csaw: ChIP-seq analysis with windows User's Guide

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Chapter 1

Introduction

1.1 Scope

This document gives an overview of the Bioconductor package csaw for detecting differential binding (DB) in ChIP-seq experiments. Specifically, csaw uses sliding windows to identify significant changes in binding patterns for transcription factors (TFs) or histone marks across different biological conditions. However, it can also be applied to any sequencing technique where reads represent coverage of enriched genomic regions. The statistical methods described here are based upon those in the edgeR package [Robinson et al., 2010]. Knowledge of edgeR is useful but not a necessary prerequesite in this guide.

1.2 How to get help

Most questions about csaw will (hopefully) be answered by the documentation. Every function mentioned in this guide has its own help page. For example, a detailed description of the arguments and output of the windowCounts function can be obtained by typing ?windowCounts or help(windowCounts) at the R prompt. The relevant references are also supplied in each help page.

The authors of the package always appreciate receiving reports of bugs in the package functions or in the documentation. The same goes for well-considered suggestions for improvements. Other questions about how to use csaw are best sent to the Bioconductor mailing list bioconductor@stat.math.ethz.ch. To subscribe to the mailing list, see https://stat.ethz.ch/mailman/listinfo/bioconductor. Please send requests for general assistance and advice to the mailing list rather than to the individual authors.

Users posting to the mailing list for the first time may find it helpful to read the helpful posting guide at http://www.bioconductor.org/doc/postingGuide.html.

1.3 Quick start

A typical ChIP-seq analysis would look something like the code described below. This assumes that a vector of BAM file paths is provided in bam.files and a design matrix in design.

```
> require(csaw)
> data <- windowCounts(bam.files, ext=110)
> binned <- windowCounts(bam.files, bin=TRUE, width=10000)
> normfacs <- normalizeChIP(binned$counts, binned$totals)
> require(edgeR)
> y <- DGEList(data$counts, lib.size=data$totals, norm.factors=normfacs)
> y <- estimateDisp(y, design)
> results <- glmQLFTest(y, design, robust=TRUE)
> merged <- mergeWindows(data$region, tol=1000L)
> tabcom <- combineTests(merged$id, results$table)</pre>
```

For anyone still reading, the csaw analysis pipeline can be summarized into several steps:

- 1. Loading in data from BAM files.
- 2. Calculating normalization factors.
- 3. Filtering out uninteresting tests.
- 4. Identifying DB windows.
- 5. Correcting for multiple hypothesis tests.

Each step will be demonstrated in this guide with some publicly available data. This particular dataset looks for changes in NFYA binding between embryonic stem cells and terminal neurons [Tiwari et al., 2012].

```
> bam.files <- c("es_1.bam", "es_2.bam", "tn_1.bam", "tn_2.bam")
> design <- model.matrix(~factor(c('es', 'es', 'tn', 'tn')))
> colnames(design) <- c("intercept", "cell.type")</pre>
```

A comprehensive listing of the datasets used in this guide is provided in Section 8.1, along with instructions on how to obtain and process them for entry into the csaw pipeline.

Chapter 2

Converting reads to counts

Hello, reader. A little box like this will be present at the start of each chapter. It's intended to tell you which objects from previous chapters are needed to get the code in the current chapter to work. At this point, all we need are the bam.files we defined in the introduction above.

2.1 Counting reads into windows

The windowCounts function counts fragments throughout the genome for a set of libraries using a sliding window approach. For single-end data, the fragment corresponding to a read is estimated by directionally extending each read to the average fragment length. For paired-end data, each fragment is defined as the interval spanned by the two reads in a proper pair. In both cases, the number of fragments overlapping a genomic window is counted. This is repeated after sliding the window along the genome to a new position. A count is then obtained for each window in each library. Sorted and indexed BAM (i.e. binary SAM) files are required as input into windowCounts.

```
> frag.len <- 110
> data <- windowCounts(bam.files, ext=frag.len, width=10)
> head(data$counts)

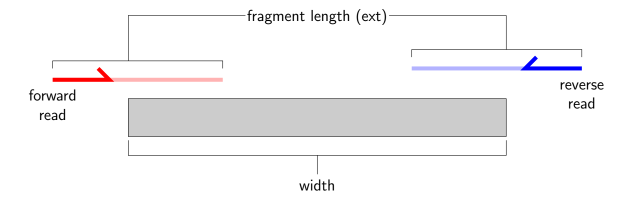
[,1] [,2] [,3] [,4]
[1,] 6 7 4 4
[2,] 1 2 9 13
[3,] 0 2 9 13
```

^{[4,] 13 5 4 1} [5,] 16 5 5 1 [6,] 10 13 21 5

> head(data\$region)

```
GRanges with 6 ranges and 0 metadata columns:
      seqnames
                            ranges strand
         <Rle>
                         <IRanges>
                                     <Rle>
  [1]
          chr1 [3003751, 3003760]
  [2]
          chr1 [3012951, 3012960]
  [3]
          chr1 [3013001, 3013010]
  [4]
          chr1 [3043151, 3043160]
  [5]
          chr1 [3043201, 3043210]
          chr1 [3065501, 3065510]
  [6]
  seqlengths:
                                         chr10 ... chrY_JH584303_random
                    chr1
              195471971
                                     130694993 ...
                                                                   158099
```

For single-end data, suitable values for the average fragment length in ext can be estimated from the primary peak in a cross-correlation plot (see Section 2.3). Alternatively, the length can be estimated from wet lab diagnostics such as post-fragmentation gel electrophoresis images. Typical values range from 100 to 300 bp, depending on the efficiency of sonication and the use of size selection steps in library preparation.



The specified width of each window controls the compromise between spatial resolution and count size. Larger windows will lead to greater counts which (when considered in isolation) will provide more power for DB detection. However, spatial resolution is also lost for large windows. Reads from a DB site may be counting alongside reads from a non-DB site (e.g., non-specific background) or even those from a DB site in the opposite direction. This leads to loss of detection power.

The window size can be interpreted as a measure of the width of the binding site. Thus, TF analyses will usually use a window size of several base pairs. This also maximizes spatial resolution which is critical for punctate enrichment. For histone marks, widths of at least 150 bp are recommended [Humburg et al., 2011]. This corresponds to the length of DNA wrapped up in each nucleosome, i.e., the smallest relevant unit for histone mark enrichment.

For diffuse marks, the sizes of enriched regions are more variable and so the compromise between resolution and power is more arbitrary. Analyses with multiple width values can be combined to provide a comprehensive picture of differential binding at all resolutions.

2.1.1 Filtering out low-quality reads

Reads which have been marked as PCR duplicates in the SAM flag can be ignored by setting dedup=TRUE. This can reduce the variability between replicates caused by inconsistent duplication. However, it also caps the number of reads at each position. This can lead to loss of power in high abundance regions and false positives when the same upper bound is applied to libraries of varying size. Thus, duplicate removal is generally not recommended for routine DB analyses.

Reads can also be filtered based on the minimum mapping score with the minq argument. Low mapping scores are indicative of incorrectly and/or non-uniquely aligned sequences. Removal of these reads may provide more reliable results. The exact value of the threshold depends on the range of scores provided by the aligner. The subread aligner [Liao et al., 2013] was used for this dataset so a value of 100 is appropriate.

```
> demo <- windowCounts(bam.files, ext=frag.len, minq=100, dedup=TRUE)
```

On a more subjective note, reads on particular chromosomes can be specifically counted by specifying the chromosomes of interest in restrict. This avoids the need to load up reads on unassigned contigs or uninteresting chromosomes (e.g., mitochondrial genome). Similarly, it allows windowCounts to work with limited memory by analyzing only one chromosome at a time.

Finally, reads lying in certain regions can also be removed by specifying those regions in discard. This is intended to remove reads that are wholly aligned within known repeat regions. Repeats are problematic as changes in repeat copy number between conditions can lead to spurious DB. Removal of reads can avoid detection of biologically irrelevant differences.

```
> repeats <- GRanges("chr1", IRanges(3000001, 3002128))
> demo <- windowCounts(bam.files, ext=frag.len, discard=repeats,
+ restrict=c("chr1", "chr10", "chrX"))</pre>
```

Other potential uses include the removal of reads overlapping known binding sites when attempting to discover new sites. Needless to say, any DB sites within the discarded regions will not be detected. Some caution is therefore required when specifying the regions of disinterest.

2.1.2 Increasing speed and memory efficiency

The spacing parameter controls the distance with which windows are shifted to the next position in the genome. Using a higher value will reduce the computational load as fewer count combinations are extracted for analysis. This may be useful when memory is limited on the machine. Of course, this sacrifices spatial resolution as adjacent positions are not counted and thus cannot be distinguished.

```
> demo <- windowCounts(bam.files, spacing=100, ext=frag.len)</pre>
> head(demo$region)
GRanges with 6 ranges and 0 metadata columns:
      seanames
                            ranges strand
         <Rle>
                         <IRanges>
                                     <Rle>
  [1]
          chr1 [3013001, 3013001]
          chr1 [3043201, 3043201]
  [2]
  [3]
          chr1 [3065501, 3065501]
          chr1 [3089401, 3089401]
  [4]
  [5]
          chr1 [3241701, 3241701]
          chr1 [3254601, 3254601]
  [6]
  seqlengths:
                                         chr10 ... chrY_JH584303_random
                    chr1
               195471971
                                     130694993 ...
                                                                   158099
```

For analyses with large windows, it is also worth increasing the spacing to a fraction of the specified width. This reduces the computational work by decreasing the number of windows and extracted counts. Any loss in spatial resolution due to a larger spacing interval is negligible when compared to that already lost by using a large window width.

Windows with a low sum of counts across all libraries can be filtered out using the filter argument. This improves memory efficiency by discarding the majority of low-abundance windows corresponding to uninteresting background regions. The default filter value is set as the number of libraries multiplied by 5. This roughly corresponds to the minimum average count required for accurate downstream modelling. Note that more sophisticated filtering is recommended and can be applied later (see Chapter 4).

```
> demo <- windowCounts(bam.files, ext=frag.len, filter=30)
> head(demo$counts)
     [,1] [,2] [,3] [,4]
[1,]
       10
                  21
             13
[2,]
       10
             12
                  18
                         4
[3,]
        9
              8
                  10
                         3
[4,]
       22
             18
                  21
                        12
[5,]
       42
             38
                  40
                        29
[6,]
             23
                  20
                        17
```

A special case occurs when bin=TRUE. This will set spacing=width and only use the 5' end of each read for counting. Reads are then counted into contiguous bins across the genome. However, no filtering will be performed. Users should set width to a reasonably large value, otherwise reads will be counted and reported for every single base in the genome.

```
> demo <- windowCounts(bam.files, width=1000, bin=TRUE)
> head(demo$region)
GRanges with 6 ranges and 0 metadata columns:
      seqnames
                           ranges strand
         <Rle>
                       <IRanges>
                                   <Rle>
  [1]
          chr1 [3000001, 3001000]
          chr1 [3001001, 3002000]
  [2]
  [3]
          chr1 [3002001, 3003000]
  [4]
          chr1 [3003001, 3004000]
  [5]
          chr1 [3004001, 3005000]
  [6]
          chr1 [3005001, 3006000]
  seglengths:
                                        chr10 ... chrY_JH584303_random
                   chr1
              195471971
                                    130694993 ...
```

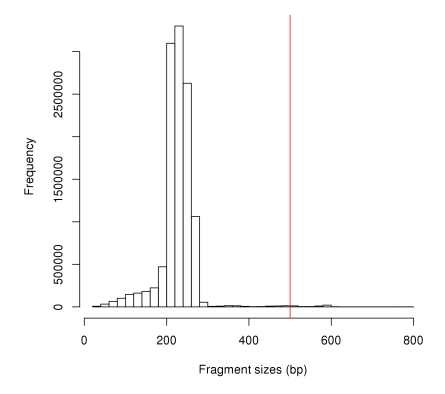
2.2 Experiments involving paired-end tags

ChIP experiments with paired-end sequencing can be accommodated by setting pet=TRUE in the windowCounts function call. Read extension is not required for ChIP-PET data as the genomic interval spanned by the originating fragment is explicitly defined by the positions of the paired reads. By default, only proper pairs are counted whereby the two reads are on the same chromosome, face inward and are no more than max.frag apart.

```
> petBamFile <- "example-pet.bam"
> demo <- windowCounts(petBamFile, max.frag=500, pet="both")</pre>
```

A suitable value for max.frag can be chosen by examining the distribution of fragment sizes using the getPETSizes function. In this example, a user might use a value of around 500 bp as it covers most of the fragment size distribution. The plot can also be used to examine the quality of the PET sequencing procedure. The location of the peak should be consistent with the fragmentation and size selection steps in library preparation.

```
> out <- getPETSizes(petBamFile)
> frag.sizes <- out$sizes[out$sizes<=800]
> hist(frag.sizes, breaks=50, xlab="Fragment sizes (bp)", ylab="Frequency", main="")
> abline(v=500, col="red")
```



The number of fragments exceeding the maximum size can be recorded for quality control. The getPETSizes function also returns the number of invalid pairs, interchromosomal pairs, pairs with one unmapped reads and single reads. A non-negligible proportion of these reads indicates that there may be some problems with the paired-end alignment or sequencing.

> c(out\$diagnostics, too.large=sum(out\$sizes > 500))

total	single	unoriented	mate.unmapped	inter.chr
30854223	0	254594	888175	2928351
too.large				
176116				

In cases where there are a non-negligible proportion of invalid pairs, the reads can be rescued by setting rescue.pairs=TRUE. For each invalid read pair, the read with the highest mapping quality score will be directionally extended to ext to form a fragment. Counting will then be performed with these fragments in addition to those from the valid pairs. A value of ext can be chosen by examining the distribution above.

> demo <- windowCounts(petBamFile, max.frag=500, pet="both", ext=200, rescue.pairs=TRUE)

Paired-end data can also be treated as single end data by specifiying pet="first" or pet="second". This will only take the first or second read of each pair as defined in the SAM flags. Unlike rescue.pairs, the use of one read is done for all read pairs regardless of validity. This may be useful for comparing paired-end with single-end data, or for truly disastrous situations where paired-end sequencing has failed.

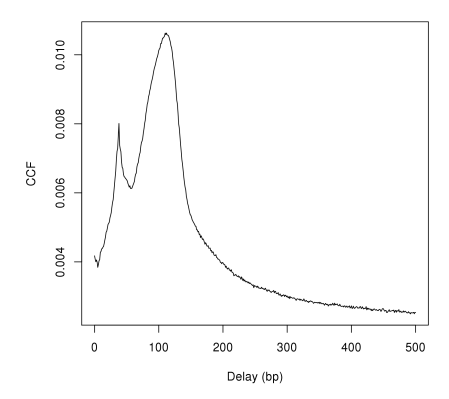
```
> demo <- windowCounts(petBamFile, max.frag=500, pet="first")</pre>
```

Note that the paired-end methods in csaw depend on the synchronisation of mate information for each alignment in the BAM file. Any file manipulations that might break this synchronisation should be corrected prior to the use of csaw.

2.3 Estimating the average fragment length

Cross-correlation plots can be generated directly from BAM files using the correlateReads function. This provides a measure of the immunoprecipitation (IP) efficiency of an experiment [Kharchenko et al., 2008]. Efficient IP should yield a smooth peak at a distance corresponding to the average fragment length. This reflects the strand-dependent bimodality of reads around TF binding sites. The location of the peak thus provides an estimate for the average fragment length (~ 110 bp below) for use in read extension.

```
> max_distance <- 500
> x <- correlateReads(bam.files, max_distance, dedup=TRUE, cross=TRUE, minq=100)
> plot(0:max_distance, x, type="1", ylab="CCF", xlab="Delay (bp)")
```

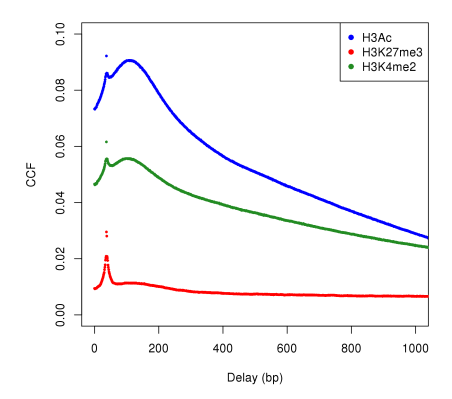


A sharp spike may also observed in the plot at a distance corresponding to the read length. This is an artefact which is thought to be a result of the preference of aligners for uniquely mapped reads. The size of the smooth peak can be compared to the height of the spike to assess the signal-to-noise ratio of the data [Landt et al., 2012]. Poor IP efficiency will result in a smaller or absent peak as bimodality is less pronounced. Note that duplicate removal is required here to reduce the size of the read length spike. Otherwise, the fragment length peak will not be visible as a separate entity.

Cross-correlation plots can also be used for fragment length estimation of punctate histone marks such as histone acetylation and H3K4 methylation. However, they are less effective for diffuse enrichment where bimodality is not obvious (e.g. H3K27 trimethylation).

```
> n <- 10000
> h3ac <- correlateReads("h3ac.bam", n, dedup=TRUE, cross=TRUE)
> h3k27me3 <- correlateReads("h3k27me3.bam", n, dedup=TRUE, cross=TRUE)
> h3k4me2 <- correlateReads("h3k4me2.bam", n, dedup=TRUE, cross=TRUE)
> plot(0:n, h3ac, col="blue", ylim=c(0, 0.1), xlim=c(0, 1000),
+ xlab="Delay (bp)", ylab="CCF", pch=16, cex=0.5)
> points(0:n, h3k27me3, col="red", pch=16, cex=0.5)
> points(0:n, h3k4me2, col="forestgreen", pch=16, cex=0.5)
```

```
> legend("topright", col=c("blue", "red", "forestgreen"),
+ c("H3Ac", "H3K27me3", "H3K4me2"), pch=16)
```



2.4 Ensuring synchronisation

In practice, most ChIP-seq analysis pipelines based on csaw will involve multiple window-Counts calls. Users should supply the same values for the bam.files, dedup, pet, minq, ext and (for paired-end data) max.frag parameters at each call. This ensures that the same reads are being used for counting throughout the analysis. For complex analyses, this synchronisation can be easily maintained by constructing a parameter list and calling the countWindows wrapper function.

```
> param <- list(bam.files=bam.files, dedup=TRUE, minq=100, pet="none", ext=frag.len)
> demo <- countWindows(param, filter=50)
> demo <- countWindows(param, bin=TRUE, width=1000)</pre>
```

This strategy avoids the need for repeated manual specification of non-default arguments at each function call. Global changes can then be implemented by altering the contents of

param directly. That said, for the sake of simplicity, the examples in the rest of this guide will use the windowCounts function. This is equivalent to read counting with the default values for each of the critical parameters previously described.

Chapter 3

Calculating normalization factors

This next chapter will need the bam.files vector again. If you bother to keep reading, you'll notice that a number of other files are called in this chapter. However, these are just for demonstration purposes and aren't necessary for the main example.

3.1 Eliminating composition biases

As the name suggests, composition biases are formed when there are differences in the composition of sequences across libraries. Highly enriched regions consume more sequencing resources and thereby suppress the representation of other regions. Differences in the magnitude of suppression can lead to spurious DB calls. Scaling by library size fails to correct for this as composition biases can still occur in libraries of the same size.

To remove composition biases in csaw, reads are counted in large bins and the counts are used for normalization with the normalizeChIP wrapper function. This uses the trimmed mean of M-values (TMM) method [Robinson and Oshlack, 2010] to correct for any systematic fold change in the coverage of the bins. The assumption here is that most bins represent non-DB background regions so any consistent difference across bins must be spurious.

```
> binned <- windowCounts(bam.files, bin=TRUE, width=10000)
> normfacs <- normalizeChIP(binned$counts, lib.size=binned$totals)
> normfacs
```

[1] 0.9843100 0.9594416 1.0334848 1.0245790

The TMM method produces normalization factors for use in edgeR. The size of each library is scaled by the corresponding factor to obtain an effective library size for modelling. A larger normalization factor results in a larger effective library size. This is conceptually

equivalent to scaling each individual count downwards, as the ratio of that count to the library size will be smaller. Check out the user's guide in edgeR for more detail.

Binning is necessary to increase the size of counts for TMM normalization. This avoids zero counts that lead to undefined M-values. Adding a prior count is only a superficial solution as the chosen prior will have undue influence on the estimate of the normalization factor when many counts are low. The variance of the fold change distribution is also higher for low counts. This reduces the effectiveness of the trimming procedure during normalization.

Of course, this strategy requires the user to supply a bin size. Excessively large bins are problematic as background and enriched regions will be included in the same bin. This makes it difficult to trim away enriched bins during the TMM procedure. Obviously, bins which are too small will have read counts which are too low. A value around 10000 bp is usually suitable. Testing multiple bin sizes is recommended to ensure that the estimates are robust to any changes.

```
> demo <- windowCounts(bam.files, bin=TRUE, width=5000)
> normalizeChIP(demo$counts, lib.size=demo$total)

[1] 0.9840325 0.9611444 1.0318386 1.0246845

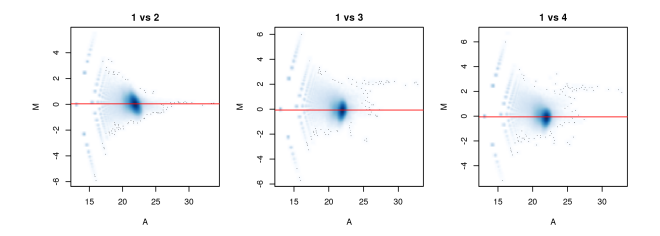
> demo <- windowCounts(bam.files, bin=TRUE, width=15000)
> normalizeChIP(demo$counts, lib.size=demo$total)

[1] 0.9848029 0.9578794 1.0344662 1.0247632
```

Note that normalizeChIP skips the precision weighting step in the TMM method. Weighting increases the contribution of bins with high counts. However, these bins are more likely to contain binding sites and thus are more likely to be DB. If any DB regions should survive trimming (e.g. those with subtle fold changes), upweighting them would then be counterproductive. Users may wish to explicitly filter out such bins and run TMM only on putative background regions, as shown below.

```
> ab <- aveLogCPM(binned$counts, lib.size=binned$total)
> keep <- ab <= quantile(ab, p=0.75)
> normalizeChIP(binned$counts[keep,], lib.size=binned$total)
[1] 0.9904911 0.9267174 1.0572098 1.0304832
```

The effectiveness of normalization can be checked using a MA plot. A single main cloud of points should be present that represents the background regions. Separation into multiple discrete points indicates that the counts are too low and that larger bin sizes should be used. Composition biases manifest as a vertical shift in the position of this cloud. Ideally, the log-ratios of the corresponding normalization factors should correspond to the centre of this mass. This indicates that undersampling has been corrected.



3.2 Eliminating efficiency biases

Efficiency biases are commonly observed in ChIP-seq data. This refers to fold changes in enrichment that are introduced by variability in IP efficiencies between libraries. These technical differences are of no biological interest and must be removed. This can be done by assuming some top percentage of bins or windows with highest abundances contain binding sites. The TMM method can then be applied to eliminate systematic differences across those bins.

```
> me.demo <- windowCounts(c("h3k4me3_mat.bam", "h3k4me3_pro.bam"), bin=TRUE, width=10000L)
> ab <- aveLogCPM(me.demo$counts, lib.size=me.demo$total)
> keep <- rank(ab)>0.99*length(ab)
> me.norm <- normalizeChIP(me.demo$count[keep,], lib.sizes=me.demo$totals)
> me.norm

[1] 0.7108908 1.4066859

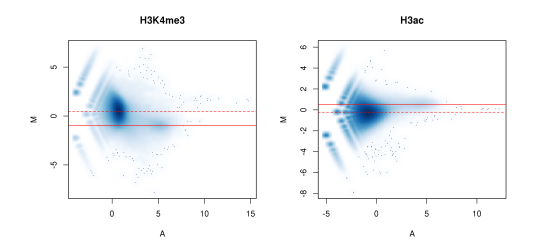
> ac.demo <- windowCounts(c("h3ac.bam", "h3ac_2.bam"), bin=TRUE, width=2000L)
> ab <- aveLogCPM(ac.demo$counts, lib.size=ac.demo$total)</pre>
```

```
> keep <- rank(ab)>0.99*length(ab)
> ac.norm <- normalizeChIP(ac.demo$count[keep,], lib.sizes=ac.demo$totals)
> ac.norm
[1] 1.1937090 0.8377251
```

This method assumes that most high-abundance bins are not DB. Specifically, any changes must be due to differences in IP efficiency (or some other uninteresting technical effect). Genuine biological differences may be removed when the assumption of a non-DB majority does not hold, i.e., overall enrichment is truly lower for one condition. Also, the percentage of bins to use may not be obvious if there is no obvious demarcation on a MA plot.

Speaking of which, the results of normalization can be visualized with MA plots. Of particular interest is the cloud at high A-values. This represents a systematic fold change in bound regions, either due to genuine DB or variable IP efficiency. Note the difference in the normalization factors from removal of efficiency bias (full) against that of composition bias (dashed).

```
> par(mfrow=c(1,2))
> adjc <- cpm(me.demo$counts, log=TRUE)
> smoothScatter(x=rowMeans(adjc), y=adjc[,1]-adjc[,2], xlab="A", ylab="M", main="H3K4me3")
> abline(h=log2(me.norm[1]/me.norm[2]), col="red")
> me.base <- normalizeChIP(me.demo$counts)
> abline(h=log2(me.base[1]/me.base[2]), col="red", lty=2)
> adjc <- cpm(ac.demo$counts, log=TRUE)
> smoothScatter(x=rowMeans(adjc), y=adjc[,1]-adjc[,2], xlab="A", ylab="M", main="H3ac")
> abline(h=log2(ac.norm[1]/ac.norm[2]), col="red")
> ac.base <- normalizeChIP(ac.demo$counts)
> abline(h=log2(ac.base[1]/ac.base[2]), col="red", lty=2)
```



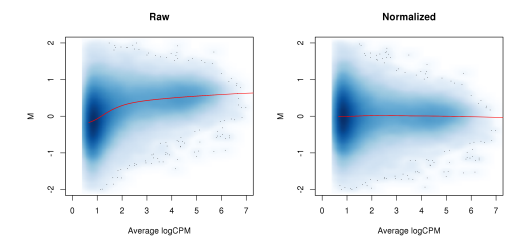
3.3 Dealing with trended biases

In more extreme cases, the bias may vary with the average abundance to form a trend. This may reflect the fact that changes in IP efficiency have little effect at low abundances (e.g., background regions) and more effect at high abundances (i.e., actual binding sites). The trend is difficult to correct with scaling methods as any one scaling factor cannot be optimal at all abundances. Rather, non-linear methods are required such as cyclic loess or quantile normalization.

One such implementation is provided in normalizeChIP. This is based on the fast loess algorithm [Ballman et al., 2004] with adaptation to count data. It can be applied after some filtering to remove low-abundance regions where the loess curve fitting is not accurate. An offset matrix is then produced that contains the log-effective library sizes for each supplied bin/window. This can then be supplied to the statistical methods in edgeR, assuming that said bins or windows are also the ones to be tested for DB.

Filtering on the average count computed by aveLogCPM is strongly recommended prior to using this method (see Chapter 4 for more detail on choices of filter value). Specifically, an average count threshold serve as a vertical cutoff in the plots below. This avoids introducing any spurious trends that might affect normalization.

```
> par(mfrow=c(1,2))
> aval <- ab[keep]
> o <- order(aval)
> adjc <- cpm(ac.demo$counts[keep,], log=TRUE, lib.size=ac.demo$total)
> mval <- adjc[,1]-adjc[,2]
> fit <- loessFit(x=aval, y=mval)
> smoothScatter(aval, mval, ylab="M", xlab="Average logCPM",
+ main="Raw", ylim=c(-2,2), xlim=c(0, 7))
> lines(aval[o], fit$fitted[o], col="red")
> re.adjc <- log2(ac.demo$counts[keep,]+0.5) - ac.off/log(2)
> mval <- re.adjc[,1]-re.adjc[,2]
> fit <- loessFit(x=aval, y=mval)
> smoothScatter(ab[keep], re.adjc[,1]-re.adjc[,2], ylab="M",
+ xlab="Average logCPM", main="Normalized", ylim=c(-2,2), xlim=c(0, 7))
> lines(aval[o], fit$fitted[o], col="red")
```



Note that all non-linear methods assume that most bins/windows are not DB throughout the covariate range. This tends to be a stronger assumption that the equivalent for scaling methods. Removal of the trend may not be appropriate if that trend represents some genuine biological phenomenon.

3.4 A word on other biases

No normalization is performed to adjust for differences in mappability or sequencability between different regions of the genome. Region-specific biases are assumed to be constant between libraries. This is generally reasonable as the biases depend on fixed properties of the genome sequence such as GC content. Thus, biases should cancel out during DB comparisons. Any variability in the bias between samples will be absorbed into the dispersion estimate.

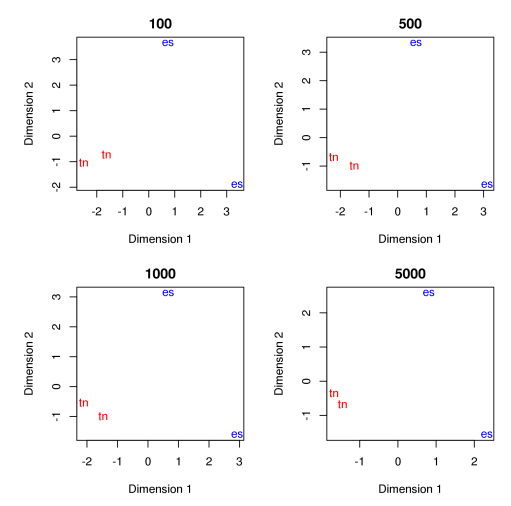
That said, explicit normalization to correct these biases can improve results for some datasets. This is due to decreases in the observed variability when systematic differences between libraries are removed. Of course, this also assumes that the targeted differences have no biological relevance. Detection power may be lost if this is not true. For example, differences in the GC content distribution can be driven by technical bias as well as biology, e.g., when protein binding is associated with a specific GC composition.

3.5 Examining replicate similarity with MDS plots

On a related note, the binned counts can be used to examine the similarity of replicates through multi-dimensional scaling (MDS) plots. The distance between each pair of libraries is computed as the square root of the mean squared log-fold change across the top set of bins with the highest absolute log-fold changes. A small top set visualizes the most extreme

differences whereas a large set visualizes overall differences. Again, binning is necessary as fold changes are being used.

```
> par(mfrow=c(2,2), mar=c(5,4,2,2))
> binned <- windowCounts(bam.files, bin=TRUE, width=2000L)
> adj.counts <- cpm(binned$counts, log=TRUE)
> for (top in c(100, 500, 1000, 5000)) {
+    out <- plotMDS(adj.counts, main=top, col=c("blue", "blue", "red", "red"),
+    labels=c("es", "es", "tn", "tn"), top=top)
+ }</pre>
```



Replicates from different groups should form separate clusters in the plot. This indicates that the reproducibility is high and that the effect sizes are large. Mixing between replicates of different conditions indicates that the biological difference has no effect on protein binding, or the data is too variable for any effect to manifest. Any outliers should also be noted as their presence may confound the downstream analysis. In the worst case, the removal of the corresponding libraries may be necessary to obtain sensible results.

Chapter 4

Filtering prior to correction

This chapter will require frag.len and data defined in Chapter 2. We will also need the normfacs vector from Chapter 3.

4.1 Independent filtering for count data

Many of the low abundance windows in the genome correspond to background regions in which DB is not expected. At the most extreme, windows with low counts will not provide enough evidence against the null to obtain low p-values. Similarly, some statistical approximations will fail at low counts. Removing such uninteresting tests reduces the severity of the multiple testing correction, increases detection power amongst the remaining tests and reduces computational work.

Filtering is valid so long as it is independent of the test statistic under the null hypothesis [Bourgon et al., 2010]. In the negative binomial (NB) framework, this corresponds to filtering on the overall mean (based on an analogy to the normal case). Row sums can also be used for datasets where the effective library sizes are not hugely different, or where the counts are assumed to be Poisson-distributed.

```
> abundances <- aveLogCPM(data$counts, lib.size=data$totals*normfacs)
> length(abundances)

[1] 473281
> summary(abundances)

Min. 1st Qu. Median Mean 3rd Qu. Max.
-2.110 -2.029 -1.890 -1.660 -1.542 12.470
```

For demonstration purposes, an arbitrary threshold of -1 is used here to filter the window abundances. Users can then restrict the dataset to the filtered values. While filtering can be performed at any stage of the analysis, doing so at earlier steps is recommended to reduce computational work. Downstream estimates of various statistics are also more relevant when restricted to the regions of interest. Of course, one should retain enough points for successful information sharing (see Chapter 5).

```
> keep <- abundances > -1
> demo <- data
> demo$counts <- demo$counts[keep,]
> demo$region <- demo$region[keep]
> sum(keep)
[1] 52369
```

The exact choice of filter threshold may not be immediately obvious. A filter which is too conservative will be ineffective whereas a filter which is too aggressive may reduce power by removing false nulls. In some respects, the filter value is necessarily arbitrary as it reflects prior expectations of the abundances of the features of interest. Nonetheless, several strategies for defining the filter threshold are described below.

4.2 By proportion

One approach is to to assume that only a certain proportion - say, 0.1% - of the genome is genuinely bound. The number of windows corresponding to these bound regions can be calculated based on the known spacing interval used in windowCounts. The top set of windows with the highest abundances can then be selected such that the size of the set is equal to the computed number.

```
> spacing <- 50
> desired <- 0.001
> genome.windows <- sum(seqlengths(data$region)/spacing)
> keep <- length(abundances) - rank(abundances) + 1 < genome.windows*desired
> sum(keep)

[1] 54617
```

This approach is simple and has the practical advantage of maintaining a constant number of windows for the downstream analysis. However, it may not adapt well to different datasets where the relative proportion of binding can vary. Assuming an inappropriate percentage of binding will result in loss of potential DB regions or excessive inclusion of background regions.

4.3 By global enrichment

An alternative approach involves choosing a filter based on enrichment over non-specific background. Specifically, the median (or any other robust average) of the binned abundances can be used as an estimate of the global background coverage in the dataset. Binning is necessary here to increase the size of the counts. This ensures that precision is maintained when estimating the average background abundance.

```
> bin.size <- 2000L
> binned <- windowCounts(bam.files, bin=TRUE, width=bin.size)
> bin.ab <- aveLogCPM(binned$counts, lib.size=binned$total*normfacs)
> threshold <- median(bin.ab)</pre>
```

As the threshold is computed for large bins, it needs to be scaled for comparison to windows. This assumes a uniform distribution of reads in background regions. This means that the threshold can be directly scaled down based on differences in the size of the read counting interval. For windows with ext > 0, the size of the interval is equal to ext + width due to read extension on each strand.

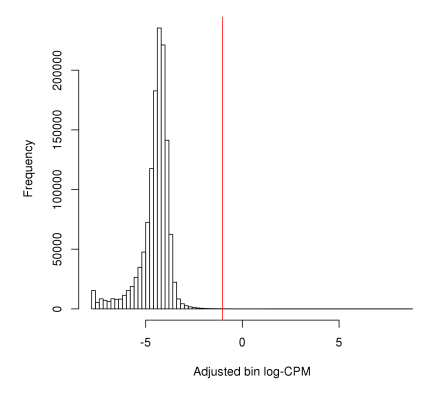
```
> width <- median(width(data$region))
> eff.win.size <- width+frag.len
> adjustment <- log2(bin.size/eff.win.size)
> threshold <- threshold-adjustment</pre>
```

Windows are then filtered based on some minimum required fold change over the global background. Here, a fold change of 10 over the background coverage is deemed necessary for a window to be considered as containing a binding site. This approach has an intuitive and experimentally relevant interpretation which adapts to the level of non-specific enrichment in the dataset.

```
> log.min.fc <- log2(10)
> threshold <- threshold+log.min.fc
> keep <- abundances >= threshold
> sum(keep)
[1] 54662
```

The effect of filtering can also be visualized with a histogram. This allows users to confirm that the bulk of (assumed) background windows are discarded upon filtering. Note that windows containing genuine binding sites will usually fail to appear on such plots due to the dominance of the background windows throughout the genome.

```
> hist(bin.ab-adjustment, xlab="Adjusted bin log-CPM", breaks=100, main="")
> abline(v=threshold, col="red")
```



Users should be aware that the pre-specified minimum fold change may be too aggressive when binding is weak. For TF data, a large cut-off works well as narrow binding sites will have high read densities and are unlikely to be filtered out. Smaller minimum fold changes are recommended for regions of diffuse enrichment where the difference to background is less obvious.

4.4 By local enrichment

4.4.1 Without negative controls

Local background estimators can also be constructed with a little imagination. This avoids inappropriate filtering due to differences in the sequencing coverage across the genome. Here, the 2 kbp region surrounding each window will be used as the "neighbourhood" over which a local estimate of non-specific enrichment for that window can be obtained. The counts for this region can be obtained with the aptly-named regionCounts function.

```
> surrounds <- 2000
```

> neighbour <- suppressWarnings(resize(data\$region, width(data\$region)+surrounds, fix="center"))

> wider <- regionCounts(bam.files, regions=neighbour, ext=frag.len)

Counts for each window are subtracted from the counts for its neighbourhood. This ensures that any enriched regions or binding sites inside the window do not interfere with estimation of its local background. Again, some adjustment for width is required for a valid comparison. Read extension is used for the neighbourhood counts, so note the addition of frag.len. The adjustment must also account for the subtraction of the window counts.

```
> neighbour.counts <- wider$counts- data$counts
> adjustment <- log2((surrounds + frag.len - eff.win.size)/eff.win.size)</pre>
```

Enrichment is then defined as the fold change of the average window abundance over the local neighbourhood. Filtering can then be performed using a quantile- or fold change-based threshold on the enrichment values. This roughly mimics the behaviour of single-sample peak-calling programs such as MACS [Zhang et al., 2008].

```
> neighbour.ab <- aveLogCPM(neighbour.counts, lib.size=wider$totals*normfacs)-adjustment
> en.ab <- abundances - neighbour.ab
> summary(en.ab)

Min. 1st Qu. Median Mean 3rd Qu. Max.
-9.524 1.445 1.930 1.872 2.330 12.340
```

Note that this procedure also assumes that no other enriched regions are present in each neighbourhood. Otherwise, the local background will be overestimated and windows may be incorrectly filtered out. This may be problematic for diffuse histone marks or TFBS clusters where enrichment may be observed in both the window and its neighbourhood.

If this seems to complicated, a simpler alternative is to identify locally enriched regions using peak-callers like MACS. Filtering can then be performed to retain only windows within called peaks. However, peak calling must be done independently of the DB status of each window. If libraries are of similar size, reads should be pooled into a single library for single-sample peak calling. This is equivalent to filtering on the average count and avoids distortion of the type I error rates.

4.4.2 With negative controls

Negative controls for ChIP-seq refer to input or IgG libraries where the IP step has been skipped or compromised, respectively. In particular, IgG controls provide a measure of non-specific enrichment over the genome. These libraries are largely irrelevant when testing for DB. However, they can be used for filtering whereby windows must have average ChIP counts with some fold change above the control. The example below requires a 5-fold increase.

```
> in.demo <- windowCounts(c(bam.files, "IgG.bam"), ext=frag.len)
> chip <- aveLogCPM(in.demo$counts[,1:4], lib.size=in.demo$totals[1:4])
> control <- aveLogCPM(in.demo$counts[,5], lib.size=in.demo$totals[5])
> keep <- chip > control + log2(5)
```

4.5 By prior information

When only a subset of genomic regions are of interest, power can be improved by removing windows lying outside of these regions. This could include promoters, enhancers, gene bodies or exons. The example below retrieves the coordinates of the broad gene bodies from the mouse genome after extending each 5' end by 3 kbp to include the putative promoter region.

```
> require(org.Mm.eg.db)
> suppressWarnings(anno <- select(org.Mm.eg.db, keys=keys(org.Mm.eg.db),
     col=c("CHRLOC", "CHRLOCEND"), keytype="ENTREZID"))
> anno <- anno[!is.na(anno$CHRLOCCHR),]</pre>
> extension <- 3000
> coord5 <- ifelse(anno$CHRLOC > 0, anno$CHRLOC-extension, -anno$CHRLOC)
> coord3 <- ifelse(anno$CHRLOC > 0, anno$CHRLOCEND, -anno$CHRLOCEND+extension)
> broads <- GRanges(paste0("chr", anno$CHRLOCCHR), IRanges(coord5, coord3), names=rownames(anno))
> head(broads)
GRanges with 6 ranges and 1 metadata column:
      segnames
                               ranges strand |
         <Rle>
                            <IRanges> <Rle> | <character>
  [1]
          chr9 [ 21062393, 21070093]
  [2]
          chr7 [ 84940169, 84967009]
                                                          4
  [3]
         chr10 [ 77708457.
                            777120097
                                                          5
                                                          6
  [4]
         chr11 [ 45805083, 45842878]
  [5]
          chr4 [144157556, 144165651]
                                                          9
  [6]
          chr4 [134745412, 134771004]
                                                         12
  seqlengths:
                                         chr7 ...
                                                        chrUn_JH584304
                   chr9
                                           NA ...
```

Windows can then be filtered to only retain those which overlap with the regions of interest. Discerning users may wish to distinguish between full and partial overlaps, though this should not be a significant issue for small windows. This can also be done in conjunction with abundance filtering to obtain windows overlapping binding sites in the regions of interest.

```
> suppressWarnings(keep <- overlapsAny(data$region, broads))
> sum(keep)
```

[1] 202254

Any information used here should be independent of the DB status under the null in the current dataset. For example, DB calls from a separate dataset and/or independent annotation can be used without problems. However, using DB calls from the same dataset to filter regions would violate the null assumption and compromise type I error control.

Chapter 5

Testing differential binding

For this next section, we'll be needing the data list from Chapter 2 and filtered in Chapter 4. Just let me assign the filtered list back to data, because I put it in a dummy variable:

```
> original <- data
> data <- demo</pre>
```

You'll also need the normfacs vector from Chapter 3, as well as the design matrix from the introduction.

5.1 Introduction to edgeR

Low counts per window are typically observed in ChIP-seq datasets, even for genuine binding sites. The statistical analysis must be able to handle discreteness in the data. Software packages using count-based models are ideal for this purpose. In this guide, the quasi-likelihood (QL) framework in the edgeR package is used [Lund et al., 2012]. Counts are modelled using NB distributions that account for overdispersion between biological replicates [Robinson and Smyth, 2008].

Before proceeding with the analysis, a DGEList object must be formed from the count matrix. Additional information like the library size and the normalization factors can (and should) be added. For this analysis, the normfacs vector from TMM normalization of background bins is used. If an offset matrix is necessary (e.g., from non-linear normalization), this can be assigned into y\$offset for use in downstream functions.

> y <- DGEList(data\$counts, lib.size=data\$totals, norm.factors=normfacs)

The experimental design is described by a design matrix. In this case, the only relevant factor is the cell type of each sample. A generalized linear model (GLM) will then be fitted to the counts for each window using the specified design matrix [McCarthy et al., 2012]. This provides a general framework for the analysis of complex experiments with multiple factors. Readers are referred to the user's guide in edgeR for more details on parametrization.

> design

It is worth pointing out that any method can be used if it is able to accept a count matrix and a vector of normalization factors (or more generally, a matrix of offsets). The choice of edgeR is primarily motivated by its performance relative to alternatives [Law et al., 2014], though the author's desire to increase his h-index is also a factor.

5.2 Estimating the dispersions

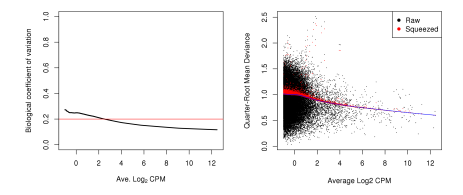
Under the QL framework, both the QL and NB dispersions are used to model biological variability in the data. The former ensures that the NB mean-variance relationship is properly specified with appropriate contributions from Poisson and gamma components. The latter accounts for variability and uncertainty in the dispersion estimate. However, limited replication in most ChIP-seq experiments means that each window does not contain enough information for precise estimation of either value.

This problem is overcome in edgeR by sharing information across windows. For the NB dispersions, a mean-dispersion trend is fitted across all windows to model the mean-variance relationship [McCarthy et al., 2012]. The raw QL dispersion for each window is estimated using the trended NB dispersion, and another trend is fitted to the QL estimates against the mean. An empirical Bayes (EB) strategy is then used to stabilize the raw QL dispersion estimates by shrinking them towards the second trend [Lund et al., 2012]. The ideal amount of shrinkage is determined from the heteroskedasticity of the data.

```
> par(mfrow=c(1,2))
> y <- estimateDisp(y, design)
> o <- order(y$AveLogCPM)
> plot(y$AveLogCPM[o], sqrt(y$trended.dispersion[o]), type="1", lwd=2,
```

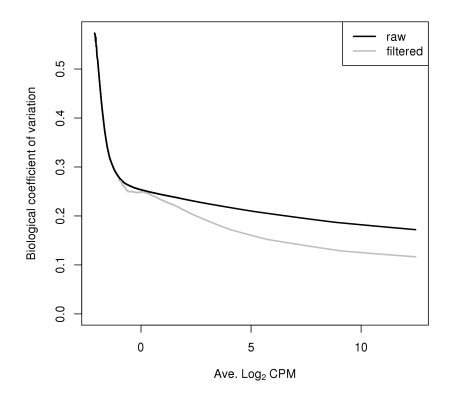
```
+ ylim=c(0, 1), xlab=expression("Ave."~Log[2]~"CPM"),
+ ylab=("Biological coefficient of variation"))
> abline(h=0.2, col="red")
> results <- glmQLFTest(y, design, robust=TRUE, plot=TRUE)</pre>
```

The effect of EB stabilisation can then be visualized by examining the biological coefficient of variation (for the NB dispersion) and the quarter-root deviance (for the QL dispersion). These plots can also be used to decide whether the fitted trend is appropriate. Sudden irregulaties may be indicative of an underlying structure in the data which cannot be modelled with the mean-dispersion trend. Discrete patterns in the raw dispersions may be observed for low counts and suggest that more aggressive filtering is required.



A strong trend may also be observed where the dispersion drops sharply with increasing mean. This is due to the disproportionate impact of artifacts such as mapping errors and PCR duplicates at low counts. It is difficult to accurately fit an empirical curve to these strong trends. This results in increased errors in the fitted values, typically manifesting as overestimation of dispersions at high abundances. Users should check whether removal of the low abundance regions affects the dispersion estimate. Large differences imply that further filtering may be desirable (see Chapter 4).

```
> yo <- DGEList(original$counts, lib=original$total, norm=normfacs)
> yo <- estimateDisp(yo, design)
> oo <- order(yo$AveLogCPM)
> plot(yo$AveLogCPM[oo], sqrt(yo$trended.dispersion[oo]), type="l", lwd=2,
+ ylim=c(0, max(sqrt(yo$trended))), xlab=expression("Ave."~Log[2]~"CPM"),
+ ylab=("Biological coefficient of variation"))
> lines(y$AveLogCPM[o], sqrt(y$trended[o]), lwd=2, col="grey")
> legend("topright", c("raw", "filtered"), col=c("black", "grey"), lwd=2)
```



5.2.1 Modelling heteroskedasticity

The heteroskedasticity of the data is modelled in edgeR by the prior degrees of freedom. A large prior degrees of freedom represents strong shrinkage and low heteroskedasticity. More EB shrinkage can be performed to reduce uncertainty and maximize power when the dispersions are not variable. However, strong shrinkage is not appropriate if the dispersions are highly variable. Lower prior degrees of freedom (and less shrinkage) is required to maintain type I error control.

> summary(results\$df.prior)

```
Min. 1st Qu. Median Mean 3rd Qu. Max. 0.5323 63.7300 63.7300 62.6000 63.7300 63.7300
```

On occasion, the estimated prior degrees of freedom will be infinite. This is indicative of a strong batch effect where the dispersions are large with minimal variability. A typical example would involve uncorrected differences in IP efficiency across replicates. In severe cases, the trended dispersion may fail to pass through the bulk of points as the variability is too low to be properly modelled in the QL framework. These batch effects are often caused by efficiency biases and can be removed with appropriate normalization.

It is worth pointing out that the prior degrees of freedom should be robustly estimated [Phipson et al., 2013]. Obviously, this protects against large positive outliers (e.g. highly variable windows) but it also protects against near-zero dispersions at low counts. These will manifest as large negative outliers after a log transformation step during estimation [Smyth, 2004]. Without robust estimation, incorporation of these outliers will inflate the observed variability in the dispersions and reduce power.

5.3 Testing for DB windows

The effect of specific factors can then be tested to identify windows with significant differential binding. In the QL framework, p-values are computed using the F-test [Lund et al., 2012]. This is more appropriate than using the likelihood ratio test as the F-test accounts for uncertainty in the dispersion estimates. Associated statistics such as log-fold changes and log-counts per million are also computed for each window.

```
> results <- glmQLFTest(y, design, robust=TRUE, contrast=c(0, 1))
> head(results$table)

logFC logCPM F PValue
1 -0.5078927 -0.5402696 1.0563315 0.30782063
2 -0.4598138 0.4265961 1.2652509 0.27607407
3 -0.5350752 -0.3695510 1.2712313 0.26363466
4 -0.4692340 -0.4043727 0.9678305 0.32882914
5 -0.8460653 -0.4803792 2.9451508 0.09084544
6 -0.2987352 -0.5264553 0.3757897 0.54198044
```

The null hypothesis here is that the cell type has no effect. The contrast argument in the glmQLFTest function specifies which factors are of interest. In this case, a contrast of c(0, 1) defines the null hypothesis as 0*intercept + 1*cell.type = 0 so changes between cell types can be identified. Specification of the contrast is explained in greater depth in the edgeR user's manual.

As a side note, the glmQLFTest function will perform both the dispersion estimation and the hypothesis testing. As such, it only needs to be called once. Multiple calls are only shown here and in Section 5.2 for demonstration purposes.

Chapter 6

Correction for multiple testing

All right, we're almost there. This chapter needs the results list and from the last chapter. You'll also need the filtered data list and the frag.len value from Chapter 2, as well as the the broads object from Chapter 4.

6.1 Problems with false discovery control

The false discovery rate (FDR) is usually the most appropriate measure of error for high-throughput experiments. Control of the FDR can be provided by applying the Benjamini-Hochberg (BH) method [Benjamini and Hochberg, 1995] to a set of p-values. This is less conservative than the alternatives (e.g., Bonferroni) yet still provides some measure of error control. The most obvious approach is to apply the BH method to the set of p-values across all windows. This will control the FDR across the set of putative DB windows.

However, the FDR across all detected windows is not necessarily the error of interest. Interpretation of ChIP-seq experiments is more concerned with regions of the genome at which (differential) protein binding is found. In other words, the FDR across all detected regions is usually desired. This is not equivalent to that across windows as each region will often consist of multiple overlapping windows. Control of one will not guarantee control of the other [Lun and Smyth, 2014].

To illustrate this difference, consider an analysis where the FDR across all window positions is controlled at 10%. In the results, there are 18 adjacent window positions forming one cluster and 2 windows forming a separate cluster. Each cluster represents a region. The first set of windows is a truly DB region whereas the second set is a false positive. A window-based interpretation of the FDR is correct as only 2 of the 20 window positions are false positives. However, a region-based interpretation results in an actual FDR of 50%.

Problems from misinterpretation can be avoided by applying the BH method to a p-value

from each region. Windows can be clustered together into regions with a number of strategies. Simes' method can then be used to compute a combined p-value for each cluster based on the p-values the constituent windows [Simes, 1986]. This tests the joint null hypothesis that no enrichment is observed across any sites within the region. The combined p-values are then converted to q-values using the BH method to control the region-level FDR.

6.2 Clustering with external information

Combined p-values can be computed for a predefined set of regions based on the windows overlapping those regions. The most obvious source of predefined regions is that of annotated features such as promoters or gene bodies. Alternatively, independently called peaks can be used though some care is required to avoid data snooping. In either case, the find-Overlaps function from the GenomicRanges package can be used to identify all windows in or overlapping each specified region.

```
> olap <- findOverlaps(broads, data$region)</pre>
> olap
Hits of length 26071
queryLength: 27294
subjectLength: 52369
      queryHits subjectHits
       <integer>
                    <integer>
                         36290
 1
 2
               11
                         48150
 3
               18
                         39194
 4
               18
                         39195
 5
               18
                         39196
              . . .
                         35634
 26067
            27283
 26068
            27283
                         35635
            27286
                         36047
 26069
 26070
            27286
                         36048
 26071
            27286
                         36049
```

The combineTests function can then be used to combine the p-values for all windows in each region. This provides a single combined p-value (and the corresponding q-value) for each region. The row names of the output table correspond to the value of the integer identifiers supplied in ids. These should, in turn, correspond to the regions of interest in broads. The average log-CPM and log-FC across all windows in each region are also computed.

```
> ids <- queryHits(olap)
> wids <- subjectHits(olap)
> tabprom <- combineTests(ids, results$table[wids,,drop=FALSE])
> head(tabprom)
```

```
logFC logCPM PValue FDR
7 2.95211689 -0.7893144 1.987679e-05 0.0002377056
11 0.95217836 -0.8375883 8.860506e-02 0.1451496065
18 1.56736941 0.1369136 9.613185e-04 0.0043771167
22 0.98826601 -0.9671633 9.477974e-02 0.1536700621
23 1.10365009 -0.2706485 4.965306e-02 0.0916712846
25 0.08376962 -0.4331375 9.041186e-01 0.9346915676
```

At this point, one might imagine that it would be simpler to just collect and analyze counts over the predefined regions. This is a valid strategy but can lead to quite different results from the one presented here. Consider a promoter region containing two separate peaks which are identically DB in opposite directions. Counting reads across the promoter will give equal counts for each group so changes within the promoter will not be detected. For peaks, imprecise boundaries can lead to loss of DB detection power due to "contamination" by reads from background regions. Window-based methods are more robust as each interval of the region is examined separately.

6.3 Quick and dirty clustering

Clustering can also be quickly performed inside csaw with a simple single-linkage algorithm implemented in the mergeWindows function. This approach can be useful as it avoids potential problems with the other clustering methods, e.g., peak-calling errors, incorrect or incomplete annotation. Briefly, all windows which are less than some distance apart - say, 1 kbp - are put in the same cluster. This reflects some arbitrary minimum distance at which two binding events are considered to be separate.

```
> merged <- mergeWindows(data$region, tol=1000L)</pre>
> merged$region
GRanges with 13665 ranges and 0 metadata columns:
          seqnames
                                   ranges strand
              <Rle>
                                <IRanges>
                                            <Rle>
      [1]
                      [3695601, 3695760]
               chr1
      [2]
               chr1
                      [3981801, 3981810]
      [3]
                      [5072001, 5072010]
               chr1
      [4]
               chr1
                      [7397901, 7398110]
                      [7860851, 7860860]
      [5]
               chr1
                . . .
               chrY [90782851, 90782910]
  [13661]
  [13662]
               chrY
                    [90784051, 90784110]
               chrY [90805151, 90805210]
  [13663]
  [13664]
               chrY [90808851, 90808860]
               chrY [90812101, 90813560]
  [13665]
  seqlengths:
                    chr1
                                          chr10 ... chrY_JH584303_random
               195471971
                                     130694993 ...
                                                                    158099
```

Combined p-values are computed for each cluster as previously described. Application of the BH method then controls the FDR across all detected clusters. Like before, the row names of the output table point towards the corresponding coordinates of the clusters in merged\$regions.

```
> tabcom <- combineTests(merged$id, results$table)
> head(tabcom)

logFC logCPM PValue FDR
1 -0.4930039 -0.2218993 0.32882914 0.58915041
2 -0.8460653 -0.4803792 0.09084544 0.26014312
3 -0.2987352 -0.5264553 0.54198044 0.77730506
4 1.0222183 0.4770579 0.01119654 0.05846416
5 -1.0761098 -0.9931622 0.06697002 0.21188824
6 1.3899478 -0.5090193 0.01215433 0.06215900
```

If many overlapping windows are present, very large clusters may be formed which are difficult to interpret. A simple check can be used to determine whether most clusters are of an acceptable size. Many huge clusters indicate that more aggressive filtering from Chapter 4 is required. This mitigates chaining effects by increasing the average spacing between each window on the genome.

> summary(width(merged\$region))

```
Min. 1st Qu. Median Mean 3rd Qu. Max.
10 10 60 269 160 51410
```

More generally, this problem can be avoided by setting the max.width parameter to restrict the sizes of the merged intervals. The chosen value should be small enough so as to be separate differentially bound regions from unchanged neighbours, yet large enough to avoid difficulties in interpretation from many adjacent windows. A value from 2000 to 10000 bp is recommended.

```
> merged <- mergeWindows(data$region, tol=1000L, max.width=5000L)
> summary(width(merged$region))

Min. 1st Qu. Median Mean 3rd Qu. Max.
10.0 10.0 60.0 263.8 160.0 4960.0
```

There are also provisions for clustering based on the sign of the log-fold change. The idea is that clusters will be broken up wherever the sign changes. This will separate binding sites that are close together but are changing in opposite directions. The mergeWindows function can do this if a vector is supplied in sign indicating whether each window has a positive log-fold change.

```
> merged <- mergeWindows(data$region, tol=1000L, sign=(results$table$logFC > 0))
> summary(width(merged$region))
```

```
Min. 1st Qu. Median Mean 3rd Qu. Max. 10.0 10.0 60.0 166.8 110.0 49860.0
```

However, sign-based filtering is not recommended as the sign of windows within each cluster is not independent of their DB status. A spurious/genuine DB region will form one cluster (consistent sign) whereas non-DB regions will form many cluster (inconsistent sign, as the log-fold change is small). This results in more large p-values and conservativeness. Furthermore, any attempt to filter away small clusters will cause liberalness if too many large p-values are lost.

Chapter 7

Post-processing steps

This is where we bring it all together. We'll need the merged list and the tabcom table from the previous chapter. There's a bit about visualization at the end where we need the y object from Chapter 5 and the bam.files that we started off with.

7.1 Adding gene-based annotation

Annotation can be added to a given set of regions using the detailRanges function. This will identify overlaps between the regions and annotated features such as exons, introns and promoters. Here, promoters are defined as some interval 3 kbp up- and 1 kbp downstream of the TSS. Any exonic features within dist on the left or right side of the region will also be reported.

```
> require(TxDb.Mmusculus.UCSC.mm10.knownGene)
> sig.bins <- merged$region[as.integer(rownames(tabcom))]</pre>
> anno <- detailRanges(sig.bins, txdb=TxDb.Mmusculus.UCSC.mm10.knownGene,
     orgdb=org.Mm.eg.db, promoter=c(3000, 1000), dist=5000)
> head(anno$overlap)
[1] ""
                             "Rgs20|0|-" ""
> head(anno$left)
[1] ""
                       11 11
                                          "Rgs20|1|-[1716]" ""
[5] ""
> head(anno$right)
[1] ""
                                                          11 11
[5] ""
                      "Rrs1|1|+[3898]"
```

Character vectors of compact string representations are provided that describe the features overlapped by each supplied region. Each pattern represents <code>GENE|EXONS|STRAND</code> for the overlapped exons of that gene. Promoters are labelled as exon <code>0</code> whereas introns are labelled as <code>I</code>. For <code>left</code> and <code>right</code>, an additional <code>DISTANCE</code> field is included which indicates the gap between the annotated feature and the supplied region.

While the string representation is space-saving, it is not easy to work with. If the annotation needs to manipulated directly, users can obtain it from the detailRanges command by not specifying the regions of interest. This can then be used, e.g., to identify the genes containing DB sites overlapping the promoter.

> anno.ranges <- detailRanges(txdb=TxDb.Mmusculus.UCSC.mm10.knownGene, orgdb=org.Mm.eg.db)
> head(anno.ranges)

GRanges with 6 ranges and 2 metadata columns:

0		O					
	s	eqnames		ranges	strand	symbol	exon
		<rle></rle>		Ranges	<rle></rle>	<pre><character></character></pre>	<pre><integer></integer></pre>
1000	09600	chr9	[21062393,	21062717]	-	Zglp1	5
1000	09600	chr9	[21062894,	21062987]	-	Zglp1	4
1000	09600	chr9	[21063314,	21063396]	-	Zglp1	3
1000	09600	chr9	[21066024,	21066377]	-	Zglp1	2
1000	09600	chr9	[21066940,	21067925]	-	Zglp1	1
1000	09609	chr7	[84940169,	84941088]	-	Vmn2r65	6
seqlengths:							
-	chr1		ır1	cl	nr2	chrUn_JH	H584304
		1954719	971	1821132	224		114452

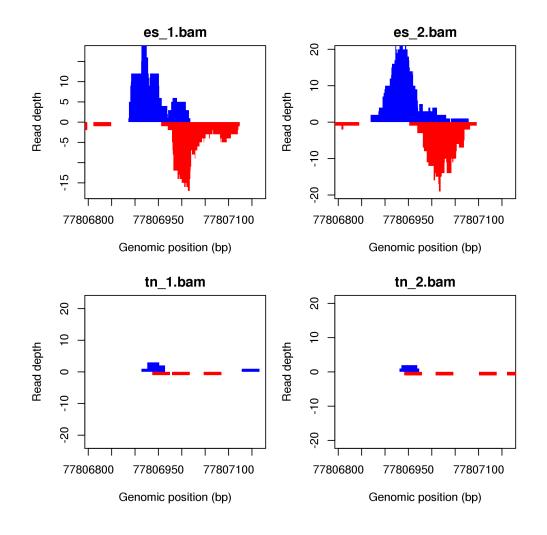
7.2 Saving the results to file

It is then a simple matter to save the results. This is done here in the *.tsv format where all detail is preserved. Compression is used to reduce the file size. Of course, other formats can be used depending on the purpose of the file, e.g., exporting to BED files through the rtracklayer package for visual inspection of the data with genomic browsers.

```
> ofile <- gzfile("clusters.gz", open="w")
> write.table(data.frame(chr=as.character(seqnames(sig.bins)), start=start(sig.bins),
+ end=end(sig.bins), tabcom, anno), file=ofile,
+ row.names=FALSE, quote=FALSE, sep="\t")
> close(ofile)
```

7.3 Simple home-made visualization

A quick and dirty inspection of the read depth around interesting features is also possible within. The blue and red tracks represent the coverage on the forward and reverse strands, respectively. The height of each plot is adjusted according to the library sizes to avoid any misrepresentation of read depth.



Chapter 8

Epilogue

Congratulations on getting to the end. Here's a poem for your efforts.

There once was a man named Will Who never ate less than his fill. He ate meat and bread Until he was fed But died when he saw the bill.

8.1 Datasets

8.1.1 Obtaining the FastQ files

The main NFYA dataset used throughout the guide was first mentioned in Section 1.3. This was generated by Tiwari et al. [2012] and is available from the NCBI Gene Expression Omnibus (GEO) with the accession number GSE25532. FastQ files can be obtained from the Sequence Read Archive (SRA) with accession numbers of SRR074398 for es_1.bam, SRR074399 for es_2.bam, SRR074417 for tn_1.bam and SRR074418 for tn_2.bam.

The paired-end dataset used in Section 2.2 was generated by Pal et al. [2013] and is available from the NCBI GEO under the accession GSE43212. FastQ files can be obtained from the SRA with the accession SRR642390 for example-pet.bam.

All libraries used in Section 2.3 were generated by Zhang et al. [2012] and are available from the NCBI GEO under the accession GSE31233. FastQ files can be obtained from the SRA under the accessions SRR330784 and SRR330785 for h3ac.bam; SRR330800 and SRR330801 for h3k4me2.bam; and SRR330814, SRR330815 and SRR330816 for h3k27me3.bam. Each set of FastQ files represents technical replicates that are merged into a single BAM file.

Finally, the H3K4me3 dataset in Section 3.2 was generated by Revilla-I-Domingo et al.

[2012] and is available from the NCBI GEO under the accession GSE38046. FastQ files can be obtained from the SRA under the accessions SRR499732 and SRR499733 for h3k4me3_pro.bam, and SRR499716 and SRR499717 for h3k4me3_mat.bam. Again, each set of FastQ files represents technical replicates. For H3ac normalization, the FastQ file at SRR330786 was also downloaded and used as h3ac_2.bam.

8.1.2 Alignment and processing to produce BAM files

Technically, the downloaded files are in the SRA format. These can be unpacked to form FastQ files using the fastq-dump command from the SRA Toolkit. For the paired-end library, users will need to specify fastq-dump -split-files to ensure that two separate files are produced, i.e., one containing sequences from each end.

The reads in the FastQ files can then be mapped with the aligner of choice. For this particular guide, all reads were aligned to the mm10 build of the mouse genome using the subread program [Liao et al., 2013]. This can be obtained from Bioconductor as Rsubread or as a standalone C program from http://subread.sourceforge.net/. Default settings were used with the exception of the consensus threshold, which was lowered to 2 to accommodate the short read lengths. Paired-end data was aligned by supplying both FastQ files to subread in the same run.

Once aligned, SAM files were converted to BAM files using the samtools suite [Li et al., 2009]. BAM files were position-sorted with the samtools sort command, and duplicate reads were marked using the MarkDuplicates command from the Picard suite (http://picard.sourceforge.net/). Any technical replicates were merged together using the samtools merge command to form a single library. Indexing was performed using the samtools index command.

8.2 Session information

> sessionInfo()

```
R version 3.1.0 (2014-04-10)
Platform: x86_64-unknown-linux-gnu (64-bit)
locale:
 [1] LC_CTYPE=en_US.UTF-8
                                LC_NUMERIC=C
 [3] LC_TIME=en_US.UTF-8
                                LC_COLLATE=en_US.UTF-8
 [5] LC_MONETARY=en_US.UTF-8
                                LC_MESSAGES=en_US.UTF-8
 [7] LC PAPER=en US.UTF-8
                                LC NAME=C
 [9] LC_ADDRESS=C
                                LC_TELEPHONE=C
[11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
attached base packages:
[1] splines parallel stats
                                  graphics grDevices utils
                                                                 datasets
```

[8] methods

```
other attached packages:
 [1] TxDb.Mmusculus.UCSC.mm10.knownGene_2.14.0
 [2] GenomicFeatures_1.16.2
 [3] org.Mm.eg.db_2.14.0
 [4] RSQLite_0.11.4
 [5] DBI_0.2-7
 [6] AnnotationDbi_1.26.0
 [7] Biobase_2.24.0
 [8] statmod_1.4.20
 [9] locfit_1.5-9.1
[10] edgeR_3.6.2
[11] limma_3.20.6
[12] csaw_0.99.0
[13] GenomicRanges_1.16.3
[14] GenomeInfoDb_1.0.2
[15] IRanges_1.22.9
[16] BiocGenerics_0.10.0
```

loaded via a namespace (and not attached):

[1]	BatchJobs_1.2	BBmisc_1.6	BiocParallel_0.6.1
[4]	biomaRt_2.20.0	Biostrings_2.32.0	bitops_1.0-6
[7]	brew_1.0-6	BSgenome_1.32.0	codetools_0.2-8
[10]	digest_0.6.4	fail_1.2	foreach_1.4.2
[13]	<pre>GenomicAlignments_1.0.1</pre>	grid_3.1.0	iterators_1.0.7
[16]	KernSmooth_2.23-12	lattice_0.20-29	plyr_1.8.1
[19]	Rcpp_0.11.2	RCurl_1.95-4.1	Rsamtools_1.16.1
[22]	rtracklayer_1.24.2	sendmailR_1.1-2	stats4_3.1.0
[25]	stringr_0.6.2	tools_3.1.0	XML_3.98-1.1
[28]	XVector_0.4.0	zlibbioc_1.10.0	

8.3 References

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