# Assessing ChIP-seq sample quality with ChIPQC

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### 1 Introduction

This vignette illustrates how to compute quality metrics for aligned data from ChIP-seq experiments using the *ChIPQC* package. *ChIPQC* will automatically compute a number of quality metrics, and provides simple ways to generate a ChIP-seq experiment quality report which can be examined to asses the absolute and relative quality of individual ChIP-seq samples (and their associated controls, as well as overall quality of the experimental data.)

Two examples of ChIP-seq experiments comprising multiple samples are included in the vignette. The first contains some problematic samples to demonstrate how quality issues can be detected., while the second is a complete experiment of reasonable quality. There is another example showing how to process a single sample.

First is a summary of the minimal set of commands needed to generate a QC report using *ChIPQC*.

### 1.1 Summary: Generating a QC report for a ChIP-seq Experiment

Given a sample sheet samples, a report can be generated in two commands. The first constructs a ChIPQCexperiment object, including calculating all the QC metrics; the second (optional) step shows a quick summary of the primary QC metrics; and the third writes out a summary report that can be viewed in a browser, as follows:

- > experiment = ChIPQC(samples)
- > experiment
- > ChIPQCreport(experiment)

By default, the HTML report and associated figures will be written in a sub directory named ChIPQCreport, with the main html file named ChIPQC.html.

An example report (corresponding to the second example below) is available for examination at http://ChIPQC.starkhome.com/Reports/tamoxifen/ChIPQC.html.

### 1.2 Summary: Generating a QC report for a ChIP-seq sample

QC reports can be generated for single samples. This example checks a single bam file, chip.bam:

```
> sample = ChIPQCsample("chip.bam")
```

```
> sample
```

> ChIPQCreport(sample)

The following examples explore this in more depth.

# 2 Example 1: ENCODE data set with problematic samples

The first example, derived using ENCODE data for three transcription factors (CTCF, CMYC, and E2F1) [1]. Each ChIP has been performed in replicate, for a total of six samples (there are no controls in this example, see the following example for working with controls).

The actual reads and peak for this experiment are not included with this vignette as they are too large. The sample sheet (see below) include the SRA SRR numbers for each of the experiments, so you can download reads and peaks to fully run the vignette. If you do so, you should put the reads in a sub directory called reads and the peaks in a sub directory called peaks. We intend to make all the vignette data more easily obtainable; email the package authors if you are interested.

### 2.1 Experiment sample sheet

The first step is to construct a sample sheet describing the ChIP-seq experiment. This can be passed as a data frame, or saved in a csv file. The experiment can also be represented by a DBA object constructed using the *DiffBind* package, which accepts the same sample sheets as *ChIPQC*. A csv sample sheet has been included for this data set, and can be accessed as follow:

```
> library(ChIPQC)
> samples = read.csv(file.path(system.file("extdata", package="ChIPQC"),
                                             "example_QCexperiment.csv"))
+
> samples
  SampleID Tissue Factor Replicate
                                                bamReads
              A549
                                   1 reads/SRR568129.bam
1
    CTCF_1
                     CTCF
2
    CTCF_2
              A549
                     CTCF
                                   2 reads/SRR568130.bam
3
    cMYC_1
              A549
                     cMYC
                                   1 reads/SRR568131.bam
4
    cMYC_2
              A549
                     cMYC
                                   2 reads/SRR568132.bam
5
    E2F1_1 HeLa-S3
                     E2F1
                                   1 reads/SRR502355.bam
6
    E2F1_2 HeLa-S3
                     E2F1
                                   2 reads/SRR502356.bam
                             Peaks
1 peaks/SRR568129_chr22_peaks.bed
2 peaks/SRR568130_chr22_peaks.bed
3 peaks/SRR568131_chr22_peaks.bed
4 peaks/SRR568132_chr22_peaks.bed
5 peaks/SRR502355_chr22_peaks.bed
6 peaks/SRR502356_chr22_peaks.bed
```

This sample sheet details the metadata for the experiment. It also contains file paths to the aligned reads and previously called peaks. Note that if you have the sample sheet in a .csv file, you do not have to first load it into a data frame – the filename can be passed directly to ChIPQC.

### 2.2 Constructing a ChIPQCexperiment object

The main entry point for assessing experiments is ChIPQC. This takes a sample sheet and some other optional parameters and computes quality metrics for each of the individual samples. It does this using the *BiocParallel* package, which by default will run in parallel, using all available cores on your machine. For each unique sample (including controls), a ChIPQCsam ple object us constructed and added to a list. This list of samples forms the core of the ChIPQCexperiment object, which is constructed and returned by ChIPQC.

The parameters for ChIPQC include an annotation indicating what assembly of what genome is being analyzed. The chromosomes parameter specifies which chromosomes to examine when computing quality metrics. Restricting the chromosomes analysed can greatly increase the speed of the computations, as well as reducing the size of the resulting ChIPQCexperiment object. For this example we will limit the analysis to chromosome 18, the only chromosome for which data is included.

Other parameters include mapQCth, which represents a threshold for filtering on mapping (alignment) quality, which is set to 15 by default. An optional blacklist is a file or GRanges object with genomic intervals whereby reads in those regions will also be filtered out. Use of blacklists can be very important in ChIP-seq data processing ([?]), and should be applied for peak calling as well.

For this example, the main experiment is analyzed as follows:

```
> exampleExp = ChIPQC(samples,annotaiton="hg19")
> exampleExp
```

By default, the "hg19" annotation will use a default blacklist, and only the first chromosome (in this case "chr22") will be examined.

For the purposes of this vignette, we can load a pre-computed version of this object:

```
> data(example_QCexperiment)
> exampleExp
Samples: 6 : CTCF_1 CTCF_2 ... E2F1_1 E2F1_2
       Tissue Factor Replicate Peaks
CTCF_1
                CTCF
                              1 1118
          A549
CTCF_2
          A549
                CTCF
                              2
                                 648
cMYC_1
          A549
                              1
                                 253
                 cMYC
cMYC_2
          A549
                 cMYC
                              2
                                  159
                              1
                                 325
E2F1_1 HeLa-S3
                E2F1
E2F1_2 HeLa-S3
                E2F1
                              2
                                  249
       Reads Map% Filt%
                          Dup% ReadL FragL RelCC SSD RiP% RiBL%
CTCF_1 341055 100 26.3 16.600
                                  28
                                       131 2.740 2.53 31.20 1.33
CTCF_2 303856 100
                   28.4 7.320
                                   28
                                       131 2.690 1.43 12.80
                                                              0.00
cMYC_1 287462 100
                   31.0 13.600
                                  28
                                        97 0.347 1.47 6.59 1.78
cMYC_2 317537
               100
                   29.9 4.540
                                   28
                                       129 0.386 1.09
                                                       2.79
                                                              0.00
E2F1_1 223580
              100
                   31.7
                         1.010
                                   28
                                       101 0.701 1.29
                                                       7.80
                                                              2.00
E2F1_2 194919
              100 31.8 0.663
                                   28
                                       107 0.303 1.40 5.36 2.79
```

### 2.3 Quality metrics summary

Looking at the output, the first line indicates how many samples there are. The next section shows the metadata for the six ChIP samples (cell line, transcription factor ChIP'ed, replicate number, and number of peaks called).

The final section (also retrievable as a *matrix* by calling QCmetrics(exampleExp)) shows a summary of the main QC metrics. These include the total number of reads in the bam file for each sample, the percentage of these that were successfully mapped (aligned), and the percentage of reads having a mapping quality score less than or equal to the mapQCth (in this case, 15). The percentage of reads that map to the exact position in the genome as at least one other read is then reported. In this example, we can see substantial variance in the duplication rates, although none are extremely high. Indeed, good quality ChIPs for narrowly binding transcriptions factors are expected to have regions of very high enrichment, which will correctly include fragments that originate at the same location. Sequencing of such "duplicate" fragments is expected and biologically meaningful where the factor binds with high affinity. The extremely low duplication rates for the E2F1 samples are flags of potential problems with these ChIPs.

Next the read length, derived from the data in the bam files, is reported, along with the estimated mean fragment length. The fragment length is estimated by methods implemented in the *chipseq* package by systematically shifting the reads on each strand towards each other until the minimum genome coverage is achieved. The RelativeCC metric is calculated by comparing the maximum cross coverage peak (at the shift size corresponding to the fragment length) to the cross coverage at a shift size corresponding to the read length, with higher scores (generally 1 or greater) indicating good enrichment. In this example, the RelativeCC score for the all CTCF samplesis greater than 1, indicating these samples are of high quality.

The SSD score, as implemented in *htSeqTools*, is another indication of evidence of enrichment. It is computed by looking at the standard deviation of signal pile-up along the genome normalised to the total number of reads. An enriched sample typically has regions of significant pile-up so a higher SSD is more indicative of better enrichment. SSD scores are dependent on the degree of total genome wide signal pile-up and so are sensitive to regions of high signal found with Blacklisted regions as well as genuine ChIP enrichment. Here the first CTCF replicate shows the highest SSD score and so greatest enrichment for depth of signal.

The final two metrics report the percentage of reads in different regions of interest. The first reports the percentage of reads that overlap called peaks (also known as FRIP). This is another good indication of how "enriched" the sample is, and can be considered a "signal-to-noise" measure of what proportion of the library consists of fragments from binding sites vs. background reads. RiP% values for ChIPs around 5% or higher generally reflect successful enrichment.

The final measure reports the percentage of reads that overlapped blacklisted regions (RiBL). The signal from blacklisted has been shown to contribute to confound peak callers and fragment length estimation as well as contribute to the read length peak in cross coverage and cross coverage read length peak ([2]). The RiBL score then may act as a guide for the level of background signal in a ChIP or input and is found to be correlated with SSD in input samples and the read length cross coverage score in both input and ChIP samples ([2]).

# 2.4 Generating a summary QC report for experimental sample groups

The easiest way to get a visual overview of the QC metrics for the experiment is to generate a summary HTML report. This is accomplished by calling the ChIPQCreport method:

#### > ChIPQCreport(exampleExp)

By default, a sub directory will be created named ChIPQCreport (this is configurable using the reportFolder parameter), which will contain each of the plots. The main interactive HTML page (built using the R package *Nozzle*) is named ChIPQC.html by default. This can be loaded into a browser. This report is available at http://ChIPQC.starkhome.com/ Reports/exampleExp/ChIPQC.html.

The report is interactive in several ways. The individual sections can be expanded or compressed to focus in on specific areas of the assessment. Table can be sorted by clicking on the column labels. Individual plots can be enlarged or shrunk by clicking on them.

#### 2.4.1 Report: Overview

The first section include a textual overview of the report contents.

#### 2.4.2 Report: QC Summary table

The second section, *QC Summary*, contains a table with the key QC metrics for each sample, similar to the table output by <u>QCmetrics</u>.

#### 2.4.3 Report: QC Results plots

The third section contains the plots themselves, divided into three sections. Samples are grouped together using the experimental metadata; by default, samples sharing the same Tissue and Factor values are considered sample groups (this is controllable by the user using the facetBy parameter with ChIPQCreport() function).

The first section, *Mapping, Filtering, and Duplication Rate,* focuses on read mapping. begins with a table showing how the sequencing reads can be filtered: how many are mappable (align to a position in the genome) and how many reads pass the mapping quality filter and are not duplicates. Also included are the overall duplication rate for each sample, the percentage that pass the mapping quality filter, and the percentage of total reads that pass both the mapping quality filter and are not duplicates. Next is a plot showing the effect of blacklisting, with the proportion of reads that do and do not overlap with blacklisted regions. The final plot in this section uses the genomic annotation to show where reads map in terms of genomic features. This is represented as a heatmap showing the enrichment of reads compared to the background levels of the feature.

The second section, *ChIP Signal Distribution and Structure*, looks at the inherent "peakiness" of the samples. The first plot is a coverage histogram. The X-axis represent the read pileup height at a basepair position, and the Y-axis represents how many positions have this pileup height. This is on a log scale. A ChIP sample with good enrichment should have a reasonable "tail". This can be captured using the *SSD* metric, which is the standard deviation of this histogram (normalized to library depth so as to allow for sample-to-sample comparisons.) Samples with low enrichment, consisting of mostly background reads and genome wide low pile-up (such as in controls), should have lower SSD values than efficient ChIPs. Equivalence

between ChIP and control samples may reflect either low enrichment for ChIP or, when SSD is high for controls, the presense of large regions of high depth, aberrant signal in control samples and hence a flag for further blacklisting of genomic regions.

The other plot in this section show the cross-coverage scores (*CC Score*) of reads along the two strands at a range of successive shifts. There is usually a distinctive peak around the read length and a second peak should be at the expected fragment length. The read length peak is excluded from identification of the the shift with maximum cross coverage score, region of exclusion shown in red, to allow for the prediction of sensible fragment length scores. Failure to show a clear peak at the fragment length is indicative of a non-enriched sample. This can be seen in the *Relative CC* metric, which is the maximum CC Score (estimated fragment size) divided by the read length CC Score; low values tend to indicate a lack of enrichment.

In this example, both these plots suggest that the CTCF sample group is more enriched than the other two sets of samples, with the first replicate showing particularly high enrichment. The coverage histogram stays higher above the X-axis for the CTCF samples. Likewise, the CTCF samples show a clear peak at the fragment length in the cross-coverage plot, while the other samples do not show a clear peak other than at the read length.

The final set of plots, *Peak Profile and ChIP Enrichment*, are based on metric computed using the supplied peaks if available. The first plot shows average peak profiles, centered on the summit (point of highest pileup) for each peak. These profiles can vary depending on what type of mark is being studied – transcription factor, histone mark, or other DNAbinding protein such as a polymerase – but similar marks usually have a distinctive profile in successful ChIPs.

Next is a bar chart of the Reads in Peaks. ChIP samples with good enrichment will have a higher proportion of their reads overlapping called peaks. The final plot shows these data in a different way, offering a box plot of the distribution of how many reads are in each peak.

Finally, plots showing how the sample cluster are presented. The correlation heatmap is based on correlation values for all the peak scores for each sample. The other plot shows the first two principal component values for each sample. In this example, the replicates do cluster by replicate, which is a positive sign. the CTCF replicates are highly correlated and form their own cluster. The E2F1 replicates are also highly correlated with each other, and anti-correlated with the CTCF samples, probably due to their being from different cell lines.

#### 2.4.4 Report: Files and Versions

The final section of the report gives some version details as to the software run.

# 3 Example 2: ER binding in tamoxifen resistant data set

The next example demonstrates the use of *ChIPQC* with a larger experiment with good overall quality. This includes 11 ChIPs for the transcription factor ER in five cell lines, with at least two replicates for each ChIP, as well as 5 associated input DNA controls (one for each cell line). This data set is derived from [3]. The example only includes data from chromosome 18 (chr18) for time and space reasons. The actual aligned reads for the samples, in the form of BAM files, are not included with the package as they are approximately 200MB. The peaks (called using MACS [4]) are also not included. To run all aspects of the vignette, you will

need to download these separately – inquire of the package maintainers if you are interested in how to do this. Note that this data set is the same at that used for the vignette and example for the *DiffBind* package, which provides tools for further analyzing ChIP-seq experiments.

### 3.1 Experimental sample sheet

The first step is to construct a sample sheet describing the ChIP-seq experiment. This can be passed as a data frame, or saved in a csv file. The experiment can also be represented by a DBA object constructed using the *DiffBind* package, which accepts the same sample sheets as *ChIPQC*. A csv sample sheet has been included for the tamoxifen data set, and can be accessed as follow:

```
> library(ChIPQC)
> samples = read.csv(file.path(system.file("extdata", package="ChIPQC"),
+
                                             "tamoxifenQC.csv"))
> samples
   SampleID Tissue Factor Condition Treatment Replicate
1
     BT4741 BT474
                       ER Resistant Full-Media
                                                          1
     BT4742 BT474
                                                          2
2
                       ER Resistant Full-Media
3
      MCF71
              MCF7
                       ER Responsive Full-Media
                                                          1
4
      MCF72
                       ER Responsive Full-Media
              MCF7
                                                          2
5
      MCF73
              MCF7
                       ER Responsive Full-Media
                                                          3
6
      T47D1
              T47D
                       ER Responsive Full-Media
                                                          1
7
      T47D2
              T47D
                       ER Responsive Full-Media
                                                          2
8
      TAMR1
              MCF7
                       ER Resistant Full-Media
                                                          1
9
                                                          2
      TAMR2
              MCF7
                       ER Resistant Full-Media
10
      ZR751
              ZR75
                        ER Responsive Full-Media
                                                          1
11
      ZR752
              ZR75
                        ER Responsive Full-Media
                                                          2
                      bamReads ControlID
                                                           bamControl
1
   reads/Chr18_BT474_ER_1.bam
                                  BT474c reads/Chr18_BT474_input.bam
2
   reads/Chr18_BT474_ER_2.bam
                                  BT474c reads/Chr18_BT474_input.bam
3
    reads/Chr18_MCF7_ER_1.bam
                                   MCF7c
                                          reads/Chr18_MCF7_input.bam
4
    reads/Chr18_MCF7_ER_2.bam
                                   MCF7c
                                          reads/Chr18_MCF7_input.bam
5
    reads/Chr18_MCF7_ER_3.bam
                                   MCF7c
                                          reads/Chr18_MCF7_input.bam
6
    reads/Chr18_T47D_ER_1.bam
                                   T47Dc
                                                 reads/T47D_input.bam
7
    reads/Chr18_T47D_ER_2.bam
                                   T47Dc
                                                 reads/T47D_input.bam
8
    reads/Chr18_TAMR_ER_1.bam
                                   TAMRc
                                                 reads/TAMR_input.bam
9
          reads/TAMR_ER_2.bam
                                   TAMRc
                                                 reads/TAMR_input.bam
10
    reads/Chr18_ZR75_ER_1.bam
                                   ZR75c
                                                 reads/ZR75_input.bam
11
    reads/Chr18_ZR75_ER_2.bam
                                   ZR75c
                                                 reads/ZR75_input.bam
                      Peaks PeakCaller
1
   peaks/BT474_ER_1.bed.gz
                                   bed
2
   peaks/BT474_ER_2.bed.gz
                                   bed
3
    peaks/MCF7_ER_1.bed.gz
                                   bed
4
                                   bed
    peaks/MCF7_ER_2.bed.gz
5
    peaks/MCF7_ER_3.bed.gz
                                   bed
6
    peaks/T47D_ER_1.bed.gz
                                   bed
7
    peaks/T47D_ER_2.bed.gz
                                   bed
8
    peaks/TAMR_ER_1.bed.gz
                                   bed
9
    peaks/TAMR_ER_2.bed.gz
                                   bed
10
    peaks/ZR75_ER_1.bed.gz
                                   bed
```

#### 11 peaks/ZR75\_ER\_2.bed.gz bed

This sample sheet details the metadata for the experiment, including how controls are matches with ChIP samples. It also contains file paths to the aligned reads and previously called peaks. Note that in this case the paths are relative to the current working directory. If you download the associated data to run all step of the vignette, you will need to unzip the "reads" and "peaks" directories as sub directories of where the sample sheet is.

If you have the sample sheet in a .csv file, you do not have to first load it into a data frame – the filename can be passed directly to ChIPQC.

#### 3.2 Constructing a ChIPQCexperiment object

The main entry point for assessing experiments is ChIPQC. This takes a sample sheet and some other optional parameters and computes quality metrics for each of the individual samples. It does this using the *BiocParallel* package, which by default will run in parallel, using all available cores on your machine. For each unique sample (including controls), a ChIPQCsam ple object us constructed and added to a list. This list of samples forms the core of the ChIPQCexperiment object, which is constructed and returned by ChIPQC.

The parameters for ChIPQC include an annotation indicating what assembly of what genome is being analyzed. The chromosomes parameter specifies which chromosomes to examine when computing quality metrics. Restricting the chromosomes analysed can greatly increase the speed of the computations, as well as reducing the size of the resulting ChIPQCexperiment object. For this example we will limit the analysis to chromosome 18, the only chromosome for which data is included.

Other parameters include mapQCth, which represents a threshold for filtering on mapping (alignment) quality, which is set to 15 by default. An optional blacklist is a file or GRanges object with genomic intervals whereby reads in those regions will also be filtered out. Use of blacklists can be very important in ChIP-seq data processing ([?]), and should be applied for peak calling as well.

For this example, some other features of ChIPQC are used as well. By setting consensus=TRUE, a consensus peak set will be derived, so that even the control samples can be compared to determine their relative enrichment in these regions. With bCounts=TRUE, the peak scores will be computed by counting how many reads overlap these regions for every sample (include controls). This feature relies on the dba.count of the *DiffBind* package. By passing an extra parameter, summits=250, through to dba.count, the peaks will include 250 basepairs upstream and downstream of the peak summit.

Note that only the sample sheet actually needed to be specified; by default, if the annotation parameter is missing, it will default to "hg19", while a missing chromosomes parameter will result in the first chromosome that has peak being examined (in this case there re only peaks for "chr18", so that would be the default), and if the "hg19" is used, and the blacklist is missing, it will default to the one supplied with the package.

For the purposes of this vignette, we can load a pre-computed version of this object:

> data(tamoxifen_QC) > tamoxifen							
Samples: 16 : BT4741 BT4742 TAMRc ZR75c							
Tissue			Treatment	Renlic	ato I	Doaks	
BT4741 BT474		sistant Fu		Neptit	1	2844	
BT4742 BT474		sistant Fu			2	2844	
MCF71 MCF7		ponsive Fu			1	2844	
MCF72 MCF7		ponsive Fu			2	2844	
MCF73 MCF7		ponsive Fu			3	2844	
T47D1 T47D		ponsive Fu			1	2844	
T47D2 T47D		ponsive Fu			2	2844	
TAMR1 MCF7		sistant Fu			1	2844	
TAMR2 MCF7		sistant Fu			2	2844	
ZR751 ZR75	ER Res	ponsive Fu	ull-Media		1	2844	
ZR752 ZR75	ER Res	ponsive Fu	ull-Media		2	2844	
BT474c BT474	Control Re	sistant Fu	ull-Media		c1	2844	
MCF7c MCF7	Control Res	ponsive Fu	ull-Media		c2	2844	
T47Dc T47D	Control Res	ponsive Fu	ull-Media		c3	2844	
TAMRc MCF7	Control Re	sistant Fu	ull-Media		c4	2844	
ZR75c ZR75	Control Res	ponsive Fu	ull-Media		c5	2844	
Read	s Map% Filt%	Dup% Rea	adL FragL	RelCC	SSD	RiP%	RiBL%
BT4741 77635	3 100 15.8	8.42	28 153	2.110	1.88	14.90	1.75
BT4742 78241	9 100 15.1	10.30	28 147	2.010	1.63	13.80	1.68
MCF71 43899	4 100 20.9	21.30	28 134	2.300	2.52	26.50	1.70
MCF72 46570	0 100 20.8	4.84	28 155	2.090	1.65	16.10	2.17
MCF73 57727	3 100 19.1	10.30	28 153	2.460	2.18	21.80	1.83
T47D1 50749	2 100 21.1	7.86	28 153	1.520	1.59	9.62	2.24
T47D2 183176	6 100 19.3	9.56	28 162	1.530	2.30	5.70	2.26
TAMR1 74761	0 100 17.4	15.50	28 151	2.490	2.15	19.30	2.05
TAMR2 72860	1 100 18.4	5.84		1.850			1.56
ZR751 80442	7 100 12.0	15.80	28 158	2.950	3.52	30.70	1.39
ZR752 291854	9 100 11.6	23.80	28 160	3.140	4.45	20.70	1.22
BT474c 59801				0.202			1.72
MCF7c 48519				0.108			2.47
T47Dc 40039		31.60		0.268			8.56
TAMRc 77910				0.267			2.06
ZR75c 102398	7 100 26.2	20.10	36 220	0.573	5.36	1.50	5.35

### 3.3 Quality metrics summary

Looking at the output, the first line indicates how many samples there are (16, including 11 ChIPs and 5 controls). The next section shows the metadata for the 11 ChIP samples. All samples have the same number of peaks as a consensus set (derived from merging overlapping peaks, and keeping peaks that were identified in at least two samples) was used.

The final section (also retrievable as a *matrix* by calling QCmetrics(tamoxifen)) shows a summary of the main QC metrics. These include the total number of reads in the bam file for each sample, the percentage of these that were successfully mapped (aligned), and the percentage of reads that were filtered out due to having a mapping quality score less than or equal to the mapQCth (in this case, 15). The percentage of reads that map to the exact

position in the genome as at least one other read is then reported. In this example, we can see that three of the controls have very low duplication rates, with the T47D and ZR75 having duplication rates above 20%. This suggest it owuld be reasonable to filter duplciates from those two control. The ChIPs show substantial variance in their duplication rates, several above 10%. Higher duplication rates are expected in the ChIPs, as they should have regions of very high enrichment, which will correctly include fragments that originate at the same location. This is why we expect to see the controls samples exhibiting lower duplication rates (<5%) with the ChIP samples having higher rates (15%-20%), but not too high (>50%). A very high duplication rate (>50%) can result from a sample that has been biased by uneven PCR duplication of some fragments. This example shows some variance in duplication rates amongst the samples, including between replicates of the same condition. For example the third MCF7 replicate has a duplication rate double that of the first replicat (although neither rate is so high as to indicate a massive PCR duplication issue). The second replicate has a very low rate, which should draw our attention to other metrics that indicate peak enrichment, such as SSD, Retlative CC, and the peak profile plot.

Next the read length, derived from the data in the bam files, is reported, along with the estimated mean fragment length. The fragment length is estimated from the data by systematically shifting the reads on each strand towards each other until the highest degree of cross-coverage is obtained. The RelativeCC metric is calculated by comparing the maximal cross coverage peak (at the shift size corresponding to he fragment length) to the cross coverage at a shift size corresponding to the read length, with higher scores (generally 2 or greater) indicating good enrichment. In this example, the RelativeCC scores for the controls are very low, as expected, while the scores for the ChIPs are all greater than 1.5, with most above 2, indicating good enrichment and reliable fragment length estimation.

The SSD value is another indication of evidence of enrichment. It is computed by looking at the density of positions with different pileup values. An enriched sample typically has a range of pileup values, with a high standard deviation, so a higher SSD is more indicative of better enrichment. SSD values close to 1 are generally correlated of poorly enriched samples, while successful ChIPs can expect values around 1.5, with highly enriched samples having SSD values of 2 or higher. In this example, the **RelativeCC** scores for the ChIPs are greater than 1.5, with several above 2, while most of the controls are closer to 1, indicating background sampling without clear peaks where reads along the two strands converge in peaks. The two controls with high duplication rates also have high SSD scores, probably driven by false peaks driven by PCR duplication. Several of the plots below, such as the coverage histogram, cross-coverage, and peak profiles show that the signals for these controls do not show meaningful enrichment when duplicates are removed.

The final two metrics report the percentage of reads in different regions of interest. The first reports the percentage of reads that overlap called peaks (also known as FRIP). This is another good indication of how "enriched" the sample is, and can be considered a "signal-to-noise" measure of what proportion of the library consists of fragments from binding sites vs. background reads. RiP% values for ChIPs around 15% or higher generally reflect successful enrichment. In this case,where a consensus peak set was used (instead of peak sets unique to each sample), samples that originally had fewer peaks called can be expected to have lower RiP values (for example the T47D samples). Use of the consensus set does allow enrichment in the controls to be directly compared; in this example, the controls have much lower RiP% values, indicating that most of the the fragment sequenced in them were not located at peak sites.

The final measure report the percentage of reads that overlapped blacklisted regions (RiBL).

### 3.4 Generating a summary HTML report

As in the previous example, a summary HTML report can be generated for this example experiment by invoking ChIPQCreport. The only difference is that, in this case, the sample groups should be derived using the Tissue and Condition metadata values, as follows:

> ChIPQCreport(tamoxifen,facetBy=c("Tissue","Condition"))

The resulting report is not included in this vignette due to space limitations, but you are invited to generate it yourself and have a look. It is also available for examination at http://ChIPQC.starkhome.com/Reports/tamoxifen/ChIPQC.html.

### 3.5 Plotting QC metrics for experimental sample groups

As an alternative to generating a complete summary report, plotting methods are provided to generate specific plots, as follows:

#### 3.5.1 Plotting Coverage Histograms

The coverage histogram plot is generated as follows:

```
> plotCoverageHist(tamoxifen,facetBy=c("Tissue","Condition"))
```

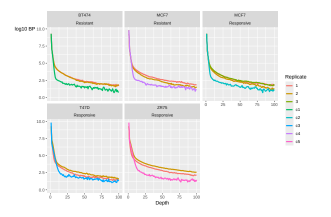


Figure 1: Example Coverage Histogram plot. Generated by plotCoverage
Hist(tamoxifen,facetBy=c("Tissue","Condition"))

As shown in Figure 1, this plots the distribution of pileup values at each basepair. The first thing to looks for is that the controls are "below" the ChIPs in the plots. Enriched ChIP samples will tail off less quickly than input controls consisting of background reads, indicating positions with high pileup values (ultimately corresponding to peaks).

#### 3.5.2 Plotting Cross-Coverage

A cross-coverage plot can be generated as follows:

```
> plotCC(tamoxifen,facetBy=c("Tissue","Condition"))
```

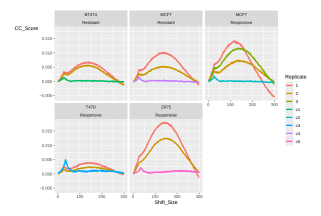
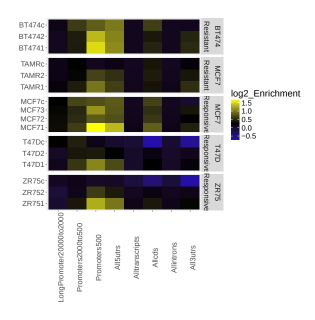


Figure 2: Example Cross-coverage plot. Generated by plotCC(tamoxifen,facetBy=c("Tissue", "Condition"))

As shown in Figure 2, this plots the cross-coverage values over a range of shift sizes. The region up to the read length is highlighted, and a small peak in cross-coverage is expected to be found here. Notice that the controls do not have another higher peak, while ChIP samples with good enrichment have another peak in cross-coverage value at the fragment length, as shifting the reads on both strand should increase coverage at peak sites.

#### 3.5.3 Plotting Relative Enrichment of reads in Genomic Intervals

A heatmap plot showing relative enrichment of reads around annotated genomic features can be generated as follows:



> plotRegi(tamoxifen, facetBy=c("Tissue", "Condition"))

**Figure 3:** Example Relative Enrichment of Genomic features heatmap plot. Generated by plotRegi(tamoxifen,facetBy=c("Tissue","Condition"))

As shown in Figure 3, samples with more reads than expected around certain features can be seen. Note that these genomic features tend to be enriched (and background genomic DNA depleted) in ChIP and control samples, as it is more likely that the chromatin is open near coding genes, and hence fragmentation can occur there more frequently.

#### 3.5.4 Plotting Peak Profiles

Plots of composite peak profile can be generated as follows:

> plotPeakProfile(tamoxifen, facetBy=c("Tissue", "Condition"))

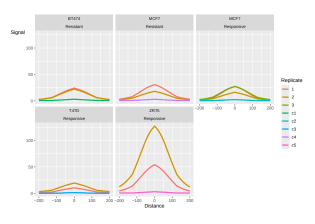


Figure 4: Example Peak Profile plot. Generated by plotPeakProfile(tamoxifen,facetBy=c("Tissue","Condition"))

As shown in Figure 4, each peak is centered on its summit (point of highest pileup after extending the reads to the calculated fragment length), and the pileup values at bases in a window upstream and downstream of the summits is computed and averaged for all peaks in the sample. Good ChIPs will show distinctive patterns of enrichment in these peaks, while associated controls will be relatively flat. The exact shape depends on the specific protein being ChIPed and the biological conditions. This example shows a profile for a transcription factor binding directly to the DNA.

#### 3.5.5 Plotting Reads overlapping Peaks and the Blacklist

Barplots of the relative number of reads that overlap peaks vs. background reads, as well and the proportion of reads overlapping blacklisted regions, can be generated as follows:

```
> plotRap(tamoxifen,facetBy=c("Tissue","Condition"))
```

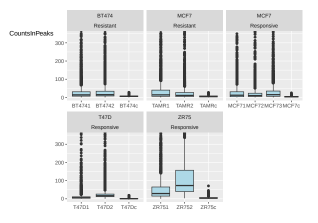
> plotFribl(tamoxifen,facetBy=c("Tissue","Condition"))

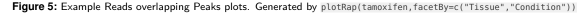
These plots are shown for the example data set in Figures 5. and 6.

#### 3.5.6 Plotting Sample Clustering

It can be useful when assessing the overall success of an experiment to see how the samples cluster. Two plots to aid in this can be generated as follows:

> plotCorHeatmap(tamoxifen,attributes=c("Tissue","Factor","Condition","Replicate"))





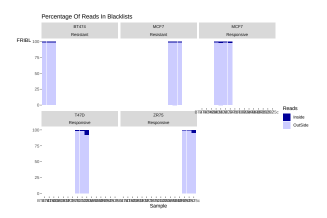


Figure 6: Example Reads overlapping Blacklist plots. Generated by plot
Fribl(tamoxifen,facetBy=c("Tissue","Condition"))

> plotPrincomp(tamoxifen,attributes=c("Tissue","Condition"))

Figure 7 shows the clustering in a correlation heatmap, where the scores for every peak are computed for each sample, and then scores for each pair of samples are used to calculate a correlation value. In this example, the controls form a distinct cluster, while the replicates each cluster together.

Additional insight can be gained from viewing the results of a principal components analysis using the peak scores. Figure 8 plots each sample along the first two components. The replicates, each plotted with the same color, mostly plot close to each other. The two controls, with very low peak scores (as evidenced by their low RiP% values), end up in almost the exact same point (with the yellow dot superimposed over a red dot) along the first component to the right, while the T47D samples, which are the least enriched of the ChIPs (lower RiP%), separate in the second component, occupying the top portion (green dots).

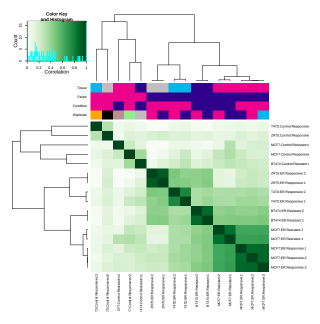


Figure 7: Example Correlation Heatmap with Clustering. Generated by plot CorHeatmap(tamoxifen,attributes=c("Tissue","Condition"))

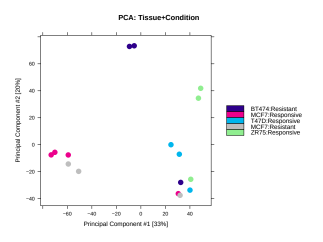


Figure 8: Example Principal Components plot. Generated by plotPrin comp(tamoxifen,attributes=c("Tissue","Condition"))

# 4 Example 3: Single sample assessment

While the vignette has focused on using *ChIPQC* to assess experiment consisting of multiple ChIP and control samples, it can also be used to examine samples individually without requiring the full experimental structure. This section briefly describes how to work with individual samples.

### 4.1 Constructing a ChIPQsample object and retrieving QC metrics

Given only a bam file (or a set of aligned reads in a *GappedAlignemnt* object), the QC metrics can be easily calculated (note the peak file is optional, but if missing, no peak-related metrics can be calculated):

> CTCF1 = ChIPQCsample("reads/SRR568129.bam", peaks="peaks/SRR568129\_chr22\_peaks.bed")

As the source data is not included with this vignette, the same effect can be simulated by retrieving the sample object we would have generated from the example experiment:

> CTCF1 = QCsample(exampleExp,1)

This returns a ChIPQCsample object, that can be examined as follows:

> CTCF1

	Proportion0	fCounts		
Peaks	0.4	2345591		
BlackList	0.0	1807147		
LongPromoter200001	to2000 0.3	9820917		
Promoters2000to500	9 0.0	5128542		
Promoters500	0.0	2985514		
All5utrs	0.0	1987424		
Alltranscripts	0.7	7778972		
Allcds	0.0	3430038		
Allintrons	0.7	1927332		
All3utrs	0.0	3180118		
GRanges object wit	th 1118 ranges and	2 metada	ta columns:	
seqnames	ranges	strand	Counts	bedRangesSummitsTemp
<rle></rle>	<iranges></iranges>	<rle>  </rle>	<integer></integer>	<numeric></numeric>
[1] chr22	16084142-16084556	*	29	16084371
[2] chr22	16156879-16157272	*	29	16157010
[3] chr22	16166244-16166805	*	464	16166577
[4] chr22	16185962-16186461	*	36	16186189
[5] chr22	16192141-16193098	*	65	16192792
[1114] chr22	51129810-51130449	*	105	51130147
[1115] chr22	51165140-51165330	*	14	51165225
[1116] chr22	51170467-51171033	*	133	51170738
[1117] chr22	51195483-51195886	*	24	51195608
[1118] chr22	51213448-51214119	*	164	51213729
seqinfo: 1 seque	ence from an unspe	cified ge	nome; no se	qlengths

This shows some summary statistics and the *GRanges* object containing the examined peaks.

A simpler way to see the key QC metrics is as follows:

<pre>&gt; QCmetrics(CTCF1)</pre>							
Reads 341055.00 RiP%	Map% 100.00 RiBL%	Filt% 26.30	Dup% 16.60	ReadL 28.00	FragL 131.00	RelCC 2.74	SSD 2.53

#### 31.20 1.33

tamThis returns a *vector* containing 10 metrics: the total number of reads in the bam file, the percentage that are aligned to the genome, the percentage that fall below the default mapping quality filter score of 15, the percentage of reads that align to exactly the same starting position as another read, the length of reads, the calculated fragment length, the relative cross-coverage score (based on the cross-coverage score when shifting by the fragment length vs. unshifted), the SSD (based on the standard deviation of the distribution of coverage pileup scores), and the percentage of reads that overlap peaks and blacklisted regions.

### 4.2 Plotting QC metrics for a sample

Plots can be generated for individual samples. For example, here is the cross-coverage plot:

#### > plotCC(CTCF1)

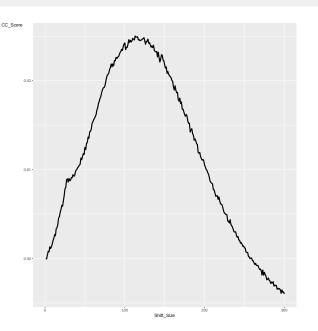


Figure 9: Sample-level cross-coverage plot. Generated by figSampCC(CTCF)

shown as Figure 9

### 4.3 Combining multiple samples into an Experiment

Samples that have been analyzed separately as *ChIPQCsample* objects may be combined into an *ChIPQCexperiment* object, without requiring the QC metrics to be re-computed, by combining them with a list and supplying a sample sheet with the experimental metadata.

For example, suppose the samples in the tamoxifen resistance example had been assessed separately and been combined in a list. We can get equivalent of this using the <code>QCsample</code> method on the experiment:

```
> sampleList = QCsample(tamoxifen)
> class(sampleList)
```

```
[1] "list"
> length(sampleList)
[1] 16
> names(sampleList)
 [1] "BT4741" "BT4742" "MCF71"
                                "MCF72" "MCF73"
                                                  "T47D1"
                                                           "T47D2"
                                                                    "TAMR1"
 [9] "TAMR2" "ZR751" "ZR752" "BT474c" "MCF7c"
                                                  "T47Dc"
                                                           "TAMRc"
                                                                    "7R75c"
> class(sampleList[[1]])
[1] "ChIPQCsample"
attr(,"package")
[1] "ChIPQC"
Suppose we also have constructed a matching samplesheet as a data.frame:
> sampleSheet = read.csv(file.path(system.file("extdata", package="ChIPQC"),
+
                                           "tamoxifenQC.csv"))
> sampleSheet
   SampleID Tissue Factor Condition Treatment Replicate
                       ER Resistant Full-Media
1
     BT4741 BT474
                                                        1
2
     BT4742 BT474
                       ER Resistant Full-Media
                                                        2
3
     MCF71
            MCF7
                       ER Responsive Full-Media
                                                        1
4
      MCF72
             MCF7
                       ER Responsive Full-Media
                                                        2
5
     MCF73
              MCF7
                       ER Responsive Full-Media
                                                        3
6
     T47D1
             T47D
                       ER Responsive Full-Media
                                                        1
7
     T47D2
             T47D
                       ER Responsive Full-Media
                                                        2
8
     TAMR1
              MCF7
                       ER Resistant Full-Media
                                                        1
9
      TAMR2
              MCF7
                       ER Resistant Full-Media
                                                        2
10
      ZR751
              ZR75
                       ER Responsive Full-Media
                                                        1
11
      ZR752
             ZR75
                       ER Responsive Full-Media
                                                        2
                     bamReads ControlID
                                                         bamControl
1 reads/Chr18_BT474_ER_1.bam
                                 BT474c reads/Chr18_BT474_input.bam
2
   reads/Chr18_BT474_ER_2.bam
                                 BT474c reads/Chr18_BT474_input.bam
3
    reads/Chr18_MCF7_ER_1.bam
                                  MCF7c reads/Chr18_MCF7_input.bam
4
    reads/Chr18_MCF7_ER_2.bam
                                  MCF7c
                                         reads/Chr18_MCF7_input.bam
5
    reads/Chr18_MCF7_ER_3.bam
                                  MCF7c
                                         reads/Chr18_MCF7_input.bam
6
                                  T47Dc
    reads/Chr18_T47D_ER_1.bam
                                               reads/T47D_input.bam
7
    reads/Chr18_T47D_ER_2.bam
                                  T47Dc
                                               reads/T47D_input.bam
8
    reads/Chr18_TAMR_ER_1.bam
                                  TAMRc
                                               reads/TAMR_input.bam
9
          reads/TAMR_ER_2.bam
                                  TAMRc
                                               reads/TAMR_input.bam
10
    reads/Chr18_ZR75_ER_1.bam
                                  ZR75c
                                               reads/ZR75_input.bam
11
    reads/Chr18_ZR75_ER_2.bam
                                  ZR75c
                                               reads/ZR75_input.bam
                     Peaks PeakCaller
1 peaks/BT474_ER_1.bed.gz
                                  bed
2 peaks/BT474_ER_2.bed.gz
                                  bed
3
                                  bed
    peaks/MCF7_ER_1.bed.gz
4
                                  bed
    peaks/MCF7_ER_2.bed.gz
5
                                  bed
    peaks/MCF7_ER_3.bed.gz
6
    peaks/T47D_ER_1.bed.gz
                                  bed
7
    peaks/T47D_ER_2.bed.gz
                                  bed
```

8	peaks/TAMR_ER_1.bed.gz	bed
9	peaks/TAMR_ER_2.bed.gz	bed
10	peaks/ZR75_ER_1.bed.gz	bed
11	peaks/ZR75_ER_2.bed.gz	bed

we can then re-construct the original *ChIPQCexperiment* object without having to re-compute the QC metrics:

```
> tamoxifen = ChIPQC(sampleSheet, samples=sampleList)
```

```
> tamoxifen
```

We could also supply a sample sheet that refers to only a subset of the *ChIPQCsample* objects to form a smaller experiment:

```
> BT474s = ChIPQC(sampleSheet[1:2,], samples=sampleList)
```

# 5 Customising QC plots

*ChIPQC* allows for some customisation of plots produced by the use of the additional parameters of facetBy, colourBy and lineBy. facetBy accepts a character vector corresponding to metadata column names and will group plots according to those supplied. In this example we choose to group by sample names to produce individual plots per sample (Figure 10).

```
> plotCC(exampleExp,facetBy="Sample")
```

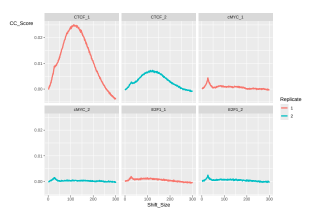


Figure 10: Cross-coverage plot grouped by Sample ID. Generated by plotCC(exampleExp,facetBy="Sample")

We can also use the colourBy and lineBy parameters for the plotCoverageHist(), plot PeakProfile() and plotCC() functions to alter line colours and types used. In Figure 11) we colour by factor and set the line type to be by Tissue.

> plotCC(exampleExp, facetBy="Sample", colourBy="Factor", lineBy="Tissue")

In addition to these options, extra metadata information can be passed to the plotting functions and this used by colourBy, lineBy and facetBy. We illustrate this by showing thr relationship between SSD and cross-coverage scores by supplying the QCmetrics as additional metadata using the addMetaData parameter.

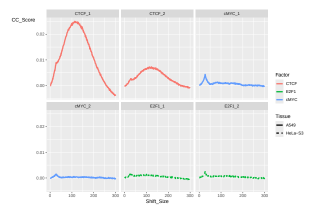


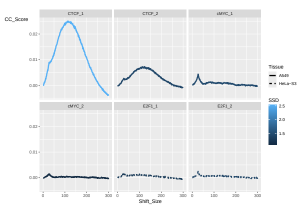
Figure 11: Cross-coverage plot grouped by Sample ID. Generated by
plotCC(exampleExp,facetBy="Sample",colourBy="Factor",lineBy="Tissue")

```
> extraMetadata <- data.frame(Sample = rownames(QCmetrics(exampleExp)),</pre>
```

- + FRiBL = QCmetrics(exampleExp)[,("RiBL%")],
- + SSD =QCmetrics(exampleExp)[,("SSD")]

```
+ )
```

> plotCC(exampleExp,facetBy="Sample",lineBy="Tissue",addMetaData=extraMetadata,colourBy="SSD")
>





*ChIPQC* uses the *ggplot2* framework to make plots and the resulting gg object can be further customised after the production of the plot. Additional *ggplot2* aesthetics can be added and this instance we change the colour scale used.

> plotCC(exampleExp,facetBy="Sample",lineBy="Tissue",addMetaData=extraMetadata,colourBy="SSD")
> + scale\_color\_gradient2(high="red",mid="black",low="white",midpoint=1.5)

# 6 Using ChIPQC and DiffBind together

*ChIPQC* and *DiffBind* are both packages that help manage and analyze ChIP-seq experiments, and are designed to be used together.

If you already have a project in *DiffBind*, the main constructor ChIPQC can accept a *DBA* object in place of the sample sheet.

Once a *ChIPQCexperiment* object has been constructed, it can be used in place of a *DBA* object in most calls to *DiffBind*. All plotting, counting, and analysis functions are available from *DiffBind*.

It is also possible to extract a *DBA* object from a *ChIPQCexperiment* object using the QCdba method. The resulting *DBA* object can be used in *DiffBind* without restriction, although neither it nor *DBA* objects based on it can be re-attached to the original *ChIPQCexperiment* object (although they can be used in lieu of a sample sheet when creating a new one.)

In a typical workflow, the first step would be to run a *ChIPQC* analysis before peak calling to assess library quality and establish what filtering should be done at the read level (mapping quality, duplicates, and blacklists). Next peaks would be called externally, and read into a new *ChIPQCexperiment* object to assess peak-based metrics, such as FRIP, peak profiles, and clustering.

At this point, *DiffBind* could be used to perform occupancy analysis, derive consensus peak sets, re-count reads to form a binding matrix, and set up contrasts to carry out full differential binding analyses using the *edgeR* and *DESeq2* packages, along with plotting and reporting functions.

### 7 Acknowledgements

We would like to acknowledge our co-authors, Wei Liu and Ines de Santiago, as well as Gordon Brown and everyone in the Bioinformatics Core at Cancer Research UK's Cambridge Institute at the University of Cambridge.

# 8 Session Info

> toLatex(sessionInfo())

- R version 4.4.0 RC (2024-04-16 r86468), x86\_64-pc-linux-gnu
- Locale: LC\_CTYPE=en\_US.UTF-8, LC\_NUMERIC=C, LC\_TIME=en\_GB, LC\_COLLATE=C, LC\_MONETARY=en\_US.UTF-8, LC\_MESSAGES=en\_US.UTF-8, LC\_PAPER=en\_US.UTF-8, LC\_NAME=C, LC\_ADDRESS=C, LC\_TELEPHONE=C, LC\_MEASUREMENT=en\_US.UTF-8, LC\_IDENTIFICATION=C
- Time zone: America/New\_York
- TZcode source: system (glibc)
- Running under: Ubuntu 22.04.4 LTS
- Matrix products: default
- BLAS: /home/biocbuild/bbs-3.20-bioc/R/lib/libRblas.so

- LAPACK: /usr/lib/x86\_64-linux-gnu/lapack/liblapack.so.3.10.0
- Base packages: base, datasets, grDevices, graphics, methods, stats, stats4, utils
- Other packages: Biobase 2.65.0, BiocGenerics 0.51.0, BiocParallel 1.39.0, ChIPQC 1.41.0, DiffBind 3.15.0, GenomeInfoDb 1.41.0, GenomicRanges 1.57.0, IRanges 2.39.0, MatrixGenerics 1.17.0, S4Vectors 0.43.0, SummarizedExperiment 1.35.0, ggplot2 3.5.1, matrixStats 1.3.0
- Loaded via a namespace (and not attached): AnnotationDbi 1.67.0, BSgenome 1.73.0, BiocIO 1.15.0, BiocManager 1.30.22, BiocStyle 2.33.0, Biostrings 2.73.0, DBI 1.2.2, DelayedArray 0.31.0, GenomeInfoDbData 1.2.12, GenomicAlignments 1.41.0, GenomicFeatures 1.57.0, GreyListChIP 1.37.0, KEGGREST 1.45.0, KernSmooth 2.23-22, MASS 7.3-60.2, Matrix 1.7-0, Nozzle.R1 1.1-1.1, R6 2.5.1, RColorBrewer 1.1-3, RCurl 1.98-1.14, RSQLite 2.3.6, Rcpp 1.0.12, Rsamtools 2.21.0, S4Arrays 1.5.0, SQUAREM 2021.1, ShortRead 1.63.0, SparseArray 1.5.0, TxDb.Celegans.UCSC.ce6.ensGene 3.2.2, TxDb.Dmelanogaster.UCSC.dm3.ensGene 3.2.2, TxDb.Hsapiens.UCSC.hg18.knownGene 3.2.2, TxDb.Hsapiens.UCSC.hg19.knownGene 3.2.2, TxDb.Mmusculus.UCSC.mm10.knownGene 3.10.0, TxDb.Mmusculus.UCSC.mm9.knownGene 3.2.2, TxDb.Rnorvegicus.UCSC.rn4.ensGene 3.2.2, UCSC.utils 1.1.0, XML 3.99-0.16.1, XVector 0.45.0, abind 1.4-5, amap 0.8-19, apeglm 1.27.0, ashr 2.2-63, bbmle 1.0.25.1, bdsmatrix 1.3-7, bit 4.0.5, bit64 4.0.5, bitops 1.0-7, blob 1.2.4, caTools 1.18.2, cachem 1.0.8, chipseq 1.55.0, cli 3.6.2, coda 0.19-4.1, codetools 0.2-20, colorspace 2.1-0, compiler 4.4.0, crayon 1.5.2, curl 5.2.1, deldir 2.0-4, digest 0.6.35, dplyr 1.1.4, emdbook 1.3.13, evaluate 0.23, fansi 1.0.6, farver 2.1.1, fastmap 1.1.1, generics 0.1.3, ggrepel 0.9.5, glue 1.7.0, gplots 3.1.3.1, grid 4.4.0, gtable 0.3.5, gtools 3.9.5, htmltools 0.5.8.1, htmlwidgets 1.6.4, httr 1.4.7, hwriter 1.3.2.1, interp 1.1-6, invgamma 1.1, irlba 2.3.5.1, jpeg 0.1-10, jsonlite 1.8.8, knitr 1.46, labeling 0.4.3, lattice 0.22-6, latticeExtra 0.6-30, lifecycle 1.0.4, limma 3.61.0, locfit 1.5-9.9, magrittr 2.0.3, memoise 2.0.1, mixsqp 0.3-54, munsell 0.5.1, mvtnorm 1.2-4, numDeriv 2016.8-1.1, parallel 4.4.0, pillar 1.9.0, pkgconfig 2.0.3, plyr 1.8.9, png 0.1-8, pwalign 1.1.0, reshape2 1.4.4, restfulr 0.0.15, rjson 0.2.21, rlang 1.1.3, rmarkdown 2.26, rtracklayer 1.65.0, scales 1.3.0, statmod 1.5.0, stringi 1.8.3, stringr 1.5.1, systemPipeR 2.11.0, tibble 3.2.1, tidyselect 1.2.1, tools 4.4.0, truncnorm 1.0-9, utf8 1.2.4, vctrs 0.6.5, withr 3.0.0, xfun 0.43, yaml 2.3.8, zlibbioc 1.51.0

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