# Differential analysis of RNA-Seq data at the gene level using the DESeq2 package

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#### 1 Introduction

In this lab, you will learn how to analyse a count table, such as arising from a summarised RNA-Seq experiment, for differentially expressed genes.

## 2 Input data

## 2.1 Experiment data

We read in a prepared *SummarizedExperiment*, which was generated from publicly available data from the article by Felix Haglund et al., "Evidence of a Functional Estrogen Receptor in Parathyroid Adenomas", J Clin Endocrin Metab, Sep 2012, http://www.ncbi.nlm.nih.gov/pubmed/23024189. Details on the generation of this object can be found in the vignette for the *parathyroidSE* package, http://bioconductor.org/packages/release/data/experiment/html/parathyroidSE.html.

The purpose of the experiment was to investigate the role of the estrogen receptor in parathyroid tumors. The investigators derived primary cultures of parathyroid adenoma cells from 4 patients. These primary cultures were treated with diarylpropionitrile (DPN), an estrogen receptor  $\beta$  agonist, or with 4-hydroxytamoxifen (OHT). RNA was extracted at 24 hours and 48 hours from cultures under treatment and control. The blocked design of the experiment allows for statistical analysis of the treatment effects while controlling for patient-to-patient variation.

We first load the *DESeq2* package and the data package *parathyroidSE*, which contains the example data set.

```
library("DESeq2")
library("parathyroidSE")
```

The data command loads a data object.

```
data("parathyroidGenesSE")
```

The information in a *SummarizedExperiment* object can be accessed with accessor functions. For example, to see the actual data, i.e., here, the read counts, we use the assay function. (The head function restricts the output to the first few lines.)

```
head(assay(parathyroidGenesSE))
##
                    [,1] [,2] [,3] [,4] [,5] [,6] [,7] [,8] [,9] [,10] [,11] [,12]
## ENSG00000000003
                     792 1064
                                444
                                     953
                                           519
                                                855
                                                      413
                                                           365
                                                                278
                                                                      1173
                                                                             463
                                                                                    316
   ENSG00000000005
                                  2
                                             3
                                                                         0
                                                                               0
                                                                                      0
                             1
                                                  1
                                                             1
                          282
   ENSG00000000419
                     294
                                164
                                     263
                                           179
                                                217
                                                      277
                                                           204
                                                                189
                                                                       601
                                                                             257
                                                                                    183
   ENSG00000000457
                                 93
                                            75
                                                122
                                                      228
                                                           171
                                                                 116
                                                                       422
                                                                                    122
                     156
                           184
                                     145
                                                                             182
                                                173
                                                      611
                     396
                           207
                                     212
                                                           199
                                                                426
                                                                      1391
                                                                             286
   ENSG00000000460
                                210
                                           221
                                                                                    417
##
   ENSG00000000938
                       3
                             8
                                  2
                                       5
                                             0
                                                  4
                                                       13
                                                            22
                                                                   3
                                                                        38
                                                                              13
                                                                                     10
                    [,13] [,14] [,15] [,16] [,17]
                                                           [,19] [,20] [,21] [,22]
                                                     [,18]
##
##
  ENSG00000000003
                      987
                             424
                                   305
                                          391
                                                586
                                                       714
                                                             957
                                                                    346
                                                                          433
   ENSG00000000005
                      0
                               0
                                     0
                                            0
                                                  0
                                                         0
                                                                      0
                                                                            0
                                                                                   0
                                                               1
  ENSG00000000419
                      588
                             275
                                   263
                                          281
                                                406
                                                       568
                                                             764
                                                                    288
                                                                          259
                                                                                 250
   ENSG00000000457
                      441
                             211
                                   131
                                          115
                                                196
                                                       266
                                                             347
                                                                    133
                                                                          168
                                                                                 148
## ENSG0000000460 1452
                             238
                                   188
                                          102
                                                389
                                                       294
                                                             778
                                                                   162
                                                                           85
                                                                                 339
  ENSG00000000938
                       26
                              13
                                     7
                                            3
                                                 10
                                                       18
                                                              15
                                                                    7
                                                                            8
                                                                                   7
                    [,23] [,24] [,25] [,26] [,27]
##
## ENSG00000000003
                      277
                             511
                                   366
                                          271
                                                492
                                            0
  ENSG00000000005
                       0
                               0
                                     0
                                                  0
  ENSG00000000419
                             271
                                   227
                                          197
                                                363
                      147
## ENSG0000000457
                       83
                             184
                                   136
                                                195
                                          118
## ENSG0000000460
                       75
                             154
                                   314
                                          117
                                                233
  ENSG00000000938
                      5
                              13
```

In this count table, each row represents an Ensembl gene, each column a sequenced RNA library, and the values give the raw numbers of sequencing reads that were mapped to the respective gene in each library.

Question 1: For how many genes are there counts in this table?

#### Answer 1:

```
nrow(parathyroidGenesSE)
## [1] 63193
```

We also have metadata on each of the samples (the "columns" of the count table):

```
colData(parathyroidGenesSE)
## DataFrame with 27 rows and 8 columns
##
              run experiment patient treatment
                                                    time submission
                                                                        study
##
                   <factor> <factor> <factor> <factor>
      <character>
                                                           <factor> <factor>
## 1
        SRR479052 SRX140503
                                 1
                                        Control
                                                     24h SRA051611 SRP012167
## 2
        SRR479053 SRX140504
                                    1
                                        Control
                                                     48h SRA051611 SRP012167
## 3
        SRR479054 SRX140505
                                    1
                                            DPN
                                                     24h SRA051611 SRP012167
## 4
        SRR479055 SRX140506
                                            DPN
                                                     48h SRA051611 SRP012167
                                    1
## 5
        SRR479056 SRX140507
                                    1
                                            OHT
                                                     24h SRA051611 SRP012167
##
## 23
        SRR479074 SRX140523
                                    4
                                            DPN
                                                     48h SRA051611 SRP012167
        SRR479075 SRX140523
                                            DPN
## 24
                                    4
                                                     48h SRA051611 SRP012167
```

```
## 25 SRR479076 SRX140524
                                           OHT
                                                   24h SRA051611 SRP012167
## 26 SRR479077 SRX140525
                                   4
                                                   48h SRA051611 SRP012167
                                           OHT
                                           OHT
     SRR479078 SRX140525
                                                   48h SRA051611 SRP012167
##
        sample
##
      <factor>
## 1 SRS308865
## 2
     SRS308866
## 3
     SRS308867
## 4
      SRS308868
## 5 SRS308869
## ...
## 23 SRS308885
## 24 SRS308885
## 25 SRS308886
## 26 SRS308887
## 27 SRS308887
```

## **Question 2**: What are the metadata for the genes (the "rows" of the count table)? **Answer 2**:

```
rowData(parathyroidGenesSE)
## GRangesList of length 63193:
## $ENSG0000000000
## GRanges with 17 ranges and 2 metadata columns:
##
       seqnames ranges strand | exon_id
                                                                              exon_name
          <Rle> <IRanges> <Rle> | <integer>
                                                                         <character>
##
               X [99883667, 99884983]
                                               - | 664095 ENSE00001459322
##
       [1]
       [2]
                   X [99885756, 99885863]
##
                                                              664096 ENSE00000868868
##
       [3]
                                                   - | 664097 ENSE00000401072
                  X [99887482, 99887565]
                 X [99887538, 99887565]
                                                   - | 664098 ENSE00001849132
##
       [4]
      [5]
                 X [99888402, 99888536]
##
                                                               664099 ENSE00003554016
     [13] X [99890555, 99890743] - | 664106 ENSE00003512331

[14] X [99891188, 99891686] - | 664108 ENSE00001886883

[15] X [99891605, 99891803] - | 664109 ENSE00001855382

[16] X [99891790, 99892101] - | 664110 ENSE00001863395

[17] Y [09894942 99894988] - | 664111 ENSE00001828996
##
##
##
##
##
##
##
## <63192 more elements>
## ---
## seqlengths:
                                            2 ...
##
                                                               LRG 98
                                                                                     LRG 99
                       1
              249250621
                                   243199373 ...
                                                                 18750
                                                                                      13294
```

## 2.2 Collapsing technical replicates

There are a number of samples which were sequenced in multiple runs. For example, sample SRS308873 was sequenced twice. To see, we list the respective columns of the colData. (The

use of as.data.frame forces R to show us the full list, not just the beginning and the end as before.)

```
as.data.frame(colData(parathyroidGenesSE)[, c("sample", "patient", "treatment", "time")])
##
        sample patient treatment time
## 1 SRS308865 1 Control 24h
## 2 SRS308866
                    1 Control 48h
## 3 SRS308867
                    1
                          DPN 24h
## 4 SRS308868 1
## 5 SRS308869 1
                             DPN 48h
                             OHT 24h
## 6 SRS308870
                            OHT 48h
## 7 SRS308871
                    2 Control 24h
## 8 SRS308872 2 Control 48h
## 9 SRS308873 2 DPN 24h
                  2
2
2
2
## 10 SRS308873
                             DPN 24h
## 11 SRS308874
                             DPN 48h
## 12 SRS308875
                             OHT 24h
## 13 SRS308875
                             OHT 24h
## 14 SRS308876
                    2
                           OHT 48h
                    3 Control 24h
## 15 SRS308877
## 16 SRS308878 3 Control 48h
## 17 SRS308879 3 DPN 24h
## 18 SRS308880 3 DPN 48h
## 19 SRS308881
## 20 SRS308882
                   3
                             OHT 24h
                  3 Uni
4 Control 48h
DPN 24h
## 21 SRS308883
## 22 SRS308884
## 23 SRS308885
                   4
                             DPN 48h
                  4
## 24 SRS308885
                             DPN 48h
## 25 SRS308886
                   4
                             OHT 24h
## 26 SRS308887
                   4
                             OHT 48h
## 27 SRS308887
                             OHT 48h
```

We recommend to first add together technical replicates (i.e., libraries derived from the same samples), such that we have one column per sample.

**Fixme**: Present the 'pedestrian' approach of chunk poormanssum first, only then (and as 'optional') the sapply-magic.

As is often the case, this preparatory step looks more complicated than the subsequent actual analysis. In fact, the following operations are not specific to *DESeq2*, but are specific preparations needed for this data set. To understand the general ideas of *DESeq2*, you could now skip to Section 3. What you will learn in the rest of this section is an example of a typical preparatory data manipulation task done with elementary R functions. Details on these can be found in general textbooks on R; also consider reading the help pages of the functions used.

We first use the function split to see which columns need to be collapsed.

```
allColSamples <- colData(parathyroidGenesSE)$sample
sp <- split(seq(along = allColSamples), allColSamples)</pre>
```

Using sapply, we loop over the elements of sp, which correspond to the distinct samples, construct subtables of the count table (i.e., assay(parathyroidGenesSE)) corresponding only to the current sample considered, and add up across rows if there is more than one column. The result of the sapply call is a new table, in which each column now corresponds to a different sample.

```
countdata <- sapply(sp, function(columns) rowSums(assay(parathyroidGenesSE)[, columns,
    drop = FALSE]))
head(countdata)
##
                    SRS308865 SRS308866 SRS308867 SRS308868 SRS308869 SRS308870
                          792
                                    1064
                                               444
                                                          953
                                                                    519
## ENSG00000000003
                                                 2
  ENSG00000000005
                           4
                                                           3
                                                                     3
                                      1
                                                                                 1
  ENSG00000000419
                          294
                                    282
                                                          263
                                                                    179
                                               164
                                                                               217
                                                                     75
  ENSG00000000457
                          156
                                    184
                                                93
                                                          145
                                                                               122
                                    207
  ENSG00000000460
                          396
                                               210
                                                          212
                                                                    221
                                                                               173
  ENSG00000000938
                          3
                                      8
                                                 2
                                                           5
                                                                      0
                                                                                 4
                    SRS308871 SRS308872 SRS308873 SRS308874 SRS308875 SRS308876
##
## ENSG0000000003
                          413
                                    365
                                              1451
                                                          463
                                                                   1303
                                                                               424
  ENSG00000000005
                           0
                                      1
                                                 0
                                                           0
                                                                      0
                                                                                 0
## ENSG00000000419
                          277
                                    204
                                               790
                                                          257
                                                                    771
                                                                               275
## ENSG0000000457
                          228
                                    171
                                               538
                                                          182
                                                                    563
                                                                               211
## ENSG0000000460
                          611
                                     199
                                              1817
                                                          286
                                                                   1869
                                                                               238
## ENSG00000000938
                                     22
                                                          13
                                                                     36
                          13
                                                41
                                                                                13
##
                    SRS308877 SRS308878 SRS308879 SRS308880 SRS308881 SRS308882
                          305
                                    391
## ENSG0000000003
                                               586
                                                          714
                                                                    957
                                                                               346
## ENSG0000000005
                          0
                                      0
                                                 0
                                                           0
                                                                     1
                                                                                 0
## ENSG0000000419
                          263
                                    281
                                               406
                                                          568
                                                                    764
                                                                               288
  ENSG00000000457
                                    115
                                               196
                                                          266
                                                                    347
                                                                               133
                          131
                          188
                                    102
                                               389
                                                          294
                                                                    778
                                                                               162
  ENSG00000000460
  ENSG00000000938
                          7
                                      3
                                                10
                                                          18
                                                                     15
                                                                                 7
##
                    SRS308883 SRS308884 SRS308885 SRS308886 SRS308887
## ENSG00000000003
                          433
                                    402
                                               788
                                                          366
                                                                    763
                                      0
## ENSG00000000005
                          0
                                                 0
                                                           0
                                                                      0
## ENSG00000000419
                          259
                                    250
                                               418
                                                          227
                                                                    560
## ENSG0000000457
                          168
                                    148
                                               267
                                                          136
                                                                    313
## ENSG0000000460
                           85
                                    339
                                               229
                                                          314
                                                                    350
## ENSG0000000938
                                                18
                                                                     15
```

Novice users might find the preceding two code chunks difficult. Of course, there is a much easier way to add up the columns, namely by explicitly specifying the indices of the columns we want to use as is and the columns we want to add up, and using cbind to bind all the columns to a matrix:

```
a <- assay(parathyroidGenesSE)
countdata2 <- cbind(a[, 1:8], a[, 9] + a[, 10], a[, 11], a[, 12] + a[, 13], a[, 14:22],
        a[, 23] + a[, 24], a[, 25], a[, 26] + a[, 27])
all(countdata == countdata2)
## [1] TRUE</pre>
```

While this is simpler to understand, it is more error-prone. Mistakes can easily happen when determining the column indices, and it is tedious to update the code if the input data changes, for instance, if at a later time you would like to add more replicates to your data set. Hence, if you are a beginner in R and want to improve your R skills, try to understand how the split and the sapply calls above work, because only learning to master such expressions will give you the skills to make full use of R.

Having reduced our count data table to only one column per sample, we next need to subset the column metadata accordingly, as we now have less columns. We also now use the sample names as names for the column data rows:

```
coldata <- colData(parathyroidGenesSE)[sapply(sp, `[`, 1), ]</pre>
rownames(coldata) <- coldata$sample</pre>
coldata
## DataFrame with 23 rows and 8 columns
            run experiment patient treatment time submission
    <character> <factor> <factor> <factor> <factor> <factor>
##
## SRS308865 SRR479052 SRX140503 1 Control 24h SRA051611
## SRS308866 SRR479052 SRX140505 1 Control 48h SRA051611
## SRS308866 SRR479054 SRX140505 1 DPN 24h SRA051611
## SRS308868 SRR479055 SRX140506 1 DPN 48h SRA051611
## SRS308869 SRR479056 SRX140507 1 OHT 24h SRA051611
## SRS308883 SRR479072 SRX140521 4 Control 48h SRA051611
## SRS308884 SRR479073 SRX140522 4 DPN 24h SRA051611
## SRS308885 SRR479074 SRX140523 4 DPN 48h SRA051611
## SRS308886 SRR479076 SRX140524 4 DPN 48h SRA051611
## SRS308886 SRR479076 SRX140524 4 DPN 48h SRA051611
## SRS308885 SRR479074 SRX140523 4 DPN 48h SRA051611
## SRS308886 SRR479076 SRX140524 4 OHT 24h SRA051611
## SRS308887 SRR479077 SRX140525 4 OHT 48h SRA051611
## study sample
## <factor> <factor>
## SRS308865 SRP012167 SRS308865
## SRS308866 SRP012167 SRS308866
## SRS308867 SRP012167 SRS308867
## SRS308868 SRP012167 SRS308868
## SRS308869 SRP012167 SRS308869
## SRS308883 SRP012167 SRS308883
## SRS308884 SRP012167 SRS308884
## SRS308885 SRP012167 SRS308885
## SRS308886 SRP012167 SRS308886
## SRS308887 SRP012167 SRS308887
```

**Question 3**: What do the quotation marks in the expression '[' do? What happens if you omit them?

#### Answer 3:

The function sapply expects an R function as its second argument. Here, we want to provide it with the function for vector subsetting (as in a[1]), and the name of this function is [. However, if we provide that name without the quotation marks, the R interpreter gets confused and complains about the unexpected symbol (try this out). Hence we need to quote the function name

in our call to sapply.

To unclutter the output in the subsequent steps, we only keep the column data columns that we actually need for our analysis.

```
coldata <- coldata[, c("patient", "treatment", "time")]</pre>
head(coldata)
## DataFrame with 6 rows and 3 columns
##
      patient treatment time
   <factor> <factor> <factor>
##
## SRS308865 1 Control 24h
## SRS308866
                 1 Control
## SRS308867 1 DPN
## SRS308868 1 DPN
## SRS308869 1 OHT
                                 24h
                                  48h
                                  24h
## SRS308870 1 OHT
                                  48h
```

Our *SummarizedExperiment* object also contains metadata on the rows, which we can simply keep unchanged:

```
rowdata <- rowData(parathyroidGenesSE)</pre>
rowdata
## GRangesList of length 63193:
## $ENSG0000000003
## GRanges with 17 ranges and 2 metadata columns:
## seqnames ranges strand | exon_id exon_name
## <Rle> <IRanges> <Rle> | <integer> <character>
     [1] X [99883667, 99884983] - | 664095 ENSE00001459322
[2] X [99885756, 99885863] - | 664096 ENSE00000868868
##
##
     [3]
                                        - | 664097 ENSE00000401072
##
             X [99887482, 99887565]
   ##
##
##
##
##
##
##
##
##
## ...
## <63192 more elements>
## ---
## seqlengths:
##
                  1
                                    2 ...
                                                  LRG 98
                                                                    LRG 99
                            243199373 ...
                                                                     13294
           249250621
                                                    18750
```

We now have all the ingredients to prepare our data object in a form that is suitable for analysis, namely:

• countdata: a table with the read counts, with technical replicates summed up,

- coldata: a table with metadata on the count table's columns, i.e., on the samples,
- rowdata: a table with metadata on the count table's rows, i.e., on the genes, and
- a design formula, which tells which factors in the column metadata table specify the experimental design and how these factors should be used in the analysis. We specify patient + treatment, which means that we want to test for the effect of treatment (the last factor), controlling for the effect of patient (the first factor). You can use R's formula notation to express any experimental design that can be described within an ANOVA-like framework.

To now construct the data object from the matrix of counts and the metadata table, we use:

```
ddsFull <- DESeqDataSetFromMatrix(countData = countdata, colData = coldata, design = "patient +
    treatment, rowData = rowdata)
## Usage note: the following factors have 3 or more levels:
##
## patient, treatment
##
## For DESeq2 versions < 1.3, if you plan on extracting results for
## these factors, we recommend using betaPrior=FALSE as an argument
## when calling DESeq().
## As currently implemented in version 1.2, the log2 fold changes can
## vary if the base level is changed, when extracting results for a
## factor with 3 or more levels. A solution will be implemented in
## version 1.3 which allows for the use of a beta prior and symmetric
## log2 fold change estimates regardless of the selection of base level.
ddsFull
## class: DESeqDataSet
## dim: 63193 23
## exptData(0):
## assays(1): counts
## rownames(63193): ENSG00000000003 ENSG0000000005 ... LRG_98 LRG_99
## rowData metadata column names(0):
## colnames(23): SRS308865 SRS308866 ... SRS308886 SRS308887
## colData names(3): patient treatment time
```

## 3 Running the DESeq2 pipeline

Here we will analyze a subset of the samples, namely those taken after 48 hours, with either control or DPN treatment, taking into account the multifactor design.

## 3.1 Preparing the data object for the analysis of interest

First we subset the relevant columns from the full dataset:

```
dds <- ddsFull[, colData(ddsFull)$treatment %in% c("Control", "DPN") & colData(ddsFull)$time ==
    "48h"]</pre>
```

Sometimes it is necessary to "refactor" the factors, in case that levels have been dropped. (Here, for example, the treatment factor still contains the level "OHT", but no sample to this level.)

```
dds$patient <- factor(dds$patient)
dds$treatment <- factor(dds$treatment)</pre>
```

It will be convenient to make sure that Control is the first level in the treatment factor, so that the  $\log_2$  fold changes are calculated as treatment over control. The function relevel achieves this:

```
dds$treatment <- relevel(dds$treatment, "Control")</pre>
```

A guick check whether we now have the right samples:

```
colData(dds)
## DataFrame with 8 rows and 3 columns
## patient treatment time
## <factor> <factor> <factor>
## SRS308866 1 Control 48h
## SRS308868
               1 DPN
                            48h
## SRS308872 2 Control
## SRS308874 2 DPN
                            48h
              2 DPN
## SRS308874
                            48h
             3 Control
3 DPN
## SRS308878
                            48h
## SRS308880
                            48h
## SRS308883 4 Control
                             48h
## SRS308885
```

## 3.2 Running the pipeline

With the data object prepared, the *DESeq2* analysis can now be run with a single call to the function DESeq:

```
dds <- DESeq(dds)

## estimating size factors
## estimating dispersions</pre>
```

```
## gene-wise dispersion estimates
## mean-dispersion relationship
## final dispersion estimates
## 1 rows did not converge in dispersion, labelled in mcols(object)$dispConv. Use larger maxit argument
with estimateDispersions
## fitting model and testing
```

## 3.3 Inspecting the results table

The results for the last variable in the design formula, in our case the treatment variable, can be extracted using the results function.

```
res <- results(dds)
res
## DataFrame with 63193 rows and 6 columns
               baseMean log2FoldChange lfcSE stat
##
                                                        pvalue
##
                <numeric> <numeric> <numeric> <numeric> <numeric>
## ENSG0000000000 623.42
                            -0.0205 0.0585 -0.350 0.72664
## ENSG0000000005
                  0.68
                             0.0210 0.2053
                                                0.102 0.91848
## ENSG00000000419 299.75
                             -0.0148 0.0742 -0.200 0.84148
## ENSG0000000457 183.50
                             -0.0892 0.1065 -0.837 0.40252
## ENSG0000000460 200.46
                             0.3539 0.1109 3.192 0.00141
## ...
                                               . . .
                    . . .
                                 . . .
                                          . . .
## LRG_94
                     0
                                  NA
                                         NA
                                                  NA
                                                           NA
## LRG_96
                     0
                                 NA
                                         NA
                                                  NA
                                                           NA
## LRG_97
                     0
                                         NA
                                                  NA
                                 NA
                                                           NΑ
                                         NA
NA
## LRG_98
                      0
                                  NA
                                                   NA
## LRG_99
                      0
                                  NA
                                                   NA
                                                           NA
##
                   padj
##
             <numeric>
## ENSG0000000000 0.9057
## ENSG0000000005
## ENSG0000000419 0.9483
## ENSG0000000457 0.7455
## ENSG0000000460 0.0283
## ...
## LRG_94
                     NA
## LRG_96
                     NA
## LRG_97
                     NA
## LRG_98
                     NA
## LRG_99
                      NA
```

As res is a DataFrame object, it carries metadata with information on the meaning of the columns:

```
mcols(res)
## DataFrame with 6 rows and 2 columns
```

```
##
             type
                                                      description
##
      <character>
                                                      <character>
## 1 intermediate
                                      the base mean over all rows
## 2
         results log2 fold change (MAP): treatment DPN vs Control
## 3
         results
                         standard error: treatment DPN vs Control
## 4
         results
                         Wald statistic: treatment DPN vs Control
         results
## 5
                      Wald test p-value: treatment DPN vs Control
## 6
         results
                                             BH adjusted p-values
```

The first column, baseMean, is a just the average of the normalized count values, taken over all samples. The remaining four columns refer to a specific *contrast*, namely the comparison of the levels *DPN* versus *Control* of the factor variable *treatment*. See the help page for results (by typing ?results) for information on how to obtain other contrasts.

The column log2FoldChange is the effect size estimate. It tells us how much the gene's expression seems to have changed due to treatment with DPN in comparison to control. This value is reported on a logarithmic scale to base 2: for example, a  $log_2$  fold change of 1.5 means that the gene's expression is increased by a factor of  $2^{1.5} \approx 2.82$ .

Of course, this estimate has an uncertainty associated with it, which is available in the column lfcSE, the standard error estimate for the log2 fold change estimate. We can also express the uncertainty of a particular effect size estimate as the result of a statistical test. The purpose of a test for differential expression is to test whether the data provides sufficient evidence to conclude that this value is really different from zero (and that the sign is correct). DESeq2 performs for each gene a *hypothesis test* to see whether evidence is sufficient to decide against the *null hypothesis* that there is no effect of the treatment on the gene and that the observed difference between treatment and control was merely caused by experimental variability (i. e., the type of variability that you can just as well expect between different samples in the same treatment group). As usual in statistics, the result of this test is reported as a *p value*, and it is found in the column pvalue. (Remember that a p value indicates the probability that a fold change as strong as the observed one, or even stronger, would be seen under the situation described by the null hypothesis.)

Finally, we note that a subset of the p values in res are NA ("not available"). This is DESeq's way of reporting that all counts for this gene were zero, and hence not test was applied.

Question 4: How could you check to see if the baseMean is the mean of raw counts or the mean of normalized counts?

**Answer 4**: The raw counts and normalized counts of a *DESeqDataSet* object are available via the accessor function counts, which has an argument normalized, which defaults to FALSE.

```
all.equal(res$baseMean, rowMeans(counts(dds)))
## [1] "names for current but not for target"
## [2] "Mean relative difference: 0.0582"
all.equal(res$baseMean, rowMeans(counts(dds, normalized = TRUE)))
## [1] "names for current but not for target"
```

## 3.4 Multiple testing

Novices in high-throughput biology often assume that thresholding these p values at 0.05, as is often done in other settings, would be appropriate – but it is not. We briefly explain why:

There are 2115 genes with a p value below 0.05 among the 30434 genes, for which the test succeeded in reporting a p value:

```
sum(res$pvalue < 0.05, na.rm = TRUE)

## [1] 2115

table(is.na(res$pvalue))

## FALSE TRUE
## 30434 32759</pre>
```

Now, assume for a moment that the null hypothesis is true for all genes, i.e., no gene is affected by the treatment with DPN. Then, by the definition of p value, we expect up to 5% of the genes to have a p value below 0.05. This amounts to 1522 genes. If we just considered the list of genes with a p value below 0.05 as differentially expressed, this list should therefore be expected to contain up to 1522/2115 = 72% false positives!

DESeq2 uses the so-called Benjamini-Hochberg (BH) adjustment; in brief, this method calculates for each gene an adjusted p value which answers the following question: if one called significant all genes with a p value less than or equal to this gene's p value threshold, what would be the fraction of false positives (the false discovery rate, FDR) among them (in the sense of the calculation outlined above)? These values, called the BH-adjusted p values, are given in the column padj of the results object.

Hence, if we consider a fraction of 10% false positives acceptable, we can consider all genes with an *adjusted* p value below 10%=0.1 as significant. How many such genes are there?

```
sum(res$padj < 0.1, na.rm = TRUE)
## [1] 882</pre>
```

We subset the results table to these genes and then sort it by the log2-fold-change estimate to get the significant genes with the strongest down-regulation

```
resSig <- res[which(res$padj < 0.1), ]
head(resSig[order(resSig$log2FoldChange), ])

## DataFrame with 6 rows and 6 columns
## baseMean log2FoldChange lfcSE stat pvalue
## <numeric> <numeric> <numeric> <numeric> <numeric> <numeric> <numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric></numeric>
```

```
## ENSG0000163631
                     269
                                 -0.942
                                          0.1033
                                                   -9.11 8.09e-20
## ENSG0000169239
                     1548
                                -0.753 0.1065
                                                   -7.07 1.50e-12
## ENSG0000041982
                    1493
                                -0.686 0.0922
                                                   -7.44 9.96e-14
## ENSG0000145244
                    173
                                -0.676 0.2556
                                                   -2.65 8.15e-03
                                -0.674 0.1075 -6.27 3.56e-10
                    183
## ENSG0000119946
## ENSG00000155111
                     588
                                -0.665 0.0857
                                                   -7.75 8.96e-15
##
                     padj
##
               <numeric>
## ENSG00000163631 6.39e-17
## ENSG00000169239 5.91e-10
## ENSG00000041982 4.97e-11
## ENSG00000145244 9.21e-02
## ENSG00000119946 7.68e-08
## ENSG00000155111 5.00e-12
```

#### and with the strongest upregulation

```
tail(resSig[order(resSig$log2FoldChange), ])
## DataFrame with 6 rows and 6 columns
##
                         baseMean log2FoldChange
                                                               lfcSE
                                                                              stat
                                                                                            pvalue
                         <numeric> <numeric> <numeric> <numeric> <numeric>
## ENSG0000005189 227
                                              0.641 0.229 2.80 5.18e-03
## ENSG00000156414 137
## ENSG00000103257 168
## ENSG00000101255 285
## ENSG00000135069 136

      0.727
      0.138
      5.27
      1.40e-07

      0.780
      0.151
      5.16
      2.51e-07

      0.797
      0.186
      4.29
      1.76e-05

      0.843
      0.321
      2.63
      8.66e-03

                                                 0.869 0.151
                                                                              5.76 8.33e-09
## ENSG00000092621
                                594
##
                                padj
##
                         <numeric>
## ENSG0000005189 6.88e-02
## ENSG00000156414 1.53e-05
## ENSG00000103257 2.41e-05
## ENSG00000101255 9.98e-04
## ENSG00000135069 9.60e-02
## ENSG00000092621 1.32e-06
```

**Question 5**: What is the proportion of down- and up-regulation among the genes with adjusted p value less than 0.1?

#### Answer 5:

```
table(sign(resSig$log2FoldChange))

##
## -1   1
## 459 423
```

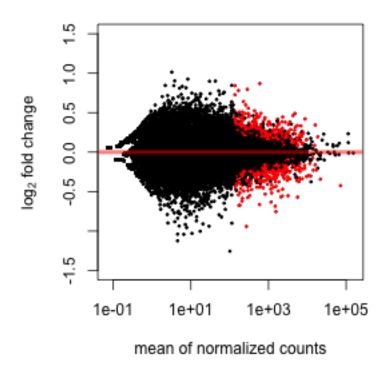


Figure 1: The MA-plot shows the  $\log_2$  fold changes from the treatment over the mean of normalized counts, i.e. the average of counts normalized by size factor. The *DESeq2* package incorporates a prior on  $\log_2$  fold changes, resulting in moderated estimates from genes with low counts and highly variable counts, as can be seen by the narrowing of spread of points on the left side of the plot.

## 3.5 Diagnostic plots

A so-called MA plot provides a useful overview for an experiment with a two-group comparison:

```
plotMA(dds, ylim = c(-1.5, 1.5))
```

The plot (Fig. 1) represents each gene with a dot. The x axis is the average expression over all samples, the y axis the  $\log_2$  fold change between treatment and control. Genes with an adjusted p value below a threshold (here 0.1, the default) are shown in red.

This plot demonstrates that only genes with an average normalized count above 10 contain sufficient information to yield a significant call, and only above about 300 counts can smaller fold-changes become significant.

Also note *DESeq2*'s shrinkage estimation of log fold changes (LFCs): When count values are too low to allow an accurate estimate of the LFC, the value is "shrunken" towards zero to avoid

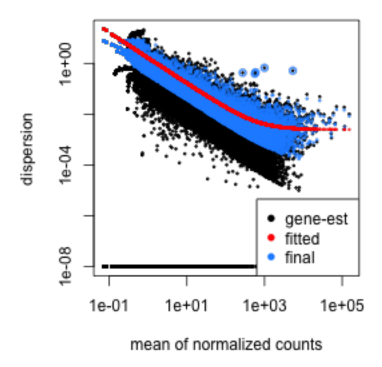


Figure 2: Plot of dispersion estimates. See text for details.

that these values, which otherwise would frequently be unrealistically large, dominate the top-ranked log fold changes.

Whether a gene is called significant depends not only on its LFC but also on its within-group variability, which DESeq2 quantifies as the dispersion. For strongly expressed genes, the dispersion can be understood as a squared coefficient of variation: a dispersion value of 0.01 means that the gene's expression tends to differ by typically  $\sqrt{0.01} = 10\%$  between samples of the same treatment group. For weak genes, the Poisson noise is an additional source of noise, which is added to the dispersion.

The function plotDispEsts visualizes *DESeq2*'s dispersion estimates:

#### plotDispEsts(dds)

The black dots are the dispersion estimates for each gene as obtained by considering the information from each gene separately. Unless one has many samples, these values fluctuate strongly around their true values. Therefore, we fit the red trend line, which shows the dispersions' dependence on the mean, and then shrink each gene's estimate towards the red line to obtain the final estimates (blue circles) that are then used in the hypothesis test.

Question 6: How could you change the MA-plot so as to color those genes with adjusted

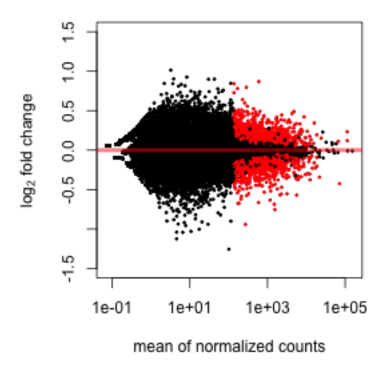


Figure 3: The MA-plot with red points indicating adjusted p value less than 0.5.

p-value less than 0.5 instead of 0.1?

#### Answer 6:

```
plotMA(dds, pvalCutoff = 0.5, ylim = c(-1.5, 1.5))
```

#### See Figure 3.

Another useful diagnostic plot is the histogram of the p values (Fig. 4).

```
hist(res$pvalue, breaks = 100)
```

**Question 7**: Revisit the discussion about p values and multiple testing in the previous section. Which part of the histogram is caused by genes that are called significant? And which part is caused by those that are truly significant? Why are there "spikes" at intermediate values?

**Answer 7**: Genes that are not differentially expressed have p values that are approximately uniformly distributed between 0 and 1. This gives rise to the floor of bars of equal heights. The truly differentially expressed genes give rise to the tall bar(s) at the very left – but only to that part of the bars that raises above the uniform floor. Of course, we cannot know which of the genes in these tall bars are true ones and which are not. When only looking at the bars to the

## Histogram of res\$pvalue

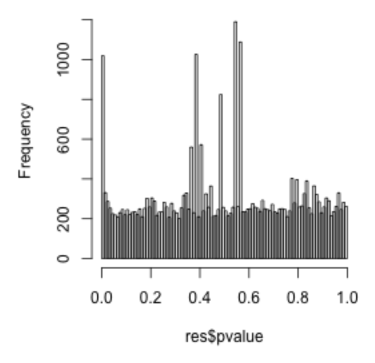


Figure 4: Histogram of the p values returned by the test for differential expression.

left of our chosen p value cut-off, the ratio of "floor" area to total area provides an estimate of the false discovery rate. This is a graphical way of understanding FDR.

The rule that p values from null cases are uniform is true only for continuous test statistics. However, for genes with low counts, the fact that we are working with integer counts becomes noticeable, and gives rise to the spikes at intermediate p values.

## 4 Independent filtering

The MA plot (Figure 1) highlights an important property of RNA-Seq data. For weakly expressed genes, we have no chance of seeing differential expression, because the low read counts suffer from so high Poisson noise that any biological effect is drowned in the uncertainties from the read counting. The MA plot suggests that for genes with less than one or two counts per sample, averaged over all samples, there is no real inferential power. We loose little if we filter out these genes:

```
filterThreshold <- 2
keep <- rowMeans(counts(dds, normalized = TRUE)) > filterThreshold
```

```
table(keep)

## keep
## FALSE TRUE
## 43939 19254
```

Note that none of the genes below the threshold had a significant adjusted p value

```
min(res$padj[!keep], na.rm = TRUE)

## Warning: no non-missing arguments to min; returning Inf

## [1] Inf
```

At first sight, there may seem to be little benefit in filtering out these genes. After all, the test found them to be non-significant anyway. However, these genes have an influence on the multiple testing adjustment, whose performance improves if such genes are removed. Compare:

```
table(p.adjust(res$pvalue, method = "BH") < 0.1)

##
## FALSE TRUE
## 29863 571

table(p.adjust(res$pvalue[keep], method = "BH") < 0.1)

##
## FALSE TRUE
## 18545 709</pre>
```

By removing the weakly-expressed genes from the input to the FDR procedure, we have found more genes to be significant among those which we kept, and so improved the power of our test. This approach is known as *independent filtering*.

The term *independent* highlights an important caveat. Such filtering is permissible only if the filter criterion is independent of the actual test statistic [?]. Otherwise, the filtering would invalidate the test and consequently the assumptions of the BH procedure. This is why we filtered on the average over *all* samples: this filter is blind to the assignment of samples to the treatment and control group and hence independent.

**Question 8**: Redo the histogram as in Figure 4, now only using the genes that passed the filtering. What happened to the spikes at intermediate values?

Answer 8: Run

```
hist(res$pvalue[keep], breaks = 100)
```

## Histogram of res\$pvalue[keep]

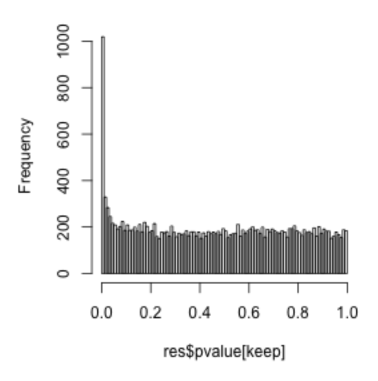


Figure 5: Histogram of the p values returned by the test for differential expression.

See Figure 5. As explained before, the spikes were caused by genes with low counts. Having removed these, our p value histogram now looks smoother.

In this vignette, we have determined the value for filterThreshold, 2, by looking at Figure 1. More formal, automatable ways exist; if you are interested, please have a look at the vignette *Diagnostics for independent filtering* in the *genefilter* package.

## 4.1 Adding gene names

Our result table only uses Ensembl gene IDs, but gene names may be more informative. Bio-conductor's annotation packages help with mapping various ID schemes to each other.

We load the annotation package *org.Hs.eg.db*:

```
library("org.Hs.eg.db")
```

This is the organism annotation package ("org") for *Homo sapiens* ("Hs"), organized as an *AnnotationDbi* package ("db"), using Entrez Gene IDs ("eg") as primary key.

To get a list of all available key types, use

```
cols(org.Hs.eg.db)
## Warning: 'cols' has been deprecated and replaced by 'columns' for versions of
## Bioc that are higher than 2.13. Please use 'columns' anywhere that
## you previously used 'cols'
                       "PFAM"
                                      "IPI"
  [1] "ENTREZID"
                                                      "PROSITE"
                                                                     "ACCNUM"
##
                       "CHR"
                                                     "CHRLOCEND"
                                                                     "ENZYME"
##
   [6] "ALIAS"
                                      "CHRLOC"
## [11] "MAP"
                       "PATH"
                                      "PMID"
                                                     "REFSEQ"
                                                                     "SYMBOL"
## [16] "UNIGENE"
                       "ENSEMBL"
                                      "ENSEMBLPROT"
                                                     "ENSEMBLTRANS" "GENENAME"
## [21] "UNIPROT"
                       "GO"
                                                     "ONTOLOGY"
                                                                     "GOALL"
                                      "EVIDENCE"
   [26] "EVIDENCEALL" "ONTOLOGYALL" "OMIM"
                                                     "UCSCKG"
```

Converting IDs with the native functions from the *AnnotationDbi* package is currently a bit cumbersome, so we provide the following convenience function (without explaining how exactly it works):

```
convertIDs <- function(ids, fromKey, toKey, db, ifMultiple = c("putNA", "useFirst")) {
    stopifnot(inherits(db, "AnnotationDb"))
    ifMultiple <- match.arg(ifMultiple)
    suppressWarnings(selRes <- AnnotationDbi::select(db, keys = ids, keytype = fromKey,
        cols = c(fromKey, toKey)))
    if (ifMultiple == "putNA") {
        duplicatedIds <- selRes[duplicated(selRes[, 1]), 1]
        selRes <- selRes[!selRes[, 1] %in% duplicatedIds, ]
    }
    return(selRes[match(ids, selRes[, 1]), 2])
}</pre>
```

This function takes a list of IDs as first argument and their key type as the second argument. The third argument is the key type we want to convert to, the fourth is the *AnnotationDb* object to use. Finally, the last argument specifies what to do if one source ID maps to several target IDs: should the function return an NA or simply the first of the multiple IDs?

To convert the Ensembl IDs in the rownames of res to gene symbols and add them as a new column, we use:

```
res$symbol <- convertIDs(row.names(res), "ENSEMBL", "SYMBOL", org.Hs.eg.db)
## DataFrame with 63193 rows and 7 columns
                baseMean log2FoldChange
                                           lfcSE
                                                    stat
                                                              pvalue
##
                 <numeric>
                             <numeric> <numeric> <numeric> <numeric>
## ENSG0000000003
                    623.42
                                -0.0205 0.0585
                                                  -0.350 0.72664
## ENSG0000000005
                    0.68
                                 0.0210
                                           0.2053
                                                    0.102 0.91848
## ENSG0000000419
                    299.75
                                 -0.0148
                                           0.0742
                                                    -0.200 0.84148
## ENSG0000000457
                    183.50
                                 -0.0892
                                           0.1065
                                                    -0.837 0.40252
## ENSG0000000460
                                  0.3539
                                           0.1109
                                                    3.192 0.00141
                    200.46
                                                       . . .
## LRG_94
                         0
                                     NA
                                              NA
                                                        NA
                                                                 NA
```

```
## LRG_96
                                                              NA
                           0
                                                    NA
                                                                         NA
                           0
## LRG_97
                                          NA
                                                    NA
                                                              NA
                                                                         NΑ
## LRG_98
                           0
                                          NA
                                                    NA
                                                              NA
                                                                         NA
## LRG_99
                           0
                                                    NA
                                                              NA
                                                                         NA
                                          NA
##
                        padj
                                  symbol
##
                   <numeric> <character>
## ENSG00000000003
                      0.9057
                                  TSPAN6
## ENSG0000000005
                          NA
                                    TNMD
## ENSG0000000419
                      0.9483
                                    DPM1
## ENSG0000000457 0.7455
                                   SCYL3
## ENSG0000000460
                    0.0283
                              Clorf112
## LRG_94
                          NA
## LRG_96
## LRG_97
                          NA
                                      NΑ
## LRG_98
                          NA
## LRG_99
                          NA
                                       NA
```

Finally, we note that you can easily save the results table in a CSV file, which you can then load with a spreadsheet program such as Excel:

```
write.csv(as.data.frame(res), file = "results.csv")
```

## 5 Downstream analyses

A list of gene names is no final result. We demonstrate two possible further analysis steps.

## 5.1 Gene set enrichment analysis

Do the genes with a strong up- or down-regulation have something in common? We perform next a gene-set enrichment analysis (GSEA) to examine this question.

We use the gene sets in the Reactome database

```
library("reactome.db")
```

This database works with Entrez IDs, so we add a column with such IDs, using our convertIDs function:

```
res$entrez <- convertIDs(row.names(res), "ENSEMBL", "ENTREZID", org.Hs.eg.db)</pre>
```

Next, we subset the results table, res, to only those genes for which the Reactome database has data (i.e, whose Entrez ID we find in the respective key column of reactome.db) and for which the test gave a p value that was not NA.

```
res2 <- res[res$entrez %in% keys(reactome.db, "ENTREZID") & !is.na(res$pvalue), ]
head(res2)
## DataFrame with 6 rows and 8 columns
                 baseMean log2FoldChange
                                                            pvalue
                                          lfcSE
                                                   stat
##
                 <numeric>
                          <numeric> <numeric> <numeric> <numeric>
## ENSG0000000419
                   299.7
                               -0.01483 0.0742 -0.2000
                                                            0.841
## ENSG0000000938
                    12.1
                              -0.00483 0.3205 -0.0151
                                                             0.988
## ENSG0000000971
                    19.7
                              -0.38555 0.2648 -1.4559
                                                            0.145
                               0.04504
                                        0.0945
                                                  0.4765
## ENSG0000001084
                    323.5
                                                             0.634
                                                            0.601
## ENSG0000001167
                    412.4
                               -0.05387
                                         0.1031
                                                 -0.5225
## ENSG0000001626
                    11.0
                               0.15267 0.2988 0.5109
                                                            0.609
##
                     padj
                             symbol
                                        entrez
##
                 <numeric> <character> <character>
## ENSG0000000419
                   0.948
                                DPM1
                                           8813
## ENSG0000000938
                                FGR
                                           2268
## ENSG0000000971
                      NA
                                 CFH
                                           3075
## ENSG0000001084
                    0.870
                                GCLC
                                           2729
## ENSG0000001167
                    0.859
                                NFYA
                                           4800
## ENSG0000001626
                                CFTR
                                           1080
```

Using select, a function from *AnnotationDbi* for querying database objects, we get a table with the mapping from Entrez IDs to Reactome Path IDs

```
reactomeTable <- AnnotationDbi::select(reactome.db, keys = res2$entrez, keytype = "ENTREZID",
    columns = c("ENTREZID", "REACTOMEID"))
## Warning: 'select' and duplicate query keys resulted in 1:many mapping between
## keys and return rows
head(reactomeTable)
    ENTREZID REACTOMEID
## 1
        8813
                 162699
## 2
        8813
                 163125
## 3
        8813
                 392499
## 4
         8813
                 446193
## 5
         8813
                  446203
                 446219
```

The next code chunk transforms this table into an *incidence matrix*. This is a boolean matrix with one row for each Reactome Path and one column for each gene in res2, which tells us which genes are members of which Reactome Paths. (If you want to understand how this chunk exactly works, read up about the tapply function.)

```
incm <- do.call(rbind, with(reactomeTable, tapply(ENTREZID, factor(REACTOMEID), function(x) res2$entrez %i
    x)))
colnames(incm) <- res2$entrez
str(incm)</pre>
```

```
## logi [1:1435, 1:5494] FALSE FALSE FALSE FALSE FALSE FALSE ...
## - attr(*, "dimnames")=List of 2
## ..$ : chr [1:1435] "1059683" "109581" "109582" "109606" ...
## ..$ : chr [1:5494] "8813" "2268" "3075" "2729" ...
```

We remove all rows corresponding to Reactome Paths with less than 5 assigned genes.

```
incm <- incm[rowSums(incm) >= 5, ]
```

To test whether the genes in a Reactome Path behave in a special way in our experiment, we perform t-tests to see whether the average of the genes'  $\log_2$  fold change values are different from zero. If so, we can say that our treatment tends to upregulate (or downregulate) the genes in the category. To facilitate the computations, we define a little helper function:

```
testCategory <- function(reactomeID) {
   isMember <- incm[reactomeID, ]
   data.frame(reactomeID = reactomeID, numGenes = sum(isMember), avgLFC = mean(res2$log2FoldChange[isMember)
        strength = sum(res2$log2FoldChange[isMember])/sqrt(sum(isMember)), pvalue = t.test(res2$log2FoldChange[isMember])
}</pre>
```

The function can be called with a Reactome Path ID:

```
testCategory("109581")

## reactomeID numGenes avgLFC strength pvalue reactomeName
## 1 109581 148 -0.00865 -0.105 0.356 Homo sapiens: Apoptosis
```

As you can see the function not only performs the t test and returns the p value but also lists other useful information such as the number of genes in the category, the average log fold change, a "strength" measure (see below) and the name with which Reactome describes the Path.

We call the function for all Paths in our incidence matrix and collect the results in a data frame:

```
reactomeResult <- do.call(rbind, lapply(rownames(incm), testCategory))</pre>
```

As we performed many tests, we should again use a multiple testing adjustment.

```
reactomeResult$padjust <- p.adjust(reactomeResult$pvalue, "BH")</pre>
```

This is a list of Reactome Paths which are significantly differentially expressed in our comparison of DPN treatment with control, sorted according to sign and strength of the signal:

```
reactomeResultSignif <- reactomeResult[reactomeResult$padjust < 0.05, ]</pre>
reactomeResultSignif[order(reactomeResultSignif$strength), ]
##
        reactomeID numGenes avgLFC strength
                                               pvalue
## 878
                        20 -0.1442
                                      -0.645 4.54e-04
            445355
                                      -0.546 2.05e-04
## 1137
            913531
                        140 -0.0462
## 104
           1280215
                        246 -0.0314
                                      -0.492 3.84e-04
## 69
           1169408
                        64 -0.0572
                                     -0.457 6.47e-04
## 70
          1169410
                        64 -0.0572
                                      -0.457 6.47e-04
           2032785
                        10 -0.1175
## 552
                                      -0.372 2.34e-04
## 1066
            74159
                       137 -0.0307
                                      -0.359 2.85e-04
                        96 -0.0365
## 1044
            73857
                                     -0.358 2.56e-05
## 651
           2468052
                        11 -0.1011
                                     -0.335 2.87e-04
## 1079
            75071
                       124 -0.0301
                                      -0.335 3.23e-05
## 1013
                       101 -0.0324
                                     -0.325 2.33e-05
            72163
## 1015
            72172
                      101 -0.0324
                                     -0.325 2.33e-05
                       105 -0.0316
## 1018
            72203
                                      -0.324 2.72e-05
## 439
            189445
                        14 0.0972
                                       0.364 2.26e-04
## 467
           191273
                        22 0.1012
                                       0.475 1.42e-04
## 928
                       176 0.0367
                                       0.486 7.30e-04
           597592
                        20 0.1112
                                       0.497 6.37e-04
## 169
           156590
           2024096
                        20 0.1115
                                       0.499 1.38e-04
## 539
                        49 0.0730
## 165
           156580
                                       0.511 5.04e-04
## 505
           196849
                        53 0.0737
                                       0.537 6.85e-04
## 506
           196854
                        53 0.0737
                                       0.537 6.85e-04
## 731
           381070
                        46
                            0.0829
                                       0.563 2.51e-04
## 130
          1430728
                       1319 0.0201
                                       0.730 2.11e-05
##
                                                            reactomeName padjust
## 878
                                 Homo sapiens: Smooth Muscle Contraction 0.03386
## 1137
                                      Homo sapiens: Interferon Signaling 0.02443
## 104
                       Homo sapiens: Cytokine Signaling in Immune system 0.03055
## 69
                                 Homo sapiens: ISG15 antiviral mechanism 0.03711
## 70
               Homo sapiens: Antiviral mechanism by IFN-stimulated genes 0.03711
## 552
          Homo sapiens: YAP1- and WWTR1 (TAZ)-stimulated gene expression 0.02443
## 1066
                                             Homo sapiens: Transcription 0.02443
## 1044
                           Homo sapiens: RNA Polymerase II Transcription 0.00641
## 651
                Homo sapiens: Establishment of Sister Chromatid Cohesion 0.02443
## 1079
                                           Homo sapiens: mRNA Processing 0.00641
## 1013
                             Homo sapiens: mRNA Splicing - Major Pathway 0.00641
## 1015
                                             Homo sapiens: mRNA Splicing 0.00641
## 1018
           Homo sapiens: Processing of Capped Intron-Containing Pre-mRNA 0.00641
## 439
                                  Homo sapiens: Metabolism of porphyrins 0.02443
## 467
                                  Homo sapiens: Cholesterol biosynthesis 0.02113
## 928
                   Homo sapiens: Post-translational protein modification 0.03783
## 169
                                   Homo sapiens: Glutathione conjugation 0.03711
## 539
                                        Homo sapiens: HS-GAG degradation 0.02113
## 165
                                      Homo sapiens: Phase II conjugation 0.03536
## 505
        Homo sapiens: Metabolism of water-soluble vitamins and cofactors 0.03711
## 506
                      Homo sapiens: Metabolism of vitamins and cofactors 0.03711
## 731
                     Homo sapiens: Activation of Chaperones by IRE1alpha 0.02443
## 130
                                                Homo sapiens: Metabolism 0.00641
```

Note that such lists need to be interpreted with care, and a grain of salt. Which of these

## 5.2 Nearest peak to a differentially expressed gene

The RNA-Seq experiment analyzed above provides a list of genes which have responded to a selective estrogen-receptor-beta agonist. We can investigate whether we find estrogen receptor binding sites in the vicinity of the gene with the highest fold induction. In order to match differentially expressed genes to other experiment data, we will use annotated binding sites of estrogen receptor alpha from the ENCODE project. It is not necessarily the case that these annotated binding sites are actually functional in the cell lines of the RNA-Seq experiment or biologically relevant as the alpha and beta subtypes are distinct proteins transcribed from different genes; here we only use these binding site data for demonstration purposes.

Let us consider a particular gene with a low p value. The rowData function provides us with all the information about the gene model; each of the exons is represented as a *GRanges*, and these are tied together as a *GRangesList*. We use the function range to extract the entire range of the gene, from the start of the left-most exon to the end of the right-most exon. This is all the information we need in order to find the nearest binding site.

```
deGeneID <- "ENSG00000099194"
res[deGeneID, ]
## DataFrame with 1 row and 8 columns
     baseMean log2FoldChange lfcSE stat pvalue
##
       ##
## ENSG00000099194 8794 0.421 0.0193
## padj symbol entrez
## <numeric> <character> <character>
## ENSG00000099194 4.69e-102 SCD 6319
deGene <- range(rowData(dds[deGeneID, ])[[1]])</pre>
names(deGene) <- deGeneID</pre>
deGene
## GRanges with 1 range and 0 metadata columns:
  seqnames ranges strand
<Rle> <IRanges> <Rle>
##
##
  ENSG00000099194 10 [102106881, 102124591]
##
##
## seqlengths:
                        2 ...
                                           LRG_98
                                                          LRG_99
##
          249250621
                                            18750
                                                           13294
##
                         243199373 ...
```

We would like to compare the location of this gene with the location of annotated estrogen receptor binding sites, provided by the UCSC Genome Browser. We must first alter the sequence name (the chromosome name) of the differentially expressed gene, as the Ensembl gene annotation does not use the "chr" prefix, which the UCSC chromosomes are annotated

with. (Note that we ignore here another complication, which is that the Ensembl sequence "MT" corresponds to the UCSC's sequence "chrM".) We use the paste0 function, which concatenates the character vectors provided without using any separating characters. We then create a range which is 10 Mb to the left and right of the start of the deGene object.

```
as.character(seqnames(deGene))
## [1] "10"
ucscChrom <- paste0("chr", as.character(seqnames(deGene)))</pre>
ucscRanges <- ranges(flank(deGene, width = 1e+07, both = TRUE))
subsetRange <- GRanges(ucscChrom, ucscRanges)</pre>
subsetRange
## GRanges with 1 range and 0 metadata columns:
##
                seqnames ranges strand
                   <Rle> <IRanges> <Rle>
##
## ENSG00000099194 chr10 [92106881, 112106880]
##
## seqlengths:
## chr10
##
        NA
```

We now provide code which would download a track from the UCSC Genome Browser, in our case a track containing transcription factor binding sites obtained from ChIP-Seq experiments across various cell lines, generated by the ENCODE project.

The track names and table names must match a track name provided by the UCSC Genome Browser. For more information on these steps, see the detailed instructions in the vignette of the useful Bioconductor package *rtracklayer*.

Here we use a locally cached copy of ucscTable:

```
ucscTableFile <- system.file("extdata/localUcscTable.csv.gz", package = "BiocBrazil2014")
ucscTable <- read.csv(gzfile(ucscTableFile), stringsAsFactors = FALSE)</pre>
```

We now can use the downloaded table of annotated estrogen receptor peaks. Whether to use a cutoff on the provided peak scores at this step, or what scores cutoff to use, depends on your

experience with the specific transcription factor and the ChIP-Seq experiments used to define these peaks. It often makes sense to visualize tracks in a genome browser in order to get a sense of the qualitative difference between peaks of different scores.

We create a *GRanges* object, peaks, from the table obtained from UCSC, and then we convert the chromosome names back to the Ensembl style using the global substitute function, gsub. Finally, we enforce that the sequence levels of the peaks match the sequence levels of the differential expressed gene, which is necessary for performing the nearest matching in the following code chunk.

```
peaks <- with(ucscTable, GRanges(chrom, IRanges(chromStart, chromEnd), score = score))
seqlevels(peaks) <- gsub("chr(.+)", "\\1", seqlevels(peaks))
seqlevels(peaks) <- seqlevels(deGene)</pre>
```

Now we have two *GRanges* objects, defined over the same chromosomes, so we can use the distanceToNearest function from the package *GRanges*. This provides a *Hits* object, which contains the matches between the "query" and the "subject", the first and second arguments to the function, as well as the distance from the query to the subject. As we only have a single query, there should only be one nearest range in the subject. See the documentation via <code>?distanceToNearest</code> and <code>?Hits</code> for more information on the options for the this matching step.

```
d2nearest <- distanceToNearest(deGene, peaks)
```

**Question 9**: What is the distance from the differentially expressed gene to all the peaks? **Answer 9**:

```
distance(deGene, peaks)
```

```
[1] 9475047 9429128 9425728 9416450 9378087 9377697 9292445 9147971 9052350
##  [10] 8985898 8790054 8757486 8746708 8742787 8675294 8674780 8664979 8656386
## [19] 8482847 8223783 8221925 8135509 8001210 7455663 7085829 6942799 6925411
## [28] 6901313 6888089 6884998 6881075 6880706 6864676 6780255 6775784 6605487
## [37] 6599859 6596686 6588697 6583718 6571499 6420064 6351948 6351146 6329893
## [46] 6319663 6310500 6310175 6308781 6304333 6293607 6288353 6271339 6268796
## [55] 6262968 5922099 5213835 5117195 5058165 5057764 5038070 4950763 4846639
   [64] 4840470 4690192 4604930 4150399 4075280 3867684 3837681 3750152 3728198
## [73] 3713689 3672012 3545140 3489164 3483060 3482656 3374532 3145464 3009849
## [82] 3009475 2945584 2938228 2898743 2780415 2775156 2772196 2766543 2697779
## [91] 2134525 2099928 2093216 2080614 2079696 2078669 2077381 2074629 2052830
## [100] 2050846 2044996 2044079 2043235 2032452 1972277 1970323 1864919 1843269
## [109] 1397487 951766 778762 566972 415890 336981 235041
                                                                  45754
## [118]
             44 518646 536372 624809 727175 759664 988971 1236678 1267088
## [127] 1482084 1528951 1573655 1600301 1752606 1755604 1767986 1810760 1842340
## [136] 1962210 2003350 2039054 2048291 2137914 2293613 2342835 2345825 2347756
## [145] 2399792 2400158 2404602 2409590 2489595 2791121 2803606 2806523 2866416
## [154] 2867474 4107727 4108577 4109757 4112806 4115469 4116070 4198736 7246349
## [163] 9538145 9592618 9593558 9704719 9860874 9909237
```

We can now examine the object d2nearest. This tells us that the nearest peak is 44 base pairs from the differential expressed gene.

```
## Hits of length 1
## queryLength: 1
## subjectLength: 168
## queryHits subjectHits distance
## <integer> <integer> <integer>
## 1 1 118 44
```

The function subjectHits is used to extract the index of the closest hit in the peaks object.

```
deGene
## GRanges with 1 range and 0 metadata columns:
    seqnames ranges strand
<Rle> <IRanges> <Rle>
##
##
## ENSG00000099194 10 [102106881, 102124591]
##
## seqlengths:
          1 2 ...
249250621 243199373 ...
##
                                            LRG_98
                                                            LRG_99
                                              18750
                                                             13294
##
peaks[subjectHits(d2nearest)]
## GRanges with 1 range and 1 metadata column:
## seqnames ranges strand |
                <IRanges> <Rle> | <integer>
##
      <Rle>
## [1] 10 [102124636, 102124912] * | 76
    seqlengths:
##
##
                  1
                                2 ...
                                             LRG_98
                                                            LRG_99
##
                 NΑ
                               NA ...
```

Is 44 base pairs unexpectedly close? Here we make a simple plot of the starting points of the peaks and gene along the chromosome, to get a sense of the distribution of peaks and how surprised we should be with the distance of the nearest. To identify the nearest peak, we construct a logical vector <code>peakNearest</code>, which can be used to change the y value and the color of the point corresponding to the nearest peak.

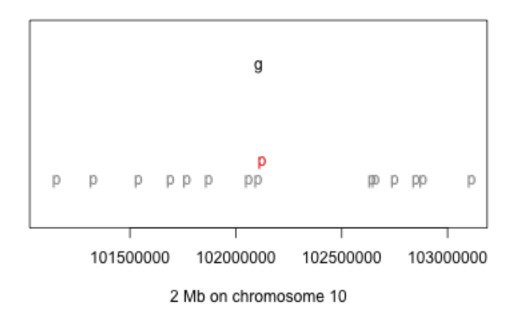


Figure 6: A 2 Mb genomic range showing the location of the differentially expressedgene (labelled 'g'), and the peaks (labelled 'p').

Again, the biological relevance of the distances between peaks and genes is another matter, especially considering the data are from different sources. An important consideration when investigating the distribution of distances between two sets of genomic features, is how the individual sets cluster along the genome.

**Question 10**: Are the peaks relatively uniformly distributed?

#### Answer 10:

We can answer this question by investigating the inter-peak distances. As all of our peaks are on the same chromosome, we just sort the peak starts and subtract the 2nd from the 1st, the 3rd from the 2nd, etc. Then we call the summary function which provides the mean and median. Note that the mean is constricted: it must be equal to the total span divided by the number of inter-peak distances. The median distance is about one quarter of the mean, so the peaks tend to cluster. You can also verify this by plotting the histogram of peakDists.

```
peakDists <- diff(sort(start(peaks)))
summary(peakDists)

## Min. 1st Qu. Median Mean 3rd Qu. Max.
## 364 5110 32500 116000 103000 3050000

mean(peakDists)</pre>
```

## 6 Working with rlog-transformed data

## 6.1 The rlog transform

Many common statistical methods for exploratory analysis of multidimensional data, especially methods for clustering and ordination (e.g., principal-component analysis and the like), work best for (at least approximately) homoskedastic data; this means that the variance of an observable (i.e., here, the expression strength of a gene) does not depend on the mean. In RNA-Seq data, however, variance grows with the mean. For example, if one performs PCA directly on a matrix of normalized read counts, the result typically depends only on the few most strongly expressed genes because they show the largest absolute differences between samples. A simple and often used strategy to avoid this is to take the logarithm of the normalized count values; however, now the genes with low counts tend to dominate the results because, due to the strong Poisson noise inherent to small count values, they show the strongest relative differences between samples.

As a solution, DESeq2 offers the regularized-logarithm transformation, or rlog for short. For genes with high counts, the rlog transformation differs not much from an ordinary  $\log_2$  transformation. For genes with lower counts, however, the values are shrunken towards the genes' averages across all samples. Using an empirical Bayesian prior in the form of a ridge penality, this is done such that the rlog-transformed data are approximately homoskedastic.

The function rlogTransform returns a *SummarizedExperiment* object which contains the rlog-transformed values in its *assay* slot:

```
rld <- rlogTransformation(dds)</pre>
## you had estimated gene-wise dispersions, removing these
## you had estimated fitted dispersions, removing these
head(assay(rld))
##
                 SRS308866 SRS308868 SRS308872 SRS308874 SRS308878 SRS308880
## ENSG0000000000 9.716 9.687 9.133 9.190 8.960 8.871
                                   -0.629
## ENSG0000000005
                   -0.679
                          -0.475
                                             -0.782
                                                      -0.783
                                                               -0.795
                                              8.290
## ENSG0000000419
                   8.100
                            8.110
                                     8.240
                                                       8.302
                                                                8.307
## ENSG0000000457
                   7.444
                            7.302
                                     7.816
                                             7.709
                                                       7.222
                                                                7.343
                            7.671
                                             8.171
## ENSG0000000460
                   7.573
                                     7.987
                                                       7.135
                                                                7.453
## ENSG0000000938 3.235
                          3.075
                                     4.180
                                             3.672
                                                       2.966
                                                                3.403
##
               SRS308883 SRS308885
## ENSG0000000000 9.094
                          9.114
## ENSG0000000000 -0.783 -0.793
```

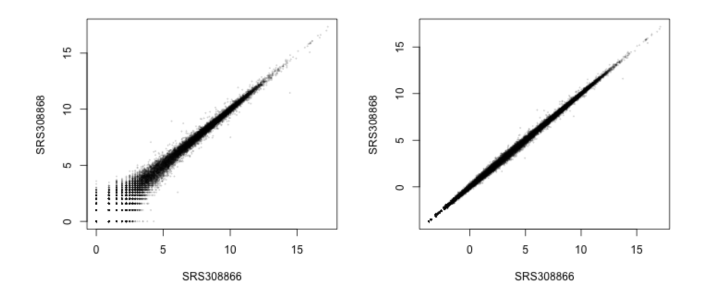


Figure 7: Scatter plot of sample 2 versus sample 1. Left: using an ordinary  $\log_2$  transformation. Right: Using the rlog transformation.

```
## ENSG00000000419 8.264 8.172

## ENSG00000000457 7.601 7.495

## ENSG00000000460 7.017 7.375

## ENSG000000000938 3.368 3.500
```

To show the effect of the transformation, we plot the first sample against the second, first simply using the log2 function (after adding 1, to avoid taking the log of zero), and then using the rlog-transformed values.

Note that, in order to make it easier to see where several points are plotted on top of each other, we set the plotting color to a semi-transparent black (encoded as #00000020) and changed the points to solid disks (pch=20) with reduced size (cex=0.3)<sup>1</sup>.

In Figure 7, we can see how genes with low counts seem to be excessively variable on the ordinary logarithmic scale, while the rlog transform compresses differences for genes for which the data cannot provide good information anyway.

<sup>&</sup>lt;sup>1</sup>The function heatscatter from the package *LSD* offers a colourful alternative.

## 6.2 Sample distances

A useful first step in an RNA-Seq analysis is often to assess overall similarity between samples: Which samples are similar to each other, which are different? Does this fit to the expectation from the experiment's design?

We use the R function dist to calculate the Euclidean distance between samples. To avoid that the distance measure is dominated by a few highly variable genes, and have a roughly equal contribution from all genes, we use it on the rlog-transformed data:

```
sampleDists <- dist(t(assay(rld)))</pre>
sampleDists
##
           SRS308866 SRS308868 SRS308872 SRS308874 SRS308878 SRS308880 SRS308883
## SRS308868
              27.6
              70.4
## SRS308872
                        72.7
                      71.5
## SRS308874
              70.3
                               27.8
              91.8
                      94.4
## SRS308878
                                 91.8
                                         93.9
              91.8
                      93.8
                                         94.3
                                                  26.8
## SRS308880
                                92.9
## SRS308883
              79.9
                      82.7
                                76.6
                                         78.4
                                                  69.4
                                                           69.9
## SRS308885
              78.0
                      79.8
                                75.2
                                         76.1
                                                  71.0
                                                           70.0
                                                                    27.4
```

Note the use of the function t to transpose the data matrix. We need this because dist calculates distances between data *rows* and our samples constitute the columns.

We visualize the distances in a heatmap, using the function heatmap. 2 from the gplots package.

```
sampleDistMatrix <- as.matrix(sampleDists)
rownames(sampleDistMatrix) <- paste(colData(rld)$treatment, colData(rld)$patient,
    sep = "-")
library("gplots")
heatmap.2(sampleDistMatrix, trace = "none")</pre>
```

Note that we have changed the row names of the distance matrix to contain treatment type and patient number instead of sample ID, so that we have all this information in view when looking at the heatmap (Fig. 8).

**Question 11**: Some people find the colour scheme used in Figure 8 ugly. Make a better version. *Hint:* Look at the sequential colour schemes in the *RColorBrewer* package and at the colorRampPalette function.

#### Answer 11:

```
library("RColorBrewer")
colours = colorRampPalette(rev(brewer.pal(9, "Blues")))(255)
heatmap.2(sampleDistMatrix, trace = "none", col = colours)
```

See Figure 9.

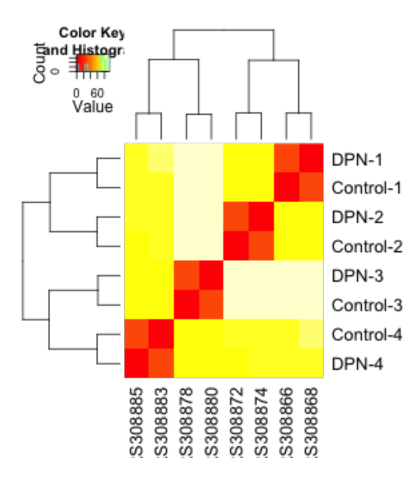


Figure 8: Heatmap of Euclidean sample distances after rlog transformation.

Another way to visualize sample-to-sample distances is a principal-components analysis (PCA). In this ordination method, the data points (i.e., here, the samples) are projected onto the 2D plane such that they spread out optimally (Fig. 10).

```
print(plotPCA(rld, intgroup = c("patient", "treatment")))
```

Here, we have used the function plotPCA which comes with *DESeq2*. The two terms specified as intgroup are column names from our sample data; they tell the function to use them to choose colours.

From both visualizations, we see that the differences between patients is much larger than the difference between treatment and control samples of the same patient. This shows why it was important to account for this paired design ("paired", because each treated sample is paired with one control sample from the *same* patient). We did so by using the design formula !~ patient treatment! when setting up the data object in the beginning. Had we used an unpaired analysis, by specifying only ~ treatment, we would not have found many hits, because then, the patient-to-patient differences would have drowned out any treatment effects.

Here, we have performed this sample distance analysis towards the end of our analysis. In practice, however, this is a step suitable to give a first overview on the data. Hence, one will

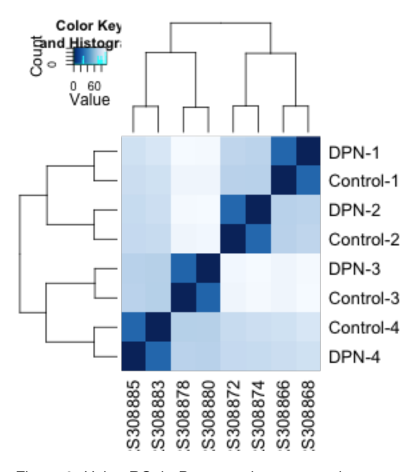


Figure 9: Using RColorBrewer to improve our heatmap

typically carry out this analysis as one of the first steps in an analysis. To this end, you may also find the function <code>arrayQualityMetrics</code>, from the equinymous package, useful.

## 6.3 Gene clustering

In the heatmap of Fig. 8, the dendrogram at the side shows us a hierarchical clustering of the samples. Such a clustering can also be performed for the genes.

Since the clustering is only relevant for genes that actually carry signal, one usually carries it out only for a subset of most highly variable genes. Here, for demonstration, let us select the 35 genes with the highest variance across samples:

```
library("genefilter")
topVarGenes <- head(order(rowVars(assay(rld)), decreasing = TRUE), 35)</pre>
```

The heatmap becomes more interesting if we do not look at absolute expression strength but rather at the amount by which each gene deviates in a specific sample from the gene's average across all samples. Hence, we center and scale each genes' values across samples, and plot a heatmap.

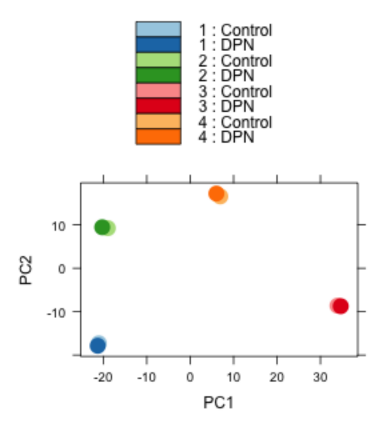


Figure 10: The same heatmap as in Figure 8 but with better colours

We can now see (Fig. 11) blocks of genes which covary across patients. Often, such a heatmap is insightful, even though here, seeing these variations across patients is of limited value because we are rather interested in the effects between the two samples from each patient.

## 7 Advanced Questions

For these questions, we provide (and probably have) no solutions, advanced readers are encouraged to explore them.

1. DESeq2 performs the shrinkage of the dispersion estimates by fitting a parametric curve on the mean of normalized counts (cf. Figure 2). However, one could argue that the biological variability of genes should not be a function of counts, but of counts per gene length (i. e., expression level), and that regression on that covariate should lead to a better fit. Write your own version of the estimateDispersions function to explore this question.

2. What is the contribution of UTR length variations to the between-replicates variability modelled by *DESeq2*? The read counting script (available in the vignette of *parathyroidSE*) uses all exons of the genes, which includes UTRs. Would detection power be increased –or would we preferentially detect different phenomena– if we left out UTRs from the counting (i. e. count reads that fall on coding exons only); or indeed, if we looked only at UTRs?

## References

[1] Richard Bourgon, Robert Gentleman, and Wolfgang Huber. Independent filtering increases detection power for high-throughput experiments. *PNAS*, 107(21):9546–9551, 2010.

## 9 Session Info

As last part of this document, we call the function <code>sessionInfo</code>, which reports the version numbers of R and all the packages used in this session. It is good practice to always keep such a record as it will help to trace down what has happened in case that an R script ceases to work because a package has been changed in a newer version.

```
## R version 3.0.2 Patched (2014-01-22 r64855)
## Platform: x86_64-apple-darwin10.8.0 (64-bit)
## locale:
## [1] C
## attached base packages:
## [1] grid parallel stats graphics grDevices utils
                                                                  datasets
## [8] methods base
##
## other attached packages:
## [1] rtracklayer_1.21.12
## [2] reactome.db_1.44.0
## [3] randomForest_4.6-7
## [4] pheatmap_0.7.7
## [5] org.Hs.eg.db_2.10.1
## [6] gplots_2.11.3
## [7] gtools_3.1.0
## [8] ggplot2_0.9.3.1
## [9] genefilter_1.43.0
## [10] gdata_2.13.2
## [11] devtools_1.3
## [12] caTools_1.14
## [13] VariantAnnotation_1.7.47
## [14] TxDb.Hsapiens.UCSC.hg19.knownGene_2.10.1
## [15] ShortRead_1.19.13
## [16] SRAdb_1.15.0
## [17] graph_1.39.3
## [18] SNPlocs.Hsapiens.dbSNP.20120608_0.99.9
## [19] Rsamtools_1.13.46
## [20] RSQLite_0.11.4
## [21] RCurl_1.95-4.1
## [22] bitops_1.0-6
## [23] RColorBrewer_1.0-5
## [24] MASS_7.3-29
## [25] KernSmooth_2.23-10
## [26] Gviz_1.5.15
## [27] GenomicFeatures_1.13.43
## [28] GEOquery_2.28.0
## [29] DBI_0.2-7
## [30] BiocBrazil2014_1.0
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## [31] parathyroidSE_1.0.4
## [32] DESeq2_1.2.9
## [33] RcppArmadillo_0.3.910.0
## [34] Rcpp_0.10.4
## [35] minfiData_0.4.2
## [36] IlluminaHumanMethylation450kanno.ilmn12.hg19_0.2.1
## [37] IlluminaHumanMethylation450kmanifest_0.4.0
## [38] minfi_1.8.9
## [39] bumphunter_1.1.17
## [40] locfit_1.5-9.1
## [41] iterators_1.0.6
## [42] foreach_1.4.1
## [43] reshape_0.8.4
## [44] plyr_1.8
## [45] lattice_0.20-24
## [46] BSgenome. Hsapiens. UCSC. hg19_1.3.19
## [47] BSgenome_1.29.1
## [48] Biostrings_2.29.19
## [49] GenomicRanges_1.13.45
## [50] XVector_0.1.4
## [51] IRanges_1.19.38
## [52] AnnotationDbi_1.24.0
## [53] ALL_1.4.14
## [54] Biobase_2.21.7
## [55] BiocGenerics_0.7.5
## [56] knitr_1.5
## [57] BiocInstaller_1.12.0
##
## loaded via a namespace (and not attached):
                                                     XML_3.95-0.2
## [1] Hmisc_3.12-2
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## [4] annotate_1.39.0
                              base64_1.1
                                                     beamplot_1.1
## [7] biomaRt_2.17.3
                              biovizBase_1.9.4
                                                     cluster_1.14.4
## [10] codetools_0.2-8
                              colorspace_1.2-3
                                                     compiler_3.0.2
## [13] dichromat_2.0-0
                              digest_0.6.3
                                                     doRNG_1.5.5
## [16] evaluate_0.5
                              formatR_0.9
                                                     gtable_0.1.2
## [19] highr_0.2.1
                              httr_0.2
                                                     hwriter_1.3
## [22] illuminaio_0.3.11
                              itertools_0.1-1
                                                     labeling_0.2
## [25] latticeExtra_0.6-26
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                                                     matrixStats_0.8.12
## [28] mclust_4.2
                              memoise_0.1
                                                     multtest_2.17.0
## [31] munsell_0.4.2
                              nlme_3.1-113
                                                     nor1mix_1.1-4
## [34] pkgmaker_0.17.4
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## [37] registry_0.2
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                                                     rngtools_1.2.3
## [40] rpart_4.1-4
                              scales_0.2.3
                                                     siggenes_1.35.0
                              stats4_3.0.2
## [43] splines_3.0.2
                                                     stringr_0.6.2
## [46] survival_2.37-7
                              tools_3.0.2
                                                     whisker_0.3-2
## [49] xtable_1.7-1
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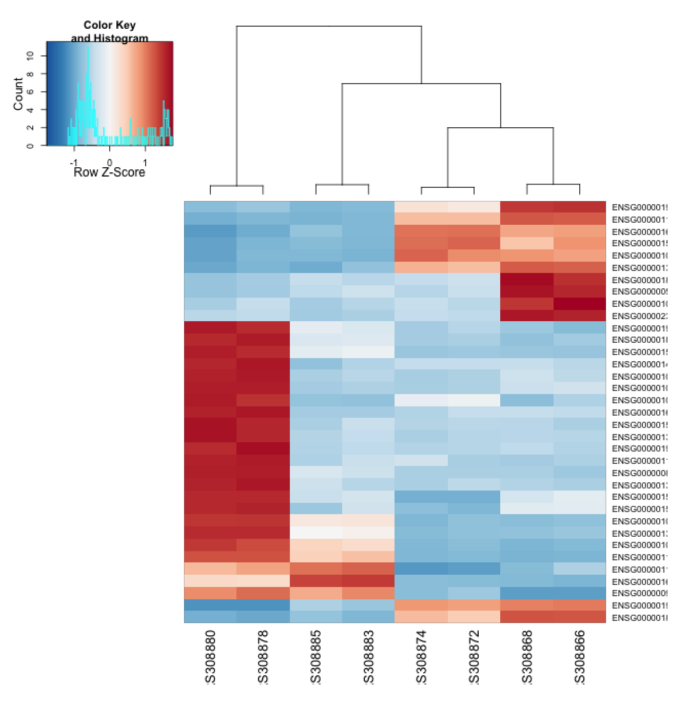


Figure 11: Heatmap with gene clustering.