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Basic DESeq2 results exploration

Project: PDF report.

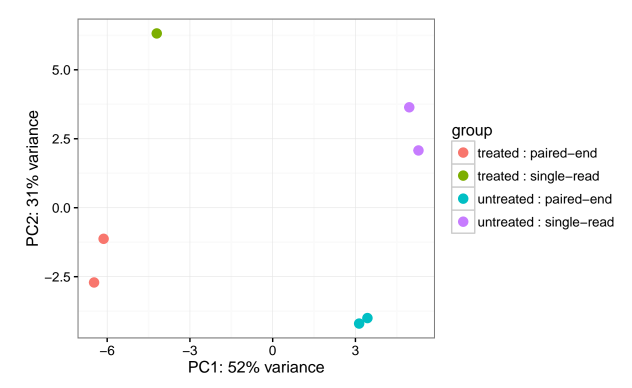
Introduction

This report is meant to help explore DESeq2 (Love, Huber, and Anders, 2014) results and was generated using the regionReport (Collado-Torres, Jaffe, and Leek, 2015) package. While the report is rich, it is meant to just start the exploration of the results and exemplify some of the code used to do so. If you need a more in-depth analysis for your specific data set you might want to use the customCode argument. This report is based on the vignette of the DESeq2 (Love, Huber, and Anders, 2014) package which you can find here.

Code setup

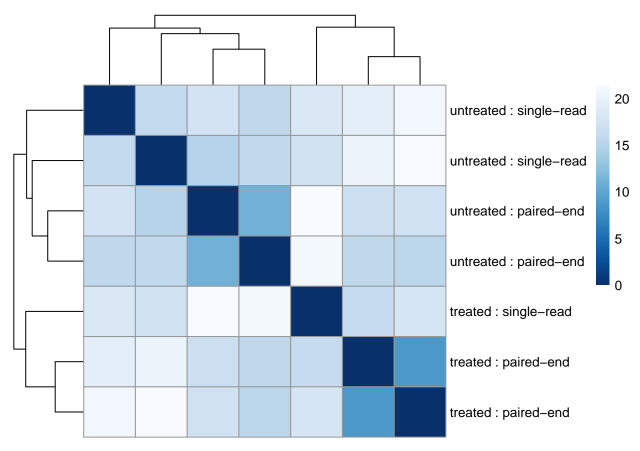
This section contains the code for setting up the rest of the report.

PCA



The above plot shows the first two principal components that explain the variability in the data using the regularized log count data. If you are unfamiliar with principal component analysis, you might want to check the Wikipedia entry or this interactive explanation. In this case, the first and second principal component explain 52 and 31 percent of the variance respectively.

Sample-to-sample distances

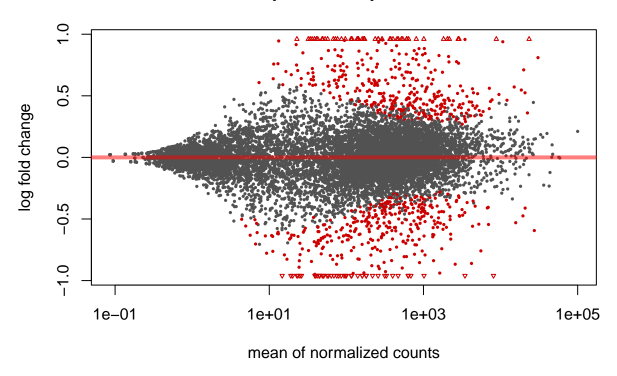


This plot shows how samples are clustered based on their euclidean distance using the regularized log transformed count data. This figure gives an overview of how the samples are hierarchically clustered. It is a complementary figure to the PCA plot.

MA plots

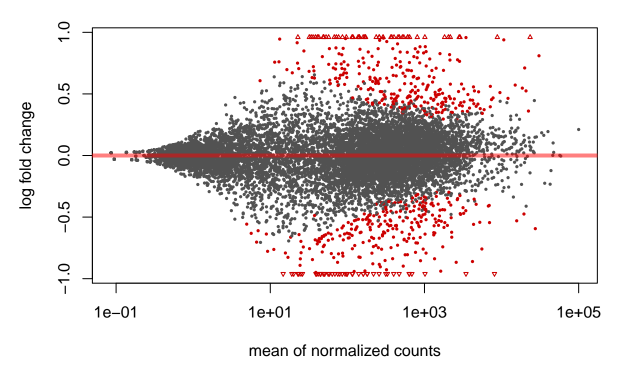
This section contains three MA plots (see Wikipedia) that compare the mean of the normalized counts against the log fold change. They show one point per feature. The points are shown in red if the feature has an adjusted p-value less than alpha, that is, the statistically significant features are shown in red.

MA plot with alpha = 0.1



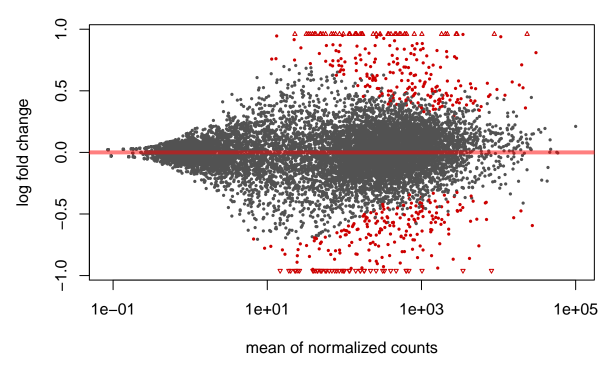
This first plot shows uses alpha = 0.1, which is the alpha value used to determine which resulting features were significant when running the function DESeq2::results().

MA plot with alpha = 0.05



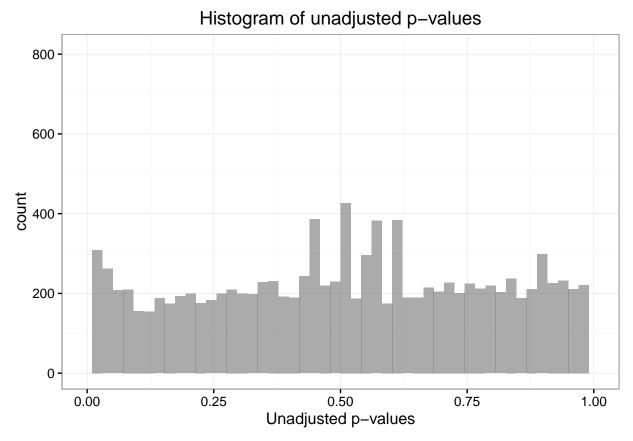
This second MA plot uses alpha = 0.05 and can be used agains the first MA plot to identify which features have adjusted p-values between 0.05 and 0.1.

MA plot for top 500 features



The third and final MA plot uses an alpha such that the top 500 features are shown in the plot. These are the features that whose details are included in the top features interactive table.

P-values distribution



This plot shows a histogram of the unadjusted p-values. It might be skewed right or left, or flat as shown in the Wikipedia examples. The shape depends on the percent of features that are differentially expressed. For further information on how to interpret a histogram of p-values check David Robinson's post on this topic.

```
## Min. 1st Qu. Median Mean 3rd Qu. Max. NA's ## 0.0000 0.2258 0.4972 0.4795 0.7287 0.9999 2649
```

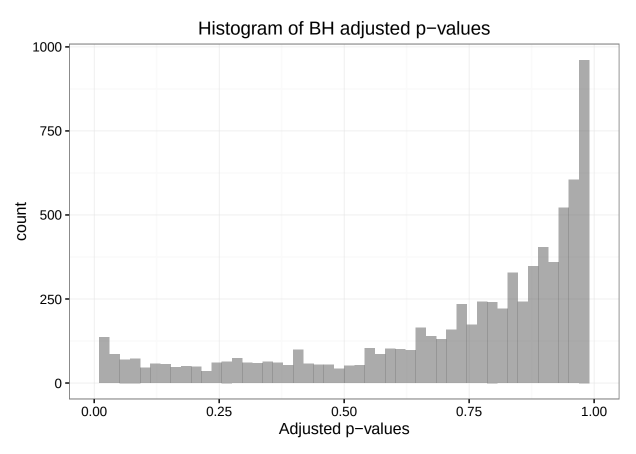
This is the numerical summary of the distribution of the p-values.

Cut	Count
0.0001	319
0.0010	468
0.0100	807
0.0250	1052
0.0500	1375
0.1000	1861
0.2000	2724
0.3000	3673
0.4000	4696
0.5000	5949
0.6000	7396
0.7000	8560
0.8000	9626
0.9000	10667

Cut	Count
1.0000	11821

This table shows the number of features with p-values less or equal than some commonly used cutoff values.

Adjusted p-values distribution



This plot shows a histogram of the BH adjusted p-values. It might be skewed right or left, or flat as shown in the Wikipedia examples.

This is the numerical summary of the distribution of the BH adjusted p-values.

Cut	Count
0.0001	194
0.0010	267
0.0100	411
0.0250	516
0.0500	631
0.1000	797
0.2000	1043

Cut	Count
0.3000	1326
0.4000	1615
0.5000	1916
0.6000	2283
0.7000	2927
0.8000	3920
0.9000	5357
1.0000	8424

This table shows the number of features with BH adjusted p-values less or equal than some commonly used cutoff values.

Top features

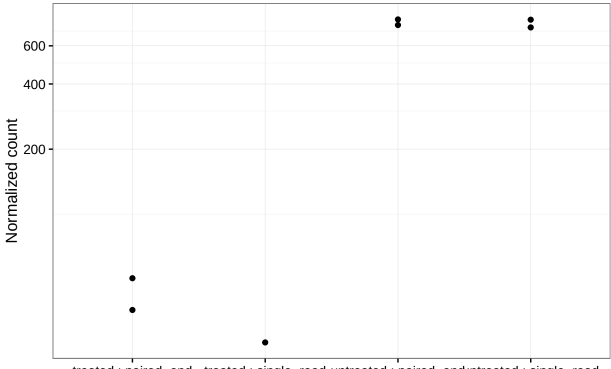
This table shows the top 500 features ordered by their BH adjusted p-values. Since the report is in PDF format, only the top 20 features are shown.

Feature	baseMean	log2FoldChange	lfcSE	stat	pvalue	padj
FBgn0039155	453.28	3.71	0.16	23.21	4.013291e-119	3.380797e-115
FBgn0029167	2165.04	2.08	0.10	20.10	6.684454e-90	2.815492e-86
FBgn0035085	366.83	2.23	0.14	16.26	1.888618e-59	5.303239e-56
FBgn0029896	257.90	2.21	0.16	13.91	5.854593e-44	1.232977e-40
FBgn0034736	118.41	2.57	0.18	13.88	8.067448e-44	1.359204 e-40
FBgn0040091	610.60	1.43	0.12	11.91	1.114552e-32	1.564831e-29
FBgn0000071	180.01	-2.14	0.18	-11.81	3.498482e-32	4.210174e-29
FBgn0011260	140.36	-1.96	0.17	-11.61	3.723734e-31	3.921092e-28
FBgn0034434	76.91	2.39	0.21	11.26	2.045205 e-29	1.914312e-26
FBgn0001226	686.72	-1.53	0.14	-11.10	1.310312e-28	1.103807e-25
FBgn0038832	167.59	2.04	0.19	10.92	9.049372e-28	6.930174 e-25
FBgn0026562	23460.08	1.90	0.18	10.58	3.761064e-26	2.640267e-23
FBgn0051092	93.33	-1.95	0.19	-10.41	2.332072e-25	1.511183e-22
FBgn0035189	115.49	-2.12	0.20	-10.37	3.376356e-25	2.031602e-22
FBgn0031191+FBgn0027279	2043.81	0.96	0.09	10.30	6.734072 e- 25	3.587332e-22
FBgn0033913	2816.22	1.10	0.11	10.30	6.813545 e-25	3.587332e-22
FBgn0023479	1823.54	1.36	0.13	10.28	8.376665 e-25	4.150884e-22
FBgn0033764	41.75	-2.23	0.22	-10.23	1.480711e-24	6.929728e-22
FBgn0003501	161.22	-1.76	0.18	-10.05	9.406230 e-24	4.170425 e-21
FBgn0039419	806.81	1.11	0.11	9.78	1.336255 e-22	5.628306e-20

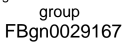
Count plots top features

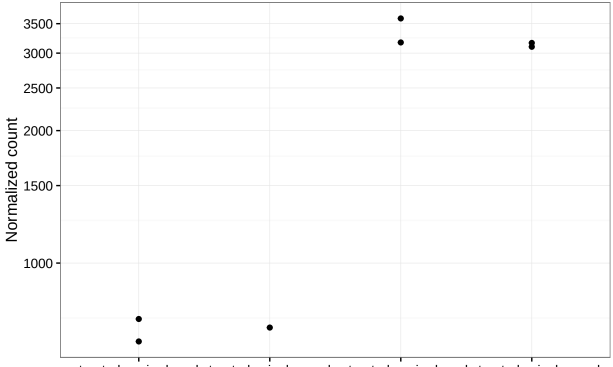
This section contains plots showing the normalized counts per sample for each group of interest. Only the best 20 features are shown, ranked by their BH adjusted p-values. The Y axis is on the log10 scale and the feature name is shown in the title of each plot.



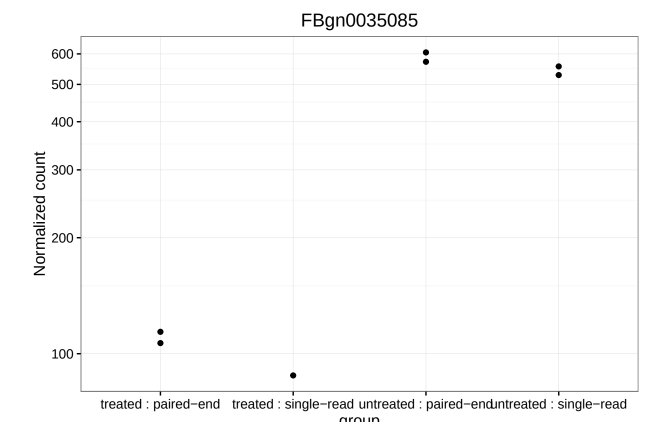


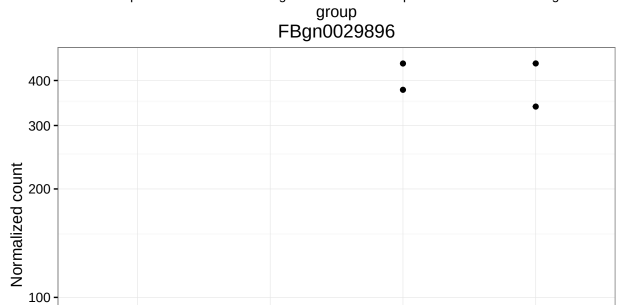
treated : paired-end treated : single-read untreated : paired-enduntreated : single-read



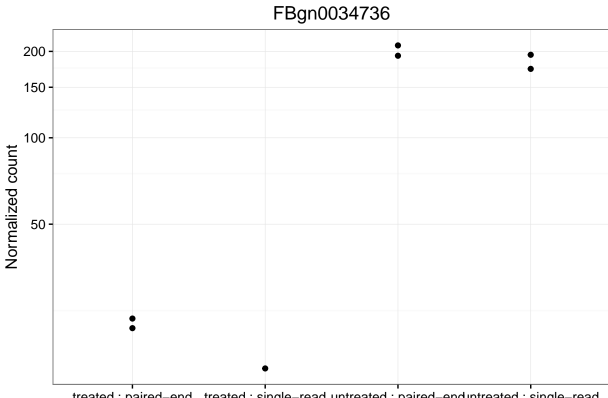


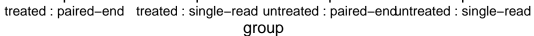
treated : paired-end treated : single-read untreated : paired-endintreated : single-read group

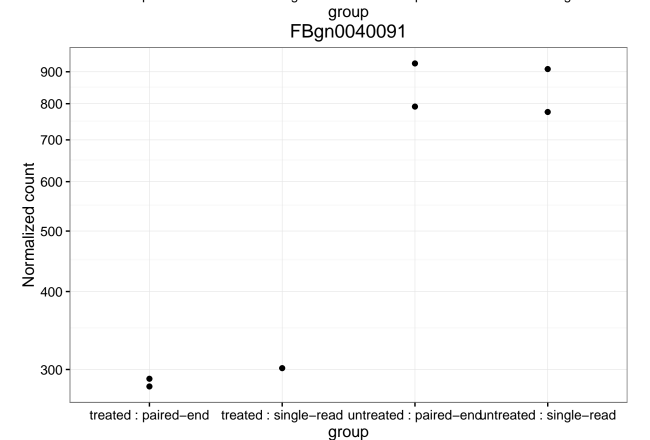


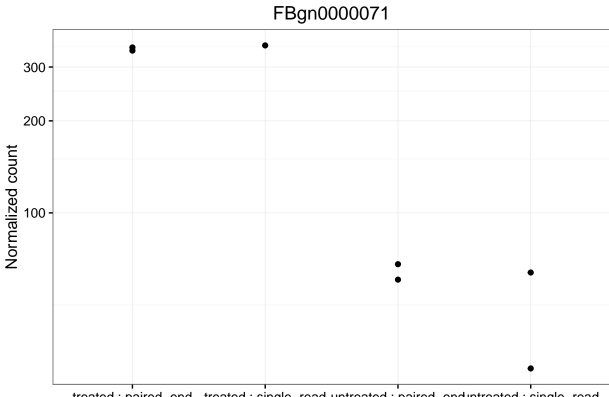


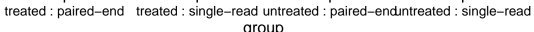
treated : paired-end treated : single-read untreated : paired-enduntreated : single-read group

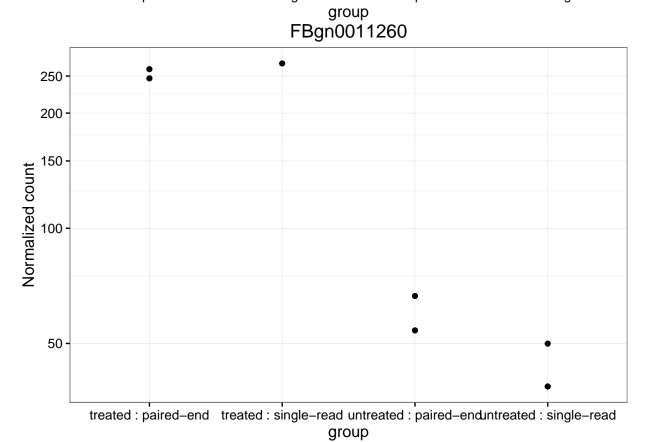


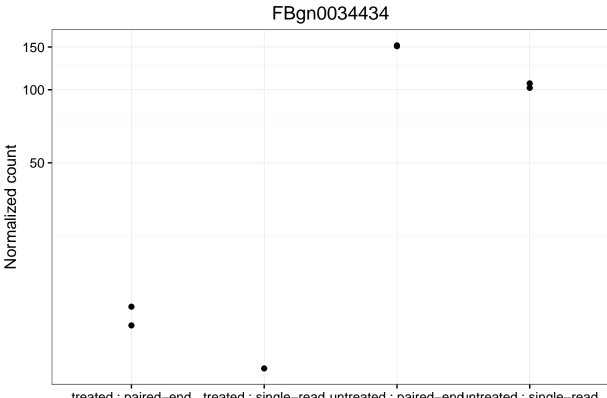


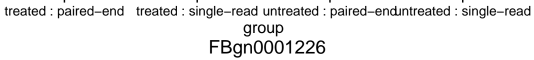


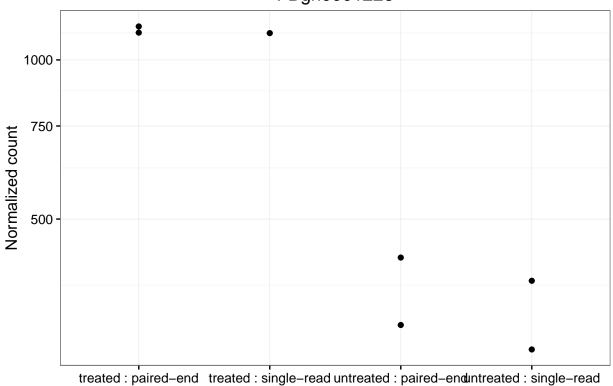






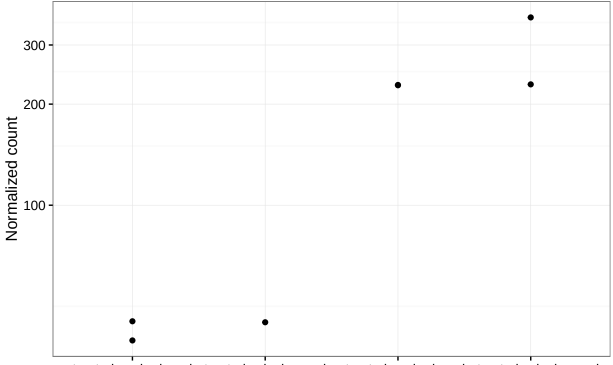






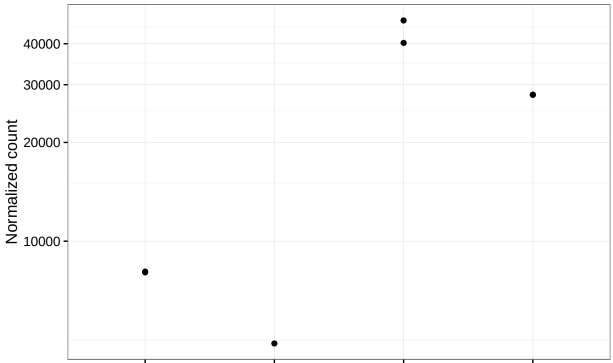
treated: paired-end treated: single-read untreated: paired-enduntreated: single-read group



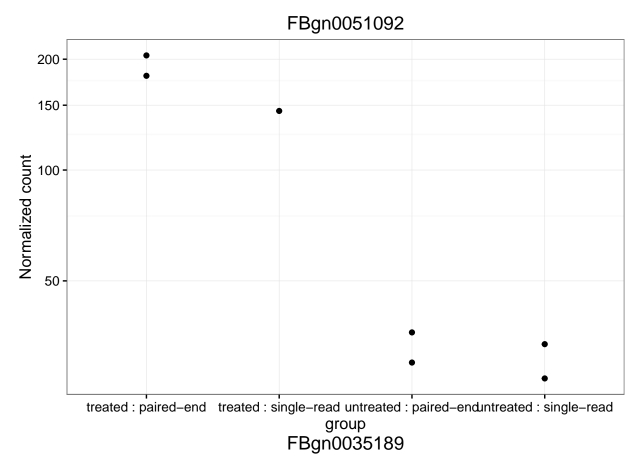


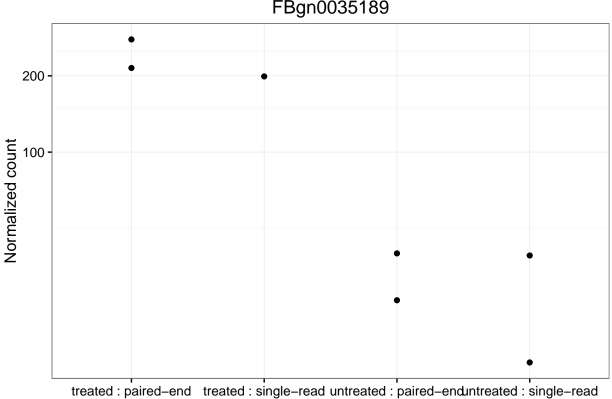
treated : paired-end treated : single-read untreated : paired-enduntreated : single-read





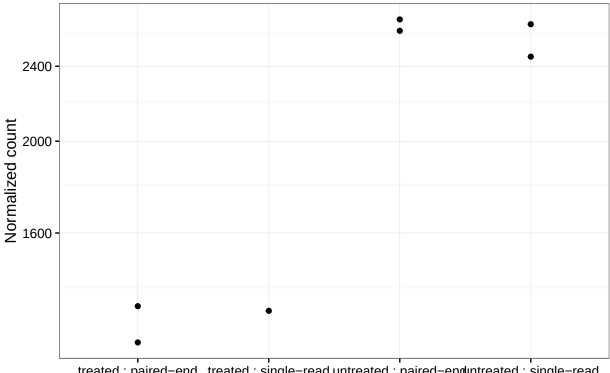
treated : paired-end treated : single-readuntreated : paired-end ntreated : single-read group



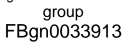


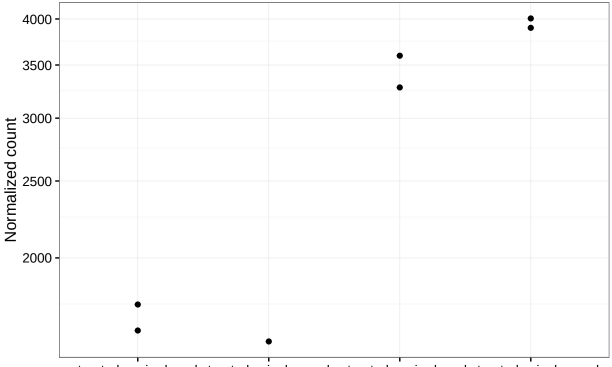
group

FBgn0031191+FBgn0027279

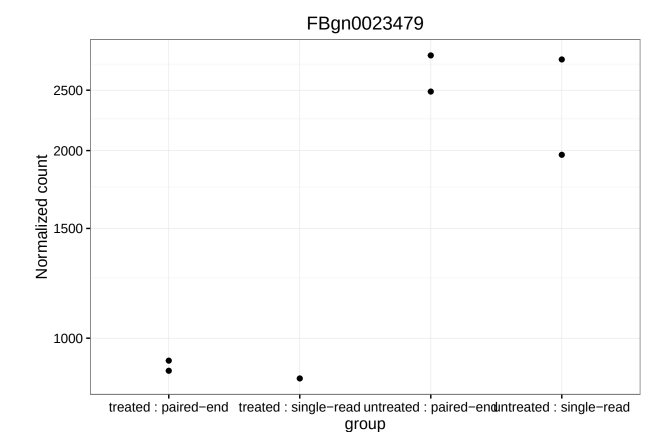


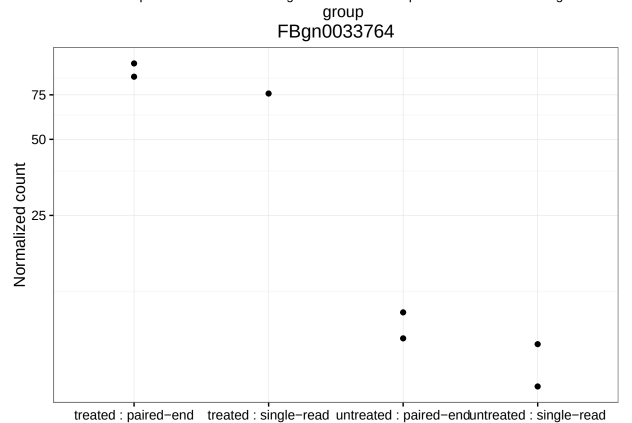
treated: paired-end treated: single-read untreated: paired-endintreated: single-read



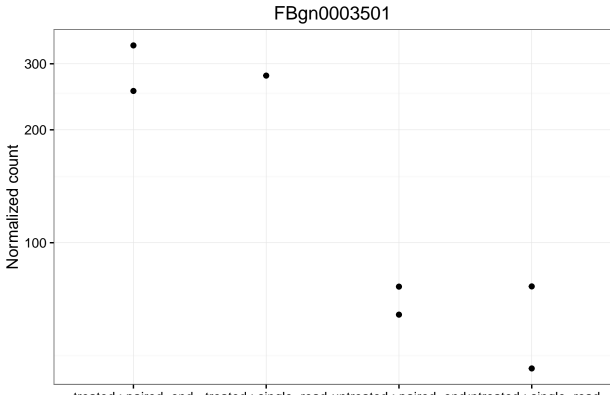


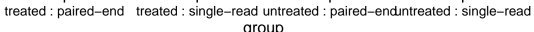
treated: paired-end treated: single-read untreated: paired-enduntreated: single-read group

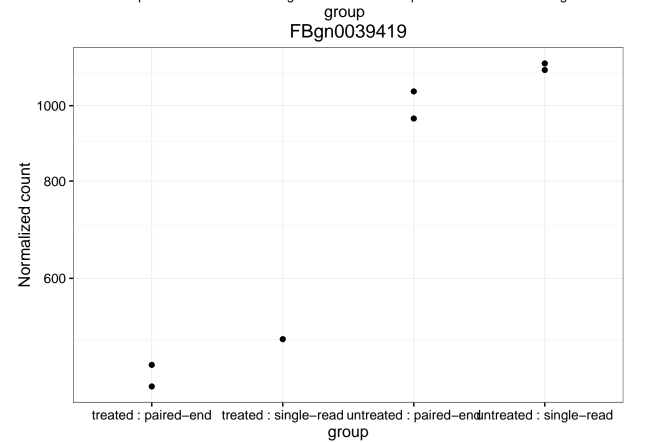




group







Reproducibility

biomaRt

bitops

BSgenome

Biostrings

bumphunter

checkmate

codetools

cluster

The input for this report was generated with DESeq2 (Love, Huber, and Anders, 2014) using version 1.11.36 and the resulting features were called significantly differentially expressed if their BH adjusted p-values were less than alpha = 0.1. This report was generated in path /Users/lcollado/Dropbox/JHSPH/Code/regionReportSupp using the following call to DESeq2Report():

```
## DESeq2Report(dds = dds, project = "PDF report", intgroup = c("condition",
      "type"), outdir = "DESeq2-example", output = "DESeq2Report",
##
     device = "pdf", theme = theme_bw(), output_format = "pdf_document")
Date the report was generated.
## [1] "2016-04-07 18:28:21 EDT"
Wallclock time spent generating the report.
## Time difference of 32.6 secs
R session information.
setting value
##
  version R version 3.3.0 alpha (2016-03-23 r70368)
## system x86_64, darwin13.4.0
          X11
## ui
## language (EN)
## collate en_US.UTF-8
## tz
          America/New_York
          2016-04-07
   date
package
                    * version date
                                       source
   acepack
                      1.3-3.3 2014-11-24 CRAN (R 3.3.0)
##
## annotate
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                              2016-02-06 Bioconductor
## AnnotationDbi
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                              2016-01-29 Bioconductor
## backports
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                              2016-03-18 CRAN (R 3.3.0)
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                              2014-12-31 CRAN (R 3.3.0)
## Biobase
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                              2016-01-14 Bioconductor
## BiocGenerics
                    * 0.17.3
                              2016-01-29 Bioconductor
## BiocParallel
                              2016-03-23 Bioconductor
                      1.5.21
```

2016-01-14 Bioconductor

2013-08-17 CRAN (R 3.3.0) 2016-02-21 Bioconductor

2016-03-29 Bioconductor

2016-03-10 CRAN (R 3.3.0)

2015-07-21 CRAN (R 3.3.0)

2015-07-15 CRAN (R 3.3.0)

2.39.12 2016-02-21 Bioconductor

2.27.2

1.0-6

1.39.4

1.11.5

1.7.3

2.0.3

0.2-14

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2015-03-11 CRAN (R 3.3.0)
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##
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                           1.5.30
                                     2016-03-25 Bioconductor
##
    derfinderHelper
                           1.5.3
                                     2016-03-23 Bioconductor
##
                           1.23.1
                                     2016-01-14 Bioconductor
    DESeq
##
   DESea2
                         * 1.11.36
                                     2016-03-31 Bioconductor
                         * 1.10.0
                                     2016-01-23 CRAN (R 3.3.0)
##
   devtools
##
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                                     2016-01-08 CRAN (R 3.3.0)
##
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                                     2014-03-07 CRAN (R 3.3.0)
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                                     2016-03-16 Bioconductor
##
    evaluate
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## foreach
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                           1.4.3
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                           0.8-66
                                     2015-08-19 CRAN (R 3.3.0)
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##
                                     2015-04-07 CRAN (R 3.3.0)
    Formula
                           1.2-1
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                                     2016-02-22 Bioconductor
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    GenomicRanges
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                         * 1.23.25
                                     2016-03-31 Bioconductor
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    gridExtra
                           2.2.1
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                           0.5.1
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                                     2016-03-24 Github (jimhester/knitrBootstrap@cdaa4a9)
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                                     2015-07-14 CRAN (R 3.3.0)
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                                     2015-12-03 CRAN (R 3.3.0)
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    magrittr
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                                     2015-04-22 CRAN (R 3.3.0)
    markdown
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##
                           1.2-4
  Matrix
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##
    munsell
                           0.4.3
                                     2016-02-13 CRAN (R 3.3.0)
##
    nnet
                           7.3 - 12
                                     2016-02-02 CRAN (R 3.3.0)
##
    pasilla
                         * 0.11.0
                                     2016-03-31 Bioconductor
##
                                     2015-12-11 CRAN (R 3.3.0)
    pheatmap
                         * 1.0.8
## pkgmaker
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                                     2014-05-14 CRAN (R 3.3.0)
##
    plyr
                           1.8.3
                                     2015-06-12 CRAN (R 3.3.0)
```

```
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##
##
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                           2.1.2
                                     2016-01-26 CRAN (R 3.3.0)
##
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   Rcpp
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##
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                                     2016-03-28 Bioconductor
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##
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##
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##
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##
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   XML
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##
##
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  XVector
                           0.11.7
                                     2016-02-13 Bioconductor
##
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                                     2014-06-12 CRAN (R 3.3.0)
                                     2016-03-19 Bioconductor
    zlibbioc
                           1.17.1
```

Bibliography

This report was created with regionReport (Collado-Torres, Jaffe, and Leek, 2015) using rmarkdown (Allaire, Cheng, Xie, McPherson, et al., 2016) while knitr (Xie, 2014) and DT (Xie, 2015) were running behind the scenes. pheatmap (Kolde, 2015) was used to create the sample distances heatmap. Several plots were made with ggplot2 (Wickham, 2009).

Citations made with knitcitations (Boettiger, 2015). The BibTeX file can be found here.

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