## seqbias

# Assessing and Adjusting for Technical Bias in High Throughput Sequencing Data

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## 1 Introduction

This package is designed as a means to assess and adjust for technical bias in high-throughput sequencing datasets, RNA-Seq being a specific target. As noted in previous studies, RNA-Seq is often subject to protocol specific bias. That is, the number of reads mapping to a particular position of the genome is dependent on the the surrounding nucleotide sequence (as well as the abundance of the RNA transcript) [1] [3]. Accounting for this bias increases uniformity of coverage and may result in more accurate quantification.

The approach implemented here trains a simple Bayesian network classifier and uses it to evaluate the per position bias. This builds off work done by Hansen, et. al. [1], available in the Genominator Bioconductor package. Another approach is taken by Li, et. al. [3] in the mseq package, available from CRAN.

For this vignette, we will use some example data taken from Mortazavi, et. al. [4] (NCBI accession number SRR001358). Because of space constraints, we have mapped the reads (using Bowtie [2]) to an artificial genome consisting of approximately 100kb of exonic DNA.

This "artificial genome" is given as,

```
> library(seqbias)
> library(Rsamtools)
> ref_fn <- system.file( "extra/example.fa", package = "seqbias" )
> ref_f <- FaFile( ref_fn )
> open.FaFile( ref_f )
   And the mapped reads,
> reads_fn <- system.file( "extra/example.bam", package = "seqbias" )</pre>
```

## 2 Assessment

As a natural first step, we would like to assess whether our sample is significantly biased. If this proves to be the case, we may wish to correct for this. A simple procedure to do so will be covered in the next section.

To assess the nucleotide frequency we will use a very simple procedure:

- 1. Generate a random sample of intervals across our reference genome.
- 2. Extract sequences for these intervals from a FASTA file.
- 3. Extract read counts across these intervals from a BAM file.
- 4. Using these sequences and counts, compute and plot nucleotide or k-mer frequencies.

## Sampling

For this step, we could use collection of known exons, but trustworthy annotations are not always available, and biasing the analysis by known exons may be a concern in some instances. Fortunately, seqbias provides a function to generate random intervals.

First, we extract a vector of sequence lengths, in the reference genome. Given an FASTA file than has been indexed with the samtools faidx command, we can use the Rsamtools package to read off the sequence lengths and to extract the sequence. First, the lengths,

```
> ref_seqs <- scanFaIndex( ref_f )</pre>
```

Once we have this, we generate 5 intervals of 100kb. It most cases we would want to generate a larger sample, but since we are working here with small reference sequence with dense coverage, we can get an accurate measurement with a few intervals.

```
> I \leftarrow random.intervals(ref_seqs, n = 5, m = 100000)
```

#### **Sequences**

Next we extract the nucleotide sequences,

```
> seqs <- scanFa( ref_f, I )</pre>
```

The scanFa function does not respect strand, so we must be sure to perform the reverse complement ourselves.

```
> neg_idx <- as.logical( I@strand == '-' )
> seqs[neg_idx] <- reverseComplement( seqs[neg_idx] )</pre>
```

#### **Counts**

Finally, we count the number of reads mapping to each position in our sampled intervals.

```
> counts <- count.reads( reads_fn, I, binary = T )</pre>
```

Unless the binary argument is FALSE, this function returns a 0-1 vector, where a position is 0 if no reads map to it, and 1 if *at least* one read maps to it. This is a more robust way to measure sequencing bias, as the frequencies can not get dominated by a few very high peaks.

## **Frequencies**

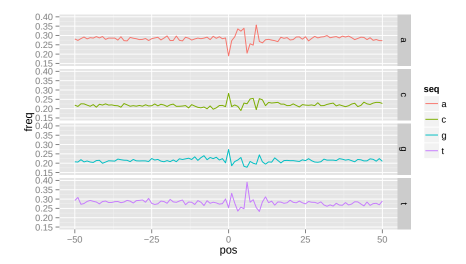
At last, we compute the k-mer frequency (where k = 1, by default).

```
> freqs <- kmer.freq( seqs, counts )</pre>
```

A nice way to plot this is with the ggplot2 package, if available.

```
> if( require(ggplot2) ) {
    P \leftarrow qplot(x = pos,
                 y = freq,
                 ylim = c(0.15, 0.4),
                 color = seq,
                 data = freqs,
                 geom = "line" )
   P <- P + facet_grid( seq ~ . )</pre>
    print(P)
+ } else {
    par(mar = c(5,1,1,1), mfrow = c(4,1))
    with( subset( freqs, seq == "a" ),
          plot( freq \tilde{} pos, ylim = c(0.15, 0.4), sub = "a", type = '1' ) )
   with( subset( freqs, seq == "c" ),
          plot( freq \tilde{} pos, ylim = c(0.15, 0.4), sub = "c", type = 'l'))
   with( subset( freqs, seq == "g" ),
          plot(freq \sim pos, vlim = c(0.15, 0.4), sub = "g", type = 'l'))
   with( subset( freqs, seq == "t" ),
          plot( freq \tilde{} pos, ylim = c(0.15, 0.4), sub = "t", type = 'l' ) )
+ }
```

Doing so will produce the following plot,



The x-axis shows the nucleotide position relative to the read start. Negative number occur in the genome to the left of mapped reads. In this set, the reads consist of positions 0-24.

We can see a clear bias here in positions 0-15, approximately. The rest of the plot looks relatively flat, as we would expect if the experiment was measuring abundance only and not a biased by the nucleotide sequence. In the next section we will adjust read counts to account for this.

# 3 Compensation

## **Training**

To begin, we must fit a seqbias model to our dataset. This done very easily with the seqbias.fit function. This will take only a few seconds, but when more reads are available a more accurate model can be trained at the expense of the training procedure taking up to several minutes.

The L and R arguments control the maximum number of positions to the left and right of the read start that may be considered. The model tries to consider only informative positions, so increasing these numbers will increase training time, but should never have a negative effect the accuracy of the model.

#### **Prediction**

Once we have trained the sequins model, we can use it to predict the sequencing bias across a set of intervals.

```
> bias <- seqbias.predict( sb, I )</pre>
```

## **Adjustment**

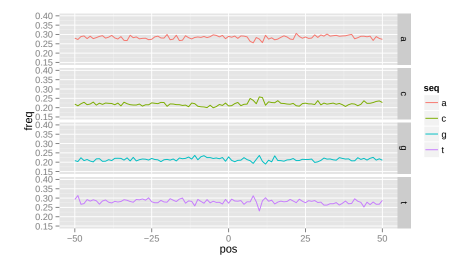
To adjust, we will can simply divide the counts vectors by the bias vectors.

```
> counts.adj <- mapply( FUN = `/`, counts, bias, SIMPLIFY = F )</pre>
```

The post-adjustment nucleotide frequencies can then be measured as before,

```
> freqs.adj <- kmer.freq( seqs, counts.adj )</pre>
  And plotted,
> if( require(ggplot2) ) {
   P \leftarrow qplot(x = pos,
                 y = freq,
                 ylim = c(0.15, 0.4),
                 color = seq,
                data = freqs.adj,
                 geom = "line" )
   P <- P + facet_grid( seq ~ . )</pre>
   print(P)
+ } else {
   par(mar = c(5,1,1,1), mfrow = c(4,1))
    with( subset( freqs.adj, seq == "a" ),
          plot( freq \tilde{} pos, ylim = c(0.15, 0.4), sub = "a", type = 'l' ) )
   with( subset( freqs.adj, seq == "c" ),
          plot(freq ~ pos, ylim = c(0.15, 0.4), sub = "c", type = 'l'))
   with( subset( freqs.adj, seq == "g" ),
          plot( freq \tilde{} pos, ylim = c(0.15, 0.4), sub = "g", type = 'l'))
    with( subset( freqs.adj, seq == "t" ),
          plot( freq \tilde{} pos, ylim = c(0.15, 0.4), sub = "t", type = '1' ) )
+ }
```

The plot below results,



Compared to the first figure, the improvement is clear.

## 4 Save / Load

If the model is fit using a large number of reads, it can take several minutes to train. To avoid repeatedly refitting the model, seqbias provides a mechanism to save and load the model to a YAML file with the seqbias.save and seqbias.load functions.

```
> seqbias.save( sb, "my_seqbias_model.yml" )
> # load the model sometime later
> sb <- seqbias.load( ref_fn, "my_seqbias_model.yml" )</pre>
```

Note when loading the model, we need to provide a reference sequence. The seqbias object keeps track of the reference sequence to make seqbias.predict more convenient.

## 5 Session Info

```
[5] LC_MONETARY=en_US.UTF-8 LC_MESSAGES=en_US.UTF-8
```

[7] LC\_PAPER=C LC\_NAME=C

[9] LC\_ADDRESS=C LC\_TELEPHONE=C

[11] LC\_MEASUREMENT=en\_US.UTF-8 LC\_IDENTIFICATION=C

#### attached base packages:

[1] parallel stats graphics grDevices utils datasets methods

[8] base

#### other attached packages:

| [1] | ggplot2_0.9.3.1   | Rsamtools_1.12.0                | seqbias_1.8.0  |
|-----|-------------------|---------------------------------|----------------|
| [4] | Biostrings_2.28.0 | <pre>GenomicRanges_1.12.0</pre> | IRanges_1.18.0 |

[7] BiocGenerics\_0.6.0

#### loaded via a namespace (and not attached):

| [1]  | MASS_7.3-26     | RColorBrewer_1.0-5 | bitops_1.0-5 | colorspace_1.2-1 |
|------|-----------------|--------------------|--------------|------------------|
| [5]  | dichromat_2.0-0 | digest_0.6.3       | grid_3.0.0   | gtable_0.1.2     |
| [9]  | labeling_0.1    | munsell_0.4        | plyr_1.8     | proto_0.3-10     |
| [13] | reshape2_1.2.2  | scales_0.2.3       | stats4_3.0.0 | stringr_0.6.2    |
|      |                 |                    |              |                  |

[17] tools\_3.0.0 zlibbioc\_1.6.0

## References

- [1] Kasper Hansen, Steven Brenner, and Sandrine Dudoit. Biases in Illumina transcriptome sequencing caused by random hexamer priming. *Nucleic acids research*, pages 1–7, April 2010.
- [2] Ben Langmead, Cole Trapnell, Mihai Pop, and Steven Salzberg. Ultrafast and memory-efficient alignment of short DNA sequences to the human genome. *Genome biology*, 10(3):R25, 2009.
- [3] Jun Li, Hui Jiang, and Wing Hung Wong. Modeling non-uniformity in short-read rates in RNA-Seq data. *Genome Biology*, 11(5):R50, 2010.
- [4] Ali Mortazavi, Brian Williams, Kenneth Mccue, Lorian Schaeffer, and Barbara Wold. Mapping and quantifying mammalian transcriptomes by RNA-Seq. *Nature Methods*, 5(7):1–8, 2008.