Differential expression of RNA-Seq data at the gene level – the DESeq package

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Abstract

A basic task in the analysis of count data from RNA-Seq is the detection of differentially expressed genes. The count data are presented as a table which reports, for each sample, the number of reads that have been assigned to a gene. Analogous analyses also arise for other assay types, such as comparative ChIP-Seq. The package *DESeq* provides methods to test for differential expression by use of the negative binonial distribution and a shrinkage estimator for the distribution's variance¹. This vignette explains the use of the package. For an exposition of the statistical method, please see our paper [1] and the additional information in Section 9.

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¹Other Bioconductor packages with similar aims are edgeR and baySeq.

10 Session Info

1 Input data and preparations

1.1 The count table

As input, the DESeq package expects count data as obtained, e.g., from RNA-Seq or another high-throughput sequencing experiment, in the form of a rectangular table of integer values. The table cell in the i-th row and the j-th column of the table tells how many reads have been mapped to gene i in sample j. Analogously, for other types of assays, the rows of the table might correspond e.g. to binding regions (with ChIP-Seq) or peptide sequences (with quantitative mass spectrometry).

The count values must be raw counts of sequencing reads. This is important for *DESeq*'s statistical model to hold, as only the actual counts allow assessing the measurement precision correctly. Hence, please do do not supply other quantities, such as (rounded) normalized counts, or counts of covered base pairs – this will only lead to nonsensical results.

Furthermore, it is important that each column stems from an independent biological replicate. For technical replicates (e.g. when the same library preparation was distributed over multiple lanes of the sequencer), please sum up their counts to get a single column, corresponding to a unique biological replicate. This is needed in order to allow *DESeq* to estimate variability in the experiment correctly.

To obtain such a count table for your own data, you will need to create it from your sequence alignments and suitable annotation. In Bioconductor, you can use the function summarizeOverlaps in the *GenomicRanges* package. See the vignette, reference [2], for a worked example. Another possibility is provided by the Bioconductor package *easyRNASeq*.

Another easy way to produce such a table from the output of the aligner is to use the htseq-count script distributed with the *HTSeq* Python package. Even though *HTSeq* is written in Python, you do not need to know any Python to use htseq-count. See http://www-huber.embl.de/users/anders/HTSeq/doc/count.html. HTSeq-count produces one count file for each sample. *DESeq* offers the function newCountDataSetFromHTSeqCount, described below, to get an analysis started from these files.

In this vignette, we will work with the gene level read counts from the pasilla data package. This data set is from an experiment on Drosophila melanogaster cell cultures and investigated the effect of RNAi knock-down of the splicing factor pasilla [3]. The detailed transcript of how we produced the pasilla count table from second generation sequencing (Illumina) FASTQ files is provided in the vignette of the data package pasilla. The table is supplied by the pasilla package as a text file of tab-separated values. The function system.file tells us where this file is stored on your computer.

```
> datafile = system.file( "extdata/pasilla_gene_counts.tsv", package="pasilla" )
> datafile
```

[1] "/home/biocbuild/bbs-3.0-bioc/R/library/pasilla/extdata/pasilla_gene_counts.tsv"

Have a look at the file with a text editor to see how it is formatted. To read this file with R, we use the function read.table.

```
> pasillaCountTable = read.table( datafile, header=TRUE, row.names=1 )
> head( pasillaCountTable )
```

	untreated1	untreated2	untreated3	untreated4	treated1	treated2	treated3
FBgn0000003	0	0	0	0	0	0	1
FBgn0000008	92	161	76	70	140	88	70
FBgn0000014	5	1	0	0	4	0	0
FBgn0000015	0	2	1	2	1	0	0
FBgn0000017	4664	8714	3564	3150	6205	3072	3334
FBgn0000018	583	761	245	310	722	299	308

Here, header=TRUE indicates that the first line contains column names and row.names=1 means that the first column should be used as row names. This leaves us with a *data.frame* containing integer count values.

1.2 The metadata

The best data are useless without metadata. *DESeq* uses familiar idioms in Bioconductor to manage the metadata that go with the count table. The metadata can be divided into three groups: information about the samples (table colums), about the features (table rows), and about the overall experiment.

First, we need a description of the samples, which we keep in a *data.frame* whose columns correspond to different types of information, and whose rows correspond to the 7 samples:

```
> pasillaDesign = data.frame(
    row.names = colnames( pasillaCountTable ),
+
    condition = c( "untreated", "untreated", "untreated",
        "untreated", "treated", "treated"),
    libType = c( "single-end", "single-end", "paired-end",
        "paired-end", "single-end", "paired-end", "paired-end"))
> pasillaDesign
          condition
                       libType
untreated1 untreated single-end
untreated2 untreated single-end
untreated3 untreated paired-end
untreated4 untreated paired-end
            treated single-end
treated1
treated2
            treated paired-end
treated3
            treated paired-end
```

Here, simply use a chunk of R code to set up this define. More often, you will read this into R from a spreadsheet table, using the R functions read.table or read.csv; or perhaps even from a relational database table using R's database access facilities.

To analyse these samples, we will have to account for the fact that we have both single-end and paired-end method. To keep things simple at the start, we defer the discussion of this to Section 4 and first demonstrate a simple analysis by using only the paired-end samples.

```
> pairedSamples = pasillaDesign$libType == "paired-end"
> countTable = pasillaCountTable[ , pairedSamples ]
> condition = pasillaDesign$condition[ pairedSamples ]
```

Now, we have data input as follows.

> head(countTable)

	untreated3	untreated4	treated2	treated3
FBgn0000003	0	0	0	1
FBgn0000008	76	70	88	70
FBgn0000014	0	0	0	0
FBgn0000015	1	2	0	0
FBgn0000017	3564	3150	3072	3334
FBgn0000018	245	310	299	308

> condition

[1] untreated untreated treated treated Levels: treated untreated

For your own data, create such a factor simply with

```
> #not run
> condition = factor( c( "untreated", "untreated", "treated", "treated" ) )
```

We can now instantiate a CountDataSet, which is the central data structure in the DESeq package:

```
> library( "DESeq" )
> cds = newCountDataSet( countTable, condition )
```

If you have used <code>htseq-count</code> (see above) to obtain your read counts, you can use the funtion <code>newCountDataSetFromHTSeqCount</code> to obtain a CountDataSe directly from the count files produced by <code>htseq-count</code>. To this end, produce a data frame, similar to <code>pasillaDesign</code> constructed above, with the sample names in the first column, the file names in the scond column, and the condition in the third column (or, if you have more than one factor in your design matrix, all these factors in the third and folling columns), and pass this data frame to the function. See the help page ("?newCountDataSetFromHTSeqCount") for further details.

The *CountDataSet* class is derived from *Biobase's eSet* class and so shares all features of this standard Bioconductor class. Furthermore, accessors are provided for its data slots². For example, the counts can be accessed with the counts function, as we will see in the next section.

1.3 Normalisation

As a first processing step, we need to estimate the effective library size. This step is sometimes also called *normalisation*, even though there is no relation to normality or a normal distribution. The effective library size information is called the *size factors* vector, since the package only needs to know the relative library sizes. If the counts of non-differentially expressed genes in one sample are, on average, twice as high as in another (because the library was sequenced twice as deeply), the size factor for the first sample should be twice that of the other sample [1, 4]. The function estimateSizeFactors estimates the size factors from the count data. (See the man page of estimateSizeFactorsForMatrix for technical details on the calculation.)

If we divide each column of the count table by the size factor for this column, the count values are brought to a common scale, making them comparable. When called with normalized=TRUE, the counts accessor function performs this calculation. This is useful, e.g., for visualization.

> head(counts(cds, normalized=TRUE))

	${\tt untreated3}$	${\tt untreated 4}$	${\tt treated2}$	${\tt treated3}$
FBgn0000003	0.00	0.00	0.0	0.897
FBgn0000008	87.05	69.27	86.1	62.803
FBgn0000014	0.00	0.00	0.0	0.000
FBgn0000015	1.15	1.98	0.0	0.000
FBgn0000017	4082.02	3116.93	3004.5	2991.238
FBgn0000018	280.61	306.75	292.4	276.335

2 Variance estimation

The inference in *DESeq* relies on an estimation of the typical relationship between the data's variance and their mean, or, equivalently, between the data's dispersion and their mean.

The dispersion can be understood as the square of the coefficient of biological variation. So, if a gene's expression typically differs from replicate to replicate sample by 20%, this gene's dispersion is $0.2^2=.04$. Note that the variance seen between counts is the sum of two components: the sample-to-sample variation just mentioned, and the uncertainty in measuring a concentration by counting reads. The latter, known as shot noise or Poisson noise, is the dominating noise source for lowly expressed genes. The former dominates for highly expressed genes. The sum of both, shot noise and dispersion, is considered in the differential expression inference.

Hence, the variance v of count values is modelled as

$$v = s\mu + \alpha s^2 \mu^2,$$

where μ is the expected normalized count value (estimated by the average normalized count value), s is the size factor for the sample under consideration, and α is the dispersion value for the gene under consideration.

To estimate the dispersions, use this command.

```
> cds = estimateDispersions( cds )
```

We could now proceed straight to the testing for differential expression in Section 3. However, let us first spend a little more time with looking at the results of estimateDispersions, in order to better understand the subsequent inference. Also, verification of the dispersion plot Figure 1, explained below, is an aspect of data quality control: how well do the data accord to the expectations of our analytic approach? Quality assessment is a crucial step of the data analysis, and we will see further quality diagnostics in Section 7.

²In fact, the object pasillaGenes from the *pasilla* package is of class *CountDataSet*, and we could do the subsequent analysis on the basis of this object. Here we re-create cds from elementary data types, a *data.frame* and a *factor*, for pedagogic effect.

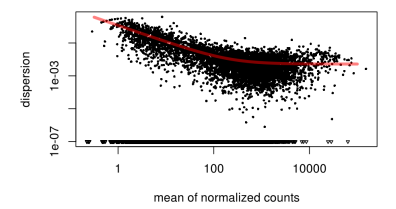


Figure 1: Empirical (black dots) and fitted (red lines) dispersion values plotted against the mean of the normalised counts.

The function estimateDispersions performs three steps. First, it estimates a dispersion value for each gene, then it fits a curve through the estimates. Finally, it assigns to each gene a dispersion value, using a choice between the per-gene estimate and the fitted value. To allow the user to inspect the intermediate steps, a fitInfo object is stored, which contains the per-gene estimate, the fitted curve and the values that will subsequently be used for inference.

It is useful to inspect the results of these steps visually. We can plot the per-gene estimates against the mean normalized counts per gene and overlay the fitted curve by using the function plotDispEsts (Fig. 1).

```
> plotDispEsts( cds )
```

As we estimated the dispersions from a small number of replicates, the estimates scatter with quite some sampling variance around their true values. An initial assumption that one could make is that the regression line shown in Figure 1 models the true underlying dispersions, and that the variation of the point estimates around simply reflects sampling variance. This is the assumption that we put forward in the first paper on DESeq [1]. However, subsequent experience with larger data sets indicates that not all of the variability of the points around the regression line seen in Figure 1 is sampling variance: some of it reflects differences of the true, underlying variance between different genes. Hence, the default behaviour of DESeq now uses a more prudent or conservative approach: if a per-gene estimates lies below the regression line, we assume that this might indeed be sampling variance, and shift the estimate upwards to the value predicted by the regression line. If, however, the per-gene estimate lies above the line, we do not shift it downwards to the line, but rather keep it as is.

The option sharingMode of the function estimateDispersions can be used to control this behaviour. The value sharingMode="maximum" corresponds to the default. If you have many replicates (where *many* starts at around 7), the choice sharingMode="gene-est-only" will typically be more adequate. If you would like to use the behaviour described in [1], this is achieved by specifying sharingMode="fit-only".

Another difference of the current *DESeq* version to the original method described in the paper is the way how the mean-dispersion relation is fitted. By default, estimateDispersions now performs a parametric fit: Using a gamma-family GLM, two coefficients α_0 , α_1 are found to parametrize the fit as $\alpha = \alpha_0 + \alpha_1/\mu$. (The values of the

two coefficients can be found in the fitInfo object, as attribute coefficients to dispFunc.) For some data sets, the parametric fit may give bad results, in which case one should try a local fit (the method described in the paper), which is available via the option fitType="local" to estimateDispersions.

In any case, the dispersion values which will be used by the subsequent testing are stored in the feature data slot of cds:

> head(fData(cds))

	${\tt disp_pooled}$
FBgn0000003	5.21332
FBgn0000008	0.02368
FBgn0000014	Inf
FBgn0000015	1.50080
FBgn0000017	0.02110
FBgn0000018	0.00928

Fixme: Why is the value for FBgn00000014 Inf and not 0 (given that all counts are 0)? You can verify that these values are indeed the maxima from the two value vectors in fitInfo(cds), which we saw on page 5. Advanced users who want to fiddle with the dispersion estimation can change the values in fData(cds) prior to calling the testing function.

3 Inference: Calling differential expression

3.1 Standard comparison between two experimental conditions

Having estimated the dispersion for each gene, it is straight-forward to look for differentially expressed genes. To contrast two conditions, e.g., to see whether there is differential expression between conditions "untreated" and "treated", we simply call the function ${\tt nbinomTest}$. It performs the tests as described in [1] and returns a data frame with the p values and other useful information.

```
> res = nbinomTest( cds, "untreated", "treated" )
```

> head(res)

	id	${\tt baseMean}$	${\tt baseMeanA}$	${\tt baseMeanB}$	${\tt foldChange}$	log2FoldChange	pval	padj
1	FBgn0000003	0.224	0.00	0.449	Inf	Inf	1.000	1.000
2	FBgn0000008	76.296	78.16	74.436	0.952	-0.0704	0.835	1.000
3	FBgn0000014	0.000	0.00	0.000	NaN	NaN	NA	NA
4	FBgn0000015	0.781	1.56	0.000	0.000	-Inf	0.416	1.000
5	FBgn0000017	3298.682	3599.47	2997.890	0.833	-0.2638	0.241	0.881
6	FBgn0000018	289.031	293.68	284.385	0.968	-0.0464	0.757	1.000

The interpretation of the columns of data.frame is as follows.

id feature identifier

baseMean mean normalised counts, averaged over all samples from both conditions

baseMeanA mean normalised counts from condition A
baseMeanB mean normalised counts from condition B
foldChange fold change from condition A to B

foldChange fold change from condition A to B log2FoldChange the logarithm (to basis 2) of the fold change

p value for the statistical significance of this change

padj p value adjusted for multiple testing with the Benjamini-Hochberg procedure (see the R function

p.adjust), which controls false discovery rate (FDR)

Let us first plot the \log_2 fold changes against the mean normalised counts, colouring in red those genes that are significant at 10% FDR (Figure 2).

> plotMA(res)

It is also instructive to look at the histogram of p values (Figure 3). The enrichment of low p values stems from the differentially expressed genes, while those not differentially expressed are spread uniformly over the range from zero to one (except for the p values from genes with very low counts, which take discrete values and so give rise to high counts for some bins at the right.)

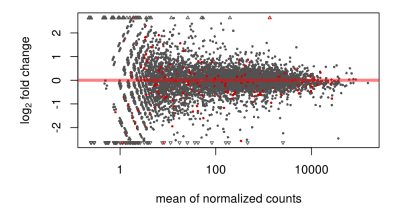


Figure 2: Plot of normalised mean versus log_2 fold change for the contrast untreated versus treated.

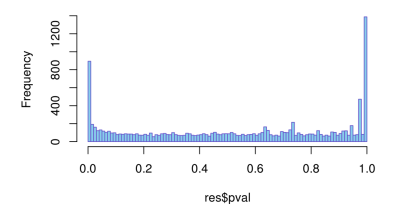


Figure 3: Histogram of p-values from the call to nbinomTest.

- > hist(res\$pval, breaks=100, col="skyblue", border="slateblue", main="")
 We can filter for significant genes, according to some chosen threshold for the false dicovery rate (FDR),
 > resSig = res[res\$padj < 0.1,]
 and list, e.g., the most significantly differentially expressed genes:</pre>
- > head(resSig[order(resSig\$pval),])

	id	baseMean	baseMeanA	baseMeanB	foldChange	log2FoldChange	pval	padj
9831	FBgn0039155	463	885	41.9	0.0474	-4.40	1.64e-124	1.89e-120
2366	FBgn0025111	1340	311	2369.3	7.6141	2.93	3.50e-107	2.01e-103
612	FBgn0003360	2544	4514	574.6	0.1273	-2.97	1.55e-99	5.95e-96
3192	FBgn0029167	2551	4211	891.7	0.2117	-2.24	4.35e-78	1.25e-74
10305	FBgn0039827	189	357	19.9	0.0556	-4.17	1.19e-65	2.74e-62
6948	FBgn0035085	447	761	133.3	0.1751	-2.51	3.15e-56	6.03e-53

We may also want to look at the most strongly down-regulated of the significant genes,

> head(resSig[order(resSig\$foldChange, -resSig\$baseMean),])

	id	baseMean	baseMeanA	baseMeanB	foldChange	log2FoldChange	pval	padj
14078	FBgn0259236	4.27	8.54	0.00	0.0000	-Inf	1.31e-03	2.60e-02
13584	FBgn0085359	36.47	71.03	1.92	0.0270	-5.21	2.40e-09	2.19e-07
9831	FBgn0039155	463.44	884.96	41.91	0.0474	-4.40	1.64e-124	1.89e-120
2277	FBgn0024288	42.56	80.89	4.24	0.0524	-4.25	5.57e-20	1.94e-17
10305	FBgn0039827	188.59	357.33	19.86	0.0556	-4.17	1.19e-65	2.74e-62
6495	FBgn0034434	82.89	155.09	10.68	0.0689	-3.86	5.94e-32	5.25e-29

or at the most strongly up-regulated ones:

```
> head( resSig[ order( -resSig$foldChange, -resSig$baseMean ), ] )
```

	id	baseMean	${\tt baseMeanA}$	${\tt baseMeanB}$	${\tt foldChange}$	${\tt log2FoldChange}$	pval	padj
6030	FBgn0033764	53.9	10.26	97.6	9.52	3.25	9.95e-18	2.93e-15
13079	FBgn0063667	11.8	2.55	21.0	8.23	3.04	1.43e-04	4.22e-03
7020	FBgn0035189	197.5	45.30	349.6	7.72	2.95	6.43e-15	1.35e-12
8499	FBgn0037290	50.5	11.66	89.2	7.65	2.94	1.04e-14	2.05e-12
2366	FBgn0025111	1340.2	311.17	2369.3	7.61	2.93	3.50e-107	2.01e-103
7264	FBgn0035539	16.0	4.19	27.8	6.63	2.73	6.76e-05	2.14e-03

To save the output to a file, use the R functions write.table and write.csv. (The latter is useful if you want to load the table in a spreadsheet program such as Excel.)

```
> write.csv( res, file="My Pasilla Analysis Result Table.csv" )
```

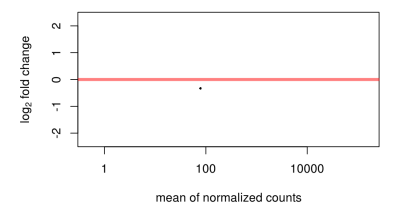


Figure 4: Plot of the log2 fold change between the untreated replicates versus average expression strength.

Note in Fig. 2 how the power to detect significant differential expression depends on the expression strength. For weakly expressed genes, stronger changes are required for the gene to be called significantly expressed. To understand the reason for this let, us compare the normalized counts between two replicate samples, here taking the two untreated samples as an example:

As one can see in Figure 4, the log fold changes between replicates are stronger for lowly expressed genes than for highly expressed ones. We ought to conclude that a gene's expression is influenced by the treatment only if the change between treated and untreated samples is stronger than what we see between replicates, and hence, the dividing line between red and black in Figure 2 mimics the shape seen in Figure 4.

3.2 Working partially without replicates

If you have replicates for one condition but not for the other, you can still proceed as before. In such cases only the conditions with replicates will be used to estimate the dispersion. Of course, this is only valid if you have good reason to believe that the unreplicated condition does not have larger variation than the replicated one.

To demonstrate, we subset our data object to only three samples:

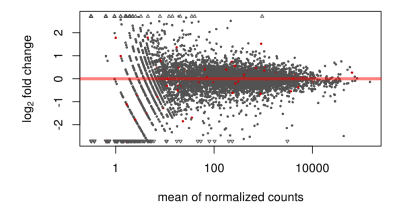


Figure 5: MvA plot for the contrast "treated" vs. "untreated", using two treated and only one untreated sample.

Now, we do the analysis as before.

```
> cdsUUT = estimateSizeFactors( cdsUUT )
> cdsUUT = estimateDispersions( cdsUUT )
> resUUT = nbinomTest( cdsUUT, "untreated", "treated" )
```

We produce the analogous plot as before, again with

```
> plotMA(resUUT)
```

Figure 5 shows the same symmetry in up- and down-regulation as in Fig. 2, but a certain asymmetry in the boundary line for significance. This has an easy explanation: low counts suffer from proportionally stronger shot noise than high counts, and since there is only one "untreated" sample versus two "treated" ones, a stronger downward fold-change is required to be called significant than for the upward direction.

3.3 Working without any replicates

Proper replicates are essential to interpret a biological experiment. After all, if one compares two conditions and finds a difference, how else can one know that this difference is due to the different conditions and would not have arisen between replicates, as well, just due to experimental or biological noise? Hence, any attempt to work without replicates will lead to conclusions of very limited reliability.

Nevertheless, such experiments are sometimes undertaken, and the *DESeq* package can deal with them, even though the soundness of the results may depend much on the circumstances.

Our primary assumption is now that the mean is a good predictor for the dispersion. Once we accept this assumption, we may argue as follows: Given two samples from different conditions and a number of genes with comparable expression levels, of which we expect only a minority to be influenced by the condition, we may take the

dispersion estimated from comparing their counts *across* conditions as ersatz for a proper estimate of the variance across replicates. After all, we assume most genes to behave the same within replicates as across conditions, and hence, the estimated variance should not be affected too much by the influence of the hopefully few differentially expressed genes. Furthermore, the differentially expressed genes will only cause the dispersion estimate to be too high, so that the test will err to the side of being too conservative.

We shall now see how well this works for our example data. We reduce our count data set to just two columns, one "untreated" and one "treated" sample:

```
> cds2 = cds[ ,c( "untreated3", "treated3" ) ]
```

Now, without any replicates at all, the estimateDispersions function will refuse to proceed unless we instruct it to ignore the condition labels and estimate the variance by treating all samples as if they were replicates of the same condition:

```
> cds2 = estimateDispersions( cds2, method="blind", sharingMode="fit-only")
```

Note the option sharingMode="fit-only". Remember that the default, sharingMode="maximum", takes care of outliers, i.e., genes with dispersion much larger than the fitted values. Without replicates, we cannot catch such outliers and so have to disable this functionality.

Now, we can attempt to find differential expression:

```
> res2 = nbinomTest( cds2, "untreated", "treated" )
```

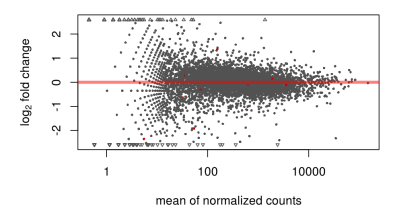


Figure 6: MvA plot, from a test using no replicates.

Unsurprisingly, we find much fewer hits, as can be seen from the plot (Fig. 6)

```
> plotMA(res2)
```

and from this table, tallying the number of significant hits in our previous and our new, restricted analysis:

```
> addmargins( table( res_sig = res$padj < .1, res2_sig = res2$padj < .1 ) )</pre>
```

```
res2_sig
res_sig FALSE TRUE Sum
FALSE 10084 1 10085
TRUE 773 48 821
Sum 10857 49 10906
```

4 Multi-factor designs

Let us return to the full pasilla data set. Remember that we started of with these data:

```
> head( pasillaCountTable )
```

	${\tt untreated1}$	${\tt untreated2}$	${\tt untreated3}$	${\tt untreated 4}$	treated1	${\tt treated2}$	treated3
FBgn0000003	0	0	0	0	0	0	1
FBgn0000008	92	161	76	70	140	88	70
FBgn0000014	5	1	0	0	4	0	0
FBgn0000015	0	2	1	2	1	0	0
FBgn0000017	4664	8714	3564	3150	6205	3072	3334
FBgn0000018	583	761	245	310	722	299	308

> pasillaDesign

```
condition libType
untreated1 untreated single-end
untreated2 untreated paired-end
untreated4 untreated paired-end
treated1 treated single-end
treated2 treated paired-end
treated3 treated paired-end
```

When creating a count data set with multiple factors, just pass a data frame instead of the condition factor:

```
> cdsFull = newCountDataSet( pasillaCountTable, pasillaDesign )
```

As before, we estimate the size factors and then the dispersions. Here, the default method (method=mpooled to estimateDispersion), means that, as before, one dispersion is computed for each gene, which is now an average over all cells (weighted by the number of samples for each cells), where the term *cell* denotes any of the four combinations of factor levels of the design. An alternative is the option method=mpooled-CRm, which uses a Cox-Reid-adjusted maximum likelihood estimator [5, 6]. The latter is slower but useful when the cell concept is not applicable (e.g. in paired designs).

```
> cdsFull = estimateSizeFactors( cdsFull )
> cdsFull = estimateDispersions( cdsFull )
```

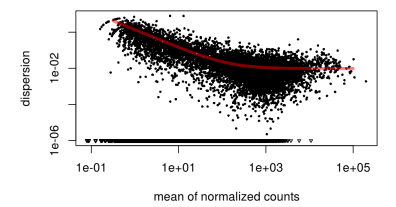


Figure 7: Estimated (black) pooled dispersion values for all seven samples, with regression curve (red).

```
We check the fit (Fig. 7):
> plotDispEsts( cdsFull )
```

For inference, we now specify two *models* by formulas. The *full model* regresses the genes' expression on both the library type and the treatment condition, the *reduced model* regresses them only on the library type. For each gene, we fit generalized linear models (GLMs) according to the two models, and then compare them in order to infer whether the additional specification of the treatment improves the fit and hence, whether the treatment has significant effect.

```
> fit1 = fitNbinomGLMs( cdsFull, count ~ libType + condition )
> fit0 = fitNbinomGLMs( cdsFull, count ~ libType )
```

These commands take a while to execute. Also, they may produce a few warnings, informing you that the GLM fit failed to converge (and the results from these genes should be interpreted with care). The "fit" objects are data frames with three columns:

Fixme: Can we add a paragraph on what to do if we only want to make specific comparisons, e.g. let fac be a factor with three levels A, B and C, and we want to test pairwise for differences between A and B (without C), between A and C (without B) etc.

The function nbinomTestGLM returned simply a vector of p values which we handed to the standard R function p.adjust to adjust for multiple testing using the Benjamini-Hochberg (BH) method.

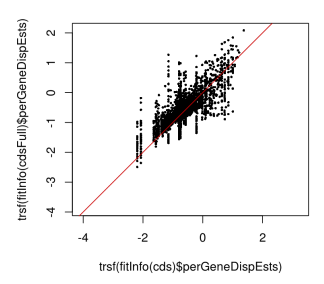


Figure 8: Comparison of per-gene estimates of the dispersion in the analysis using only the four paired-end samples (x-axis) and in the analysis using all seven samples (y-axis). For visualisation, the values are on a logarithm-like scale, defined by the function trsf, which is of the form $f(x) = \log \left((x + \sqrt{x^2 + 1})/2 \right)$. For x > 2, the values of f(x) and $\log(x)$ are approximately the same, whereas for values of x around 0, $f(x) \approx x$ is approximated by a straight line of slope 1.

Let's compare with the result from the four-samples test:

```
> tab1 = table( "paired-end only" = res$padj < .1, "all samples" = padjGLM < .1 )
> addmargins( tab1 )

all samples
paired-end only FALSE TRUE Sum
```

```
FALSE 10392 288 10680
TRUE 143 678 821
Sum 10535 966 11501
```

We see that the analyses find 678 genes in common, while 288 were only found in the analysis using all samples and 143 were specific for the *paired-end only* analysis.

In this case, the improvement in power that we gained from the additional samples is rather modest. This might indicate that the single-end samples (which are presumably older than the paired-end one) are of lower quality. This is supported by the observation that the dispersion estimates in the full data set tend to be larger than in the paired-end only one. So, despite the fact that with 7 samples we gain power from having more degrees of freedom in the estimation of the treatment effect, the additional samples also increase the dispersion (or equivalently, the variance), which loses power.

```
> table(sign(fitInfo(cds)$perGeneDispEsts - fitInfo(cdsFull)$perGeneDispEsts))
-1    1
8175 3326
See also Figure 8, which is produced by
> trsf = function(x) log( (x + sqrt(x*x+1))/2 )
> plot( trsf(fitInfo(cds)$perGeneDispEsts),
+ trsf(fitInfo(cdsFull)$perGeneDispEsts), pch=16, cex=0.45, asp=1)
> abline(a=0, b=1, col="red3")
```

Continuing with the analysis, we can now extract the significant genes from the vector padjGLM as before. To see the corresponding fold changes, we have a closer look at the object fit1.

> head(fit1)

```
(Intercept) libTypesingle-end conditionuntreated deviance converged
FBgn0000003
                -0.722
                                -31.082 -30.8575 0.452
                                                                        TRUF.
FBgn0000008
                 6.654
                                 -0.261
                                                    0.0405
                                                             2.275
                                                                        TRUE
FBgn0000014
               -30.278
                                 31.568
                                                   -0.0306
                                                             1.098
                                                                        TRUE
FBgn0000015
                -1.591
                                 -0.462
                                                    2.1028
                                                             2.759
                                                                        TRUE
FBgn0000017
                11.978
                                 -0.100
                                                    0.2562
                                                                        TRUE
                                                              2.557
FBgn0000018
                 8.568
                                  0.230
                                                    0.0647
                                                              1.585
                                                                        TRUE
```

The first three columns show the fitted coefficients, converted to a logarithm base 2 scale. The log2 fold change due to the condition is shown in the third column. As indicated by the column name, it is the effect of "untreated", i.e., the log ratio of "untreated" versus "treated". (This is unfortunately the other way round as before, due to the peculiarities of contrast coding.) Note that the library type also had noticeable influence on the expression, and hence would have increased the dispersion estimates (and so reduced power) if we had not fitted an effect for it.

The column deviance is the deviance of the fit. (Comparing the deviances with a χ^2 likelihood ratio test is how nbinomGLMTest calculates the p values.) The last column, converged, indicates whether the calculation of coefficients and deviance has fully converged. (If it is false too often, you can try to change the GLM control parameters, as explained in the help page to fitNbinomGLMs.) Fixme: This is a bit vague, 'too often' should be specified and ideally the remedy be automated.

Finally, we show that taking the library type into account was important to have good detection power by doing the analysis again using the standard workflow, as outlined earlier, and without informing *DESeq* of the library types:

```
> cdsFullB = newCountDataSet( pasillaCountTable, pasillaDesign$condition )
> cdsFullB = estimateSizeFactors( cdsFullB )
> cdsFullB = estimateDispersions( cdsFullB )
> resFullB = nbinomTest( cdsFullB, "untreated", "treated" )
> tab2 = table(
     `all samples simple` = resFullB$padj < 0.1,
     `all samples GLM`
                          = padjGLM < 0.1)
> addmargins(tab2)
                  all samples GLM
all samples simple FALSE TRUE
                                 Sum
             FALSE 11387
                           389 11776
             TRUE
                       6
                           577
                                 583
             Sum
                   11393
                           966 12359
```

We see that the two analyses find 577 genes in common, while in addition 389 genes were found by the analysis that took library type into account. The small number, namely 6, of genes that only made the FDR cutoff in the simple analysis is likely due to sampling variation.

5 Independent filtering and multiple testing

5.1 Filtering by overall count

The analyses of the previous sections involve the application of statistical tests, one by one, to each row of the data set, in order to identify those genes that have evidence for differential expression. The idea of *independent filtering* is to filter out those tests from the procedure that have no, or little chance of showing significant evidence, without even looking at their test statistic. Typically, this results in increased detection power at the same experiment-wide type I error. Here, we measure experiment-wide type I error in terms of the false discovery rate. A good choice for a filtering criterion is one that

- 1. is statistically independent from the test statistic under the null hypothesis,
- 2. is correlated with the test statistic under the alternative, and
- 3. does not notably change the dependence structure –if there is any– between the test statistics of nulls and alternatives.

The benefit from filtering relies on property 2, and we will explore it further in Section 5.2. Its statistical validity relies on properties 1 and 3. We refer to [7] for further discussion on the mathematical and conceptual background.

```
> rs = rowSums ( counts ( cdsFull ))
> theta = 0.4
> use = (rs > quantile(rs, probs=theta))
> table(use)

use
FALSE TRUE
5861 8738
> cdsFilt = cdsFull[ use, ]
```

Above, we consider as a filter criterion rs, the overall sum of counts (irrespective of biological condition), and remove the genes in the lowest 40% quantile (as indicated by the parameter theta). We perform the testing as before in Section 4.

```
> fitFilt1 = fitNbinomGLMs( cdsFilt, count ~ libType + condition )
> fitFilt0 = fitNbinomGLMs( cdsFilt, count ~ libType )
> pvalsFilt = nbinomGLMTest( fitFilt1, fitFilt0 )
> padjFilt = p.adjust(pvalsFilt, method="BH" )
```

Let us compare the number of genes found at an FDR of 0.1 by this analysis with that from the previous one (padjGLM).

```
> padjFiltForComparison = rep(+Inf, length(padjGLM))
> padjFiltForComparison[use] = padjFilt
> tab3 = table( `no filtering`
                               = padjGLM < .1,
               `with filtering` = padjFiltForComparison < .1 )
> addmargins(tab3)
           with filtering
no filtering FALSE
                    TRUE
       FALSE 11287
                     106 11393
       TRUE
                 3
                     963
                           966
             11290 1069 12359
```

The analysis with filtering found an additional 106 genes, an increase in the detection rate by about 11%, while 3 genes were only found by the previous analysis.

5.2 Why does it work?

First, consider Figure 9, which shows that among the 40–45% of genes with lowest overall counts, rs, there are essentially none that achieved an (unadjusted) p value less than 0.003 (this corresponds to about 2.5 on the $-\log_{10}$ -scale).

> plot(rank(rs)/length(rs), -log10(pvalsGLM), pch=16, cex=0.45)

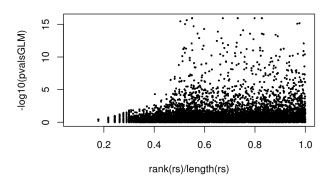


Figure 9: Scatterplot of rank of filter criterion (overall sum of counts rs) versus the negative logarithm of the test statistic pvalsGLM.

This means that by dropping the 40% genes with lowest rs, we do not loose anything substantial from our subsequent results. Second, consider the p value histogram in Figure 10. It shows how the filtering ameliorates the multiple testing problem – and thus the severity of a multiple testing adjustment – by removing a background set of hypotheses whose p values are distributed more or less uniformly in [0,1].

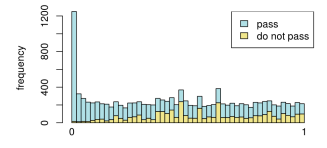


Figure 10: Histogram of p values for all tests (pvalsGLM). The area shaded in blue indicates the subset of those that pass the filtering, the area in khaki those that do not pass.

5.3 How to choose the filter statistic and the cutoff?

Please refer to the vignette Diagnostic plots for independent filtering in the genefilter package for a discussion on

- best choice of filter criterion and
- the choice of filter cutoff.

5.4 Diagnostic plots for multiple testing

The Benjamini-Hochberg multiple testing adjustment procedure [8] has a simple graphical illustration, which we produce in the following code chunk. Its result is shown in the left panel of Figure 11.

```
> orderInPlot = order(pvalsFilt)
> showInPlot = (pvalsFilt[orderInPlot] <= 0.08)
> alpha = 0.1

> plot(seq(along=which(showInPlot)), pvalsFilt[orderInPlot][showInPlot],
+ pch=".", xlab = expression(rank(p[i])), ylab=expression(p[i]))
> abline(a=0, b=alpha/length(pvalsFilt), col="red3", lwd=2)
```

Schweder and Spjøtvoll [9] suggested a diagnostic plot of the observed p-values which permits estimation of the fraction of true null hypotheses. For a series of hypothesis tests H_1, \ldots, H_m with p-values p_i , they suggested plotting

$$(1 - p_i, N(p_i)) \text{ for } i \in 1, \dots, m, \tag{1}$$

where N(p) is the number of p-values greater than p. An application of this diagnostic plot to pvalsFilt is shown in the right panel of Figure 11. When all null hypotheses are true, the p-values are each uniformly distributed in [0,1], Consequently, the cumulative distribution function of (p_1,\ldots,p_m) is expected to be close to the line F(t)=t. By symmetry, the same applies to $(1-p_1,\ldots,1-p_m)$. When (without loss of generality) the first m_0 null hypotheses are true and the other $m-m_0$ are false, the cumulative distribution function of $(1-p_1,\ldots,1-p_{m_0})$ is again expected to be close to the line $F_0(t)=t$. The cumulative distribution function of $(1-p_{m_0+1},\ldots,1-p_m)$, on the other hand, is expected to be close to a function $F_1(t)$ which stays below F_0 but shows a steep increase towards 1 as t approaches 1. In practice, we do not know which of the null hypotheses are true, so we can only observe a mixture whose cumulative distribution function is expected to be close to

$$F(t) = \frac{m_0}{m} F_0(t) + \frac{m - m_0}{m} F_1(t).$$
 (2)

Such a situation is shown in the right panel of Figure 11. If $F_1(t)/F_0(t)$ is small for small t, then the mixture fraction $\frac{m_0}{m}$ can be estimated by fitting a line to the left-hand portion of the plot, and then noting its height on the right. Such a fit is shown by the red line in the right panel of Figure 11. Here we used a value for slope of 6865, the computation can be seen in the .Rnw source code of this vignette.

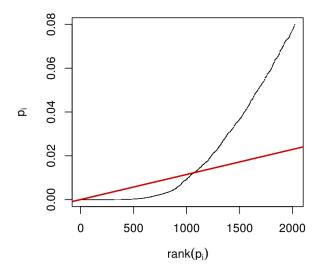
```
> plot(1-pvalsFilt[orderInPlot],
+ (length(pvalsFilt)-1):0, pch=".",
+ xlab=expression(1-p[i]), ylab=expression(N(p[i])))
> abline(a=0, b=slope, col="red3", lwd=2)
```

6 Variance stabilizing transformation

For some applications, it is useful to work with transformed versions of the count data. Maybe the most obvious choice is the logarithmic transformation. Since count values for a gene can be zero in some conditions (and non-zero in others), some advocate the use of *pseudocounts*, i.e. transformations of the form

$$y = \log_2(n+1)$$
 or more generally, $y = \log_2(n+n_0)$, (3)

where n represents the count values and n_0 is a somehow chosen positive constant. In this section, we discuss a related, alternative approach that offers more theoretical justification and a rational way of choosing the parameter equivalent to n_0 above. It is based on error modeling and the concept of variance stabilizing transformations [1,10,11]. We estimate an overall mean-dispersion relationship of the data using estimateDispersions with the argument method="blind" and call the function getVarianceStabilizedData.



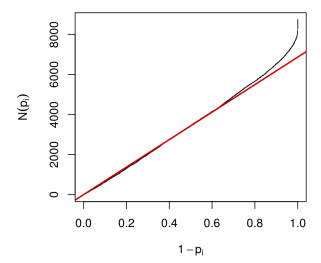


Figure 11: Left: illustration of the Benjamini-Hochberg multiple testing adjustment procedure [8]. The black line shows the p-values (y-axis) versus their rank (x-axis), starting with the smallest p-value from the left, then the second smallest, and so on. Only the first 2021 p-values are shown. The red line is a straight line with slope α/n , where n=8738 is the number of tests, and $\alpha=0.1$ is a target false discovery rate (FDR). FDR is controlled at the value α if the genes are selected that lie to the left of the rightmost intersection between the red and black lines: here, this results in 1069 genes. Right: Schweder and Spjøtvoll plot, as described in the text. For both of these plots, the p-values pvalsFilt from Section 5.1 were used as a starting point. Analogously, one can produce these types of plots for any set of p-values, for instance those from the previous sections.

```
> cdsBlind = estimateDispersions( cds, method="blind" )
> vsd = varianceStabilizingTransformation( cdsBlind )
```

Here, we have used a parametric fit for the dispersion. In this case, the a closed-form expression for the variance stabilizing transformation is used by getVarianceStabilizedData, which is derived in the document vst.pdf (you find it in the *DESeq* package alongside this vignette). If a local fit is used (option fittype="local" to estimate-Dispersion) a numerical integration is used instead. The resulting transformation is shown in Figure 12. The code that produces the figure is hidden from this vignette for the sake of brevity, but can be seen in the .Rnw or .R source file.

Figure 13 plots the standard deviation of the transformed data, across samples, against the mean, first using the shifted logarithm transformation (3), then using *DESeq*'s variance stabilising transformation. While for the latter, the standard deviation is roughly constant along the whole dynamic range, the shifted logarithm results in a highly elevated standard deviation in the lower count range.

```
> library("vsn")
> par(mfrow=c(1,2))
> notAllZero = (rowSums(counts(cds))>0)
> meanSdPlot(log2(counts(cds)[notAllZero, ] + 1), ylim = c(0,2.5))
> meanSdPlot(vsd[notAllZero, ], ylim = c(0,2.5))
```

6.1 Application to moderated fold change estimates

In the beginning of Section 3, we have seen in the data.frame res the (logarithm base 2) fold change estimate computed from the size-factor adjusted counts. When the involved counts are small, these (logarithmic) fold-change estimates can be highly variable, and can even be infinite. For some purposes, such as the clustering of samples or genes according to their expression profiles, or for visualisation of the data, this high variability from ratios between low counts tends to drown informative, systematic signal in other parts of the data. The variance stabilizing transformation offers one way to moderate the fold change estimates, so that they are more amenable to plotting or clustering.

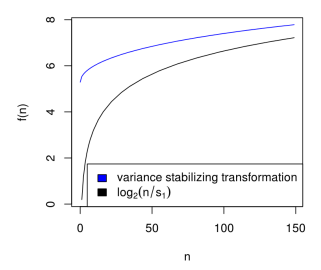


Figure 12: Graphs of the variance stabilizing transformation for sample 1, in blue, and of the transformation $f(n) = log_2(n/s_1)$, in black. n are the counts and s_1 is the size factor for the first sample.

Let us compare these to the original (\log_2) fold changes. First we find that many of the latter are infinite (resulting from division of a finite value by 0) or *not a number* (NaN, resulting from division of 0 by 0).

```
> lfc = res$log2FoldChange
> table(lfc[!is.finite(lfc)], useNA="always")
-Inf
      Inf NaN <NA>
568
      608 3098
For plotting, let us bin genes by their \log_{10} mean to colour them according to expression strength,
> logdecade = 1 + round( log10( 1+rowMeans(counts(cdsBlind, normalized=TRUE)) ) )
> lfccol = colorRampPalette( c( "gray", "blue" ) )(6)[logdecade]
and then compare their ordinary log-ratios (lfc) against the moderated log-ratios (mod_lfc) in a scatterplot.
> ymax = 4.5
 plot( pmax(-ymax, pmin(ymax, lfc)), mod_lfc,
        xlab = "ordinary log-ratio", ylab = "moderated log-ratio",
        cex=0.45, asp=1, col = 1fccol,
        pch = ifelse(lfc<(-ymax), 60, ifelse(lfc>ymax, 62, 16)))
> abline( a=0, b=1, col="red3")
```

The result is shown in Figure 14.

7 Data quality assessment by sample clustering and visualisation

Data quality assessment and quality control (i. e. the removal of insufficiently good data) are essential steps of any data analysis. Even though we present these steps towards the end of this vignette, they should typically be performed very early in the analysis of a new data set, preceding or in parallel to the differential expression testing.

We define the term *quality* as *fitness for purpose*³. Our purpose is the detection of differentially expressed genes, and we are looking in particular for samples whose experimental treatment suffered from an anormality that renders the data points obtained from these particular samples detrimental to our purpose.

³http://en.wikipedia.org/wiki/Quality_%28business%29

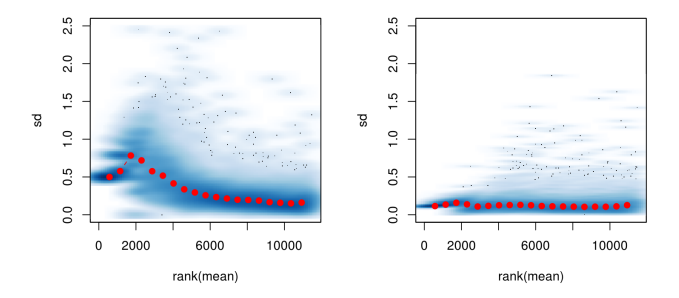


Figure 13: Per-gene standard deviation (taken across samples), against the rank of the mean, for the shifted logarithm $\log_2(n+1)$ (left) and *DESeq*'s variance stabilising transformation (right).

7.1 Heatmap of the count table

To explore a count table, it is often instructive to look at it as a heatmap. Below we show how to produce such a heatmap from the variance stabilisation transformed data for all 7 samples.

```
> cdsFullBlind = estimateDispersions( cdsFull, method = "blind" )
> vsdFull = varianceStabilizingTransformation( cdsFullBlind )
> library("RColorBrewer")
> library("gplots")
> select = order(rowMeans(counts(cdsFull)), decreasing=TRUE)[1:30]
> hmcol = colorRampPalette(brewer.pal(9, "GnBu"))(100)
> heatmap.2(exprs(vsdFull)[select,], col = hmcol, trace="none", margin=c(10, 6))
```

The left panel of Figure 15 shows the resulting heatmap for the 30 most highly expressed genes. For comparison, let us also do the same with the untransformed counts.

```
> heatmap.2(counts(cdsFull)[select,], col = hmcol, trace="none", margin=c(10,6))
```

7.2 Heatmap of the sample-to-sample distances

Another use of variance stabilized data is sample clustering. Here, we apply the dist function to the transpose of the transformed count matrix to get sample-to-sample distances.

```
> dists = dist( t( exprs(vsdFull) ) )
```

A heatmap of this distance matrix gives us an overview over similarities and dissimilarities between samples (Figure 16):

```
> mat = as.matrix( dists )
> rownames(mat) = colnames(mat) = with(pData(cdsFullBlind), paste(condition, libType, sep=" : "))
> heatmap.2(mat, trace="none", col = rev(hmcol), margin=c(13, 13))
```

The clustering correctly reflects our experimental design, i.e., samples are more similar when they have the same treatment or the same library type. (To avoid potential circularities in this conclusion, it was important to re-estimate the dispersions with method="blind" in the calculation for cdsFullBlind above, as only then, the variance stabilizing transformation is not informed about the design, and we can be sure that it is not biased towards a result supporting the design.)

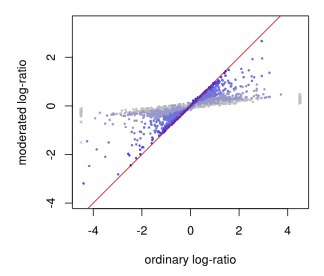


Figure 14: Scatterplot of ordinary (1fc) versus *moderated* log-ratios (mod_lfc). The points are coloured in a scale from grey to blue, representing weakly to strongly expressed genes. For the highly expressed genes (blue), lfc and mod_lfc agree. Differences arise when the involved counts are small (grey points): in this case, the moderated log-ratio is shrunk towards 0, compared to the ordinary log-ratio. The > and < symbols correspond to values of lfc whose absolute value was larger than 4.5 (including those that were infinite). All values of mod_lfc vary in a finite range.

7.3 Principal component plot of the samples

Related to the distance matrix of Section 7.2 is the PCA plot of the samples, which we obtain as follows (Figure 17).

> print(plotPCA(vsdFull, intgroup=c("condition", "libType")))

7.4 arrayQualityMetrics

Heatmaps and PCA plots similar to Figures 16 and 17 as well as some further diagnostics are also provided by the package *arrayQualityMetrics*. The package may be particularly useful for larger data set with dozens of samples. You can call its main function, which is also called arrayQualityMetrics, with an *ExpressionSet* such as vsdFull, which you will have produced from your *CountDataSet* with calls to estimateDispersions and *varianceStabilizingTransformation* as in the beginning of Section 7.1.

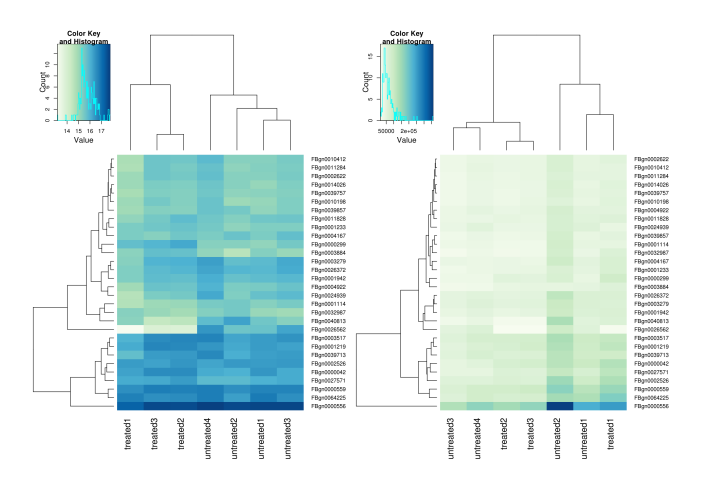


Figure 15: Heatmaps showing the expression data of the 30 most highly expressed genes. Left, the variance stabilisation transformed data (vsdFull) are shown, right, the original count data (cdsFull). In the left panel, the sample clustering aligns with the experimental factor (treated / untreated). The clustering and the colour scale in the right plot is dominated by a small number of data points with large values.

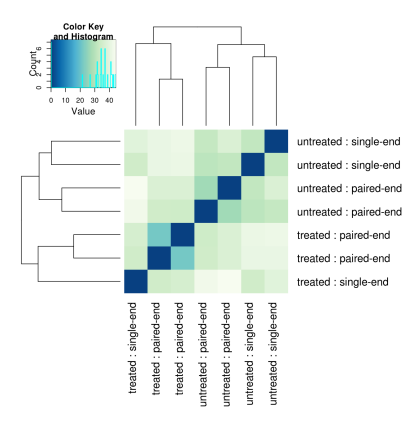


Figure 16: Heatmap showing the Euclidean distances between the samples as calculated from the variance stabilising transformation of the count data.

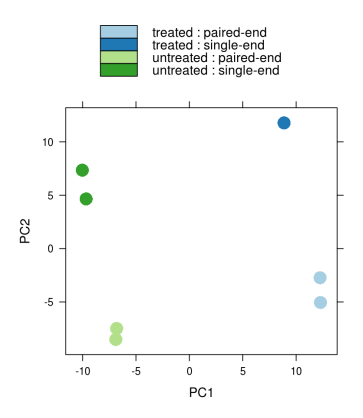


Figure 17: PCA plot. The 7 samples shown in the 2D plane spanned by their first two principal components. This type of plot is useful for visualizing the overall effect of experimental covariates and batch effects. For this data set, no batch effects besides the known effects of *condition* and *libType* are discernible.

8 Further reading

For more information on the statistical method, see our paper [1]. For more information on how to customize the *DESeq* work flow, see the package help pages, especially the help page for estimateDispersions.

9 Changes since publication of the paper

Since our paper on *DESeq* was published in Genome Biology in Oct 2010, we have made a number of changes to algorithm and implementation, which are listed here.

- nbinomTest calculates a p value by summing up the probabilities of all per-group count sums a and b that sum up to the observed count sum k_{iS} and are more extreme than the observed count sums k_{iA} and k_{iB} . Equation (11) of the paper defined more extreme as those pairs of values (a,b) that had smaller probability than the observed pair. This caused problems in cases where the dispersion exceeded 1. Hence, we now sum instead the probabilities of all values pairs that are further out in the sense that they cause a more extreme fold change $(a/s_A)/(b/s_B)$, where s_A and s_B are the sums of the size factors of the samples in conditions s_B and s_B respectively. We do this in a one-tailed manner and double the result. Furthermore, we no longer approximate the sum, but always calculate it exactly.
- We added the possibility to fit GLMs of the negative binomial family with log link. This new functionality is described in this vignette. p values are calculated by a χ^2 likelihood ratio test. The logarithms of the size factors are provided as offsets to the GLM fitting function.
- The option sharingMode='maximum' was added to estimateDispersion and made default. This change makes DESeq robust against variance outliers and was not yet discussed in the paper.
- By default, DESeq now estimates one pooled dispersion estimate across all (replicated) conditions. In the original version, we estimated a separate dispersion-mean relation for each condition. The "maximum" sharing mode achieves its goal of making DESeq robust against outliers only with pooled dispersion estimate, and hence, this is now the default. The option method='per-condition' to estimateDispersions allows the user to go back to the old method.
- In the paper, the mean-dispersion relation is fitted by local regression. Now, DESeq also offers a parametric regression, as described in this vignette. The option fitType to estimateDispersions allows the user to choose between these. If a parametric regression is used, the variance stabilizing transformation is calculated using the closed-form expression given in the vignette supplement file vst.pdf.
- Finally, instead of the term *raw squared coefficient of variance* used in the paper we now prefer the more standard term *dispersion*.

10 Session Info

> sessionInfo()

R version 3.1.1 Patched (2014-09-25 r66681) Platform: x86_64-unknown-linux-gnu (64-bit)

locale:

[7] LC_PAPER=en_US.UTF-8 LC_NAME=C LC_ADDRESS=C

[10] LC_TELEPHONE=C LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C

attached base packages:

[1] parallel stats graphics grDevices utils datasets methods base

other attached packages:

[1] gplots_2.14.2 RColorBrewer_1.0-5 vsn_3.34.0 DESeq_1.18.0

[5] lattice_0.20-29 locfit_1.5-9.1 Biobase_2.26.0 BiocGenerics_0.12.0

loaded via a namespace (and not attached):

[1]	AnnotationDbi_1.28.0	BiocInstaller_1.16.0	DBI_0.3.1	<pre>GenomeInfoDb_1.2.0</pre>
[5]	IRanges_2.0.0	KernSmooth_2.23-13	RSQLite_0.11.4	S4Vectors_0.4.0
[9]	XML_3.98-1.1	affy_1.44.0	affyio_1.34.0	annotate_1.44.0
[13]	bitops_1.0-6	caTools_1.17.1	gdata_2.13.3	genefilter_1.48.0
[17]	geneplotter_1.44.0	grid_3.1.1	gtools_3.4.1	limma_3.22.0
[21]	preprocessCore_1.28.0	splines_3.1.1	stats4_3.1.1	survival_2.37-7
[25]	tools_3.1.1	xtable_1.7-4	zlibbioc_1.12.0	

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