

erccdashboard Package Vignette

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This vignette describes the use of the erccdashboard R package to analyze External RNA Control Consortium (ERCC) spike-in control ratio mixtures in gene expression experiments. Analysis is shown for two types of samples spiked with ERCC control ratio mixtures from the SEQC project

- Rat toxicogenomics treatment and control samples for different drug treatments
- Human reference RNA samples from the MAQC I project, Universal Human Reference RNA (UHRR) and Human Brain Reference RNA (HBRR)

A subset of the large data set produced in the SEQC study is provided here as examples. The three sets of example data are:

1. Rat toxicogenomics RNA-Seq gene expression count data
2. UHRR/HBRR RNA-Seq gene expression count data
3. UHRR/HBRR Microarray gene expression fluorescent intensity data

1 Rat Toxicogenomics Example: MET (methimazole treatment) and CTL (control) Experiment

1.1 Load data and define input parameters

Load the testerccdashboard package.

```
> library( "erccdashboard" )
```

Load the package gene expression data.

```
> data(SEQC.Example)
```

The R workspace should now contain 5 objects Three of these objects are gene expression experiment expression measures:

- Lab4.array.UHRR.HBRR - Fluorescent signal data from an Illumina beadarray microarray experiment with UHRR and HBRR
- COH.RatTox.ILM.MET.CTL.countTable - RNA-Seq count data from a rat toxicogenomics experiment
- Lab5.ILM.UHRR.HBRR.countTable - RNA-Seq count data from Lab 5 in the SEQC interlaboratory study with UHRR and HBRR

The other two objects are vectors of total reads for the 2 sequencing experiments

- COH.RatTox.ILM.MET.CTL.totalReads - total sequenced reads factors for each column in the corresponding rat experiment count table

- Lab5.ILM.UHRR.HBRR.totalReads - total sequenced reads factors for each column in the corresponding UHRR/HBRR count table

Take a look at the count table for the MET experiment.

```
> head(COH.RatTox.ILM.MET.CTL.countTable)
  Feature MET_1 MET_2 MET_3 CTL_1 CTL_2 CTL_3
16499 ERCC-00002 16629 18798 26568 36600 45436 25163
16500 ERCC-00003 1347 1565 1983 3048 3447 2195
16501 ERCC-00004 4569 5570 6755 1240 1484 902
16502 ERCC-00009 811 869 1123 909 1073 537
16503 ERCC-00012 0 0 0 0 0 0
16504 ERCC-00013 3 1 2 1 5 1
```

The first column of the count table, Feature, contains unique names for all the transcripts that were quantified in this experiment. The remaining columns represent replicates of the pair of samples, in this count table the control sample is labeled CTL and the treatment sample is labeled MET. An underscore is included to separate the sample names from the replicate numbers during analysis. This column name format Sample_Rep is required for the columns of any input count table.

These RNA-Seq count tables also must be unnormalized count data, because the default differential expression testing of RNA-Seq experiments in the erccdashboard is done with the QuasiSeq package, which requires the use of integer count data. During the erccdashboard analysis the input count data will be normalized using library size factors. These factors can be provided by the user as the input argument repNormFactor, a vector of total reads. The example total reads vectors we provide here were derived from the FASTQ files associated with each column in the count tables. Alternatively, if repNormFactor is undefined (or NULL) for an RNA-Seq analysis then a vector of total mapped reads (column sums of a count table) will be calculated and used for normalization.

1.2 Quick analysis: runDashboard

To run the default analysis function runDashboard on the MET-CTL rat toxicogenomics RNA-Seq experiment, the following input arguments are required:

```
> dataType = "count" # count for RNA-Seq data, array for microarray data
> exTable = COH.RatTox.ILM.MET.CTL.countTable # the expression measure table
> filenameRoot = "COH.ILM" # user defined filename prefix for results files
> sample1Name = "MET" # name for sample 1 in the experiment
> sample2Name = "CTL" # name for sample 2 in the experiment
> erccmix = "RatioPair" # name of ERCC mixture design, "RatioPair" is default
> erccdilution = 1/100 # dilution factor used for Ambion spike-in mixtures
> spikeVol = 1 # volume (in microliters) of diluted spike-in mixture added to
>           # total RNA mass
> totalRNAmass = 0.500 # mass (in micrograms) of total RNA
> choseFDR = 0.1 # user defined false discovery rate (FDR), default is 0.05
```

For any experiment the sample spiked with ERCC Mix 1 is sample1Name and the sample spiked with ERCC Mix 2 is sample2Name. In this experiment sample1Name = MET and sample2Name = CTL. For a more robust experimental design the reverse spike-in design could be created using additional replicates of the treatment and control samples. ERCC Mix 2 would be spiked into MET samples and ERCC Mix 1 would be spiked into CTL control replicates.

The dilution factor of the pure Ambion ERCC mixes prior to spiking into total RNA samples is erccdilution. The amount of diluted ERCC mix spiked into the total RNA sample is spikeVol (units are μL). The mass of total RNA spiked with the diluted ERCC mix is totalRNAmass (units are μg).

The final required input parameter, choseFDR, is the False Discovery Rate (FDR) for differential expression testing. A typical choice would be 0.05 (5% FDR), so this is the default choseFDR value. For the rat data sets a more liberal FDR was used, choseFDR = 0.1.

The function runDashboard.R is provided for convenient default erccdashboard analysis. Execution of the runDashboard function calls the default functions for erccdashboard analysis and reports parameters and progress to the R console. The functions called within runDashboard.R are also available to the user (details provided in Section 4). All data and analysis results are stored in the list object exDat. For convenience the main diagnostic figures are saved to a pdf file and the exDat object is saved to an .RData object named using the filenameRoot provided by the user.

```
> exDat <- runDashboard(datType = "count",
  exTable = COH.RatTox.ILM.MET.CTL.countTable,
  filenameRoot = "COH.ILM", sample1Name = "MET",
  sample2Name = "CTL", erccmix = "RatioPair",
  erccdilution = 1/100, spikeVol = 1,
  totalRNAmass = 0.500, choseFDR = 0.1)

Initializing the exDat list structure...
choseFDR = 0.1
repNormFactor is NULL
Filename root is: COH.ILM.MET.CTL

Transcripts were removed with a mean count < 1 or more than 2
replicates with 0 counts.
Original data contained 16590 transcripts.
After filtering 11570 transcripts remain for analysis.
A total of 29 out of 92
ERCC controls were filtered from the data set
The excluded ERCCs are:
ERCC-00012 ERCC-00014 ERCC-00016 ERCC-00017 ERCC-00024
ERCC-00041 ERCC-00048 ERCC-00057 ERCC-00061 ERCC-00073
ERCC-00075 ERCC-00081 ERCC-00083 ERCC-00086 ERCC-00097
ERCC-00098 ERCC-00104 ERCC-00117 ERCC-00120 ERCC-00123
ERCC-00126 ERCC-00134 ERCC-00137 ERCC-00138 ERCC-00142
ERCC-00147 ERCC-00150 ERCC-00156 ERCC-00164

repNormFactor is NULL,
Using Default Upper Quartile Normalization Method - 75th percentile)

Library sizes:
438 517 473 397 546 389
Check for sample mRNA fraction differences(r_m)...

log.offset
6.082219 6.248043 6.159095 5.983936 6.302619 5.963579

Number of ERCC Controls Used in r_m estimate
63

Outlier ERCCs for GLM r_m Estimate:
None

GLM log(r_m) estimate:
```

-0.07014034

GLM log(r_m) estimate weighted s.e.:
0.1494555

Number of ERCCs in Mix 1 dyn range: 63

Number of ERCCs in Mix 2 dyn range: 63

These ERCCs were not included in the signal-abundance plot,
because not enough non-zero replicate measurements of these
controls were obtained for both samples:

ERCC-00058 ERCC-00067 ERCC-00077 ERCC-00168 ERCC-00028
ERCC-00033 ERCC-00040 ERCC-00109 ERCC-00154 ERCC-00158

Saving dynRangePlot to exDat

Starting differential expression tests

Show libe sizes

438 517 473 397 546 389
Disp = 0.0625 , BCV = 0.25
Disp = 0.06244 , BCV = 0.2499
[1] "Analyzing Gene # 2"
[1] "Analyzing Gene # 10"
[1] "Analyzing Gene # 100"
[1] "Analyzing Gene # 500"
[1] "Analyzing Gene # 1000"
[1] "Analyzing Gene # 2500"
[1] "Analyzing Gene # 5000"
[1] "Analyzing Gene # 10000"
[1] "Analyzing Gene # 2"
[1] "Analyzing Gene # 10"
[1] "Analyzing Gene # 100"
[1] "Analyzing Gene # 500"
[1] "Analyzing Gene # 1000"
[1] "Analyzing Gene # 2500"
[1] "Analyzing Gene # 5000"
[1] "Analyzing Gene # 10000"
[1] "Note: 'test.mat' not provided. Comparing each model \nfrom 'design.list' to first model in 'design.list'"
[1] "Spline scaling factor: 0.936657511021409"
[1] "Spline scaling factor: 0.933731194960756"
[1] "Analyzing Gene # 2"
[1] "Analyzing Gene # 10"
[1] "Analyzing Gene # 2"
[1] "Analyzing Gene # 10"
[1] "Note: 'test.mat' not provided. Comparing each model \nfrom 'design.list' to first model in 'design.list'"
[1] "Spline scaling factor: 0.933731194960756"
[1] "Finished DE testing"
[1] "Spline scaling factor: 0.933731194960756"

```
Finished examining dispersions
```

```
Threshold P-value  
0.007480282
```

```
Generating ROC curve and AUC statistics...
```

```
Area Under the Curve (AUC) Results:
```

	Ratio	AUC	Detected	Spiked
1	4:1	1.000	16	23
2	1:1.5	0.950	16	23
3	1:2	0.971	16	23

```
Estimating ERCC LODR
```

```
.....  
Ratio LODR Estimate 90% CI Lower Bound 90% CI Upper Bound  
1 4:1 26 19 33  
3 1:1.5 Inf <NA> <NA>  
4 1:2 280 140 410
```

```
ERCC LODR estimates are available  
Fold Ratio Count Log2Count_normalized  
1 4.000 4:1 26 -4.145050  
2 1.000 1:1 NA NA  
3 0.667 1:1.5 Inf Inf  
4 0.500 1:2 280 -0.716207
```

```
LODR estimates are available to code ratio-abundance plot
```

```
Saving main dashboard plots to pdf file...  
Saving exDat list to .RData file...  
Analysis completed.
```

1.3 Results of dashboard analysis

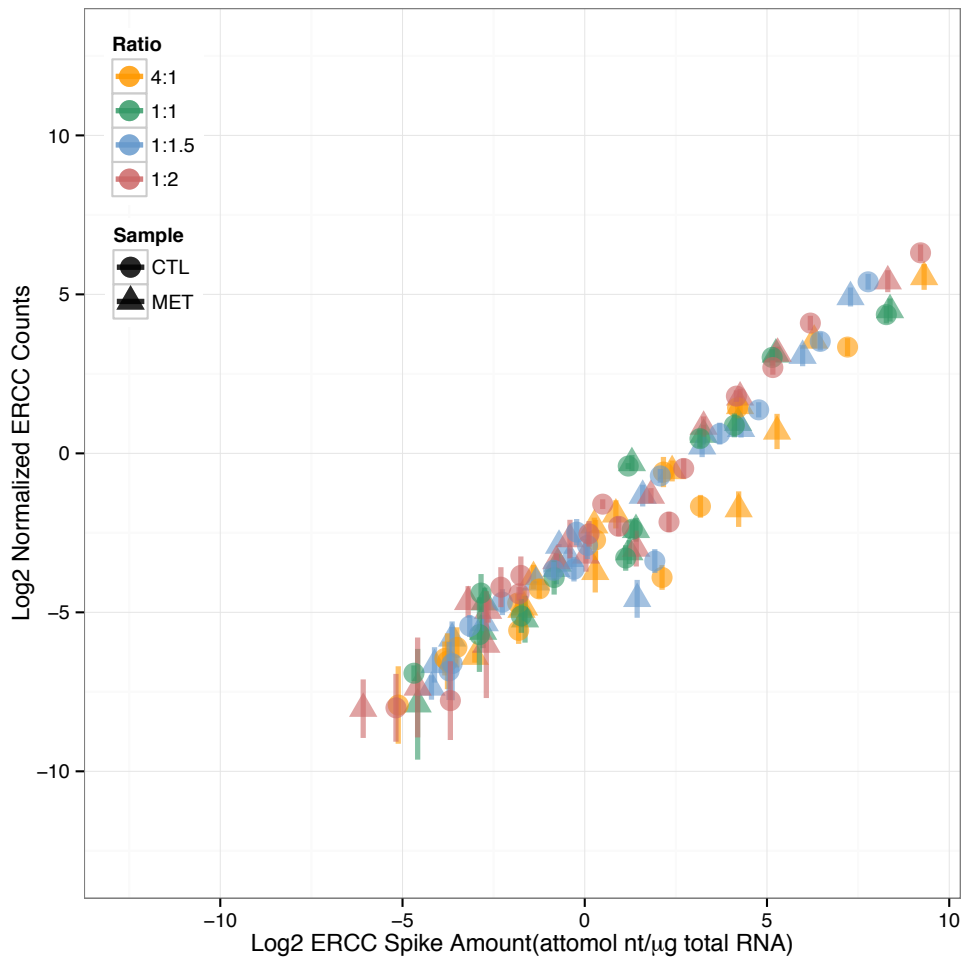
Look at the structure of exDat. The summary function will give a top level view of the list structure. The str function will give more detail. It is a good idea to set the max.level argument in the str function, because by the end of the analysis the exDat structure is quite large.

```
> summary(exDat)  
Length Class Mode  
sampleInfo 10 -none- list  
plotInfo 9 -none- list  
erccInfo 4 -none- list  
Transcripts 7 data.frame list  
designMat 3 data.frame list  
sampleNames 2 -none- character  
idCols 6 data.frame list  
normERCCDat 7 data.frame list  
libeSize 6 -none- numeric
```

```
mnLibeFactor    1      -none-      numeric
spikeFraction  1      -none-      numeric
idColsAdj      6      data.frame  list
Results         14     -none-      list
Figures        12     -none-      list
```

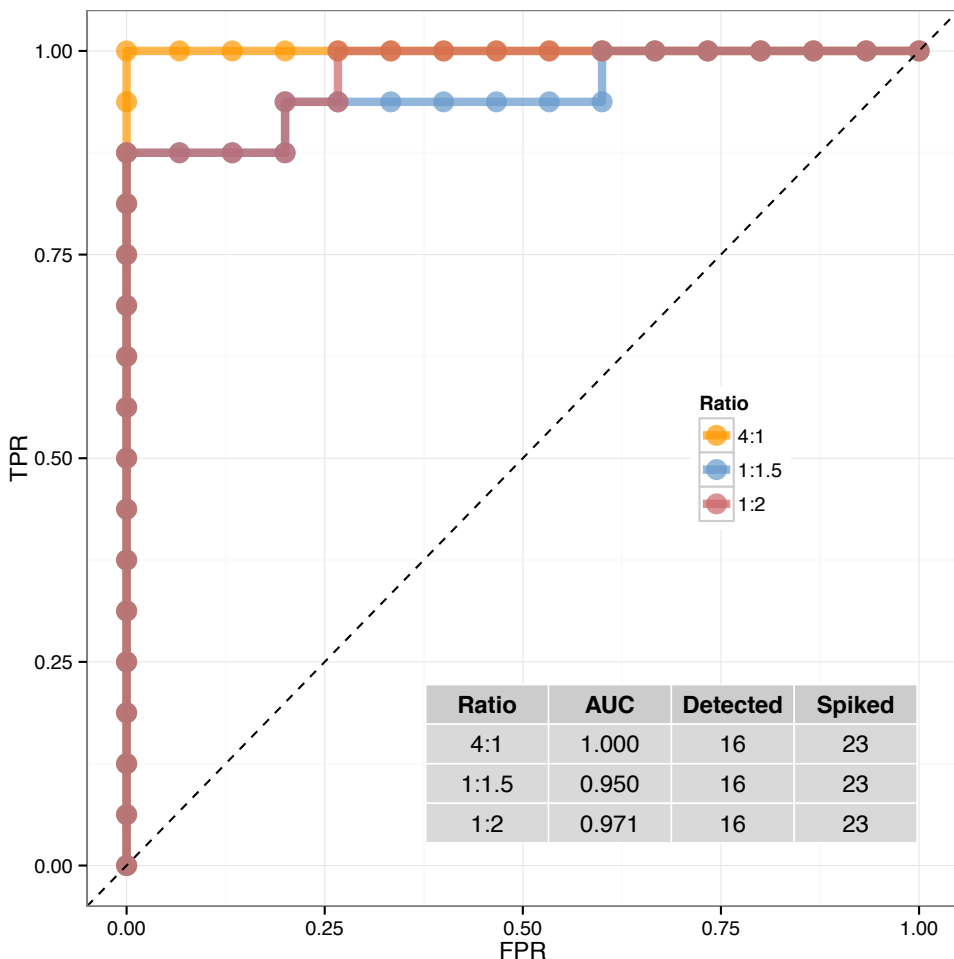
The figures from the analysis are stored in `exDat$Figures`. The four main diagnostic figures that are saved to pdf are the `dynRangePlot`, `rocPlot`, `lodrERCCPlot`, and `maPlot`.

```
> exDat$Figures$dynRangePlot
```



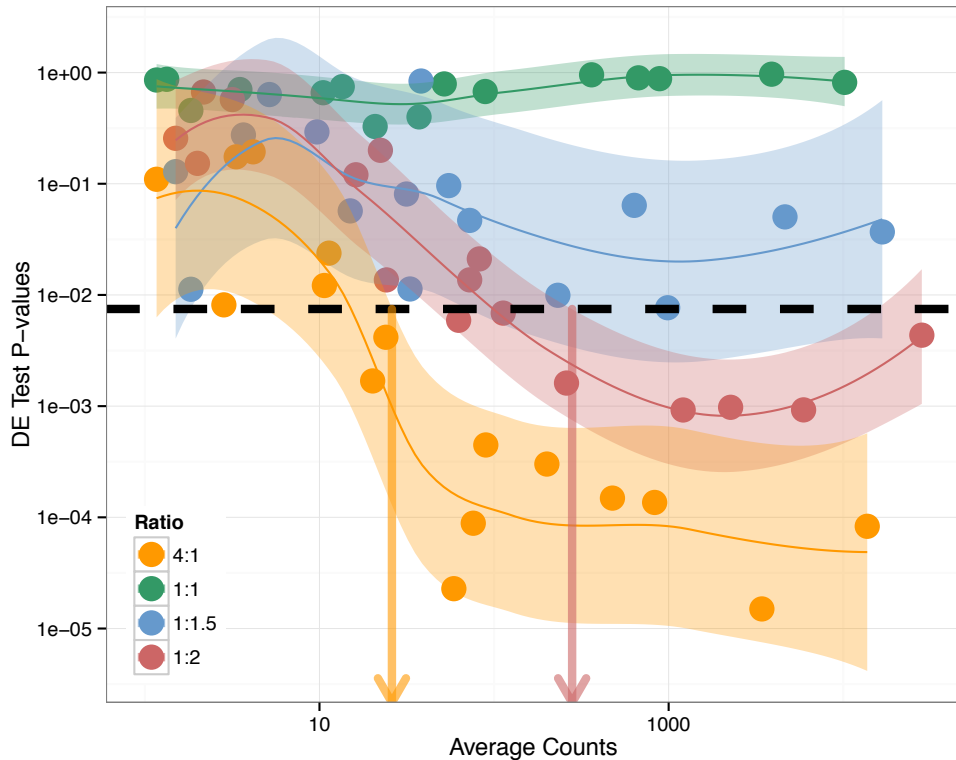
For this particular experiment the relationship between abundance and signal for the ERCC controls show that the measurement results span a 2^{15} dynamic range. These ERCC mixtures were designed to span a 2^{20} dynamic range, but there was insufficient evidence to reliably quantify ERCC transcripts at low abundances.

```
> exDat$Figures$rocPlot
```



The receiver operator characteristic (ROC) curve and the Area Under the Curve (AUC) statistic provide evidence of the diagnostic power for detecting differential expression in this rat toxicogenomics experiment. As expected with increased fold change, diagnostic power increases. The AUC summary statistic for different experiments can be used to compare diagnostic performance.

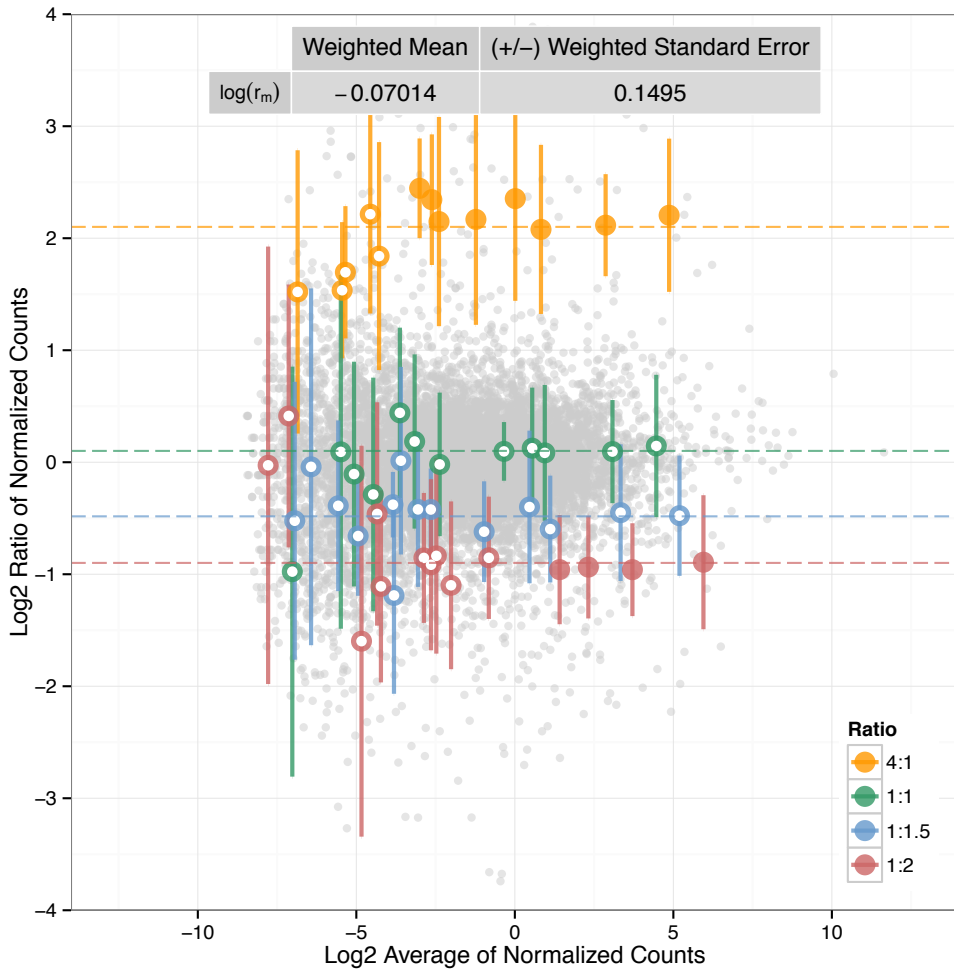
```
> exDat$Figures$lodrERCCPlot
```



Ratio	LODR Estimate	90% CI Lower Bound	90% CI Upper Bound
4:1	26	19	33
1:1.5	Inf	NA	NA
1:2	280	140	410

By modeling the relationship between average signal and p-values we can obtain Limit of Detection of Ratios (LODR) estimates for each differential fold change (or Ratio, indicated by color) and a threshold p-value, p_{thresh} , indicated by the dotted black line. LODR values can be compared between experiments to evaluate the ability to detect differences between samples as a function of transcript abundance.

```
> exDat$Figures$maPlot
```



An MA plot (Ratio of Signals vs Average Signals) shows the ratio measurements of transcripts in the pair of samples as a function of abundance. The ERCC control ratios measurements are coded to indicate which controls are above a given LODR (solid circles) or below the LODR (open circles). This plot also shows the variability in ratio measurements as a function of dynamic range and the bias in control ratio measurements (r_m), which is influenced by the mRNA fraction difference between the pair of samples.

2 SEQC Reference RNA (UHRR and HBRR) Examples

All UHRR and HBRR experiments for the SEQC interlaboratory and interplatform analysis were performed with aliquots from single large samples of UHRR and HBRR. The count data and microarray data provided here are from experiments using the same original UHRR and HBRR samples.

2.1 RNA-Seq UHRR vs. HBRR experiment analysis

```
> exDat <- runDashboard(datType="count", exTable=Lab5.ILM.UHRR.HBRR.countTable,
  filenameRoot="Lab5", sample1Name = "UHRR",
```

```
sample2Name = "HBRR", erccmix = "RatioPair",
erccdilution = 1, spikeVol = 50,
totalRNAmass = 2.5*10^(3), choseFDR = 0.01)

Initializing the exDat list structure...
choseFDR = 0.01
repNormFactor is NULL
Filename root is: Lab5.UHRR.HBRR

Transcripts were removed with a mean count < 1 or more than 2
replicates with 0 counts.
Original data contained 43919 transcripts.
After filtering 39844 transcripts remain for analysis.
A total of 2 out of 92
ERCC controls were filtered from the data set
The excluded ERCCs are:
ERCC-00057 ERCC-00083

repNormFactor is NULL,
Using Default Upper Quartile Normalization Method - 75th percentile)

Library sizes:
4048 7635 5739.25 12885 7033 6176 7059 7141.25
Check for sample mRNA fraction differences(r_m)...

log.offset
8.305978 8.940498 8.655084 9.463819 8.858369 8.728426 8.862059 8.873643

Number of ERCC Controls Used in r_m estimate
90

Outlier ERCCs for GLM r_m Estimate:
ERCC-00012 ERCC-00137 ERCC-00085 ERCC-00054 ERCC-00019

GLM log(r_m) estimate:
0.2836417

GLM log(r_m) estimate weighted s.e.:
0.03764492

Number of ERCCs in Mix 1 dyn range: 90

Number of ERCCs in Mix 2 dyn range: 90
These ERCCs were not included in the signal-abundance plot,
because not enough non-zero replicate measurements of these
controls were obtained for both samples:

ERCC-00016 ERCC-00017 ERCC-00048 ERCC-00061 ERCC-00075
ERCC-00104 ERCC-00117 ERCC-00142 ERCC-00156 ERCC-00097
ERCC-00123 ERCC-00138
```

```
Saving dynRangePlot to exDat
```

```
Starting differential expression tests
```

```
Show libe sizes
```

```
4048 7635 5739.25 12885 7033 6176 7059 7141.25
```

```
Disp = 0.00104 , BCV = 0.0322
```

```
Disp = 0.00109 , BCV = 0.033
```

```
[1] "Analyzing Gene # 2"
```

```
[1] "Analyzing Gene # 10"
```

```
[1] "Analyzing Gene # 100"
```

```
[1] "Analyzing Gene # 500"
```

```
[1] "Analyzing Gene # 1000"
```

```
[1] "Analyzing Gene # 2500"
```

```
[1] "Analyzing Gene # 5000"
```

```
[1] "Analyzing Gene # 10000"
```

```
[1] "Analyzing Gene # 15000"
```

```
[1] "Analyzing Gene # 20000"
```

```
[1] "Analyzing Gene # 25000"
```

```
[1] "Analyzing Gene # 30000"
```

```
[1] "Analyzing Gene # 35000"
```

```
[1] "Analyzing Gene # 40000"
```

```
[1] "Analyzing Gene # 2"
```

```
[1] "Analyzing Gene # 10"
```

```
[1] "Analyzing Gene # 100"
```

```
[1] "Analyzing Gene # 500"
```

```
[1] "Analyzing Gene # 1000"
```

```
[1] "Analyzing Gene # 2500"
```

```
[1] "Analyzing Gene # 5000"
```

```
[1] "Analyzing Gene # 10000"
```

```
[1] "Analyzing Gene # 15000"
```

```
[1] "Analyzing Gene # 20000"
```

```
[1] "Analyzing Gene # 25000"
```

```
[1] "Analyzing Gene # 30000"
```

```
[1] "Analyzing Gene # 35000"
```

```
[1] "Analyzing Gene # 40000"
```

```
[1] "Note: 'test.mat' not provided. Comparing each model \nfrom 'design.list' to first model in 'design.list'"
```

```
[1] "Spline scaling factor: 1.0983197686402"
```

```
[1] "Spline scaling factor: 1.09822556795109"
```

```
[1] "Analyzing Gene # 2"
```

```
[1] "Analyzing Gene # 10"
```

```
[1] "Analyzing Gene # 2"
```

```
[1] "Analyzing Gene # 10"
```

```
[1] "Note: 'test.mat' not provided. Comparing each model \nfrom 'design.list' to first model in 'design.list'"
```

```
[1] "Spline scaling factor: 1.09822556795109"
```

```
[1] "Finished DE testing"
```

```
[1] "Spline scaling factor: 1.09822556795109"
```

```
Finished examining dispersions
```

```
Threshold P-value
```

0.1517627
Threshold P-value is high for the chosen FDR of 0.01
The sample comparison indicates a large amount of
differential expression in the measured transcript
populations

Generating ROC curve and AUC statistics...

Area Under the Curve (AUC) Results:

	Ratio	AUC	Detected	Spiked
1	4:1	0.966	22	23
2	1:1.5	0.826	22	23
3	1:2	0.894	23	23

Estimating ERCC LODR

	Ratio	LODR Estimate	90% CI Lower Bound	90% CI Upper Bound
1	4:1	21	5.9	27
3	1:1.5	170	72	210
4	1:2	37	22	40

ERCC LODR estimates are available

	Fold Ratio	Count	Log2Count_normalized
1	4.000	4:1	21
2	1.000	1:1	NA
3	0.667	1:1.5	170
4	0.500	1:2	37

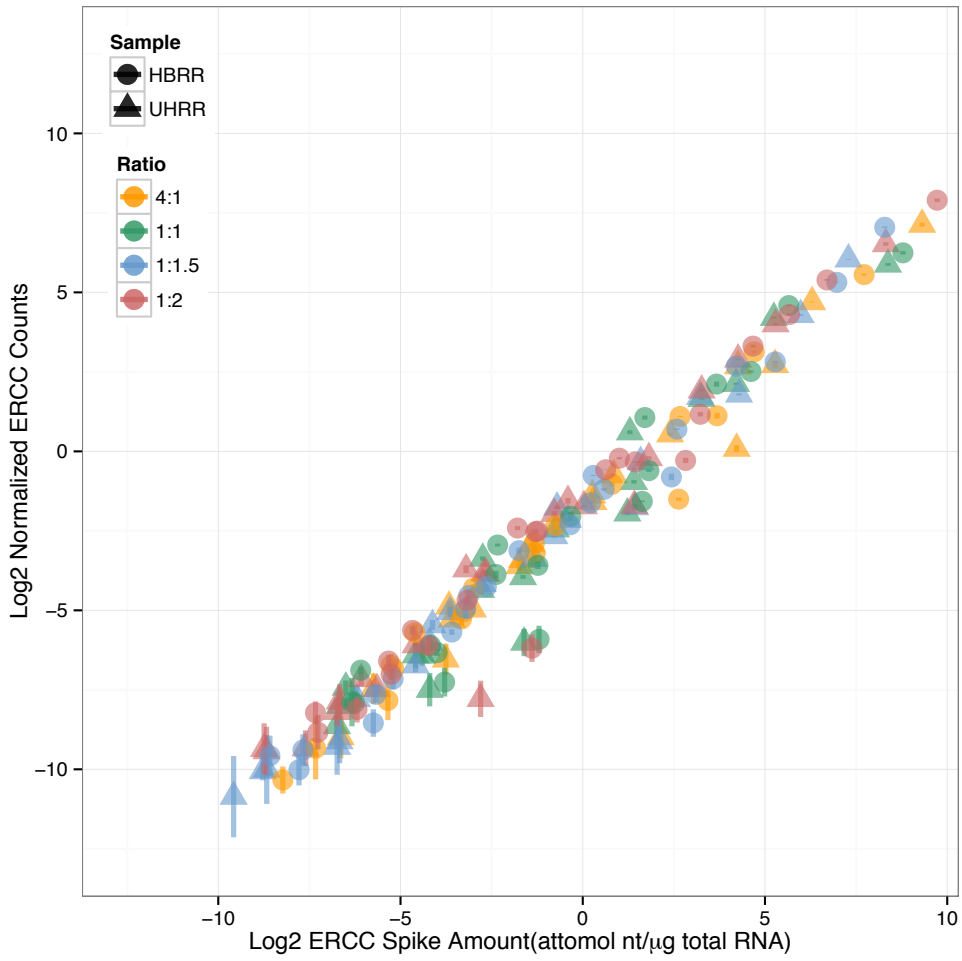
LODR estimates are available to code ratio-abundance plot

Saving main dashboard plots to pdf file...

Saving exDat list to .RData file...

Analysis completed.

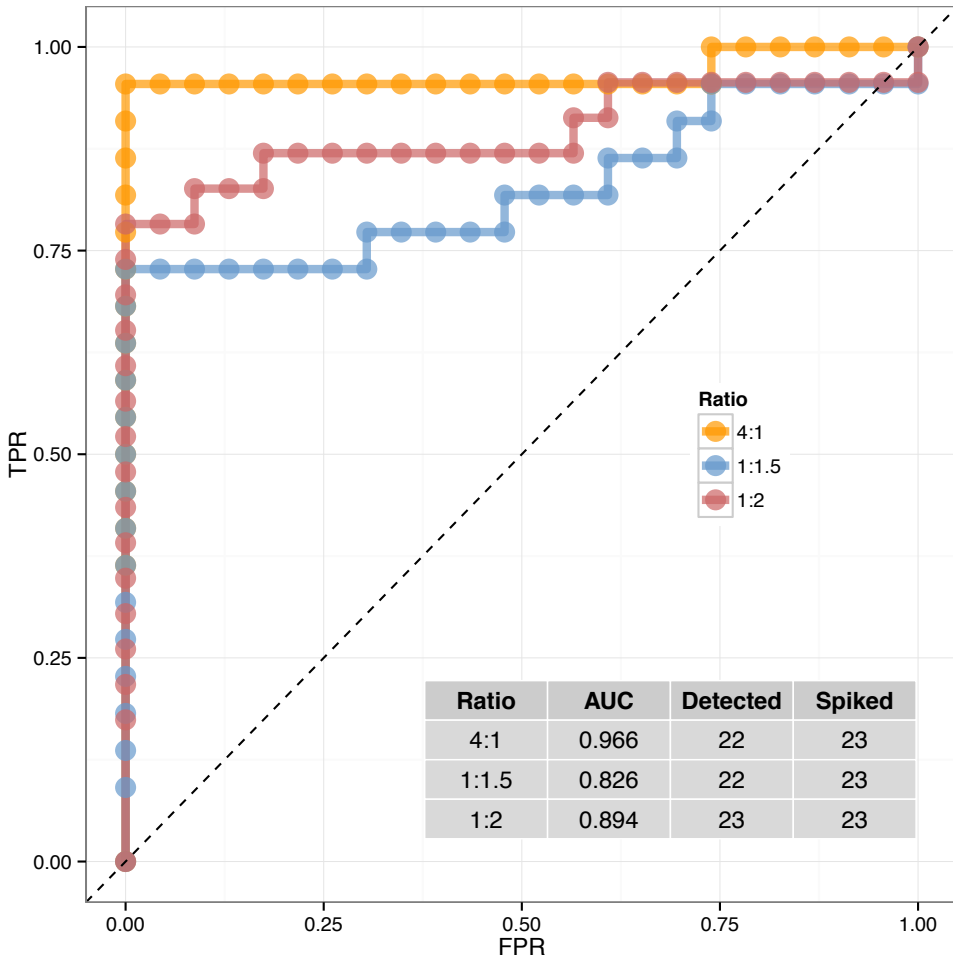
```
> exDat$Figures$dynRangePlot
```



Compared with the rat toxicogenomics experiment the UHRR/HBRR experiment from Lab 5 captures the full dynamic range of 2^{20} in the ERCC mixture design. This difference can be attributed to an much greater sequencing depth in the UHRR/HBRR experiments at each laboratory (full fluidic capacity of an instrument) compared to the MET-CTL rat toxicogenomics experiment. This difference is apparent in the total reads vectors for the two different experiments:

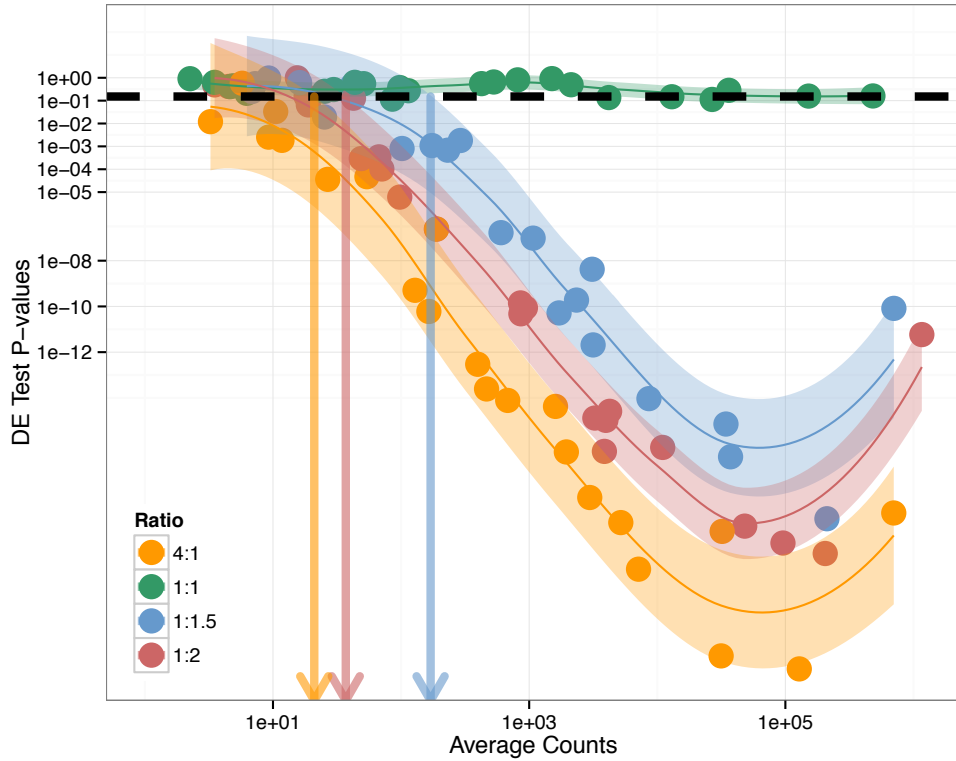
```
> COH.RatTox.ILM.MET.CTL.totalReads
[1] 41423502 46016148 44320280 38400362 47511484 33910098
> Lab5.ILM.UHRR.HBRR.totalReads
[1] 138786892 256006510 199468322 431933806 247985592
[6] 219383270 251265814 257508210
```

```
> exDat$Figures$rocPlot
```



Given the sequencing depth of the Lab 5 UHRR/HBRR experiment it is unsurprising that the ROC curve results show that almost all of the spiked ERCC transcripts were detected and tested for differential expression. At first glance, one may attempt to compare these ROC curve and AUC results to the rat toxicogenomics results, but this is ill-advised because the samples under comparison are different. Not only is the sequencing depth different for the two experiments, but the UHRR/HBRR sample transcripts also have high levels of differential expression, whereas the rat MET/CTL samples have much lower levels of differential expression. This difference in differential expression is shown in the spread and density of the points representing the endogenous transcripts in the MA plots for each experiment.

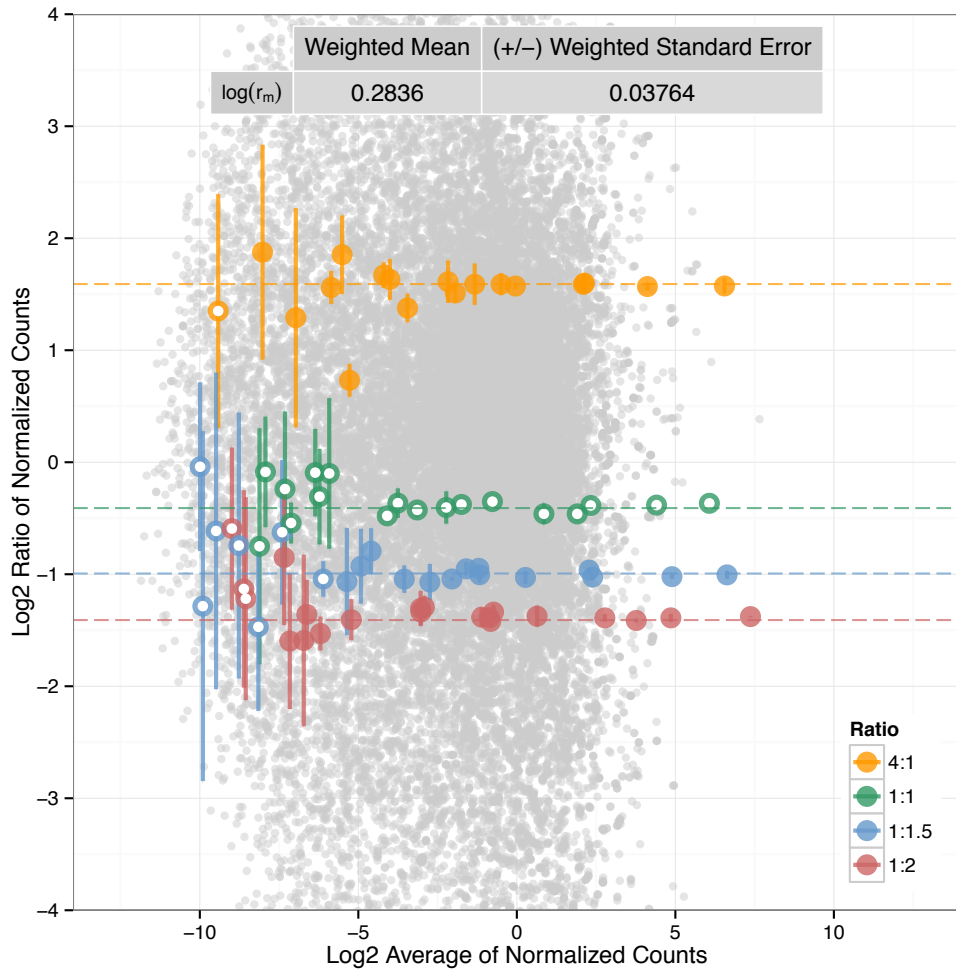
```
> exDat$Figures$lodrERCCPlot
```



Ratio	LODR Estimate	90% CI Lower Bound	90% CI Upper Bound
4:1	21	5.9	27
1:1.5	170	72	210
1:2	37	22	40

Much lower LODR estimates are found in the Lab 5 UHRR/HBRR experiment compared to the rat experiment. One can also see that the p-values in this analysis are much smaller than those in the rat experiment due to the large amount of differentially expressed transcripts in the UHRR/HBRR comparison.

```
> exDat$Figures$maPlot
```



Variability in the ratio measurements in this experiment is much lower than in the rat experiment. This is due to the nature of the replication in the two experiments. In the rat experiment the 3 replicates for each condition represent biological replicates, while in the UHRR/HBRR experiment the 4 replicates represent library preparation replicates of the same source samples and each library replicate count column was a sum of multiple fluidic replicates of the same library replicate. The bias in control ratios in this experiment is much greater than what was observed in the rat experiment. This bias is likely due to the known mRNA fraction difference between the UHRR and HBRR RNA samples.

2.2 Microarray UHRR vs. HBRR experiment analysis

Unnormalized fluorescent signals are expected for erccdashboard analysis of microarray data, the data will be log2 transformed for DE testing with the limma package. The repNormFactor variable should be NULL for microarray analysis, and repNormFactor is set to NULL if it is missing in the list of arguments for the runDashboard function. Each array replicate is normalized by the 75th quantile fluorescence intensity value of that array.

```
> exDat <- runDashboard(datType="array", exTable=Lab4.array.UHRR.HBRR,
  filenameRoot = "Lab4.array",
  sample1Name = "UHRR", sample2Name="HBRR",
  erccmix = "RatioPair", erccdilution = 1,
  spikeVol = 50, totalRNAmass = 2.5*10^(3), choseFDR=0.01)
Initializing the exDat list structure...
choseFDR = 0.01
repNormFactor is NULL
Filename root is: Lab4.array.UHRR.HBRR

Using 75th percentile (upper quartile) intensity to normalize each array

Library sizes:
694.35 758.9 705 570.55 640.25 641.3
Check for sample mRNA fraction differences(r_m)...

datType is array, using 1:1 ERCC controls for r_m estimate

log.offset
6.542976 6.63187 6.558198 6.346601 6.461859 6.463497

Number of ERCC Controls Used in r_m estimate
15

Outlier ERCCs for GLM r_m Estimate:
ERCC-00035 ERCC-00042

GLM log(r_m) estimate:
0.2070304

GLM log(r_m) estimate weighted s.e.:
0.09446045

Number of ERCCs in Mix 1 dyn range: 59

Number of ERCCs in Mix 2 dyn range: 59

datType is array, no linear model fit for ERCC specific effects

rangeResidPlot is empty

Saving dynRangePlot to exDat

Finished DE testing
```

Generating ROC curve and AUC statistics...

Area Under the Curve (AUC) Results:

	Ratio	AUC	Detected	Spiked
1	4:1	0.933	14	23
2	1:1.5	0.929	14	23
3	1:2	0.917	16	23

Estimating ERCC LODR

	Ratio	LODR Estimate	90% CI Lower Bound	90% CI Upper Bound
1	4:1	<150	<150	<150
3	1:1.5	200	<160	210
4	1:2	150	<130	160

ERCC LODR estimates are available

	Fold	Ratio	Count	Log2Count_normalized
1	4.000	4:1	150	-2.155731
2	1.000	1:1	NA	NA
3	0.667	1:1.5	200	-1.740694
4	0.500	1:2	150	-2.155731

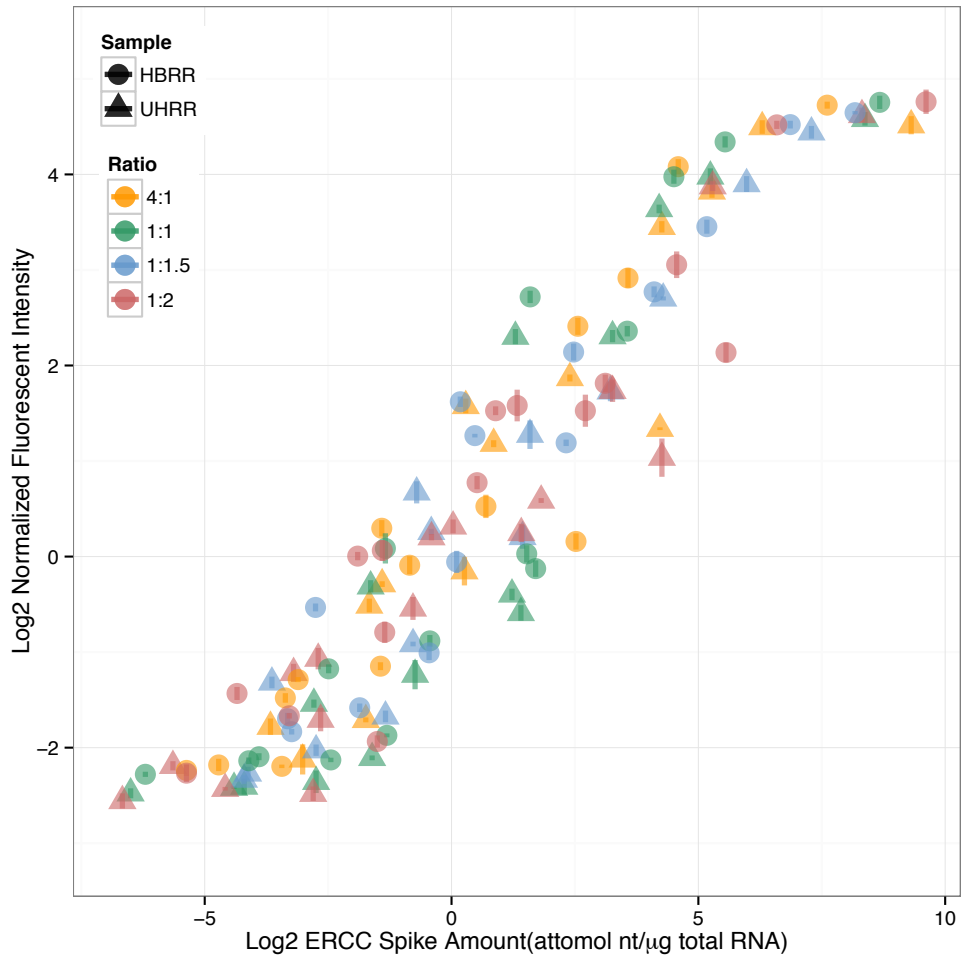
LODR estimates are available to code ratio-abundance plot

Saving main dashboard plots to pdf file...

Saving exDat list to .RData file...

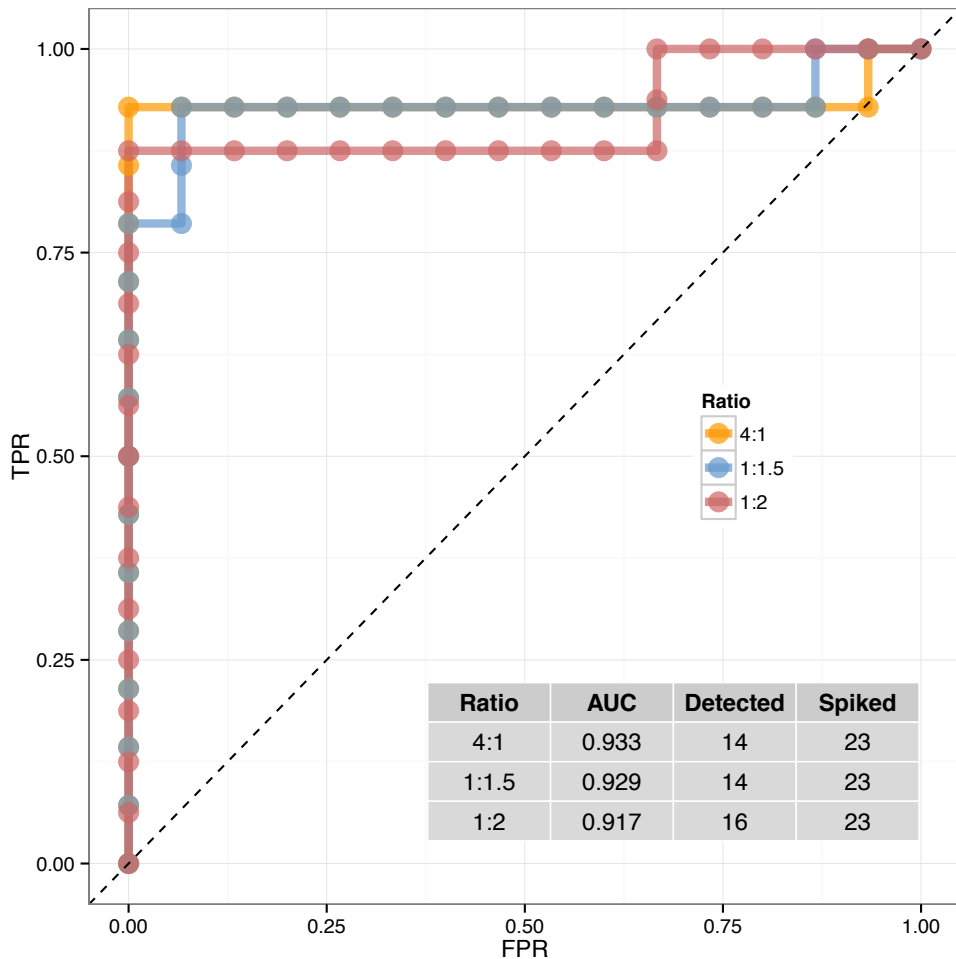
Analysis completed.

```
> exDat$Figures$dynRangePlot
```



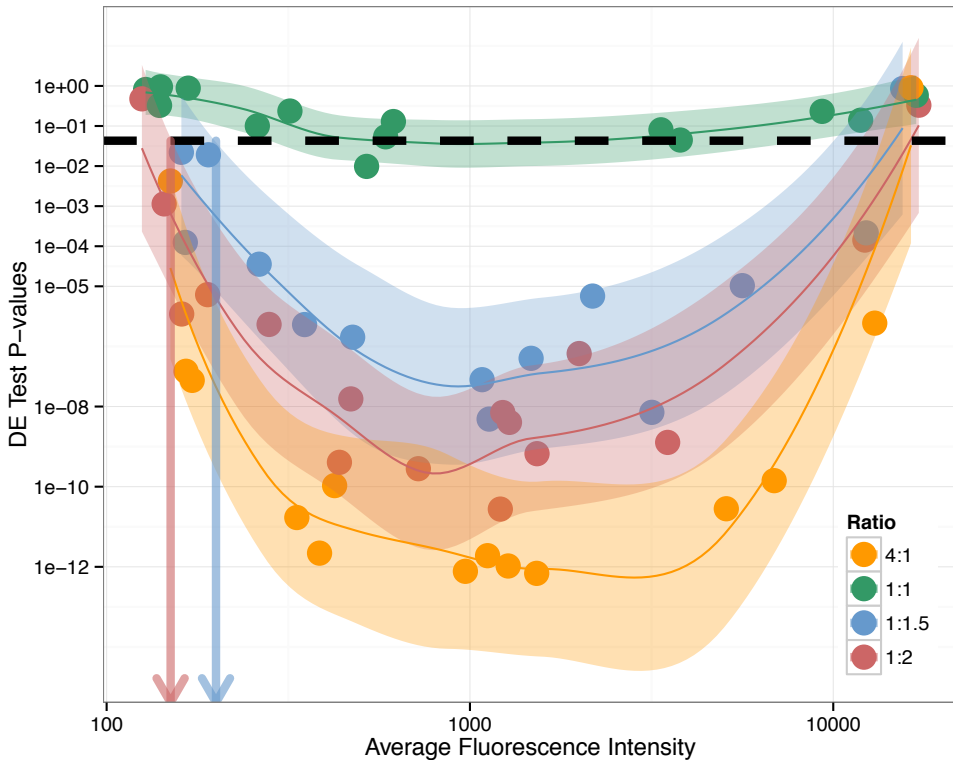
The dynamic range of the microarray experiment on reference RNA samples was 2^{10} . This figure shows the signal compression for high abundance transcripts.

```
> exDat$Figures$rocPlot
```



Comparison of these microarray ROC curves and AUC statistics to the results in the RNA-Seq UHRR/HBRR experiment from Lab 5 show similar diagnostic power for the two platforms. Fewer controls were detected above background in the microarray experiment, and this should be considered when comparing the diagnostic performance of the two platforms. More stringent filtering of RNA-Seq data would likely lead to improved diagnostic performance.

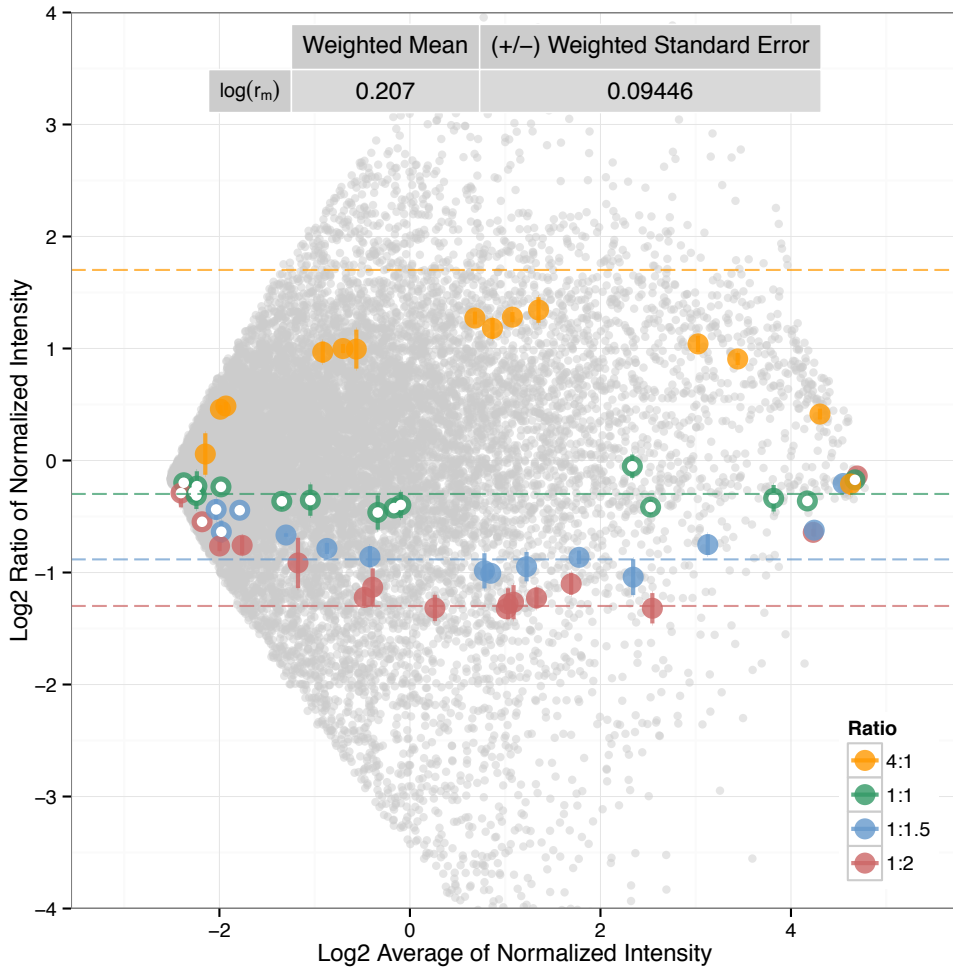
```
> exDat$Figures$lodrERCCPlot
```



Ratio	LODR Estimate	90% CI Lower Bound	90% CI Upper Bound
4:1	<150	<150	<150
1:1.5	200	<160	210
1:2	150	<130	160

LODR estimates for microarray data are derived from the intersection with the threshold p-value at the lowest signal. The same FDR was chosen for the RNA-Seq and microarray measurements of the UHRR/HBRR samples, it is interesting to note that different threshold p-values were derived for each type of experiment using the same FDR = 0.01, p.thresh = 0.1518 for RNA-Seq at Lab 5 and p.thresh = 0.0430 for the microarray experiment.

```
> exDat$Figures$maPlot
```



The variability and bias in the control ratio measurements in the microarray data measurement of UHRR/HBRR are similar to the observations in the RNA-Seq UHRR/HBRR experiment.

3 Comparison of Performance Between Experiments

The performance metrics provided here derived from measurements of ERCC ratios in gene expression experiments (AUC, LODR, r_m , and the standard deviations of the ERCC ratio measurements) can be used to assess performance between experiments within the same laboratory, or between different laboratories or technology platforms.

4 Analysis Details: Advanced Use of Functions in runDashboard

The analysis functions contained in the convenience wrapper function `runDashboard` can also be used directly by the user. Comments are provided above each analysis step included in `runDashboard` to describe the purpose and ordering constraints. View the `runDashboard` script to see comments describing the analysis functions and the ordering constraints:

```

> runDashboard
function(datType=NULL, exTable=NULL, repNormFactor=NULL,
         filenameRoot = NULL,
         sample1Name = NULL, sample2Name = NULL,
         erccmix = "RatioPair", erccdilution = 1,
         spikeVol = 1, totalRNAmass = 1, choseFDR = 0.05,
         ratioLim=c(-4,4), signalLim=c(-14,14),
         userMixFile=NULL){

  # Initialize exDat structure
  # Required for all subsequent functions
  exDat <- initDat(datType=datType, exTable=exTable,
                     repNormFactor=repNormFactor, filenameRoot=filenameRoot,
                     sample1Name=sample1Name, sample2Name=sample2Name,
                     erccmix=erccmix, erccdilution=erccdilution,
                     spikeVol=spikeVol, totalRNAmass=totalRNAmass,
                     choseFDR=choseFDR, ratioLim = ratioLim,
                     signalLim = signalLim, userMixFile=userMixFile)

  # Estimate the difference in mRNA fraction of total RNA for the two samples
  # Required for all subsequent functions
  exDat <- est_r_m(exDat)

  # Evaluate the dynamic range of the experiment (Signal-Abundance plot)
  # Not required for subsequent functions
  exDat <- dynRangePlot(exDat)

  # Evaluate pairwise ERCC control ratios within Mix1 and Mix2
  # Not required for subsequent functions
  if(erccmix == "RatioPair"){
    exDat <- withinMixRatios(exDat)
  }

  # Test for differential expression between samples
  # Required for all subsequent functions
  exDat <- geneExprTest(exDat)

  # Generate ROC curves and AUC statistics
  # Not Required for subsequent functions
  exDat <- erccROC(exDat)

  # Estimate LODR for ERCC controls
  # Required for subsequent functions
  exDat = estLODR(exDat, kind = "ERCC", prob=0.9)

  ## Estimate LODR using Simulated data from endogenous transcripts
  ## Not required for subsequent functions
  #exDat = estLODR(exDat, kind = "Sim", prob=0.9)

  # Generate MA plot (Ratio vs. Average Signal) with ERCC controls below LODR

```

```

#   annotated also flags possible False Negatives on DE gene list based on LODR
#   threshold from DE gene list
# Not required for subsequent functions
exDat <- annotLODR(exDat)

### Saving plots and results
# Convenience function to save 4 main figures to PDF
saveERCCPlots(exDat)

# Save exDat to a RData file for later use
cat("\nSaving exDat list to .RData file...")
nam <- paste(exDat$sampleInfo$filenameRoot, "exDat", sep = ".")
assign(nam, exDat)

to.save <- ls()

save(list = to.save[grepl(pattern = nam, x=to.save)],
     file=paste0(exDat$sampleInfo$filenameRoot, ".RData"))

# End analysis and return exDat to global environ. / workspace
cat("\nAnalysis completed.")
return(exDat)

}

<environment: namespace:erccdashboard>

```

4.1 Flexibility in Differential Expression Testing

The `geneExprTest` function wraps the QuasiSeq differential expression testing package for `datType = "count"` or uses the limma package for differential expression testing if `datType = "array"`. The function uses the DE testing p-value results and choseFDR parameter to select a threshold p-value for LODR estimation. In the case of count data if a correctly formatted csv file is provided with the necessary DE test results, then `geneExprTest` will bypass DE testing (with reduced runtime). The function will look for a csv file with the name `"filenameRoot.quasiSeq.res.csv"` and columns with names `"Feature"`, `"pvals"`, and `"qvals"` must be in the file.

4.2 Options for LODR Estimation

The default behavior of `runDashboard` is to use the `estLODR` function to obtain an LODR estimate using empirical data from the ERCCs and a model-based simulation using the endogenous genes in the sample. The type of LODR estimation is selected using the argument `kind` in the `estLODR` function. The other parameter that may be adjusted is the probability for the LODR estimate, in the default analysis `prob = 0.9` is selected.

4.3 Options for Printing Plots to File

The function `savePlots` will save selected figures to a pdf file. The default is the 4 manuscript figures to a single page (`plotsPerPg = "manuscript"`). If `plotsPerPg = "single"` then each plot is placed on an individual page in one pdf file. If `plotlist` is not defined (`plotlist = NULL`) then all plots in `exDat$Figures` are printed to the file.

To print 4 plots from manuscript to a single page pdf file use plotsPerPg = "manuscript" in the `saveERCCPlots` function or to create a multiplepage pdf of all plots produced use plotsPerPg = "single".

4.4 Analysis of Alternative Spike-in Designs

By default the package is configured to analyze the ERCC ratio mixtures produced by Ambion (ERCC ExFold RNA Spike-In Mixes, Catalog Number 4456739). This pair of control ratio mixtures were designed to have 1:1, 4:1, 1:1.5, and 1:2 ratios of 92 distinct RNA transcripts (23 different RNA control sequences are in each of these four ratio subpools). Alternative ERCC RNA control ratio mixture designs can be produced using the NIST DNA Plasmid Library for External Spike-in Controls (NIST Standard Reference Material 2374, <https://www-s.nist.gov/srmors/certificates/2374.pdf>). For example, a pair of RNA control mixtures could be created with a ternary ratio design, three subpools of RNA controls with either no change (1:1) or 2-fold increased (2:1) and 2-fold decreased (1:2) relative abundances between the pair of mixtures (Mix 1/Mix 2). To use alternative spike-in mixture designs with the dashboard a csv file must be provided to the package with the argument userMixFile for the `initDat` function.

If all samples from both conditions were only spiked with a single ERCC mixture (e.g. Ambion Catalog Number 4456740, ERCC RNA Spike-In Mix) a limited subset of the package functions can be used (`initDat`, `est_r_m`, and `dynRangePlot`. For `initDat` use `ERCCMixes="Single"` and `est_r_m` and `dynRangePlot` functions can then be used to examine the mRNA fraction differences for the pair of samples and evaluate the dynamic range of the experiment.

5 Notes on R version and session information

The results shown in this R vignette are the same as the results shown in our manuscript and were obtained in R version 3.0.2 with the following session information.

```
> sessionInfo()
R version 3.0.2 (2013-09-25)
Platform: x86_64-apple-darwin10.8.0 (64-bit)

locale:
[1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8

attached base packages:
[1] splines      grid        stats       graphics    grDevices
[6] utils        datasets    methods     base

other attached packages:
[1] ROCR_1.0-5          gplots_2.12.1
[3] edgeR_3.4.2         limma_3.18.12
[5] QuasiSeq_1.0-3      MASS_7.3-29
[7] reshape2_1.2.2       erccdashboard_0.9.6
[9] gridExtra_0.9.1      ggplot2_0.9.3.1

loaded via a namespace (and not attached):
[1] bitops_1.0-6          caTools_1.16
[3] colorspace_1.2-4       dichromat_2.0-0
[5] digest_0.6.4           gdata_2.13.2
[7] gtable_0.1.2           gtools_3.3.0
[9] KernSmooth_2.23-10     labeling_0.2
[11] lattice_0.20-24       locfit_1.5-9.1
```

```
[13] Matrix_1.1-2      mgcv_1.7-28
[15] munsell_0.4.2    nlme_3.1-113
[17] plyr_1.8          proto_0.3-10
[19] qvalue_1.36.0    RColorBrewer_1.0-5
[21] scales_0.2.3     stringr_0.6.2
[23] tcltk_3.0.2      tools_3.0.2
```