# NetWeAvers Vignette

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### June 24, 2013

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# 1 Introduction

This document describes the functions available in the **netweavers** package. The package may be used with proteomics mass spectrometry data to summarize quantified peptide data, fit linear models and test for differential expression, and perform network analysis on protein-protein interaction networks using protein p-values. To get started, first you need to load the package in your R session:

#### > library(netweavers)

We will use the example dataset called vanHoof, which contains mass spectrometry (MS) data from a phosphorylation study of human embryonic stem cells (hESCs), (Van Hoof

et al., 2009). The SILAC experiment measured undifferentiated hESCs and hESCs differentiated with bone morphogenetic protein 4 (BMP4) at three time points (30 minutes, 60 minutes and 240 minutes) with two biological replicates at each time point. The data were processed using PVIEW (Khan et al., 2009, 2011).

The dataset that contains the main objects required for use in this vignette is loaded by

> data(vanHoof)

# 2 Differential Expression Test

The function DEtest uses functions from the package **limma** for hypothesis testing via linear models on quantified protein- or peptide-level data. The input requires the data to be in the form of an ExpressionSet.

We will use ExpSetVH to test for differential expression at the peptide level. This ExpressionSet contains the quantified data in the assayData slot, the sequences and protein IDs in the featureData slot, and the experimental design, phenoDataVH, in the phenoData slot:

#### > ExpSetVH

```
ExpressionSet (storageMode: lockedEnvironment)
assayData: 12524 features, 6 samples
element names: exprs
protocolData: none
phenoData
sampleNames: a30min b30min ... b240min (6 total)
varLabels: sampleIDs groups
varMetadata: names labelDescription
featureData
featureNames: 1 2 ... 13096 (12524 total)
fvarLabels: Sequence Protein
fvarMetadata: labelDescription
experimentData: use 'experimentData(object)'
Annotation:
```

Note that the phenoData contains the label groups that indicates which samples belong to each of the three factor levels (time points) of the experiment:

```
> phenoData(ExpSetVH)@data
```

```
sampleIDs groups a30min min30
```

```
b30min b30min min30
a60min a60min min60
b60min b60min min60
a240min a240min min240
b240min b240min min240
```

Note that the ExpressionSet only contains 6 samples (and not 12) because with SILAC one measures the ratio of abundances of differentiated versus undifferentiated hESCs.

The function DEtest takes as input the ExpressionSet and the name of the variable subject to testing:

```
> outDE <- DEtest(ExpSetVH, label="groups")
groups (F-test)
Warning: nestedF multiple testing scheme not carried out due to missing p-values
groups (pairwise comparison): min240 - min30
groups (pairwise comparison): min240 - min60
groups (pairwise comparison): min30 - min60</pre>
```

If there are technical replicates in the data, set reps = TRUE. Note the warning: the function decideTests() with method = "nestedF" is used for multiple testing across the contrasts and proteins (or peptides); it requires complete data (see the limma User's Guide for more details). One option is to filter out peptides with too many missing values and impute the remaining NAs. The resulting object is a list containing two items:

- 1. A list of data frames comprised of the output of topTable applied to the eBayes model fit.
- 2. A data frame containing (moderated) statistics from an F-test.

#### > str(outDE)

```
List of 2
```

```
.. ..$ B
                 : num [1:12524] 0.944 0.926 0.879 0.837 0.528 ...
 .. ..$ df
                 : num [1:12524] 1 0 0 0 0 0 0 0 0 0 ...
 ..$ min240 - min60: 'data.frame':
                                         12524 obs. of 9 variables:
 ....$ Sequence : chr [1:12524] "SLDQEIARPLENENQEFLK" "VMLGETNPADSKPGTIR" "MVMIQDGPLF
 ....$ Protein : chr [1:12524] "NES" "NME1" "FUBP3" "SMARCC1" ...
 .. ..$ logFC
                 : num [1:12524] -274.7 -95.2 62.7 54.5 -37.2 ...
 ....$ AveExpr : num [1:12524] 91.6 46.3 21 32.3 13.5 ...
                : num [1:12524] -2106 -730 481 417 -285 ...
   ..$ P.Value : num [1:12524] 5.52e-07 4.02e-06 8.77e-06 1.14e-05 2.34e-05 ...
 ....$ adj.P.Val: num [1:12524] 0.00119 0.00432 0.00615 0.00615 0.01005 ...
                 : num [1:12524] 1.22 1.22 1.21 1.21 1.2 ...
                 : num [1:12524] 1 0 0 0 0 0 0 0 0 0 ...
 .. ..$ df
 ..$ min30 - min60 :'data.frame':
                                         12524 obs. of 9 variables:
 ....$ Sequence : chr [1:12524] "SLDQEIARPLENENQEFLK" "VMLGETNPADSKPGTIR" "EKPVDLQNFG
 ....$ Protein : chr [1:12524] "NES" "NME1" "SMARCC1" "AKAP8L" ...
                 : num [1:12524] -274.74 -79.75 42.31 -37.29 -4.02 ...
 .. ..$ logFC
 .. ..$ AveExpr
               : num [1:12524] 91.62 46.34 32.28 13.51 2.86 ...
 .. ..$ t
                 : num [1:12524] -2106 -611.3 324.3 -285.9 -23.4 ...
 ....$ P.Value : num [1:12524] 5.52e-07 5.60e-06 1.83e-05 2.32e-05 2.57e-05 ...
 ....$ adj.P.Val: num [1:12524] 0.00119 0.00602 0.01106 0.01106 0.01106 ...
                : num [1:12524] 1.19 1.19 1.18 1.18 3.77 ...
 .. ..$ df
                 : num [1:12524] 1 0 0 0 0 0 0 0 0 0 ...
$ resultsTable_Fpvalue:'data.frame':
                                            12524 obs. of 7 variables:
..$ Sequence : chr [1:12524] "SLDQEIARPLENENQEFLK" "VMLGETNPADSKPGTIR" "MVMIQDGPLPTGA
..$ Protein : chr [1:12524] "NES" "NME1" "FUBP3" "SMARCC1" ...
             : num [1:12524] 2956842 306746 154265 96028 54321 ...
..$ df1
              : int [1:12524] 2 2 2 2 2 2 2 2 2 2 ...
              : num [1:12524] 0 0 0 0 0 2 0 3 2 0 ...
 ..$ df2
..$ p.Val
             : num [1:12524] 8.23e-07 6.86e-06 1.31e-05 2.04e-05 3.47e-05 ...
 ..$ adj.P.Val: num [1:12524] 0.00177 0.00738 0.00937 0.01095 0.01493 ...
```

The data frames contain any original annotation and several statistics output by the function topTable including p-values and adjusted p-values. These p-values can be used as input in runNetweavers, e.g. for comparing timepoints 30 and 60 minutes you could use

> outDE\_30to60 <- outDE\$resultsTable\_limma[["min30 - min60"]]</pre>

# 3 Peptide to Protein Summarization

If your quantified MS data are at the peptide level, you will need to summarize the data to the protein level in order to do network analysis.

#### 3.1 Data Frame Summarization

One way to roll-up peptides to proteins is to input a data frame of the quantified data, along with the sequences, protein IDs and experimental design, into the function customSummarizer. The input data look like:

#### > head(dataVH)

		Sequence	Protein	a30min	b30min	a60min	b60min
1		AAAAAALQAK	RPL4	0.9390492	0.9430509	NA	NA
2		AAAAAWEEPSSGNGTAF	RCC2	NA	NA	NA	NA
3	AAAGLGHP#	SPGGSEDGPPGSEEEDAAF	ARID3A	NA	NA	NA	NA
4		AAAGEFADDPCSSVK	CTNNA1	0.8568542	NA	NA	NA
5	I	AAGQESEGPAVGPPQPLGK	ARID1A	1.8687121	NA	2.272664	NA
6	AAAI	.VDEGLDPEEHTADGEPSAK	TSHZ3	NA	0.4784046	NA	NA
	a240min	b240min					
1	0.9122702	NA					
2	0.4053151	NA					
3	NA	0.4606221					
4	0.9437617	NA					
5	1.7914869	NA					

### > phenoDataVH

```
      sampleIDs
      groups

      a30min
      a30min
      min30

      b30min
      b30min
      min30

      a60min
      a60min
      min60

      b60min
      b60min
      min60

      a240min
      a240min
      min240

      b240min
      b240min
      min240
```

NA 0.4516751

and the following shows how to specify the parameters in the function to get the desired output:

ExpressionSet (storageMode: lockedEnvironment)

assayData: 3200 features, 6 samples

element names: exprs

protocolData: none

phenoData

sampleNames: a30min b30min ... b240min (6 total)

varLabels: sampleIDs groups

varMetadata: names labelDescription

featureData

featureNames: 1 2 ... 3200 (3200 total)

fvarLabels: Protein

fvarMetadata: labelDescription

experimentData: use 'experimentData(object)'

Annotation:

The output is an ExpressionSet with the summarized, protein-level data in the assayData slot. The slot phenoData contains the phenoDataVH and the slot featureData contains the Protein field from dataVH. The output can be input into DEtest for differential expression testing.

In this example we used the default function median to aggregate all peptide values of a given protein. You may also input the function mean. Missing values in the feature data (in the case of peptides that have not been quantified in all samples) are allowed but ignored.

### 3.2 ExpressionSet Summarization

If you already have your data in an ExpressionSet, e.g. ExpSetVH, then you can simply run

```
> esetSum2 <- esetSummarizer(ExpSetVH, sumprot="median")</pre>
```

to get the summarized data. The output can be input into DEtest for differential expression analysis.

# 3.3 p-Value Summarization

Suppose you have peptide-level data and prefer to test at this level as opposed to the protein level, but still want to run the data through a network analysis. The function pvalueSummarizer allows you to do this: it summarizes peptide *p*-values (e.g. generated by DEtest) and the output may be used in runNetweavers.

The method used to aggregate p-values is Fisher's method. It returns a single p-value for each set of p-values, x, using the chi-squared distribution function on  $-2 \sum \ln(x)$  with degrees of freedom equal to the length of x.

For example, we can use p-values from outDE generated in the previous section for comparing timepoints 30 and 60 minutes:

```
> pvals3060 <- outDE_30to60[,c("Protein","P.Value")]
> names(pvals3060)[2] <- "pvalue"
> pvalsSum <- pvalueSummarizer(pvals3060)</pre>
> head(pvalsSum)
 Protein omnibus.P.Val
             0.10947111
1
     AARS
2
    ABCF2
             0.71294855
3
     ABT1
             0.30946672
4
    ACBD3
             0.36964204
5
    ACIN1
             0.05578688
6
     ACLY
             0.32122988
```

Note that Fisher's Method assumes independence, which is clearly not true for peptides from the same protein, so these p-values might be too optimistic.

#### 4 Find and Score Dense Clusters

The main function of the NetWeAvers algorithm is to perform network analysis. This section describes the core network analysis functions that find and score highly connected subgraphs within a protein-protein interaction network.

The first thing we want to do is filter an entire human network down to just those proteins of interest and their first interactors. We use **networkVH** which is the Reactome version 43 protein-protein interaction network (Matthews *et al.*, 2009):

```
> filtnet <- filterNetwork(networkVH, pvalsSum$Protein)
```

Then we find all the dense clusters of this network that have at least 4 proteins (default size) by taking random walks of length 10 (see Pons and Latapy (2006) for details):

The first item in the list is the data frame of cluster information that we will use in the following example. The second two items are special classes of the **igraph** package that can be used for visualizing the network (see Visualization section).

In order to score the clusters, we first need to calculate the protein weights and measures, which requires the use of p-values. We will use the summarized p-values from the comparison of 30 and 60 minutes (see previous section):

```
> ## change name of p-values for use in measureCalc
> names(pvalsSum)[2] <- "pvalue"
> ## calculate protein weights
> proteinWts <- weightCalc(filtnet)
> ## calculate protein measures
> proteinMsr <- measureCalc(pvalsSum, proteinWts, weightamt=3)
> head(proteinMsr)
```

	Protein	pvalue	weight	${\tt measure}$
1	A1CF	NA	0.04761905	0
2	A2M	NA	0.5000000	0
3	AARS	0.1094711	NA	NA
4	AASDHPPT	NA	1.00000000	0
5	ABCC8	NA	0.50000000	0
6	ABCF2	0.7129485	NA	NA

Note that we used a weight amount of 3, meaning we consider the weights nearly as important as p-values. Now that we have all the necessary data in proteinMsr we can score the clusters contained in outDC. The cscoremethod is the function that aggregates the protein measures per cluster and here we use the default mean:

```
> cscores <- scoreClusters(denseClusters, proteinMsr, cscoremethod="mean")
```

The larger the cluster score, the more significantly differentially regulated the proteins in the cluster are. We can use this statistic to rank our clusters, but we can also run a permutation test to see if the scores are large not by chance alone. The function permTest will do this, and here we perform 1000 permutations (B) and make sure we use the exact same cscoremethod that we did to find the cscores:

Note that we used set.seed before performing the permutation, which ensures the same results will be output from the permutation test in the next section. Finally, we can combine the cluster p-values with the cluster scores and membership information:

### 5 All-in-one: runNetweavers

The function runNetweavers is a wrapper for the functions in the NetWeAvers package. This function performs all of the steps in the NetWeAvers algorithm including filtering the protein network, finding dense clusters (only if output of findDenseClusters not already provided), and calculating protein weights, protein measures, cluster scores, and cluster p-values from a permutation test (optional). The following code runs NetWeAvers using the default values and produces the same output as above (proteinMsr, filtnet, and cdata) in a list:

```
> set.seed(1234)
> outNW <- runNetweavers(pvalsSum, filtnet, weightamt=3)
filtering network
finding dense clusters
calculating proteins weights
calculating protein measures
calculating cluster scores
calculating permutation pvalues
producing results tables
> str(outNW,list.len=5)
List of 3
 $ proteininfo:'data.frame':
                                    2363 obs. of 4 variables:
  ...$ Protein: chr [1:2363] "A1CF" "A2M" "AARS" "AASDHPPT" ...
  ..$ pvalue : num [1:2363] NA NA 0.109 NA NA ...
  ..$ weight : num [1:2363] 0.0476 0.5 NA 1 0.5 ...
  ..$ measure: num [1:2363] O O NA O O NA O NA O O ...
 $ filt_net
              :'data.frame':
                                    4823 obs. of 2 variables:
  ..$ intA: chr [1:4823] "AGPS" "AGPS" "KIF2A" "MY01C" ...
  ..$ intB: chr [1:4823] "FAR1" "FAR2" "CCNB1" "RAB4A" ...
 $ clusterinfo:'data.frame':
                                     52 obs. of 280 variables:
                         : Factor w/ 52 levels "c1", "c10", "c11", ...: 1 12 23 34 45 49 50 \,
  ..$ Cluster_ID
                         : int [1:52] 97 88 10 276 190 6 145 146 72 21 ...
  ..$ Cluster_size
  ..$ Cluster_score
                         : num [1:52] 0.246 0.6413 0.7331 0.0464 1.4122 ...
                         : num [1:52] 0.996 0.438 0.314 1 0 0.416 0.719 0.125 0.745 0.0
  ..$ Cluster_pvalue
  ..$ Cluster_symbol.X1 : chr [1:52] "EREG" "RPA1" "ACOT8" "ADCY6" ...
  .. [list output truncated]
```

### 6 Visualization

### 6.1 Writing Files for Cytoscape

Here we show how to export the data that can be used as input in Cytoscape (Shannon et al., 2003) to make the following figure. First we will create a node attributes file by adding the cluster membership to the protein list:

```
> proteininfo <- outNW$proteininfo
> clusterinfo <- outNW$clusterinfo
> proteininfo$cluster <- NA
> for(i in 1:nrow(clusterinfo)){
+  proteininfo$cluster[proteininfo[,1]%in%clusterinfo[i,-c(1:4)]] <- i
+ }</pre>
```

Then we can write the file to be uploaded into Cytoscape as node attributes. Note that Cytoscape cannot parse missing values, so we replace them with -9:

```
> ## write.table(proteininfo, "nodeAttributes.txt",
> ## sep="\t",row.names=F,quote=F,na="-9")
```

The following writes the network file:

```
> ## write.table(filtnet,"network.txt",sep="\t",row.names=F,quote=F)
```

and after a little bit of customization in Cytoscape we get, for example, the graph of cluster 5 in Figure 1.

# 6.2 Using igraph

It is also possible to visualize the resulting filtered network using the functions available in the **igraph** package. As an example, we will use the output from findDenseClusters to set up the plot (but not run, since it takes a while) of the graph with the proteins colored by cluster membership:

```
> dc_member <- outDC[['denseClus']]$membership
> dc_graph <- outDC$subGraph
> cols <- rainbow(length(unique(dc_member)))
>
> ## make the plot
> ## plot(dc_graph, vertex.color=cols, vertex.size=3, vertex.label=NA,
> ## edge.color="grey")
```

# 6.3 Using RCystocape

See RCytoscape on how to use transfer the network and attributes from R to Cytoscape.

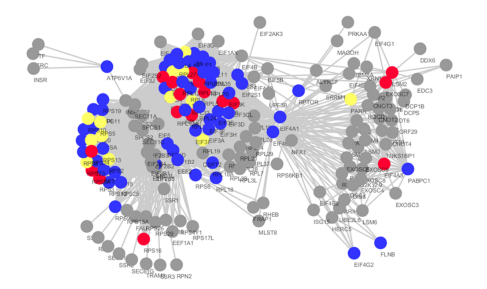


Figure 1: Image of cluster 5, generated with Cytoscape. A red node indicates a protein with a p-value less than 0.05, yellow less than 0.10, blue less than 1.00. A grey protein is one that is not in the Van Hoof dataset but is in the Reactome network.

### References

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