODULES2GENES. If no meta-information on genes is provided in GENES, then a minimal data frame will be created with one column only (ID).

Manual creation of tmod module sets: Wikipathways

Below is an example of how to use the pathway definitions from WikiPathways 7 . First, we download the data (human pathways) and clean it up:

Since each pathway is in a separate file in the zip archive we downloaded, we have to read each file separately. Below, we create a list, p2GENES, which maps the modules to the corresponding genes. To make it consistent, I decided to use gene symbols rather than the Entrez numbers (just because it makes the interpretation of results a bit easier), but actually that is not necessary: tmod does not care what gene symbols are used, as long as the mappings between genes and modules are consistent, and as long as the same identifiers are used in the lists of genes.

Furthermore, note that we filter out anything that is not an ENTREZ gene identifier. This gets rid of entities which are not genes (e.g. biochemical compounds), but also of some genes.

⁷http://www.wikipathways.org/

```
suppressMessages(library(org.Hs.eg.db))
p2GENES <- sapply( files$Name, function(fn) {
  foo <- read.csv( unz( human,
   filename= fn ), sep="\t" )
  ids <- foo$Identifier[ foo$Identifier %in% ls( org.Hs.egSYMBOL ) ]
  unique(unlist(mget(as.character(ids), org.Hs.egSYMBOL)))
})
names(p2GENES) <- files$ID</pre>
```

p2GENES is the first of three objects that we need to create. The next one is a data frame containing module definitions. We also calculate the number of associated genes and select pathways that have at least 5 associated ENTREZ genes:

```
pathways <- data.frame( ID=files$ID,
    Title=files$Title,
    stringsAsFactors=FALSE )
pathways$N <- sapply(p2GENES, length)
pathways$URL <-
    paste0("http://www.wikipathways.org/index.php/Pathway:",
    pathways$ID )
sel <- pathways$N > 4
pathways <- pathways [ sel, ]
rownames(pathways) <- pathways$ID</pre>
```

Finally, we are good to go: we can build the tmod object with the makeTmod function:

```
Hspaths <- makeTmod(modules=pathways, modules2genes=p2GENES)</pre>
```

We can now use the tmodCERNOtest to see whether it works:

```
tmodCERNOtest(tt$GENE_SYMBOL, mset=Hspaths)
```

```
## ID Title cerno N1 AUC
## WP558 WP558 Hs_Complement_and_Coagulation_Cascades 107.73082 28 0.6418746
## WP545 WP545 Hs_Complement_Activation 55.50123 11 0.7384877
## cES P.Value adj.P.Val
## WP558 1.923765 4.008176e-05 0.008938232
## WP545 2.522783 1.007705e-04 0.011235913
```

Nice – the complement pathway was also found before, when using the default data set. Unfortunately, we don't see anything else: WikiPathways are more oriented on metabolic pathways, while the blood transcriptional modules are particularly good for analyzing immune responses. However, if we were to test a specific hypothesis, we would select modules related to interferon response:

```
sel <- grep( "Interferon",
   Hspaths$MODULES$Title, ignore.case=T )
tmodCERNOtest(tt$GENE_SYMBOL, mset=Hspaths,
   modules=Hspaths$MODULES$ID[sel])</pre>
```

```
## ID Title cerno N1 AUC
## WP619 WP619 Hs_Type_II_interferon_signaling_(IFNG) 42.1566 9 0.7050031
## cES P.Value adj.P.Val
## WP619 2.342033 0.001051527 0.003154582
```

Since the number of tests is lower, the type-II interferon signalling is now significant.

Case study: Metabolic profiling of TB patients

Introduction

One of the main objectives in writing tmod was the ability to analyse metabolic profiling data and other uncommon data sets. In 2012, we have analysed metabolic profiles of serum collected from patients suffering from tuberculosis (TB) and healthy controls (Weiner 3rd et al. 2012). It turned out that there are huge differences between these two groups of individuals, involving amino acid metabolism, lipid metabolism and many others. In the course of the analysis, we found correlations between the metabolites which are not explained fully by the metabolic pathways. For example, cortisol is correlated with kynurenine due to the immunoactive function of these molecules indicating an activation of the immune system, and not because these two molecules are linked by a synthesis process. Vice versa, kynurenine and tryptophan were not directly correlated, even though these molecules are clearly linked by a metabolic process, because tryptophan is not an immune signalling molecule, while kynurenine is.

The tmod package includes both, the data set used in the Weiner et al. paper and the cluster definitions (modules) published therein. In the following, we will use these modules to analyse the metabolic profiles⁸.

First, we load the data modules and the data set to analyse.

```
data(modmetabo) ## modules
data(tbmprof)
ids <- rownames(tbmprof)
tb <- factor(gsub("\\..*", "", ids))
sex <- factor( gsub( ".*\\.([MF])\\..*", "\\1", ids))
table(tb, sex)

## sex
## tb F M
## HEALTHY 58 34
## TB 25 19</pre>
```

Differential analysis

The metabolic profiling data has not exactly a normal distribution, but that varies from one compound to another. It is possible to normalize it by ranking, but we can simply use the wilcoxon test to see differences between males and females as well as TB patients and healthy individuals.

The data frame contains the results of all tests. We can now test both the healthy/tb comparison and the male/female comparison for enrichment in metabolic profiling modules. Instead ordering the feature identifiers, we use the option "input.order" to determine the sorting.

⁸Formally, this is not correct, as the modules were derived from the data set that we are going to analyse, however it serves for demonstration purposes

```
ids <- wcx$ID
res <- list()
res$tb <- tmodCERNOtest(ids[order(wcx$pval.tb)], mset=modmetabo)</pre>
##
              ID
                                                             Title
                                                                       cerno N1
## ME.107 ME.107
                                              Amino acids cluster 104.64470 18
## ME.37
           ME.37 Kynurenines, taurocholates and cortisol cluster 116.88147 25
## MP.2
            MP.2
                                                       Amino Acid 99.16212 28
                                              adj.P.Val
                AUC
                         cES
                                   P.Value
## ME.107 0.8824576 2.906797 1.283792e-08 5.391925e-07
## ME.37 0.8837093 2.337629 2.816098e-07 5.913805e-06
          0.7060786 1.770752 3.357995e-04 4.701193e-03
res$sex <- tmodCERNOtest(ids[order(wcx$pval.sex)], mset=modmetabo)
res$sex
##
            ID
                             Title
                                       cerno N1
                                                      AUC
                                                                cES
                  Hormones cluster 62.47927 10 0.9198068 3.123963
## ME.26 ME.26
                           Steroid 60.98935 11 0.8732115 2.772243
## MS.1
          MS.1
## ME.69 ME.69 Cholesterol cluster 45.14144 11 0.8190623 2.051883
##
              P. Value
                         adj.P.Val
## ME.26 2.923803e-06 0.0001227997
## MS.1 1.593507e-05 0.0003346365
## ME.69 2.546330e-03 0.0356486262
```

Both these result tables are concordant with previous findings. The enriched modules in male vs. female comparison are what one would expect. In TB, a cluster consisting of kynurenine, bile acids and cortisol is up-regulated, while amino acids go down. We can take a closer look at it using the evidencePlot function.

Why is there a module called "Amino acid cluster" and another one called "Amino acid"? The "cluster" in the name of the module indicates that it has been build by clustering of the profiles, while the other module has been based on the biochemical classification of the molecules. This information is contained in the Category column of the MODULES data frame:

```
modmetabo$MODULES[ c("ME.107", "MP.2"), ]

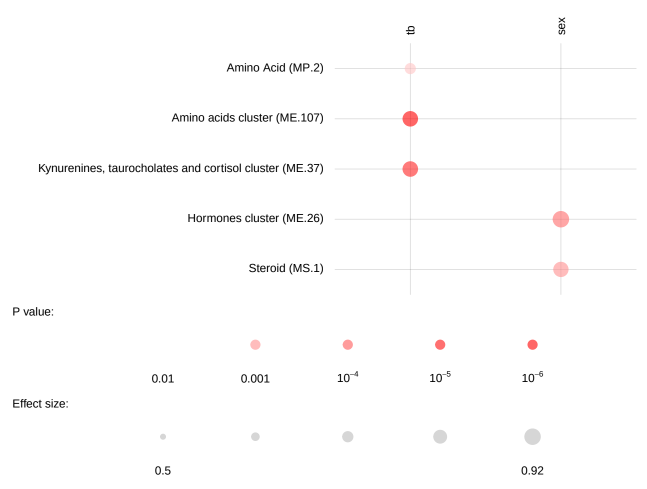
## ID Title Category

## ME.107 ME.107 Amino acids cluster Cluster

## MP.2 MP.2 Amino Acid Pathway
```

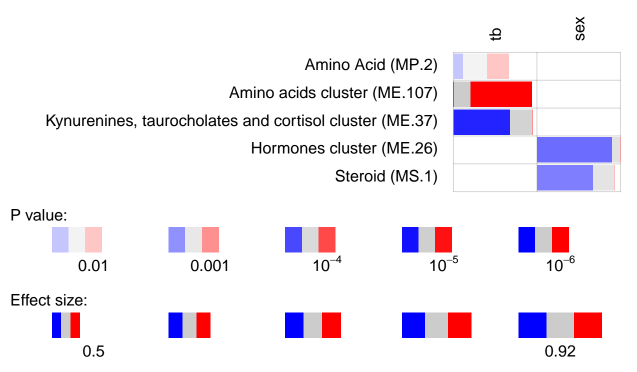
To get an overview for both of these comparisons at the same time, we can use the tmodPanelPlot function. The size of the blobs below corresponds to the AUC values from the tables above.

```
tmodPanelPlot(res)
```



This, unfortunately, does not tell us in which group the metabolites from a given modules are higher. For this, we can use the "estimate" from the wilcox.test above and a parameter for tmodPanelPlot called "pie". To create the value for this parameter – a list that describes, for each condition and for each module, how many metabolites change in one direction, and how many change in the other.

```
pie.data <- wcx[,c("E.sex", "E.tb")]
colnames(pie.data) <- c("sex", "tb")
pie <- tmodDecideTests(wcx$ID, lfc=pie.data, lfc.thr=0.2, mset=modmetabo)
tmodPanelPlot(res, pie=pie, pie.style="rug", grid="between")</pre>
```



We see now that the cortisol cluster is higher in TB, while amino acids are found at lower concentration in the patients. Also, we see that most of the steroids found (cluster ME.26 and module MS.1) are lower in females. The latter is not surprising if we inspect it closely.

```
wcx <- wcx[order(wcx$pval.sex),]</pre>
showModule(wcx[,c("E.sex", "pval.sex")], wcx$ID, "MS.1", mset=modmetabo)
##
                                pval.sex
                                                   ID
                     E.sex
               -0.86999795 3.042745e-06
## HMDB00493
                                           HMDB00493
## HMDB00365
               -0.64001269 4.033730e-05
                                           HMDB00365
## HMDB02759
               -0.61997366 1.072607e-04
                                           HMDB02759
               -0.49999995 1.486517e-04
                                             M.37186
## M.37186
## HMDB03818.1 -0.38999920 1.541563e-04 HMDB03818.1
## M.32619
               -0.35995962 3.418813e-04
                                              M.32619
## HMDB03818
               -0.45997259 4.349672e-03
                                            HMDB03818
## HMDB01032
               -0.27006303 5.284132e-03
                                            HMDB01032
## HMDB02802
               -0.10000226 8.852221e-02
                                            HMDB02802
## HMDB00063
               -0.11997203 1.552120e-01
                                            HMDB00063
  HMDB04026
               -0.07998115 3.350142e-01
                                            HMDB04026
##
##
                                                                 Name Pathway
## HMDB00493
                       5alpha-androstan-3beta,17beta-diol disulfate
                                                                        Lipid
                                             epiandrosterone sulfate
                                                                        Lipid
## HMDB00365
## HMDB02759
                                                androsterone sulfate
                                                                        Lipid
               5alpha-androstan-3alpha,17beta-diol monosulfate (1)
                                                                        Lipid
## M.37186
## HMDB03818.1
                        4-androsten-3beta,17beta-diol disulfate (2)
                                                                        Lipid
## M.32619
                                          pregn steroid monosulfate*
                                                                        Lipid
## HMDB03818
                        4-androsten-3beta,17beta-diol disulfate (1)
                                                                        Lipid
                            dehydroisoandrosterone sulfate (DHEA-S)
## HMDB01032
                                                                        Lipid
## HMDB02802
                                                                        Lipid
                                                           cortisone
## HMDB00063
                                                            cortisol
                                                                        Lipid
## HMDB04026
                                   21-hydroxypregnenolone disulfate
                                                                        Lipid
##
               Subpathway
                                HMDB
                                       KEGG MetabolonID
                  Steroid HMDB00493 C12525
                                                 M.37190
## HMDB00493
```

```
## HMDB00365
                  Steroid HMDB00365 C07635
                                                M.33973
## HMDB02759
                  Steroid HMDB02759
                                                M.31591
## M.37186
                  Steroid
                                                M.37186
## HMDB03818.1
                  Steroid HMDB03818 C04295
                                                M.37203
## M.32619
                  Steroid
                                                M.32619
                  Steroid HMDB03818 C04295
## HMDB03818
                                                M.37202
## HMDB01032
                  Steroid HMDB01032 C04555
                                                M.32425
## HMDB02802
                  Steroid HMDB02802 C00762
                                                 M.1769
## HMDB00063
                  Steroid HMDB00063 C00735
                                                 M.1712
                  Steroid HMDB04026 C05485
## HMDB04026
                                                M.46115
i <- "HMDB00493" # what is it?
modmetabo$GENES[i,]
##
                    ID
                                                                 Name Pathway
## HMDB00493 HMDB00493 5alpha-androstan-3beta,17beta-diol disulfate
                                                                        Lipid
##
             Subpathway
                             HMDB
                                     KEGG MetabolonID
                                              M.37190
                Steroid HMDB00493 C12525
## HMDB00493
par(mfrow=c(1,2))
showGene(tbmprof[,i], sex, main=modmetabo$GENES[i, "Name"],
  ylab="Relative abundance")
## now for cortisol cluster
i <- "HMDB00063"
wcx <- wcx[order(wcx$pval.tb),]</pre>
showModule(wcx[,c("E.tb", "pval.tb")], wcx$ID, "ME.37",
  mset=modmetabo)[1:10,] # only first 10!
##
                      E.tb
                                 pval.tb
                                                ID
                                                                       Name
## M.47908
             -6.999310e-01 2.665735e-14
                                           M.47908
                                                                    Unknown
## M.32599
             -7.999872e-01 2.320130e-10
                                           M.32599 glycocholenate sulfate*
## HMDB00169 -6.299770e-01 5.118925e-09 HMDB00169
                                                                    mannose
## Mx.22110 -6.448828e-05 1.379682e-08 Mx.22110
                                                       3-hydroxykynurenine
## HMDB00063 -5.399765e-01 1.990892e-08 HMDB00063
                                                                   cortisol
## HMDB00159 -2.900586e-01 2.491091e-08 HMDB00159
                                                              phenylalanine
## M.32807
             -1.219964e+00 3.577292e-08
                                           M.32807
                                                    taurocholenate sulfate
## M.46637
             -1.030041e+00 6.660175e-08
                                           M.46637
                                                                    Unknown
## M.46652
             -8.399503e-01 1.420077e-07
                                                                    Unknown
                                           M.46652
## HMDB00684 -3.100146e-01 1.788895e-07 HMDB00684
                                                                 kynurenine
                  Pathway
                                                            Subpathway
## M.47908
## M.32599
                    Lipid
                                       Secondary Bile Acid Metabolism
## HMDB00169 Carbohydrate Fructose, Mannose and Galactose Metabolism
## Mx.22110
               Amino acid
                                                Tryptophan Metabolism
## HMDB00063
                    Lipid
                                                               Steroid
## HMDB00159
               Amino Acid
                                Phenylalanine and Tyrosine Metabolism
## M.32807
                                       Secondary Bile Acid Metabolism
                    Lipid
## M.46637
## M.46652
## HMDB00684
               Amino Acid
                                                Tryptophan Metabolism
                         KEGG MetabolonID
##
                  HMDB
## M.47908
                                   M.47908
## M.32599
                                   M.32599
## HMDB00169 HMDB00169 C00159
                                     M.584
```

```
## Mx.22110
                           C02794
                                       Mx.22110
## HMDB00063 HMDB00063 C00735
                                          M.1712
## HMDB00159 HMDB00159 C00079
                                            M.64
## M.32807
                                        M.32807
## M.46637
                                        M.46637
## M.46652
                                        M.46652
## HMDB00684 HMDB00684 C00328
                                        M.15140
showGene(tbmprof[,i], tb, main=modmetabo$GENES[i, "Name"],
  ylab="Relative abundance")
                                                                                     cortisol
          5alpha-androstan-3beta,17beta-diol disulfate
                                                             3.0
    15
                                                             2.5
                                                          Relative abundance
Relative abundance
    10
                                                             2.0
```

1.5

1.0

0.5

ΤB

Functional multivariate analysis

ш

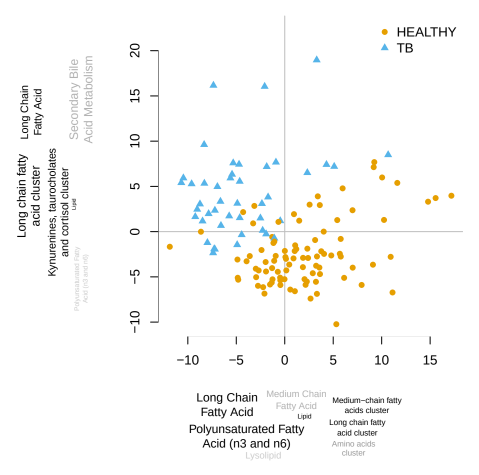
5

We can practically circumvent a gene-by-gene analysis. In fact, we are rarely interested in the p-values associated with single genes or metabolites. There is too many of them, and the statistical power is limited by the sheer number of tests and the requirement of correction for multiple testing. In case you have not read the part on FMA above, "Functional multivariate analysis", in its simplest form, is simply combining a principal component analysis (PCA) with enrichment analysis. PCA lets us explore where the variance in the data is; enrichment analysis allows us to interprete the principal components in functional terms.

Σ

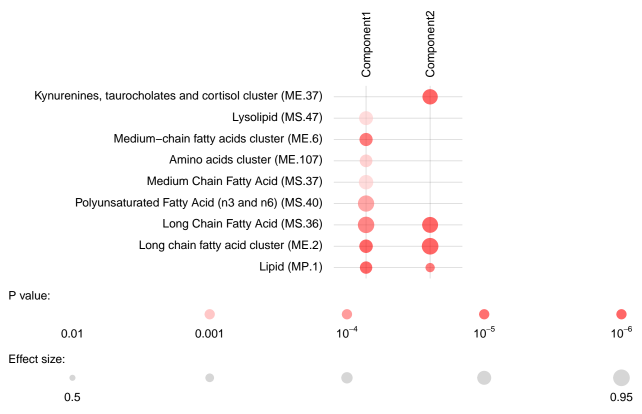
In tmod, it can be done in a few lines of code:

```
pca <- prcomp(tbmprof, scale.=T)</pre>
ret <- tmodPCA(pca, genes=colnames(tbmprof), mset=modmetabo,</pre>
  plot.params=list(group=tb, legend="topright"))
```



The ret object now contains the results of enrichments (in the ret\$enrichments member) and we can directly throw it on a panel plot:

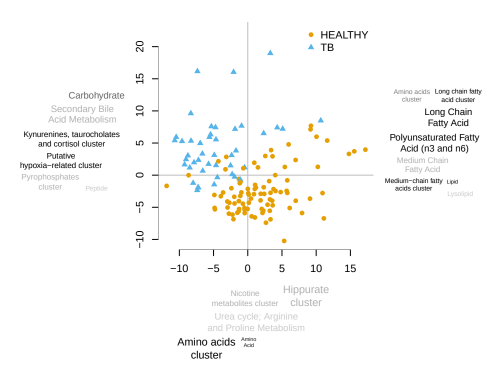
tmodPanelPlot(ret\$enrichments)



OK, but which of the terms are characteristic for TB patients? Which for the healthy controls? In the above, the enrichments were based on a list sorted by the absolute PCA weights. However, we can split it into a list ordered by signed weights ordered once from small to large values, and once from large to small values.

```
pca <- prcomp(tbmprof, scale.=T)
ret <- tmodPCA(pca, genes=colnames(tbmprof), mset=modmetabo,
    plot.params=list(group=tb, legend="topright"),
    mode="cross")</pre>
```





In essence, reading this plot is simple. First, note that this time the tag clouds on the top and the bottom correspond to the two ends of the vertical, y axis (second component); and the tag clouds at the left and right correspond to the two ends of the horizontal, x axis (first PCA component).

Now, take the amino acid cluster (bottom of the plot): it is enriched at the lower end of the y axis, which means, that features in that cluster are higher in the yellow points which are at the bottom of the plot (lower end of the y). In other words, amino acids are higher in healthy persons – a finding which corroborates the differential analysis above.

Similarly, "kynurenines" are at the left, lower side of the x axis, which means, that features from this cluster are at higher levels in TB patients.

What about the male-female differences? They probably can be found in other, less important 9 components. We could look for them manually, but we can also search which of the responses (turned to orthogonal PCA components) is best predicted by the sex factor.

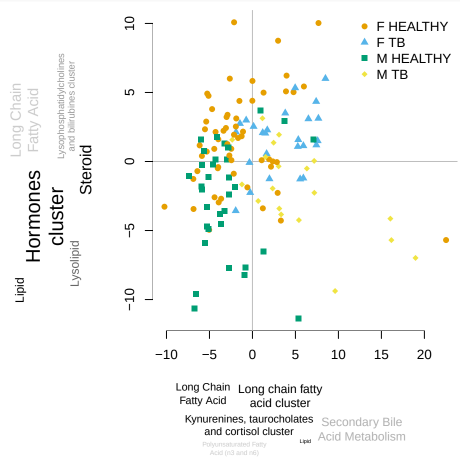
```
foo <- summary(lm(pca$x ~ sex))</pre>
foo <- t(sapply(foo,
  function(x) c(r=x$r.squared, pval=x$coefficients[2,4])))
head(foo[ order(foo[,2]), ])
##
                                      pval
                            r
## Response PC5
                  0.24569651 8.485092e-10
## Response PC10
                  0.21456193 1.359253e-08
## Response PC7
                   0.03280924 3.482826e-02
## Response PC8
                   0.02213252 8.388780e-02
## Response PC107 0.01986978 1.016566e-01
```

⁹That is, components which include a smaller fraction of the total variance in the data set

Response PC6 0.01919507 1.077110e-01

We can use the components 1 (which corresponds to TB/healthy) and components 5, which corresponds to male/female differences, as suggested by the above calculations.

```
ret <- tmodPCA(pca, genes=colnames(tbmprof), mset=modmetabo,
  plot.params=list(group=paste(sex, tb), legend="topright"),
  components=c(2,5))</pre>
```



Orange circles and blue triangles are females, located mostly in Q1 and Q2 (top half); this corresponds to differences on the y axis and the tagcloud next to it (hormone cluster, steroids etc.). On the other hand, TB patients (blue triangles and yellow circles) are in Q1 and Q4 (right-hand side), which corresponds to the TB-specific tag cloud below the y axis.

References

Banchereau, Romain, Alejandro Jordan-Villegas, Monica Ardura, Asuncion Mejias, Nicole Baldwin, Hui Xu, Elizabeth Saye, et al. 2012. "Host Immune Transcriptional Profiles Reflect the Variability in Clinical Disease Manifestations in Patients with Staphylococcus Aureus Infections." *PLoS One* 7 (4). Public Library of Science: e34390.

Chaussabel, Damien, Charles Quinn, Jing Shen, Pinakeen Patel, Casey Glaser, Nicole Baldwin, Dorothee Stichweh, et al. 2008. "A Modular Analysis Framework for Blood Genomics Studies: Application to Systemic Lupus Erythematosus." *Immunity* 29 (1). Elsevier: 150–64.

Damian, Doris, and Malka Gorfine. 2004. "Statistical Concerns About the Gsea Procedure." Nature Genetics

36 (7). Nature Publishing Group: 663-63.

Li, Shuzhao, Nadine Rouphael, Sai Duraisingham, Sandra Romero-Steiner, Scott Presnell, Carl Davis, Daniel S Schmidt, et al. 2014. "Molecular Signatures of Antibody Responses Derived from a Systems Biology Study of Five Human Vaccines." *Nature Immunology* 15 (2). Nature Publishing Group: 195–204.

Maertzdorf, Jeroen, Martin Ota, Dirk Repsilber, Hans J Mollenkopf, January Weiner, Philip C Hill, and Stefan HE Kaufmann. 2011. "Functional Correlations of Pathogenesis-Driven Gene Expression Signatures in Tuberculosis." *PloS One* 6 (10). Public Library of Science: e26938.

Smyth, Gordon K. 2005. "Limma: Linear Models for Microarray Data." In *Bioinformatics and Computational Biology Solutions Using R and Bioconductor*, edited by R. Gentleman, V. Carey, S. Dudoit, R. Irizarry, and W. Huber, 397–420. New York: Springer.

Subramanian, Aravind, Pablo Tamayo, Vamsi K Mootha, Sayan Mukherjee, Benjamin L Ebert, Michael A Gillette, Amanda Paulovich, et al. 2005. "Gene Set Enrichment Analysis: A Knowledge-Based Approach for Interpreting Genome-Wide Expression Profiles." *Proceedings of the National Academy of Sciences of the United States of America* 102 (43). National Acad Sciences: 15545–50.

Weiner 3rd, January, Shreemanta K Parida, Jeroen Maertzdorf, Gillian F Black, Dirk Repsilber, Anna Telaar, Robert P Mohney, et al. 2012. "Biomarkers of Inflammation, Immunosuppression and Stress with Active Disease Are Revealed by Metabolomic Profiling of Tuberculosis Patients." *PloS One* 7 (7). Public Library of Science: e40221.

Weiner, January. 2013. Pca3d: Three Dimensional Pca Plots.

——. 2014. Tagcloud: Tag Clouds.

Wendt, Hans W. 1972. "Dealing with a Common Problem in Social Science: A Simplified Rank-Biserial Coefficient of Correlation Based on the U Statistic." *European Journal of Social Psychology* 2 (4). Wiley Online Library: 463–65.

Yamaguchi, Ken D, Daniel L Ruderman, Ed Croze, T Charis Wagner, Sharlene Velichko, Anthony T Reder, and Hugh Salamon. 2008. "IFN- β -Regulated Genes Show Abnormal Expression in Therapy-Naïve Relapsing-remitting Ms Mononuclear Cells: Gene Expression Analysis Employing All Reported Protein-protein Interactions." Journal of Neuroimmunology 195 (1). Elsevier: 116–20.