Accounting for technical noise in single-cell RNA-seq experiments – Supplement II

All analyses in the paper were carried out in the statistical programming language *R*. This Supplement documents the complete workflow to recreate all figures and numbers shown in the paper. It is extensively commented in order to demonstrate how our suggested analysis method is performed in practice and facilitate re-implementation by the user.

Table of Contents

- 1 Preparations
 - 1.1 Packages
 - 1.2 Count table
- 2 Analysis of the GL2 cell data
 - 2.1 Preparing the count table
 - 2.2 Normalization
 - o 2.3 Plot of normalized counts
 - o 2.4 Estimating technical noise
 - 2.4.1 Plot of the fit
 - o 2.5 Testing plant genes for high variance
 - 2.5.1 Plot
 - 2.5.2 Table of highly variable genes
 - o 2.6 GO analysis
 - 2.6.1 Results:
 - o 2.7 Heatmap
 - 2.8 Read downsampling
 - o <u>2.9 Effect of transcript length</u>
- 3 Analysis of the QC cells
 - o 3.1 Preparing the count table.
 - o 3.2 Normalization
 - 3.3 Plot of normalized counts
 - o 3.4 Estimating technical noise
 - <u>3.4.1 Plot of the fit</u>
 - o 3.5 Testing plant genes for high variance
 - <u>3.5.1 Plot</u>
 - o 3.6 Table of highly variable genes
 - 3.7 GO analysis
 - 3.7.1 Results:
 - o 3.8 Heatmap
- 4 Analysis of the mouse cells
 - 4.1 Count table and normalization
 - o 4.2 Fit technical noise
 - 4.3 Test for high variance
 - o 4.4 A diagnostic plot
 - 4.5 Plot of results

1 Preparations

1.1 Packages

We load all R and Bioconductor packages that we need for this analysis:

```
library( DESeq )
library( genefilter )
library( EBImage )
library( statmod )
library( topGO )
library( org.At.tair.db )
options( max.print=300, width=100 )
```

The sessionInfo command gives the versions of R and all packages used in the present analysis run:

```
sessionInfo()
```

```
R version 2.15.3 (2013-03-01)
Platform: x86_64-pc-linux-gnu (64-bit)
locale:
[1] LC_CTYPE=en_GB.UTF-8
                              LC NUMERIC=C
                                                           LC_TIME=en_GB.UTF-8
 [4] LC_COLLATE=en_GB.UTF-8
                                LC_MONETARY=en_GB.UTF-8 LC_MESSAGES=en_GB.UTF-8
 [7] LC_PAPER=C
                                LC_NAME=C
                                                           LC_ADDRESS=C
                                LC_MEASUREMENT=en_GB.UTF-8 LC_IDENTIFICATION=C
[10] LC_TELEPHONE=C
attached base packages:
[1] stats
            graphics grDevices utils datasets methods base
other attached packages:
[1] org.At.tair.db_2.8.0 topGO_2.10.0
                                            SparseM_0.96 GO.db_2.8.0
[5] RSQLite_0.11.2 DBI_0.2-5
[9] statmod_1.4.17 EBImage_4.0.0
                                             AnnotationDbi_1.20.3 graph_1.36.2
                                              genefilter_1.40.0 DESeq_1.11.6
[13] lattice_0.20-13 locfit_1.5-8
                                             Biobase_2.18.0 BiocGenerics_0.4.0
loaded via a namespace (and not attached):
[1] abind_1.4-0 annotate_1.36.0 geneplotter_1.36.0 grid_2.15.3 [6] jpeg_0.1-2 parallel_2.15.3 png_0.1-4 RColorBrewer
                                                                                 IRanges 1.
                                                       RColorBrewer_1.0-5 splines_2.
[11] stats4_2.15.3
                       survival_2.36-14 tiff_0.1-3
                                                             tools_2.15.3
                                                                                XML_3.95-0
[16] xtable_1.7-0
```

1.2 Count table

Supplementary Table 8 contains the raw read counts for the seven GL2 samples and the six QC samples. We save this table as a CSV file and load it into R.

```
fullCountTable <- read.csv( "Supplementary_Table_8.csv", header=TRUE, row.names=1 )
head( fullCountTable )</pre>
```

```
QC1 QC2 QC3 QC4 QC5 QC6 GL2.1 GL2.2 GL2.3 GL2.4 GL2.5 GL2.6 GL2.7
               0 0 0 0 64 18 0 0 87 381
AT1G01010 0 0
AT1G01020 2 0
              7 708
                      0 0
                             422
                                  602 125
                                           58
                                               181
                                                     148
                                                          122
AT1G01030 0 0 0 0 0 0 0 0 0 0 0 0 AT1G01040 0 0 459 533 42 205 2 577 201
                                                      0 112
                                                 0
AT1G01046 0 0 0 0 0
                             0
                                   0
                                       0
                                            0
                                                 0
                                                      0
                   3 36 2 3865 5126 3804 1464 3072 3760 2112
AT1G01050 2 166 1245
```

Load the count table (new version with all seven GL2 cells), then subset to the samples from the GL2 cells (with "pGL" in the column name), and simplify the column names.

2 Analysis of the GL2 cell data

2.1 Preparing the count table

We subset the count table to only the columns referring to GL2 cells and clean up the column names.

```
countsAll <- fullCountTable[, substr( colnames(fullCountTable), 1, 3 ) == "GL2" ]
colnames(countsAll) <- gsub( "\\.", "-", colnames(countsAll) )</pre>
```

We split the count table in three sub-tables, one for the A. thalina plant genes ("At"), one for the HeLa genes ("HeLa") and one for the IVT spikes ("Sp"). The first two letter of the gene IDs (row names) are used for this categorization.

```
geneTypes <- factor( c( AT="At", pG="pGIBS", EN="HeLa", ER="ERCC" )[
   substr( rownames(countsAll), 1, 2 ) ] )
countsHeLa <- countsAll[ which( geneTypes=="HeLa" ), ]
countsAt <- countsAll[ which( geneTypes=="At" ), ]
countsSp <- countsAll[ which( geneTypes %in% c( "pGIBS", "ERCC" ) ), ]</pre>
```

We will not use the IVT spike data in this analysis. The other two tables now look as follows.

```
head( countsHeLa )
```

```
GL2-1 GL2-2 GL2-3 GL2-4 GL2-5 GL2-6 GL2-7
ENSG0000000003 581 1850 2169 392 2046 1225 166
ENSG00000000005
                         0
                                    0
                0
                    0
                               0
                                          0
                                                0
              393 1263 1411
ENSG00000000419
                               296 1754
                                         247
                                              159
ENSG00000000457
                1
                    109
                         118
                                71
                                    52
                                          0
                                                1
ENSG00000000460
                   179
                        419
                89
                                42
                                    310
                                              116
                                          16
ENSG00000000938
               0
                     0
                           0
                               0
                                    0
                                          0
                                                0
```

```
head( countsAt )
```

```
GL2-1 GL2-2 GL2-3 GL2-4 GL2-5 GL2-6 GL2-7
AT1G01010
         64 18 0 0 87 381
        422
               602
                    125
                         58
                              181
                                   148
AT1G01020
                                   0
AT1G01030
          0
             ()
                    0
                         0
                              Ω
                                        0
         205
                2
                    577
                         201
                               0
                                    0
                                        112
AT1G01040
AT1G01046
          0
                0
                    0
                         0
                               0
                                    0
AT1G01050 3865 5126 3804 1464 3072 3760 2112
```

For later use, we get a translation of gene IDs to gene symbols.

```
geneSymbolsAt <- rownames(countsAt)
names( geneSymbolsAt ) <- rownames(countsAt)
hasSymbol <- rownames(countsAt) %in% Lkeys( org.At.tairSYMBOL )
symtbl <- toTable( org.At.tairSYMBOL[ rownames(countsAt)[ hasSymbol ] ] )
symtbl <- symtbl[ !duplicated( symtbl$gene_id ), ]
geneSymbolsAt[ symtbl$gene_id ] <- symtbl$symbol
head( geneSymbolsAt )</pre>
```

```
AT1G01010 AT1G01020 AT1G01030 AT1G01040 AT1G01046 AT1G01050 "ANAC001" "ARV1" "NGA3" "ASU1" "AT1G01046" "AtPPa1"
```

2.2 Normalization

We use the function "estimateSizeFactorsForMatrix" from DESeq to get size factors. This function calculates size factors as described in the DESeq paper (Anders and Huber, 2010) and in the Online Methods of the present paper.

```
sfHeLa <- estimateSizeFactorsForMatrix( countsHeLa )
sfAt <- estimateSizeFactorsForMatrix( countsAt)</pre>
```

See the size factors and their ratios:

```
rbind( HeLa = sfHeLa, At = sfAt, ratio = sfAt / sfHeLa )
```

```
GL2-1 GL2-2 GL2-3 GL2-4 GL2-5 GL2-6 GL2-7
HeLa 0.5173383 2.022511 2.8804054 0.8552699 2.0415875 0.9551039 0.3109511
At 1.0234077 1.933931 1.7603544 1.3696320 0.8933595 0.8634580 0.5341586
ratio 1.9782176 0.956203 0.6111481 1.6014033 0.4375808 0.9040461 1.7178219
```

Divide by the size factors to get normalized counts. (Note the double use of "t" to make sure that columns, not rows, are divided by the size factors.)

```
nCountsHeLa <- t( t(countsHeLa) / sfHeLa )
nCountsAt <- t( t(countsAt) / sfAt )
```

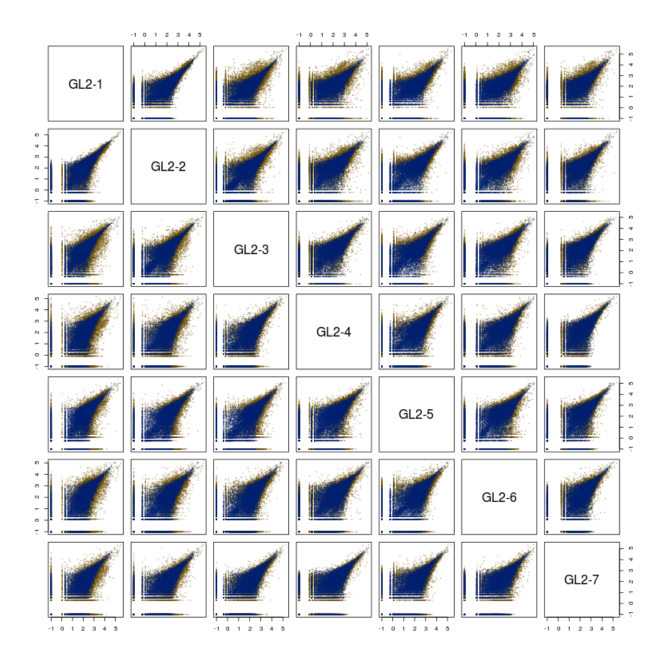
2.3 Plot of normalized counts

We use the following colours in our plots:

```
colHeLa <- "#00207040"
colAt <- "#70500040"
colAtHi <- "#B0901040"
```

Make a pairs plot of all GL-2 cells (Supplementary Figure 5).

```
pairs( log10( .1 + rbind( nCountsAt, nCountsHeLa ) ), pch=19, cex=.2,
  col = c( rep( colAt, nrow(nCountsAt) ), rep( colHeLa, nrow(nCountsHeLa) ) ) )
```

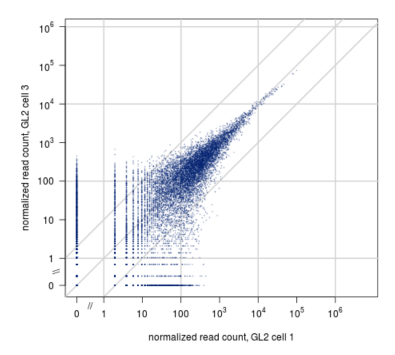


As the pairs plot is a bit large, we plot one comparsion a bit bigger, namely cells 1 vs 3 (Figures 2a and 2b).

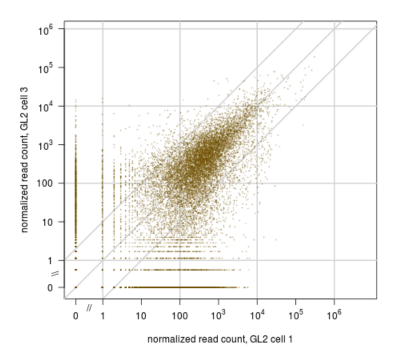
First, the following function makes nice axes etc. This function (and most of the code for plotting in the following sections) is somewhat specialized to our data. users wishing to follow this code to perform analyses of their own data may want to use simpler, more standard, R code for plotting.

Now make the two plots, first Figure 2a, then 2b:

```
geneScatterplot( nCountsHeLa[,1], nCountsHeLa[,3],
    "normalized read count, GL2 cell 1", "normalized read count, GL2 cell 3",
    colHeLa )
```



```
geneScatterplot( nCountsAt[,1], nCountsAt[,3],
   "normalized read count, GL2 cell 1", "normalized read count, GL2 cell 3",
   colAt )
```



2.4 Estimating technical noise

We start by estimating the sample moments per gene.

```
meansHeLa <- rowMeans( nCountsHeLa )
varsHeLa <- rowVars( nCountsHeLa )
cv2HeLa <- varsHeLa / meansHeLa^2
```

Next, we define a minimum mean value to exclude genes with low mean and hence high CV from fit as they would otherwise skew it downwards (Online Methods and Supplementary Notes 6 and 7).

```
minMeanForFit <- unname( quantile( meansHeLa[ which( cv2HeLa > .3 ) ], .95 ) )
minMeanForFit
```

```
[1] 417.6733
```

Perform the fit of technical noise strength on average count. We regress cv2HeLa on 1/meansForHeLa. We use the glmgam.fit function from the statmod package to perform the regression as a GLM fit of the gamma family with log link. The 'cbind' construct serves to produce a model matrix with an intercept.

```
useForFit <- meansHeLa >= minMeanForFit
fit <- glmgam.fit( cbind( a0 = 1, altilde = 1/meansHeLa[useForFit] ),
    cv2HeLa[useForFit] )
fit$coefficients</pre>
```

```
a0 altilde
0.04245659 179.66783886
```

To get the actual noise coefficients, we need to subtract Xi (see Supplementary Note 6 for the difference between altilde and al).

```
xi <- mean( 1 / sfHeLa )
```

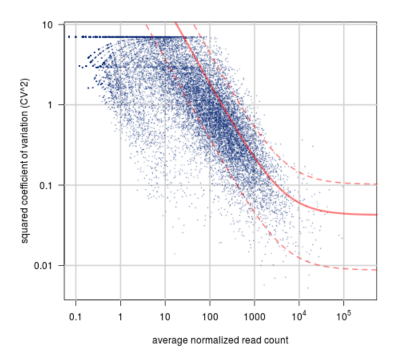
```
a0 <- unname( fit$coefficients["a0"] )
a1 <- unname( fit$coefficients["altilde"] - xi )
c( a0, a1 )</pre>
```

```
[1] 0.04245659 178.42547279
```

2.4.1 Plot of the fit

The following code produces the plot of the fit shown in Figure 2c.

```
# Prepare the plot (scales, grid, labels, etc.)
plot( NULL, xaxt="n", yaxt="n",
  log="xy", xlim = c( 1e-1, 3e5 ), ylim = c( .005, 8 ),
xlab = "average normalized read count", ylab = "squared coefficient of variation (CV^2)" axis(1, 10^{-1:5}), c("0.1", "1", "10", "1000",
 expression(10^4), expression(10^5))
axis( 2, 10^(-2:1), c( "0.01", "0.1", "1", "10" ), las=2 )
abline( h=10^(-2:1), v=10^(-1:5), col="#D0D0D0", lwd=2)
# Add the data points
points ( meansHeLa, cv2HeLa, pch=20, cex=.2, col=colHeLa )
# Plot the fitted curve
xg <- 10^seq(-2, 6, length.out=1000)
lines(xg, \frac{(xi+a1)}{xg} + a0, col="#FF000080", lwd=3)
# Plot quantile lines around the fit
df <- ncol(countsAt) - 1
lines( xg, ( (xi+a1)/xg + a0 ) * qchisq( .975, df ) / df,
  col="#FF000080", lwd=2, lty="dashed" )
lines( xg, ( (xi+a1)/xg + a0 ) * qchisq( .025, df ) / df,
      col="#FF000080", lwd=2, lty="dashed")
```



2.5 Testing plant genes for high variance

To perform the actual test (Online Methods, Supplementary Note 6), we start with again calculating the sample moments, now for the plant genes.

```
meansAt <- rowMeans( nCountsAt )
varsAt <- rowVars( nCountsAt )
cv2At <- varsAt / meansAt^2</pre>
```

The following is the term Psi + a0 * Theta, that appears in the formula for Omega (see formula in Online Methods).

```
psialtheta <- mean( 1 / sfAt ) + al * mean( sfHeLa / sfAt )
```

Now, we perform a one-sided test against the null hypothesis that the true variance is at most the technical variation plus biological variation with a CV of at most 50% (minBiolDisp = $.5^2$).

```
minBiolDisp <- .5^2
```

Calculate Omega, then perform the test, using the formula given in the Online methods and in Supplementary Note 6.

```
m <- ncol(countsAt)
cv2th <- a0 + minBiolDisp + a0 * minBiolDisp
testDenom <- ( meansAt * psialtheta + meansAt^2 * cv2th ) / ( 1 + cv2th/m )
p <- 1 - pchisq( varsAt * (m-1) / testDenom, m-1 )</pre>
```

Adjust for multiple testing with the Benjamini-Hochberg method, cut at 10%:

```
padj <- p.adjust( p, "BH" )
sig <- padj < .1
sig[is.na(sig)] <- FALSE
table( sig )</pre>
```

```
sig
FALSE TRUE
32726 876
```

2.5.1 Plot

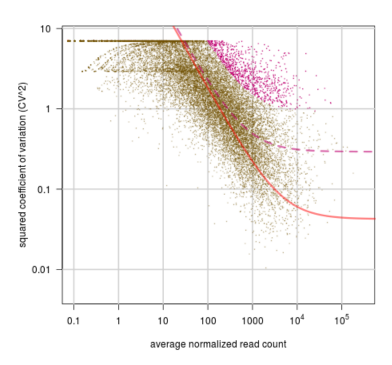
Make a plot of the plant genes, highlighting the highly variable ones (Figure 2d).

```
# Prepare plot in the same manner as before
plot( NULL, xaxt="n", yaxt="n",
    log="xy", xlim = c( 1e-1, 3e5 ), ylim = c( .005, 8 ),
    xlab = "average normalized read count", ylab = "squared coefficient of variation (CV^2)"
axis( 1, 10^(-1:5), c( "0.1", "1", "10", "100", "1000",
    expression(10^4), expression(10^5) ))
axis( 2, 10^(-2:1), c( "0.01", "0.1", "1", "10" ), las=2 )
abline( h=10^(-2:1), v=10^(-1:5), col="#DODODO", lwd=2 )

# Plot the plant genes, use a different color if they are highly variable
points( meansAt, cv2At, pch=20, cex=.2,
    col = ifelse( padj < .1, "#C0007090", colAt ) )

# Add the technical noise fit, as before
xg <- 10^seq( -2, 6, length.out=1000 )
lines( xg, (xi+a1)/xg + a0, col="#FF000080", lwd=3 )</pre>
```

```
\# Add a curve showing the expectation for the chosen biological CV^2 thershold lines( xg, psialtheta/xg + a0 + minBiolDisp, lty="dashed", col="\#C0007090", lwd=3 )
```



2.5.2 Table of highly variable genes

Most highly variable genes are strong only in one or two cells. To show this, we calculate the log ratio of the genes' expression in each cell to the mean:

```
log2RelExprAt <- log2( nCountsAt / meansAt )
```

We use this to produce a table of all significantly highly variable genes with some extra information, namely the mean normalized count, the log2 fold change of each cell to this mean, and the information which cell has the strongest expression.

```
highVarTable <- data.frame(
   row.names = NULL,
   geneID = rownames(countsAt)[ sig ],
   geneSymbol = geneSymbolsAt[ sig ],
   meanNormCount = meansAt[ sig ],
   strongest = factor( colnames( log2RelExprAt )[
      apply( log2RelExprAt[ sig, ], 1, which.max ) ] ),
   log2RelExprAt[ sig, ],
   check.names=FALSE )</pre>
head( highVarTable )
```

```
geneID geneSymbol meanNormCount strongest
                                                  GL2-1
                                                             GL2-2
                                                                       GL2-3
                                       GL2-1 1.6221783 1.0724231 -2.119163 -2.3754433
1 AT1G01100 AT1G01100
                          2534.5561
                                                              -Inf -1.293147 -0.8690356
2 AT1G01140
                CIPK9
                          1584.2303
                                       GL2-6
                                                   -Inf
3 AT1G01730 AT1G01730
                          913.3887
                                       GL2-3 -0.9406882
                                                         0.7364503 2.020811 -3.3940557
4 AT1G01740 AT1G01740
                          837.6666
                                       GL2-3 -0.5463967 1.3201552 1.876914
                                       GL2-4 -0.5788364 -5.1974315 -4.273265 2.6442089
                           208.7058
5 AT1G02230
            ANAC004
6 AT1G02780
              emb2386
                         10119.6519
                                       GL2-1 1.7777659 0.4076499 -2.532492 -1.8231270
       GL2-5
                  GL2-6
                             GL2-7
```

```
1 -0.4417918 -1.4518590 -1.7662327

2 -10.4668792 2.5764124 -3.6588172

3 -3.4629446 -9.6232830 -0.8166829

4 -Inf -4.7979905 -3.1908625

5 -Inf -Inf -Inf

6 -0.4174578 -0.7624331 -1.1493215
```

We write out the table. It is given in the Supplement as Supplementary Table 2.

```
write.csv( highVarTable, file="highly_variant_genes_GL2.tsv", row.names=FALSE )
```

2.6 GO analysis

Next, we check whether the high-variance genes are enriched in certain GO categories using TopGO (Online Methods).

To work only with genes with uniform power to detect high or low variance, we include only genes with an average count above 300.

```
minCountForEnrichment <- 300
```

The following function performs the analysis. It takes a vector with gene IDs (here: all plant genes) and then two Boolean vectors of the same length, the first indicating which genes are to be included in the analysis (here: above the mean cut-off) and the second indicating which genes are significant (i.e., highly variable).

```
topGOAnalysis <- function( geneIDs, inUniverse, inSelection )
   sapply( c( "MF", "BP", "CC" ), function( ont ) {
      alg <- factor( as.integer( inSelection[inUniverse] ) )
      names(alg) <- geneIDs[inUniverse]
      tgd <- new( "topGOdata", ontology=ont, allGenes = alg, nodeSize=5,
            annot=annFUN.org, mapping="org.At.tair.db" )
      resultTopGO <- runTest(tgd, algorithm = "elim", statistic = "Fisher" )
      GenTable( tgd, resultTopGO, topNodes=15 ) },
      simplify=FALSE )</pre>
```

We use this to perform an enrichment analysis for the list of highly variable genes:

```
goResults <-
  topGOAnalysis(
   rownames(countsAt),
  meansAt >= minCountForEnrichment & !is.na(padj),
  padj < .1 )</pre>
```

2.6.1 Results:

How to correctly adjust for multiple testing in enrichment analyses is slightly controversal, so to keep it simple, we consider categories with raw p value below 10^{-5} as clearly and below 10^{-4} as maybe significant. Categories thus deemed significant have been taken from the results below and are listed in Supplemenary Table 4.

```
goResults[["MF"]]
```

	GO.ID	Term	Annotated	Significant	Expected	re:
1	GO:0003735	structural constituent of ribosome	212	140	23.24	< :
2	GO:0015250	water channel activity	17	9	1.86	2.:
3	GO:0003677	DNA binding	391	69	42.86	2.
4	GO:0003746	translation elongation factor activity	18	7	1.97	0
5	GO:0019843	rRNA binding	7	4	0.77	0
6	GO:0016762	<pre>xyloglucan:xyloglucosyl transferase acti</pre>	8	4	0.88	0
7	GO:0016884	carbon-nitrogen ligase activity, with gl	8	4	0.88	0
8	GO:0003676	nucleic acid binding	706	123	77.39	0

```
9 GO:0003723
                                           RNA binding
                                                            229
                                                                        44
                                                                              25.10 0
10 GO:0016847 1-aminocyclopropane-1-carboxylate syntha...
                                                             6
                                                                        3
                                                                              0.66 0
                                                                        16
                                                                              9.43 0
11 GO:0004553 hydrolase activity, hydrolyzing O-glycos...
                                                             86
                                                                              10.30 0
3.07 0
12 GO:0016798 hydrolase activity, acting on glycosyl b...
                                                             94
                                                                        17
13 GO:0016684 oxidoreductase activity, acting on perox...
                                                             28
                                                                         7
                                                                               3.07 0
                                                                         7
14 GO:0004601
                                   peroxidase activity
                                                             28
                                                                               0.88 0
15 GO:0003690
                            double-stranded DNA binding
                                                                         3
                                                             8
```

```
goResults[["CC"]]
```

	GO.ID	Term	Annotated	Significant	Expected	re
1	GO:0022625	cytosolic large ribosomal subunit	92	78	10.04	<
2	GO:0022627	cytosolic small ribosomal subunit	69	48	7.53	< :
3	GO:0005730	nucleolus	161	71	17.56	5.2
4	GO:0000786	nucleosome	41	33	4.47	3.
5	GO:0009506	plasmodesma	478	108	52.15	5.
6	GO:0005618	cell wall	255	52	27.82	3.1
7	GO:0022626	cytosolic ribosome	207	139	22.58	9.
8	GO:0005576	extracellular region	126	24	13.75	0
9	GO:0005774	vacuolar membrane	334	51	36.44	0
10	GO:0005634	nucleus	919	165	100.26	0
11	GO:0009507	chloroplast	745	100	81.28	0
12	GO:0005773	vacuole	532	82	58.04	0
13	GO:0009536	plastid	784	102	85.53	0
14	GO:0048046	apoplast	102	17	11.13	0
15	GO:0005853	eukaryotic translation elongation factor	5	2	0.55	0

```
goResults[["BP"]]
```

i							
		GO.ID	Term	Annotated	Significant	Expected	re
	1	GO:0001510	RNA methylation	105	79	11.42	<
	2	GO:0006412	translation	448	155	48.74	<
	3	GO:0042254	ribosome biogenesis	108	63	11.75	7.
	4	GO:0006334	nucleosome assembly	45	34	4.90	2.
	5	GO:0009220	pyrimidine ribonucleotide biosynthetic p	53	29	5.77	3.
	6	GO:0006364	rRNA processing	39	19	4.24	4.
	7	GO:0006414	translational elongation	33	14	3.59	3.
	8	GO:0042545	cell wall modification	79	22	8.59	2.
	9	GO:0009664	plant-type cell wall organization	106	26	11.53	4.
	10	GO:0000724	double-strand break repair via homologou	9	6	0.98	0.
	11	GO:0006164	purine nucleotide biosynthetic process	50	15	5.44	0.
	12	GO:1901070	guanosine-containing compound biosynthet	5	4	0.54	0.
	13	GO:0009646	response to absence of light	20	8	2.18	0.
	14	GO:0009955	adaxial/abaxial pattern specification	12	6	1.31	0.
	15	GO:0000462	maturation of SSU-rRNA from tricistronic	6	4	0.65	0.
ı							

2.7 Heatmap

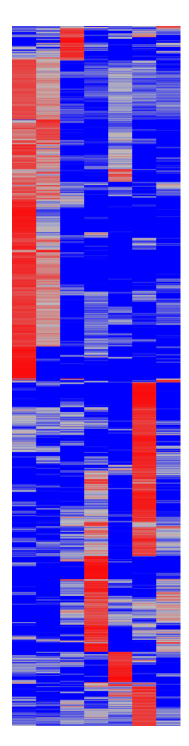
This section describes how Supplementary Figure 9a was produced.

In order to get the genes aligned to the plot's pixels, we use the following custom heatmap function instead of R's standard one. It expects a matrix with values in the range given by 'zlim' (everything outside this range will be pulled in to its margins), a number pair for zlim, a color palette, and two integers for the width and height in pixels that should be used to represent each matrix element.

```
a <- array( NA_real_, c( nrow(m)*pxWidth, ncol(m)*pxHeight, 3 ) )
for( i in 1:nrow(mn) )
    for( j in 1:ncol(mn) )
        a[ (((i-1)*pxWidth)+1) : (i*pxWidth), (((j-1)*pxHeight)+1) : (j*pxHeight), ] <-
            rep( col[ , mn[ i, j ] ] , each = pxWidth*pxHeight )
Image( a, colormode="color" )
}</pre>
```

We use this to plot the relative expressions of the highly vaiable genes.

```
writeImage( hmSig, "variableGenes_heatmap.png" )
```



We annotate this heatmap with a few hand-picked GO terms:

```
[,1] [,2]
[1,] "GO:0003735" "structural constituent of ribosome"
[2,] "GO:0001510" "RNA methylation"
[3,] "GO:0005730" "nucleolus"
[4,] "GO:0009506" "plasmodesma"
[5,] "GO:0003677" "DNA binding"
[6,] "GO:0000786" "nucleosome"
```

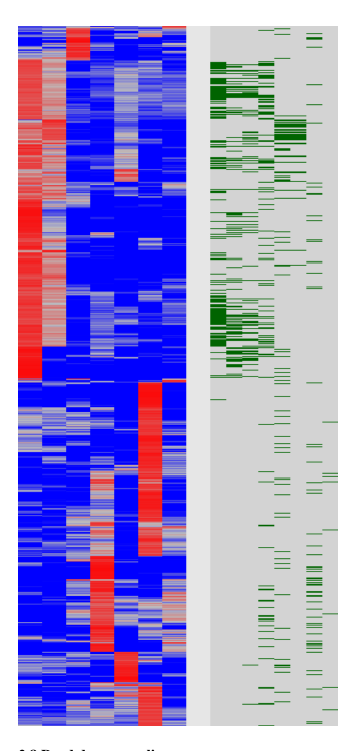
```
[7,] "GO:0012505" "endomembrane system"
[8,] "GO:0009269" "response to desiccation"
```

For these, make a second heatmap indicating the genes' membership.

```
sigInGO <- sapply( unname(someGOTerms), function(go)
  rownames(countsAt)[sig] %in% org.At.tairGO2TAIR[[ go ]] )
hmGO <- pixelHeatmap( t(sigInGO[ ord, ]), col=c( "lightgray", "darkgreen" ), pxWidth=20 )</pre>
```

Put the two heatmaps next to each other.

```
spacer <- Image( array( .9, 30 * 3 * sum(sig), dim=c( 30, sum(sig), 3 ) ), colormode="color
writeImage(
   Image( abind::abind( hmSig, spacer, hmGO, along=1 ), colormode="color" ),
   files="heatmap.png" )</pre>
```



2.8 Read downsampling

To show that we could have worked with fewer reads (as claimed in Supplementary Note 9), we downsample all read counts to 1/5 using draws from binomial distributions

```
dsfrac <- 0.2
countsAt_ds <- apply( countsAt, 1:2, function(k) rbinom( 1, k, dsfrac ) )
countsHeLa_ds <- apply( countsHeLa, 1:2, function(k) rbinom( 1, k, dsfrac ) )</pre>
```

We redo the analysis as before.

First, the normalization:

```
sfAt_ds <- estimateSizeFactorsForMatrix( countsAt_ds )
sfHeLa_ds <- estimateSizeFactorsForMatrix( countsHeLa_ds )

nCountsAt_ds <- t( t(countsAt_ds) / sfAt_ds )
nCountsHeLa_ds <- t( t(countsHeLa_ds) / sfHeLa_ds )</pre>
```

Next, the technical noise fit:

```
meansHeLa_ds <- rowMeans( nCountsHeLa_ds )
varsHeLa_ds <- rowVars( nCountsHeLa_ds )
cv2HeLa_ds <- varsHeLa_ds / meansHeLa_ds^2

minMeanForFit_ds <- unname( quantile( meansHeLa_ds[ which( cv2HeLa_ds > .3 ) ], .95 ) )
useForFit_ds <- meansHeLa_ds >= minMeanForFit_ds
fit_ds <- glmgam.fit( cbind( a0 = 1, altilde = 1/meansHeLa_ds[useForFit_ds] ),
    cv2HeLa_ds[useForFit_ds] )

xi_ds <- mean( 1 / sfHeLa_ds )
a0_ds <- unname( fit_ds$coefficients["a0"] )
a1_ds <- unname( fit_ds$coefficients["a1tilde"] - xi_ds )

c( a0_ds, a1_ds )</pre>
```

```
[1] 0.03706779 35.43892577
```

Note how this a0 stayed roughly the same, and a1 got reduced according to the downsampling fraction; compare with

```
c( a0, a1 * dsfrac )
```

```
[1] 0.04245659 35.68509456
```

Finally, the test for high variability.

```
meansAt_ds <- rowMeans( nCountsAt_ds )
varsAt_ds <- rowVars( nCountsAt_ds )

psialtheta_ds <- mean( 1 / sfAt_ds ) + al_ds * mean( sfHeLa_ds / sfAt_ds )
cv2th_ds <- a0_ds + minBiolDisp + a0_ds * minBiolDisp
testDenom_ds <- ( meansAt_ds * psialtheta_ds + meansAt_ds^2 * cv2th_ds ) / ( 1 + cv2th_ds/m

p_ds <- 1 - pchisq( varsAt_ds * (m-1) / testDenom_ds, m-1 )
padj_ds <- p.adjust( p_ds, "BH" )</pre>
```

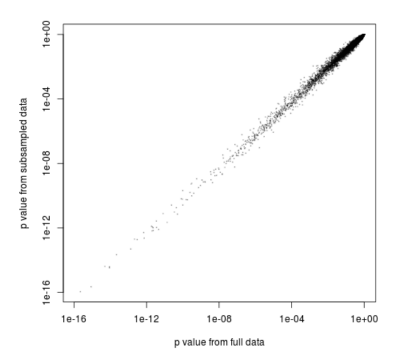
Compare the results and calculate the overlap:

```
addmargins( table( all_reads = padj < .1, subsampled = padj_ds < .1 ) )
```

```
subsampled
all_reads FALSE TRUE Sum
FALSE 17313 64 17377
TRUE 27 849 876
Sum 17340 913 18253
```

Also, a plot of the p values (Supplementary Figure 12; note that this plots always changes slightly due to the random nature of the downsampling)

```
plot( p, p_ds,
    log="xy", pch=19, cex=.2, col="#00000040",
    xlab = "p value from full data", ylab="p value from subsampled data" )
```



2.9 Effect of transcript length

Here, we test whether it is benefitial to normalize counts for transcript length.

We load a table of transcript lengths for the human genome (given in Supplementary Table 9):

```
a <- read.csv( "Supplementary_Table_9.csv" )
humanGeneLengths <- a$length
names(humanGeneLengths) <- a$geneID
head(humanGeneLengths)</pre>
```

We use this to convert the normalized counts to normalized counts per kilobase transcript length.

```
nCountsHeLaPK <- nCountsHeLa / humanGeneLengths[rownames(nCountsHeLa)] * 1000
```

We redo the fit using this data (removing a single gene for which we are missing length information):

We compare the residual variances from both fits:

```
totalVariance <- var( log( cv2HeLa[useForFit] ) )
residualVariance <- var( log( cv2HeLa[useForFit] ) - log( fitted.values(fit) ) )
totalVariancePK <- var( log( cv2HeLa[useForFitPK] ) )
residualVariancePK <- var( log( cv2HeLa[useForFitPK] ) - log( fitted.values(fitPK) ) )</pre>
```

Here are the variances and their ratios:

```
c( residualVariance, totalVariance, residualVariance / totalVariance )
c( residualVariancePK, totalVariancePK, residualVariancePK / totalVariancePK )
```

```
[1] 0.5438976 0.8333216 0.6526863
[1] 0.6610947 0.8334789 0.7931751
```

Here, the length-normalization increases the residual variance, i.e., is not benefitial.

Note: These residual variance fractions may seem large. However, bear in mind that even a perfect fit cannot explain the sampling variance of the log CV^2 estimates, which approximately have variance 2/(m-1), i.e. account for the following fraction of the total variance:

```
2/(m-1) / totalVariance
```

```
[1] 0.4000057
```

This is more than half of the residual variance fraction. Hence, if we subtract this from the tota variance to get the total "explainable" variance, we get a rather large fraction of explained variance:

```
explainedVariance <- totalVariance - residualVariance
explainableVariance <- totalVariance - 2/(m-1)
explainedVariance / explainableVariance</pre>
```

```
[1] 0.5788616
```

The same for the length-divided fit:

```
explainedVariancePK <- totalVariancePK - residualVariancePK explainableVariancePK <- totalVariancePK - 2/(m-1)
explainedVariancePK / explainableVariancePK
```

```
[1] 0.344668
```

Finally, we save the current state, for convenience.

```
save.image( "GL2_analysis_image.RData" )
```

3 Analysis of the QC cells

The analysis of the QC cells is done in exactly the same way as for the GL2 cells. Hence, we give here the code, essentially the same code as before, without much comments.

As the code here re-uses the variable names used in the previous part, we clear the global environment in order to start with a clean slate.

```
rm( list=ls() )
```

3.1 Preparing the count table.

We read in the full count table, as before, and now subset to only the columns referring to QC cells.

```
fullCountTable <- read.csv( "Supplementary_Table_8.csv", header=TRUE, row.names=1 )
countsAll <- fullCountTable[, substr( colnames(fullCountTable), 1, 2 ) == "QC" ]</pre>
```

We split the count table again into three sub-tables

```
geneTypes <- factor( c( AT="At", pG="pGIBS", EN="HeLa", ER="ERCC" )[
   substr( rownames(countsAll), 1, 2 ) ] )
countsHeLa <- countsAll[ which( geneTypes=="HeLa" ), ]
countsAt <- countsAll[ which( geneTypes=="At" ), ]
countsSp <- countsAll[ which( geneTypes %in% c( "pGIBS", "ERCC" ) ), ]</pre>
```

Again, the gene symbols:

```
geneSymbolsAt <- rownames(countsAt)
names( geneSymbolsAt ) <- rownames(countsAt)
hasSymbol <- rownames(countsAt) %in% Lkeys( org.At.tairSYMBOL )
symtbl <- toTable( org.At.tairSYMBOL[ rownames(countsAt)[ hasSymbol ] ] )
symtbl <- symtbl[ !duplicated( symtbl$gene_id ), ]
geneSymbolsAt[ symtbl$gene_id ] <- symtbl$symbol
head( geneSymbolsAt )</pre>
```

```
AT1G01010 AT1G01020 AT1G01030 AT1G01040 AT1G01046 AT1G01050 "ANAC001" "ARV1" "NGA3" "ASU1" "AT1G01046" "AtPPa1"
```

3.2 Normalization

As before:

```
sfHeLa <- estimateSizeFactorsForMatrix( countsHeLa )
sfAt <- estimateSizeFactorsForMatrix( countsAt)
```

See the size factors and their ratios:

```
rbind( HeLa = sfHeLa, At = sfAt, ratio = sfAt / sfHeLa )
```

```
QC1 QC2 QC3 QC4 QC5 QC6

HeLa 1.287133 0.8061698 0.8695366 1.206279 1.1365133 1.2384236

At 1.595883 1.2135275 1.2409527 2.315439 0.9654965 0.6946031

ratio 1.239875 1.5053001 1.4271426 1.919489 0.8495251 0.5608768
```

Divide by the size factors to get normalized counts

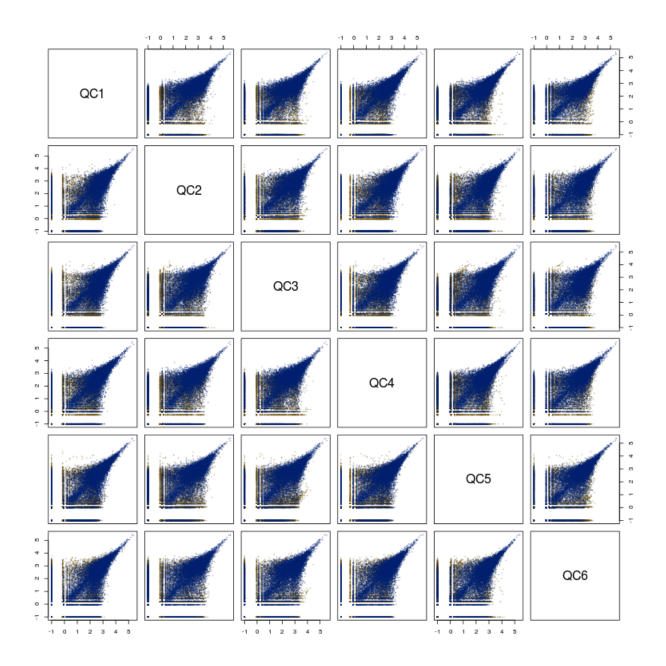
```
nCountsHeLa <- t( t(countsHeLa) / sfHeLa )
nCountsAt <- t( t(countsAt) / sfAt )
```

3.3 Plot of normalized counts

The plot shown in Supplementary Figure 6.

```
colHeLa <- "#00207040"
colAt <- "#70500040"
colAtHi <- "#B0901040"
```

```
pairs( log10( .1 + rbind( nCountsAt, nCountsHeLa ) ), pch=19, cex=.2,
    col = c( rep( colAt, nrow(nCountsAt) ), rep( colHeLa, nrow(nCountsHeLa) ) )
```



3.4 Estimating technical noise

Estimate sample moments per gene.

```
meansHeLa <- rowMeans( nCountsHeLa )
varsHeLa <- rowVars( nCountsHeLa )
cv2HeLa <- varsHeLa / meansHeLa^2
```

Find the minimum mean value for the fit.

```
minMeanForFit <- unname( quantile( meansHeLa[ which( cv2HeLa > .3 ) ], .95 ) )
minMeanForFit
```

```
[1] 1445.372
```

Perform the fit.

```
useForFit <- meansHeLa >= minMeanForFit
fit <- glmgam.fit( cbind( a0 = 1, altilde = 1/meansHeLa[useForFit] ),
    cv2HeLa[useForFit] )
fit$coefficients</pre>
```

```
a0 altilde
0.03848821 675.46699263
```

Subtract Xi.

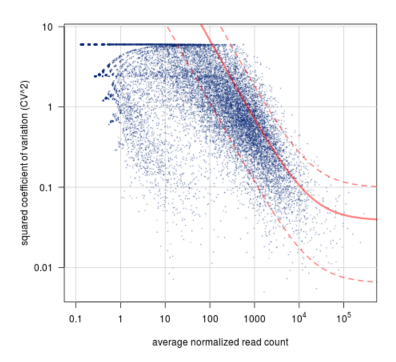
```
xi <- mean( 1 / sfHeLa )
a0 <- unname( fit$coefficients["a0"] )
a1 <- unname( fit$coefficients["altilde"] - xi )
c( a0, a1 )</pre>
```

```
[1] 0.03848821 674.51970093
```

3.4.1 Plot of the fit

A plot of the fit, code as before (Supplementary Figure 8a).

```
# Prepare the plot (scales, grid, labels, etc.)
plot( NULL, xaxt="n", yaxt="n",
  log="xy", xlim = c( 1e-1, 3e5 ), ylim = c( .005, 8 ),
  xlab = "average normalized read count", ylab = "squared coefficient of variation (CV^2)"
axis( 1, 10^{(-1:5)}, c( "0.1", "1", "10", "100", "1000",
 expression(10^4), expression(10^5))
axis( 2, 10^{(-2:1)}, c( "0.01", "0.1", "1", "10" ), las=2 )
abline( h=10^{(-2:1)}, v=10^{(-1:5)}, col="#D0D0D0")
# Add the data points
points( meansHeLa, cv2HeLa, pch=20, cex=.2, col=colHeLa )
# Plot the fitted curve
xg <- 10^seq(-2, 6, length.out=1000)
lines( xg, (xi+a1)/xg + a0, col="\#FF000080", lwd=3)
# Plot quantile lines around the fit
df <- ncol(countsAt) - 1</pre>
lines( xg, ( (xi+a1)/xg + a0 ) * qchisq( .975, df ) / df,
  col="#FF000080", lwd=2, lty="dashed")
lines( xg, ( (xi+a1)/xg + a0 ) * qchisq( .025, df ) / df,
  col="#FF000080", lwd=2, lty="dashed")
```



3.5 Testing plant genes for high variance

First the sample moments.

```
meansAt <- rowMeans( nCountsAt )
varsAt <- rowVars( nCountsAt )
cv2At <- varsAt / meansAt^2</pre>
```

Next, calculate Psi + a1 * Theta.

```
psialtheta <- mean( 1 / sfAt ) + al * mean( sfHeLa / sfAt )
```

We again test the null hypothesis that the biological CV is below 50%.

```
minBiolDisp <- .5^2
```

Calculate Omega, then perform the test

```
m <- ncol(countsAt)
cv2th <- a0 + minBiolDisp + a0 * minBiolDisp
testDenom <- ( meansAt * psialtheta + meansAt^2 * cv2th ) / ( 1 + cv2th/m )
p <- 1 - pchisq( varsAt * (m-1) / testDenom, m-1 )</pre>
```

Adjust for multiple testing, cut at 10%:

```
padj <- p.adjust( p, "BH" )
sig <- padj < .1
sig[is.na(sig)] <- FALSE
table( sig )</pre>
```

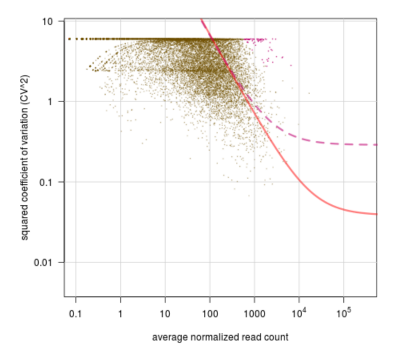
```
sig
```

```
FALSE TRUE 33538 64
```

3.5.1 Plot

Same code as before, now to produce Supplementary Figure 8b.

```
# Prepare plot in the same manner as before
plot( NULL, xaxt="n", yaxt="n",
  log="xy", xlim = c(1e-1, 3e5), ylim = c(.005, 8),
  xlab = "average normalized read count", ylab = "squared coefficient of variation (CV^2)"
axis( 1, 10^(-1:5), c( "0.1", "1", "10", "100", "1000",
 axis( 2, 10^(-2:1), c( "0.01", "0.1", "1", "10" ), las=2 )
abline ( h=10^{(-2:1)}, v=10^{(-1:5)}, col="#D0D0D0")
# Plot the plant genes, use a different color if they are highly variable
points ( meansAt, cv2At, pch=20, cex=.2,
  col = ifelse( padj < .1, "#C0007090", colAt ) )</pre>
# Add the technical noise fit, as before
xg <- 10^seq(-2, 6, length.out=1000)
lines( xg, (xi+a1)/xg + a0, col="#FF000080", lwd=3)
# Add a curve showing the expectation for the chosen biological CV^2 thershold
lines(xg, psialtheta/xg + a0 + minBiolDisp, lty="dashed", col="#C0007090", lwd=3)
```



3.6 Table of highly variable genes

Write out Supplementary Table 3.

```
log2RelExprAt <- log2( nCountsAt / meansAt )
highVarTable <- data.frame(
   row.names = NULL,</pre>
```

```
geneID = rownames(countsAt)[ sig ],
  geneSymbol = geneSymbolsAt[ sig ],
  meanNormCount = meansAt[ sig ],
  strongest = factor( colnames( log2RelExprAt )[
      apply( log2RelExprAt[ sig, ], 1, which.max ) ] ),
  log2RelExprAt[ sig, ],
  check.names=FALSE )

head( highVarTable )
```

```
geneID geneSymbol meanNormCount strongest
                                                     QC1
                                                                QC2
                                                                          QC3
                                                                                     QC4
1 AT1G02500 AtSAM1 1600.104 QC2 -1.488582 2.182097 -2.665379 -1.787801 -0
                                     QC3 -3.469101 -7.638737 2.549669 -5.054241 -6
QC3 -10.521968 -8.541857 2.577869 -10.473936 -5
2 AT1G02690
               IMPA-6
                           1478.025
            ATCSLD5
3 AT1G02730
                            1842.709
                                          QC3 -5.706594 -4.545911 2.564255 -11.565453 -5
                           1308.925
4 AT1G03780
              AtTPX2
                           4292.894
1240.020
5 AT1G06760 AT1G06760
                                         QC5 -12.742090 -3.410304 -1.912597 -2.691243 2
6 AT1G07790
                                         QC5 -3.458650 -9.555354
                                                                         -Inf -4.995580 2
                HTB1
        QC6
1 -7.7962823
2 -7.6817881
3 -7.3218736
4 -7.2434655
5 -0.6164412
6 -9.7504081
```

Write out the table

```
write.csv( highVarTable, file="highly_variant_genes_QC.tsv", row.names=FALSE )
```

3.7 GO analysis

The TopGo analysis, as before.

This time, we include only genes with an average count above 600.

```
minCountForEnrichment <- 600
```

The work function, as before.

```
topGOAnalysis <- function( geneIDs, inUniverse, inSelection )
   sapply( c( "MF", "BP", "CC" ), function( ont ) {
      alg <- factor( as.integer( inSelection[inUniverse] ) )
      names(alg) <- geneIDs[inUniverse]
      tgd <- new( "topGOdata", ontology=ont, allGenes = alg, nodeSize=5,
            annot=annFUN.org, mapping="org.At.tair.db" )
      resultTopGO <- runTest(tgd, algorithm = "elim", statistic = "Fisher" )
      GenTable( tgd, resultTopGO, topNodes=15 ) },
      simplify=FALSE )</pre>
```

The analysis.

```
goResults <-
  topGOAnalysis(
   rownames(countsAt),
  meansAt >= minCountForEnrichment & !is.na(padj),
  padj < .1 )</pre>
```

3.7.1 Results:

```
goResults[["MF"]]
```

	GO.ID	Term	Annotated	Significant	Expected	re
1	GO:0016538	cyclin-dependent protein kinase regulato	5	5	0.59	4
2	GO:0003777	microtubule motor activity	5	5	0.59	4
3	GO:0003677	DNA binding	65	19	7.66	
4	GO:0019901	protein kinase binding	7	5	0.83	0.(
5	GO:0009055	electron carrier activity	8	4	0.94	0.0
6	GO:0046906	tetrapyrrole binding	10	4	1.18	0.(
7	GO:0020037	heme binding	10	4	1.18	0.(
8	GO:0005506	iron ion binding	12	4	1.41	0.(
9	GO:0016705	oxidoreductase activity, acting on paire	10	3	1.18	0.:
10	GO:0019825	oxygen binding	6	2	0.71	0.:
11	GO:0005200	structural constituent of cytoskeleton	6	2	0.71	0.:
12	GO:0005488	binding	320	47	37.72	0.1
13	GO:0019899	enzyme binding	10	6	1.18	0.1
14	GO:0016765	transferase activity, transferring alkyl	10	2	1.18	0.:
15	GO:0032559	adenyl ribonucleotide binding	72	10	8.49	0.:

```
goResults[["CC"]]
```

	GO.ID	Term	Annotated	Significant	Expected	resul
1	GO:0000786	nucleosome	21	16	2.38	1.2e-
2	GO:0005874	microtubule	12	5	1.36	0.00
3	GO:0005634	nucleus	152	24	17.24	0.02
4	GO:0005694	chromosome	26	18	2.95	0.05
5	GO:0009579	thylakoid	9	3	1.02	0.07
6	GO:0031984	organelle subcompartment	5	2	0.57	0.10
7	GO:0043227	membrane-bounded organelle	297	38	33.68	0.13
8	GO:0043231	intracellular membrane-bounded organelle	297	38	33.68	0.13
9	GO:0045298	tubulin complex	7	2	0.79	0.18
10	GO:0044427	chromosomal part	25	17	2.84	0.29
11	GO:0031981	nuclear lumen	68	9	7.71	0.35
12	GO:0009505	plant-type cell wall	11	2	1.25	0.36
13	GO:0043233	organelle lumen	70	9	7.94	0.39
14	GO:0070013	intracellular organelle lumen	70	9	7.94	0.39
15	GO:0031974	membrane-enclosed lumen	70	9	7.94	0.39

```
goResults[["BP"]]
```

	GO.ID	Term	Annotated	Significant	Expected	re
1	GO:0006334	nucleosome assembly	23	16	2.74	3.
2	GO:0051322	anaphase	15	10	1.79	5.
3	GO:0051567	histone H3-K9 methylation	16	10	1.91	1.
4	GO:0016572	histone phosphorylation	11	8	1.31	3.
5	GO:0008283	cell proliferation	29	15	3.45	6.
6	GO:0000911	cytokinesis by cell plate formation	26	12	3.10	7.
7	GO:0000079	regulation of cyclin-dependent protein $k\dots$	5	5	0.60	2.
8	GO:0051225	spindle assembly	6	5	0.71	0.
9	GO:0007018	microtubule-based movement	12	7	1.43	0.
10	GO:0010583	response to cyclopentenone	10	6	1.19	0.
11	GO:0000087	M phase of mitotic cell cycle	10	6	1.19	0.
12	GO:0010389	regulation of $G2/M$ transition of mitotic	5	4	0.60	0.
13	GO:0006275	regulation of DNA replication	8	5	0.95	0.
14	GO:0000226	microtubule cytoskeleton organization	23	12	2.74	0.
15	GO:0006306	DNA methylation	13	6	1.55	0.

3.8 Heatmap

The code to produce the heatmap (Supplementary Figure 9b), exactly as before.

```
mn <- ( m - zlim[1] ) / ( zlim[2] - zlim[1] )
mn[ mn<0 ] <- 0
mn[ mn>1 ] <- 1
mn <- 1 + round( mn * (ncol(col)-1) )
a <- array( NA_real_, c( nrow(m)*pxWidth, ncol(m)*pxHeight, 3 ) )
for( i in 1:nrow(mn) )
    for( j in 1:ncol(mn) )
        a[ (((i-1)*pxWidth)+1) : (i*pxWidth), (((j-1)*pxHeight)+1) : (j*pxHeight), ] <-
            rep( col[, mn[ i, j ] ], each = pxWidth*pxHeight )

Image( a, colormode="color" )
}

relSig <- log2RelExprAt[ sig, ]
relSig[ relSig < -4 ] <- -4
ord <- hclust(dist(relSig))$order
hmSig <- pixelHeatmap( t( relSig[ord,] ), zlim=c( -3, 3 ), pxWidth=30, col=colorRampPalette(c("blue", "gray", "red"))(100) )</pre>
```

This time, we use these GO terms:

```
someGOTerms <- c( "GO:0009684", "GO:0003777", "GO:0016538", "GO:0051322",
    "GO:0051567", "GO:0000911", "GO:0003677", "GO:0000786")
unname( t( sapply( someGOTerms, function(x) toTable(GOTERM[ x ])[1,2:3] ) ) )</pre>
```

```
[,1] [,2]
[1,] "GO:0009684" "indoleacetic acid biosynthetic process"
[2,] "GO:0003777" "microtubule motor activity"
[3,] "GO:0016538" "cyclin-dependent protein kinase regulator activity"
[4,] "GO:0051322" "anaphase"
[5,] "GO:0051567" "histone H3-K9 methylation"
[6,] "GO:0000911" "cytokinesis by cell plate formation"
[7,] "GO:0003677" "DNA binding"
[8,] "GO:0000786" "nucleosome"
```

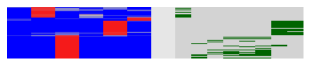
Make the seond heatmap, put it next to the first, and save the image.

```
sigInGO <- sapply( unname(someGOTerms), function(go)
    rownames(countsAt)[sig] %in% org.At.tairGO2TAIR[[ go ]] )

hmGO <- pixelHeatmap( t(sigInGO[ ord, ]), col=c( "lightgray", "darkgreen" ), pxWidth=20 )

spacer <- Image( array( .9, 30 * 3 * sum(sig), dim=c( 30, sum(sig), 3 ) ), colormode="color"

writeImage(
    Image( abind::abind( hmSig, spacer, hmGO, along=1 ), colormode="color" ),
    files="heatmap_QC.png" )</pre>
```



Save the current state.

```
save.image( "QC_analysis_image.RData" )
```

4 Analysis of the mouse cells

4.1 Count table and normalization

The count table for the mouse data can be found in Supplementary Table 5:

```
dataMouse <- read.csv( "Supplementary_Table_5.csv", row.names=1 )
dataMouse[ 1:10, 1:5 ]</pre>
```

```
length cell_01 cell_02 cell_03 cell_04
ERCC-00002
          1061 13442 39379 6800
                                     9697
ERCC-00003
                  207
                                        888
           1023
                         0
                                 33
ERCC-00004
           523
                  2762
                         6710
                                2526
                                       2375
ERCC-00009
                                217
           984
                   8
                         6125
                                         -5
                         0
ERCC-00012
           994
                    Ω
                                 0
ERCC-00013
           808
                   0
                          0
                                  0
                                          0
                          0
                    0
                                 0
ERCC-00014 1957
                                          Ω
                         0
ERCC-00016
                    0
                                  0
                                          0
           844
ERCC-00017
           1136
                    0
                                  0
                                          0
ERCC-00019
           644
                    Ω
                         1381
                                  Ω
                                          0
```

Again, we split the table into two sub-tables, one with the ERCC spikes, (countsERCC), one with the mouse genes (countsMmus). The first column, with the transcript length, is set aside.

```
geneTypes <- factor( c( Mm="Mmus", ER="ERCC" )[
   substr( rownames(dataMouse), 1, 2 ) ] )
countsMmus <- dataMouse[ which( geneTypes=="Mmus" ), -1 ]
countsERCC <- dataMouse[ which( geneTypes=="ERCC" ), -1 ]
lengthsMmus <- dataMouse[ which( geneTypes=="Mmus" ), 1 ]
lengthsERCC <- dataMouse[ which( geneTypes=="ERCC" ), 1 ]</pre>
```

Calculate size factors

```
sfMmus <- estimateSizeFactorsForMatrix( countsMmus )
sfERCC <- estimateSizeFactorsForMatrix( countsERCC )
rbind( sfMmus, sfERCC )</pre>
```

```
cell 01 cell 02 cell 03 cell 04 cell 05 cell 08 cell 09 cell 10
sfMmus 1.054806 0.8631698 1.314802 0.8392654 1.923025 2.038466 0.9484543 1.318534 0.9377162
sfERCC 1.341199 3.5780220 1.001075 1.0414057 1.258551 0.714875 0.5333393 1.178008 1.6108530
                 cell_13 cell_14 cell_15 cell_16 cell_17 cell_18 cell_19 cel.
        cell_12
sfMmus 1.0650922 0.05676978 0.7071812 1.508444 1.5083955 1.6092835 1.348358 0.7929501 1.575
sfERCC 0.4885753 3.90732773 1.0575947 1.088776 0.3442378 0.9069057 1.019297 1.6010922 0.446
        cell_21 cell_22 cell_23
                                  cell_24 cell_25 cell_26 cell_27 cell_28 cell_27 cell_28 cell_28 cell_29
sfMmus 1.1275932 0.7125047 0.2097335 1.2002740 1.2652195 0.09265965 1.6101646 1.5289089 0.6
sfERCC 0.9001471 1.9687150 1.6600806 0.5552053 0.7471373 3.73041540 0.9049005 0.9355409 2.7
        cell_30 cell_31 cell_32 cell_33 cell_34 cell_35 cell_36 cell_37 cel
sfMmus 0.8833294 1.6117294 0.9131230 0.6266095 0.4910416 1.231094 0.9456337 0.4741488 1.626
sfERCC 0.2198629 0.6530132 0.1959819 1.4602831 3.7636827 1.771898 0.1028514 2.7608761 0.694
       sfMmus 2.067016 1.5703639 2.2319662 1.531569 0.4751253 1.433638 1.4033780 1.509945 1.385977
sfERCC 1.936887 0.3855543 0.7174266 1.316699 3.2065296 0.851406 0.3767175 1.511495 0.422995
        cell_48 cell_49 cell_50 cell_51 cell_52 cell_53 cell_54
                                                                    cell_55 cell_56
sfMmus 0.9752096 1.7122014 1.395363 2.029381 0.4192704 1.733174 2.007804 1.6556897 1.551027
sfERCC 0.4137318 0.3755807 2.453767 2.088099 1.8768628 1.515678 1.754392 0.8157246 0.307815
        cell_62 cell_63
sfMmus 1.8791183 1.8109205 2.0053160 0.5212242 1.286422 1.7020164 2.2123285 0.1373123 1.71
sfERCC 0.7091403 0.6844449 0.7593432 0.7342300 1.234711 0.4970688 0.9522804 13.6460462 1.62
       cell_66 cell_67 cell_68 cell_69 cell_70
                                                   cell_71 cell_72 cell_73
sfMmus 1.725958 1.4598322 1.674938 1.5899826 1.4466890 1.0997875 0.8398078 1.734892 1.23888
sfERCC 2.499659 0.7241718 1.702510 0.4682272 0.4076763 0.1952154 6.2377984 1.656883 0.85136
                         cell_78 cell_79 cell_80 cell_81 cell_82 cell_83
        cell_76 cell_77
sfMmus 0.9557100 1.2679037 0.7024893 1.6270177 1.509954 1.6773127 1.875683 0.633388 2.58677
sfERCC 0.6596151 0.5664658 1.1481714 0.8049232 6.085711 0.5256015 2.378635 3.469949 0.90372
        cell_85 cell_86 cell_87 cell_88 cell_89 cell_90 cell_91 cell_94 cell_95
sfMmus 0.3708505 1.964748 1.7201196 2.022113 1.324386 1.131953 0.9803028 1.369537 1.71133 1
sfERCC 3.1826913 1.139850 0.9348608 1.020459 1.204492 1.308119 0.9468956 2.970329 1.65396 0
```

Normalize by them:

```
nCountsERCC <- t( t(countsERCC) / sfERCC )
nCountsMmus <- t( t(countsMmus) / sfMmus )
```

We calculate the sample moments:

```
meansERCC <- rowMeans( nCountsERCC )
varsERCC <- rowVars( nCountsERCC )
cv2ERCC <- varsERCC / meansERCC^2

meansMmus <- rowMeans( nCountsMmus )
varsMmus <- rowVars( nCountsMmus )
cv2Mmus <- varsMmus / meansMmus^2</pre>
```

Normalize the mean counts by transcrip length (i.e., "per kilobase", "PK"), too:

```
meansERCCPK <- meansERCC / lengthsERCC * 1e3
meansMmusPK <- meansMmus / lengthsMmus * 1e3
```

4.2 Fit technical noise

We perform the fit as usual. However, as we have only rather few spikes, we have to be a bit more generous with the mean cut-off, now using the 80-percentile instead of the 95-percentile.

```
minMeanForFitA <- unname( quantile( meansERCC[ which( cv2ERCC > .3 ) ], .8 ) )
useForFitA <- meansERCC >= minMeanForFitA
minMeanForFitA
table( useForFitA )
```

```
[1] 81.63606
useForFitA
FALSE TRUE
71 21
```

Afterwards, we will compare with a fit using length-normalized counts. We prepare by finding the minimum for these, too:

```
minMeanForFitB <- unname( quantile( meansERCCPK[ which( cv2ERCC > .3 ) ], .8 ) )
useForFitB <- meansERCCPK >= minMeanForFitB
minMeanForFitB
table( A=useForFitA, B=useForFitB )
```

```
[1] 100.2707

B
A FALSE TRUE
FALSE 70 1
TRUE 1 20
```

Note that the two lists overlap well.

We perform both fits.

```
fitA <- glmgam.fit( cbind( a0 = 1, altilde = 1/meansERCC[useForFitA] ),
    cv2ERCC[useForFitA] )

fitB <- glmgam.fit( cbind( a0 = 1, altilde = 1/meansERCCPK[useForFitB] ),
    cv2ERCC[useForFitB] )</pre>
```

How much variance do the two fits explain?

```
residualA <- var( log( fitted.values(fitA) ) - log( cv2ERCC[useForFitA] ) )
```

```
A B
0.7945620 0.8873808
```

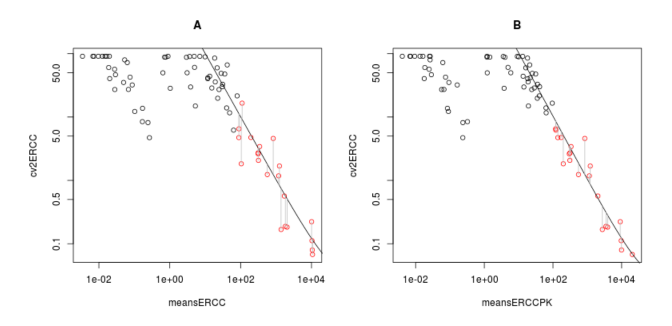
Fit B, which used the length-normalized counts, performed better.

As a second check, we plot both fits.

```
par( mfrow=c(1,2) )

plot( meansERCC, cv2ERCC, log="xy", col=1+useForFitA, main="A" )
  xg <- 10^seq( -3, 5, length.out=100 )
  lines( xg, coefficients(fitA) ["a0"] + coefficients(fitA) ["altilde"]/xg )
  segments( meansERCC[useForFitA], cv2ERCC[useForFitA],
      meansERCC[useForFitA], fitA$fitted.values, col="gray" )

plot( meansERCCPK, cv2ERCC, log="xy", col=1+useForFitB, main="B" )
  lines( xg, coefficients(fitB) ["a0"] + coefficients(fitB) ["altilde"]/xg )
  segments( meansERCCPK[useForFitB], cv2ERCC[useForFitB],
      meansERCCPK[useForFitB], fitB$fitted.values, col="gray" )</pre>
```



4.3 Test for high variance

We start with the test using fit A:

```
minBiolDisp <- .5^2

xi <- mean( 1 / sfERCC )
m <- ncol(countsMmus)
psialthetaA <- mean( 1 / sfERCC ) +
        ( coefficients(fitA)["altilde"] - xi ) * mean( sfERCC / sfMmus )
cv2thA <- coefficients(fitA)["a0"] + minBiolDisp + coefficients(fitA)["a0"] * minBiolDisp</pre>
```

```
testDenomA <- ( meansMmus * psialthetaA + meansMmus^2 * cv2thA ) / ( 1 + cv2thA/m )
pA <- 1 - pchisq( varsMmus * (m-1) / testDenomA, m-1 )
padjA <- p.adjust( pA, "BH" )
table( padjA < .1 )</pre>
```

```
FALSE TRUE
29489 1198
```

Using fit B and the length-normalized counts, we get

```
varsMmusPK <- rowVars( nCountsMmus / lengthsMmus * 1e3 )

psialthetaB <- mean( 1 / sfERCC ) +
    ( coefficients(fitB)["altilde"] - xi ) * mean( sfERCC / sfMmus )
    cv2thB <- coefficients(fitB)["a0"] + minBiolDisp + coefficients(fitB)["a0"] * minBiolDisp
    testDenomB <- ( meansMmusPK * psialthetaB + meansMmusPK^2 * cv2thB ) / ( 1 + cv2thB/m )

pB <- 1 - pchisq( varsMmusPK * (m-1) / testDenomB, m-1 )
    padjB <- p.adjust( pB, "BH" )

table( B = padjB < .1 )</pre>
```

```
B
FALSE TRUE
30137 523
```

Despite the better technical noise fit in case B (length adjusted), we get more results in case A (not length adjusted). The next section explores this.

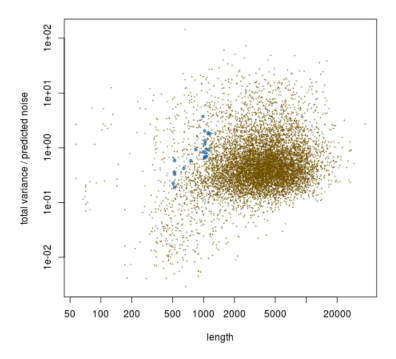
4.4 A diagnostic plot

We have a closer look at the reliability of the technical noise predictions from fit A. The variance from technical noise, predicted for a biological gene is given by Omega(0, mu), where mu is the normalized mean count for the gene, and Omega is the function defined in the Online Methods. Dividing by mu^2 to get CV^2 values (and ignoring the negligible term a0/m), we get

```
predictedNoiseCV2 <- psialthetaA / meansMmus + coefficients(fitA)["a0"]</pre>
```

We plot the ratio of observed total CV^2 to predicted technical CV^2 against transcript length, using only genes with a mean count above the cut-off also used for the fit. This is Supplementary Figure 7.

```
useInPlot <- meansMmus>minMeanForFitA
plot( lengthsMmus[useInPlot], ( cv2Mmus / predictedNoiseCV2 )[useInPlot], log="xy",
    pch=20, cex=.2, col = "#705000A0", xlab = "length", ylab="total variance / predicted noisepoints( lengthsERCC[useForFitA], cv2ERCC[useForFitA] / fitted.values(fitA), pch=20, cex=1,
```

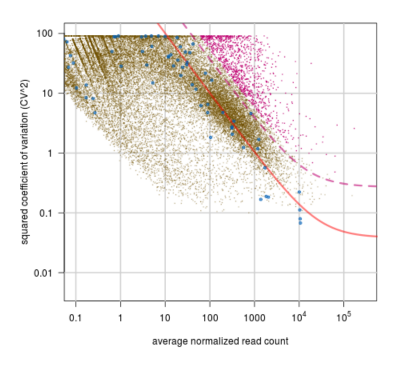


See Supplementary Note 5 for a discussion.

4.5 Plot of results

To produce Figure 3, which depicts the results of fit and test A, the following code was used

```
plot( NULL, xaxt="n", yaxt="n",
     log="xy", xlim = c( 1e-1, 3e5 ), ylim = c( .005, 100 ),
    xlab = "average normalized read count", ylab = "squared coefficient of variation (CV^2)
axis( 1, 10^(-1:5), c( "0.1", "1", "10", "100", "1000",
                    expression(10^4), expression(10^5))
axis(2, 10^{(-2:2)}, c("0.01", "0.1", "1", "10", "100"), las=2) abline(h=10^{(-2:1)}, v=10^{(-1:5)}, col="#DODODO", lwd=2)
# Plot the plant genes, use a different color if they are highly variable
points ( meansMmus, cv2Mmus, pch=20, cex=.2,
      col = ifelse(padjA < .1, "#C0007090", "#70500040"))
# Add the technical noise fit, as before
xg <- 10^seq(-2, 6, length.out=1000)
lines( xg, coefficients(fitA)["altilde"] / xg + a0, col="#FF000080", lwd=3)
# Add a curve showing the expectation for the chosen biological CV^2 thershold
lines (xg, psialthetaA/xg + coefficients(fitA)["a0"] + minBiolDisp,
   lty="dashed", col="#C0007090", lwd=3 )
# Add the normalised ERCC points
points( meansERCC, cv2ERCC, pch=20, cex=1, col="#0060B8A0" )
```



Save image:

```
save.image( "mouse_analysis_image.RData" )
```

Date: 2013-08-08 11:03:10 CEST

Author: Simon Anders

Org version 7.8.02 with Emacs version 23

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