RnBeads – Annotations

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1 Introduction

RnBeads is accompanied by four annotation packages: RnBeads.hg19, RnBeads.mm10, RnBeads.mm9, RnBeads.rn5. They contain genomic locus and Infinium probe annotation tables, as well as definition tables for tiling regions, genes, promoters and CpG islands. The RnBeads package contains routines for handling and updating these annotations. The vignette named RnBeads – Comprehensive DNA Methylation Analysis also contains examples that use functions presented here.

The data packages described above are hosted on the RnBeads website\(^1\). Note that at least one of the data packages should be installed prior to installing RnBeads.

2 Data Extraction and Processing

2.1 Sites

Currently, every data package examines cytosines in the context of CpG and contains an annotation table of all CpGs in the respective genome. CpG density and GC content are also computed for the neighborhood of length 100 base pairs centered on each locus. The total number of dinucleotides annotated in HG19 is 28,217,009 represented both on the forward and reverse DNA strands.

2.2 HumanMethylation450 Probes

The Infinium HumanMethylation450 BeadChip interrogates over 470,000 loci in the human genome. Control and methylation probe annotation is obtained from online sources, validated for consistency and enriched with locations for the SNP-associated loci obtained from dbSNP. In addition, the annotation includes information about CpG density and GC content of the sequence neighborhood around every probe target as described in the section above.

2.3 dbSNP

Single-Nucleotide Polymorphisms (SNPs) in the human genome are downloaded from the respective FTP directory of dbSNP\(^2\). RnBeads uses the provided VCF files; for mouse and rat these are one file per chromosome. The following table shows the references in dbSNP used for

\(^1\)rnbeads.mpi-inf.mpg.de
the supported assemblies.

<table>
<thead>
<tr>
<th>Package</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>RnBeads.hg19</td>
<td>GRCh37.p10</td>
</tr>
<tr>
<td>RnBeads.mm10</td>
<td>GCF_000001635.21</td>
</tr>
<tr>
<td>RnBeads.rn5</td>
<td>GCF_000001895.4</td>
</tr>
</tbody>
</table>

The data package RnBeads.mm9 does not include SNP-related information. Only SNPs that meet all of the following criteria are considered:

1. **assembly** is GRCh37.p5, reference;
2. **chrom** is the expected chromosome;
3. **allele origin** is not somatic;
4. (for HG19) major allele frequency is at most 0.95.

These SNP positions are used to enrich the annotations for CpG dinucleotides and Infinium probes.

### 2.4 Regions

Every data package defines the following sets of regions for the dedicated assembly:

**Tiling regions** Tiling regions with a window size of 5 kilobases are defined over the whole genome.

**Genes and promoters** Ensembl\(^5\) gene definitions are downloaded using the **biomaRt** package. A promoter is defined as the region spanning 1,500 bases upstream and 500 bases downstream of the transcription start site of the corresponding gene.

**CpG islands** The CpG island track is downloaded from the dedicated FTP directory of the UCSC Genome Browser\(^4\).

CpG density and GC content are computed for all region types listed above.

### 3 Annotations

#### 3.1 Genome Assemblies

Every annotation table and other data structure is specific to a genome assembly. The list of supported assemblies can be obtained by the calling the function `rnb.get.assemblies`, as shown in the example below.

```r
> rnb.get.assemblies()
```

```
[1] "hg19"  "hg38"  "mm10"  "mm9"  "rn5"
```

\(^5\)www.ensembl.org/
\(^4\)genome.ucsc.edu/
Table 1: Columns in the HumanMethylation450 control probe annotation table.

3.2 Built-in Annotation Tables

Control probe information for the Illumina 450K array is stored (and can be extracted) as a data.frame in which every row corresponds to a control probe. Every other annotation table is represented by a dedicated GRangesList object, that is, a list of consistent GRanges objects, one per chromosome. Every instance of GRanges defines the genomic positions of the corresponding sites, probes or regions. Identifiers, if present, can be obtained using the names method. Strand information is also included when applicable. Any additional annotation is stored as metadata. Please refer to the documentations of the GenomicRanges package and the GRanges class for more details.

In an R session, every annotation object can be accessed using the function rnb.get.annotation. The code snippet below shows how the control probe annotations can be extracted. Table 1 briefly describes the characteristics of these probes.

```r
> control.annotation <- rnb.get.annotation("controls450")
> head(control.annotation)
```

In a similar manner, the command below extracts the annotation of HumanMethylation450 probes. The metadata structure for the methylation and SNP-associated probes is presented in Table 2.

```r
> probe.annotation <- rnb.get.annotation("probes450")
> probe.annotation
```

Every data package integrates various region types with the CpG sites (and Infinium probes for HG19) presented in the previous section. Obtaining a region annotation information is very similar to accessing site or probe annotation. The code snippet below extracts the available gene definitions. Table 3 provides information about the metadata of these regions.

```r
> gene.annotation <- rnb.get.annotation("genes")
> attr(gene.annotation, "version")
> gene.annotation
```
<table>
<thead>
<tr>
<th>No.</th>
<th>Column Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Design</td>
<td>Probe design type.</td>
</tr>
<tr>
<td>2</td>
<td>Color</td>
<td>Color channel.</td>
</tr>
<tr>
<td>3</td>
<td>Context</td>
<td>Probe context.</td>
</tr>
<tr>
<td>4</td>
<td>Random</td>
<td>Flag indicating if the probe’s location was randomly chosen.</td>
</tr>
<tr>
<td>5</td>
<td>HumanMethylation27</td>
<td>Flag indicating if the probe is also covered by HumanMethylation27k assay.</td>
</tr>
<tr>
<td>6</td>
<td>Mismatches A</td>
<td>Number of base mismatches between the provided and expected probe sequence.</td>
</tr>
<tr>
<td>7</td>
<td>Mismatches B</td>
<td>Number of base mismatches between the provided and expected probe sequence.</td>
</tr>
<tr>
<td>8</td>
<td>CGI Relation</td>
<td>Relation to a CpG island.</td>
</tr>
<tr>
<td>9</td>
<td>CpG</td>
<td>Number of CpG dinucleotides in the sequence neighborhood of the target.</td>
</tr>
<tr>
<td>10</td>
<td>GC</td>
<td>Percentage of C and G bases in the sequence neighborhood of the target.</td>
</tr>
<tr>
<td>11</td>
<td>SNPs 3</td>
<td>Number of SNPs that overlap with the last 3 bases of a probe’s target sequence.</td>
</tr>
<tr>
<td>12</td>
<td>SNPs 5</td>
<td>Number of SNPs that overlap with the last 5 bases of a probe’s target sequence.</td>
</tr>
<tr>
<td>13</td>
<td>SNPs Full</td>
<td>Number of SNPs that overlap with the probe’s sequence.</td>
</tr>
</tbody>
</table>

Table 2: Metadata accompanying the HumanMethylation450 probe definitions.

<table>
<thead>
<tr>
<th>No.</th>
<th>Column Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>symbol</td>
<td>Gene symbols associated with this region.</td>
</tr>
<tr>
<td>2</td>
<td>entrezID</td>
<td>Entrez gene identifiers associated with this region.</td>
</tr>
<tr>
<td>3</td>
<td>CpG</td>
<td>Number of CpG dinucleotides in the region.</td>
</tr>
<tr>
<td>4</td>
<td>GC</td>
<td>Total number of C and G bases in the region.</td>
</tr>
</tbody>
</table>

Table 3: Metadata accompanying the Ensembl gene and promoter definitions.
The function `rnb.region.types` returns all supported region definitions for a given genome assembly. These currently include "cpgislands", "genes", "promoters", "tiling".

The `GRangesList` objects are well optimized for storage and fast access of genomic elements. However, exporting such objects to tables in human-readable form is not trivial. The function `rnb.annotation2data.frame` provides an easy way to convert a probe or region annotation to a single table. The code snippet below exemplifies this function by saving the gene promoter definitions to a comma-separated value (CSV) file named `promoters.csv`.

```r
> promoters <- rnb.annotation2data.frame(rnb.get.annotation("promoters"))
> write.csv(promoters, file = "promoters.csv", row.names = FALSE)
```

### 3.3 Custom Annotations

Custom region annotations can be included using the function `rnb.set.annotation`. The information is either loaded from a BED file, or it must be presented as a `data.frame` containing at least the following three columns - chromosome, start and end. Note that the included genomic regions remain available in the current R session only.

The vignette named RnBeads – Comprehensive DNA Methylation Analysis contains a detailed example for including custom annotations to an analysis.

### 4 Data Sources

The data packages accompanying RnBeads obtained data for the annotation tables from the following sources:

**Other Bioconductor packages** Methylation and control probe annotation is obtained from the package `FDb.InfiniumMethylation.hg19`. In addition, RnBeads uses the packages:

- `BSgenome.Hsapiens.UCSC.hg19`
- `BSgenome.Mmusculus.UCSC.mm9`
- `BSgenome.Mmusculus.UCSC.mm10`
- `BSgenome.Rnorvegicus.UCSC.rn5`

which validate the probe annotations and to enrich all annotation tables with sequence-based properties.

**CRAN packages** Additional control probe information is obtained from the package HumMeth27QCReport.

**dbSNP** Information on SNPs in the human genome is downloaded as BED files from the FTP site of dbSNP.

**Gene Expression Omnibus (GEO)** Methylation and control probe annotation is obtained from GEO record GPL13534 and validated for consistency with the other sources.

**Biomart** Ensembl gene definitions (Ensembl versions 73 (for hg19, mm10 and rn5), and 67 (for mm9) are downloaded using the biomaRt package.

**UCSC Genome Browser** The CpG island tracks are downloaded from the FTP site of the UCSC Genome Browser.