# Performing Some Basic Quality Checking and Analysis on Sequencing Data with Repitools

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November 30, 2010

### 1 Introduction

Repitools is a package that allows statistics of differential epigenetic marking to be calculated, as well as summaries of genome - wide trends to be visualised in a variety of formats. Some basic quality checking utilities are also available for sequencing data. The utility of Repitools comes from that most of the functionality available is implemented for both microarrays and next generation sequencing, with very similar function calls for both types of data.

In this vignette, quality checking of the sequencing data, followed by analysis and visualisation will be demonstrated. A more detailed description of the package can be found in the associated Bioinformatics Applications Note <sup>1</sup>

To start with, load the Repitools package.

> library(Repitools)

## 2 Data

A GRangesList of mapped short reads from an Illumina Genome Analyser of four samples is included with the package. GRanges objects of mapped files from many popular aligners can be created by first reading them into R with the readAligned function in the ShortRead package, then coerced with as(alnRdObj, "GRanges"). Only reads on chromosome 21 were kept, to have fast - running examples. The details of the samples are:

- > dataPath <- system.file("data", package = "Repitools")</pre>
- > load(paste(dataPath, "samplesList.Rdata", sep = .Platform\$file.sep))
- > names(samplesList)
- [1] "Cancer Input" "Cancer MBD2IP" "Normal Input" "Normal MBD2IP"

Also, an annotation of genes located on chromosome 21 is included. The annotation is based on one provided from Affymetrix with their expression arrays <sup>2</sup>.

<sup>&</sup>lt;sup>1</sup>Repitools: an R package for the analysis of enrichment-based epigenomic data

<sup>&</sup>lt;sup>2</sup>http://www.affymetrix.com/Auth/analysis/downloads/na27/wtgene/HuGene-1\_0-st-v1.na27.hg18.transcript.csv.zip

```
> geneAnno <- read.csv(paste(dataPath, "chr21genes.csv", sep = .Platform$file.sep),
      stringsAsFactors = FALSE)
> head(geneAnno)
    chr strand
                             end
                                    symbol
                  start
                                                      name
             + 33837227 33871651
                                   Clorf92
1 chr21
                                                  AK127947
             - 39636487 39642849 C15orf32 ENST00000380748
2 chr21
3 chr21
             + 13341774 13341881
                                       --- ENST00000384369
             + 13915768 13915875
                                       --- ENST00000384745
4 chr21
             + 14295118 14295292
                                       --- ENST00000330957
5 chr21
             + 14365062 14365178
                                       --- ENST00000364942
6 chr21
```

Lastly, there is matrix of gene expression difference data, with each element related to the corresponding row in the gene annotation table. These values are the t-statistics of background corrected and RMA normalised Affymetrix expression array experiments. The expression differences matrix will be used when illustrating some of the visualisation functionality later in the vignette.

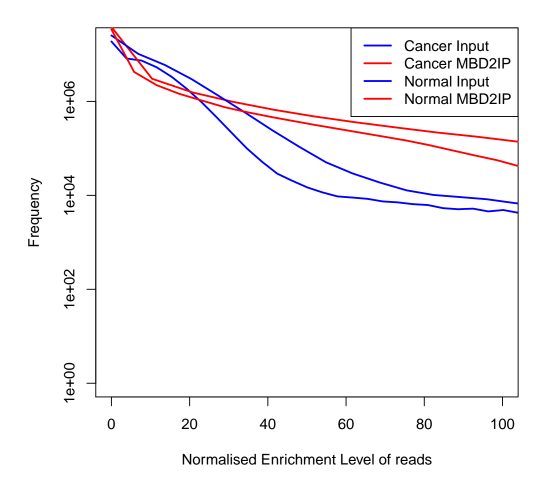
```
> load(paste(dataPath, "expr.Rdata", sep = .Platform$file.sep))
> head(expr)
```

t-stat
0.08
1.31
0.11
0.24
-1.88
-0.56

# 3 Quality Checking

Notice that two of the samples are MBD2 IPs, and two are inputs. Therefore, the IP samples should differ to the inputs in two ways. Firstly, they should be more CpG rich, since DNA methylation rarely ever occurs outside of this sequence context. Also, since DNA methylation tends to occur in peaks, rather than spread out regions, a higher frequency of bases should have high coverage of reads in the IP samples than in input samples. The enrichmentPlot and cpgDensityPlot functions allow examination of this.

#### **Enrichment Plot**

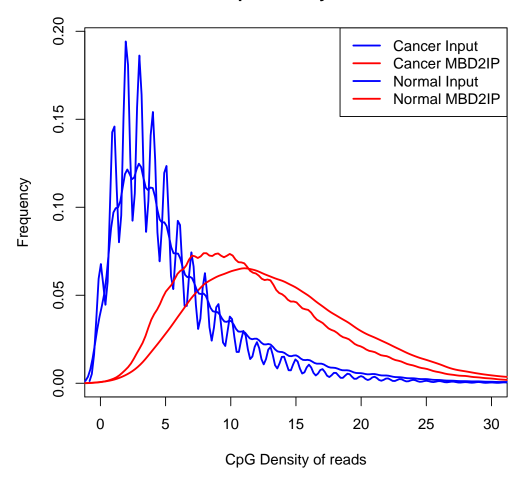


The above code uses the Hsapiens object to get the maximum base of chromosomes. The normalisation of the coverage used is to scale every coverage value by  $10 \ million/number\_of\_reads\_in\_sample$ . 300 is passed in as the seqLen parameter, because that is approximately the real length of the fragments sequenced. As expected, many more bases in the IP samples have high read coverages.

Next, the CpG density of reads is examined.

```
> cpgDensityPlot(samplesList, cols = c("blue", "red", "blue", "red"),
+ xlim = c(0, 30), wFunction = "none", organism = Hsapiens,
+ seqLen = 300, lwd = 2)
```

### **CpG Density Plot**



This time the Hsapiens annotation is required so that the 300 base DNA sequence (tags are only 36 bp long) may be fetched. The wFunction parameter allows the count of CpGs to be weighted. In this example, raw counts are used.

Notice that at lower CpG densities, the two input samples have a higher frequency of reads than the two IP samples. At higher CpG densities, this trend is reversed. This suggests that the DNA methylation IP has worked.

For more general sequencing quality checking, the FastQC <sup>3</sup> Java application has been gaining in popularity due to its speed and variety of results. A container class, FastQC, accessors, and the method readFastQC for reading in the raw Java program text file output and creating a FastQC R object, have been made to provide a framework for the fast development of quality control report generating pipelines. Higher level container classes with accessor methods are SequenceQC, which groups FastQC objects for the aligned and unaligned reads of a single sample together, and SequenceQCSet which is a collection of SequenceQC objects, perhaps of multiple sequencing samples within the same experimental run.

<sup>&</sup>lt;sup>3</sup>http://www.bioinformatics.bbsrc.ac.uk/projects/fastqc/

## 4 Analyses and Visualisations

The blocksStats function is a convenient way to do a statistical test of differential enrichment between two groups or treatments, for counts in windows surrounding some genomic landmarks, like TSSs. The function leverages the package edgeR's modelling of counts as negative binomial distributed and its adaptation of Fisher's exact test to overdispersed data.

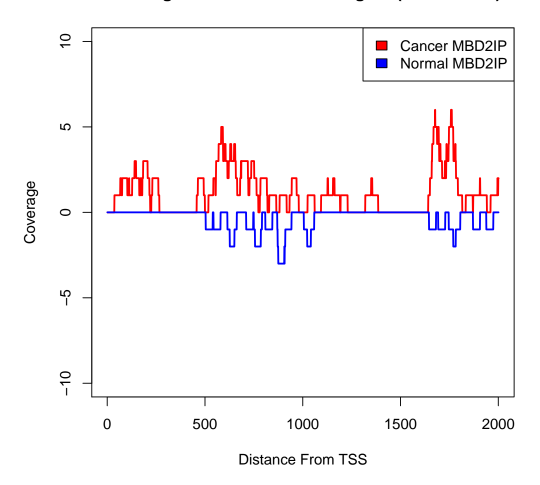
```
> designMatrix <- matrix(c(0, 1, 0, -1), dimnames = list(names(samplesList),
      "Cancer - Normal Methylation"))
> designMatrix
              Cancer - Normal Methylation
Cancer Input
                                         1
Cancer MBD2IP
Normal Input
                                         0
Normal MBD2IP
                                        -1
> stats <- blocksStats(samplesList, coordinatesTable = geneAnno,
      design = designMatrix, upStream = 2000, downStream = 0, seqLen = 300)
Generating table of counts
Counting in Cancer Input
Counting in Cancer MBD2IP
Counting in Normal Input
Counting in Normal MBD2IP
Counting successful.
Processing column 1 of design matrix
Comparison of groups:
                       1 - -1
> stats <- stats[order(stats$`FDR_Cancer - Normal Methylation`),
> head(stats)
                chr strand
                                                 symbol Cancer Input
         name
                               start
                                          end
52
     BC111558 chr21
                         - 44054086 44056875 LOC284837
                                                                   11
                                               FLJ41733
26
     AK123727 chr21
                         + 43607639 43610874
                                                                   14
198 NM_004540 chr21
                         + 21292503 21833085
                                                                   14
                                                  NCAM2
217 NM_006988 chr21
                         - 27130476 27139599
                                                                   17
                                                ADAMTS1
     BC031097 chr21
                         - 45093942 45118023
                                                PTTG1IP
                                                                   13
45
208 NM_005806 chr21
                         + 33320108 33323370
                                                  OLIG2
                                                                   22
    Cancer MBD2IP Normal Input Normal MBD2IP Cancer MBD2IP_pseudo
52
               73
                             13
                                           21
                                                           97.74692
26
               78
                             18
                                           29
                                                          104.44012
               44
                             12
                                            8
                                                           58.92638
198
217
               44
                             15
                                           12
                                                           58.92638
               26
                             12
                                            2
                                                           34.83090
45
208
               34
                              3
                                            6
                                                          45.54000
    Normal MBD2IP_pseudo logConc_Cancer - Normal Methylation
               15.664843
52
```

```
26
               21.641086
                                                      -11.40371
198
                5.953344
                                                      -12.74569
217
                8.941529
                                                      -12.45320
45
                1.470797
                                                      -14.12518
208
                4.459219
                                                      -13.13919
    logFC_Cancer - Normal Methylation PValue_Cancer - Normal Methylation
52
                              2.639043
                                                               6.125403e-16
26
                              2.268957
                                                               2.792528e-14
198
                              3.300968
                                                               3.169950e-12
217
                              2.716005
                                                               4.322107e-10
                                                               4.656613e-10
45
                              4.541976
208
                              3.344036
                                                               8.654979e-10
    FDR_Cancer - Normal Methylation
52
                        1.892749e-13
                        4.314456e-12
26
198
                        3.265049e-10
217
                        2.877787e-08
                        2.877787e-08
45
208
                        4.457314e-08
```

The example calculates statistics on TSS regions which start 2000 bases upstream of the TSS and finish at the TSS, after the reads have been extended to being 300 bases long. A coverage plot is shown to illustrate the best found region.

```
> cvgs <- lapply(samplesList[c(2, 4)], function(sampleRanges) coverage(sampleRanges)[[1]]
+ 2000), drop = TRUE])
> plot(cvgs[[1]], type = "l", ylim = c(-10, 10), main = "Coverage Plot for Best TSS Region
+ xlab = "Distance From TSS", ylab = "Coverage", col = "red",
+ lwd = 2)
> lines(-cvgs[[2]], col = "blue", lwd = 2)
> legend("topright", names(samplesList)[c(2, 4)], fill = c("red",
+ "blue"))
```

## **Coverage Plot for Best TSS Region (LOC284837)**

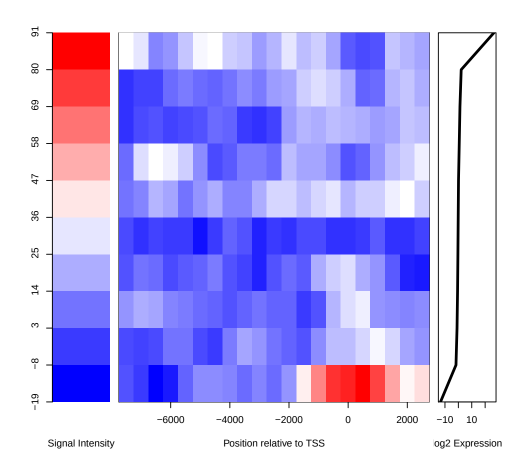


Epigenomic data is often gathered with other data, such as gene expression. It may be of interest to see the profile of epigenetic mark enrichment at a variety of distances from TSSs, and stratify this into groups by the expression of genes. The binPlots function is a convenient way to look at these interactions.

```
> binPlots(samplesList, geneAnno, design = designMatrix, by = 500,
+ bw = 500, seqLen = 300, ordering = expr, plotType = "heatmap",
+ nbins = 10)

gdata: read.xls support for 'XLS' (Excel 97-2004) files ENABLED.

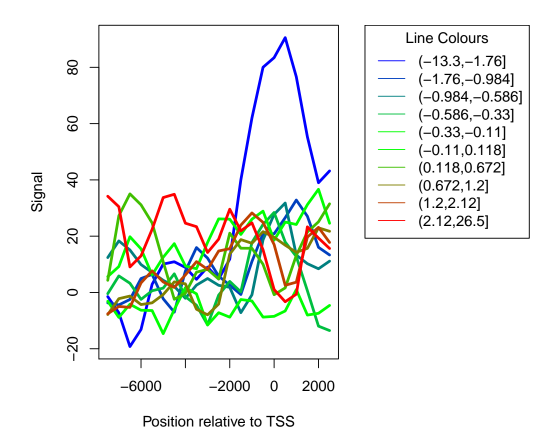
gdata: read.xls support for 'XLSX' (Excel 2007+) files ENABLED.
```



This example made counts in 500 base non - overlapping windows between -7500 bases upstream and 2500 bases downstream (the default range) for each gene, then split them into categories based on the expression difference value, and averaged over all counts for each particular window and expression category. It is encouraging to see that the lowest level of expression has a rather fine enrichment of DNA methylation about 2000 bases either side of the TSS. Apart from the heatmap visualisation, there are a number of other styles. Details can be found in the documentation of the function. More bins can be used for a nicer-looking heatmap when the full set of genes (genome-wide) is used.

To demonstrate how similar it is to generate another style of binned plot, the next example shows the same data as a line-plot. Note that the function call is the same, apart from the plotType parameter. Notice the spike in DNA methylation for the set of lowest expressed genes, which form the blue line.

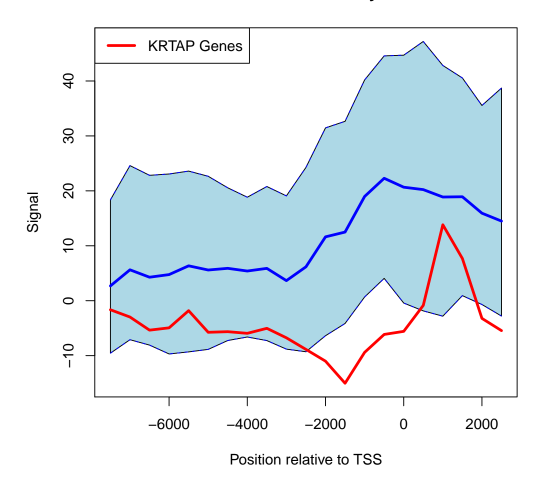
```
> binPlots(samplesList, geneAnno, design = designMatrix, by = 500,
+ bw = 500, seqLen = 300, ordering = expr, plotType = "line",
+ nbins = 10)
```



Some genes may be of interest to the researcher for some reason. This subset of genes may be known to be strongly marked with another epigenetic mark, or change in expression in the same direction strongly, or many other reasons. No matter what the reason for selecting the subset is, the profile of intensities or counts can be plotted versus the profile of randomly selected gene lists and compared with the significancePlots function. In the following example, it will be checked whether the DNA methylation profile of keratin-associated protein coding genes is any different to that of the other genes.

```
> whichKRTAP <- grep("KRTAP", geneAnno$symbol)
> significancePlots(samplesList, geneAnno, geneList = list(`KRTAP Genes` = whichKRTAP),
+ design = designMatrix, by = 500, bw = 500, seqLen = 300)
```

### **Cancer – Normal Methylation**



The blue region forms the null distribution that was created by sampling random gene lists of the same size as the user - specified gene list a number of times, as set by the nSamples parameter. By default, the null region is a between the 0.025 and 0.975 quantiles of the null distribution. In this example, it appears that the KRTAP gene set has a significant loss of methylation around from the gene TSS to around 2000 bases upstream, in comparison to random sets of other genes.

Another analysis of interest for the epigenomics research community is to find regions of the genome where epigenetics marks or changes in such marks occur in consecutive genes on a particular chromosome. The function findClusters addresses this need. The method of determining clusters is to look through the column of statistics for a set of consecutive scores in the same direction. Which potential clusters are significant is determined by randomising the ordering of the statistics column a number of times, and counting the number of clusters found in the real statistics column and the randomised statistics columns, from a loose cutoff to a tight cutoff, and choosing the cutoff to be the first cutoff that meets or is below the user - specified FDR. It will not be demonstrated, as this randomisation procedure is time consuming.

# 5 Utility Functions

These functions perform some task that is commonly made with the data, but is not a formal analysis. Often, it is required to create a set of windows covering the entire genome, for some analysis. The function genomeBlocks dies this.

> genomeBlocks(Hsapiens, "chr21", 5000)

GRanges with 9389 ranges and 0 elementMetadata values

seqnames	ranges	${ t strand}$	- 1
<rle></rle>	Ranges	<rle></rle>	
chr21	[ 1, 5000]	*	
chr21	[ 5001, 10000]	*	
chr21	[10001, 15000]	*	
chr21	[15001, 20000]	*	
chr21	[20001, 25000]	*	
chr21	[25001, 30000]	*	
chr21	[30001, 35000]	*	
chr21	[35001, 40000]	*	
chr21	[40001, 45000]	*	
chr21	[46900001, 46905000]	*	
chr21	[46905001, 46910000]	*	
chr21	[46910001, 46915000]	*	
chr21	[46915001, 46920000]	*	
chr21	[46920001, 46925000]	*	
chr21	[46925001, 46930000]	*	
chr21	[46930001, 46935000]	*	
chr21	[46935001, 46940000]	*	
chr21	[46940001, 46945000]	*	
	<pre><r1e> chr21 chr21</r1e></pre>	<pre></pre>	<pre> <rle></rle></pre>

```
seqlengths
chr21
NA
```

This example makes a GRanges object of 5 kb windows along chromosome 21.

The researcher might have a set of locations that they want to know the CpG density of.

```
> cpgDensityCalc(head(geneAnno), window = 100, organism = Hsapiens)
```

```
[1] 3.38 4.74 0.84 1.12 0.16 2.88
```

This example calculates the CpG density of a window 100 bases either side of the TSS for the first six genes on chromosome 21. By default, the CpG density is a linearly weighted count, such that a CpG at the TSS counts as one, and progressively less, towards zero, as the CpGs occur toward the edges of the region.

annotationCounts is useful to understand the counts of reads surrounding some landmarks, like TSSs. annotationBlocksCounts is the analogous function for counting in user-specified regions of the genome.

> annotationCounts(samplesList, head(geneAnno), 2000, 500, 300)

```
Counting in Cancer Input
Counting in Cancer MBD2IP
Counting in Normal Input
Counting in Normal MBD2IP
Counting successful.
```

	${\tt Cancer}$	Input	${\tt Cancer}$	MBD2IP	Normal	Input	Normal	MBD2IP
AK127947		13		3		16		9
ENST00000380748		7		4		15		19
ENST00000384369		6		0		0		0
ENST00000384745		5		1		4		0
ENST00000330957		6		0		0		2
ENST00000364942		20		51		23		60

This example counts reads that fall within 2000 bases upstream and 500 bases downstream of the first six genes on chromosome 21.

It would be good to know when seeing a lack of reads in some windows, if the mappability of the window is the cause. Some regions of the genome have low complexity sequence, and it is common practice in the analysis of sequencing data to discard reads that map to multiple locations in the genome. The function mappabilityCalc calculates the percentage of each region that can be mapped to by reads generated from the experiment. It operates on a user-created BSgenome object of a masked genome sequence. The definition of which bases are mappable and which are not depends on the fragment length of the sequencing technology used. Therefore, there is no one masked BSgenome object that can be used by all users. Note that by masking, it is meant replacing the unmappable reference sequence bases by 'N', not creating a built-in mask.

# 6 Summary

Repitools has a number of useful functions for quality checking, analysis, and comparison of trends. Many of the functions work seamlessly on array data, as well as sequencing data. Also, there are numerous utility functions, that perform some common task in the investigation of epigenomic data. Consult the package documentation for instructions on how to use functions that were not demonstrated by this vignette.

## 7 Environment

This vignette was created in:

> sessionInfo()

R version 2.12.0 Patched (2010-10-19 r53373) Platform: i386-apple-darwin9.8.0/i386 (32-bit)

locale:

[1] C/en\_CA.UTF-8/C/C/en\_CA.UTF-8/en\_CA.UTF-8

### attached base packages:

[1] grid stats graphics grDevices utils datasets methods

[8] base

#### other attached packages:

[1] gplots\_2.8.0 caTools\_1.10 [3] bitops\_1.0-4.1 gdata\_2.8.0 [5] gtools\_2.6.2 edgeR\_1.8.2 [7] BSgenome.Hsapiens.UCSC.hg18\_1.3.16 Repitools\_1.49

[9] BSgenome\_1.18.2 Biostrings\_2.18.0

[11] GenomicRanges\_1.2.1 IRanges\_1.8.3

[13] R.methodsS3\_1.2.1

loaded via a namespace (and not attached):

[1] Biobase\_2.10.0 limma\_3.6.0 tools\_2.12.0