# genomationData: R package with high throughput genomic data

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### 1 Introduction

genomationData is a Bioconductor-compliant R package that contains raw and processed data from high throughput genomics experiments. The data is originally intended for demo of the Bioconductor package genomation. genomationData holds data from CHiP Seq and Bisulphite sequencing experiments produced by Encode and Epigenetics Roadmap. All datasets come from human H1 embryonic stem cell line and are mapped to the hg19/GRCh37 version of the genome.

## 2 Getting started

To load the *genomationData* package into your R environment type:

```
library(genomationData)
```

## 2.1 Sample description

To list the available files type:

```
list.files(system.file("extdata", package = "genomationData"))
```

The package currently holds 5 CHiP Sequencing and one Bisulfite sequencing samples. Files with the .bam extension contain raw CHiP seq reads, while \*.broadPeak and \*.narrow-Peak represent processed peaks. Due to the large sizes of the samples, the \*.bam files have been restricted to human chromosome 21 (chr21).

H1.Bisulfite-Seq.combined.chr21.bedGraph.gz' contains methylation calls for each CpG dinucleotide on human chromosome 21. The original file was converted from wig into bed-Graph format using the standard UCSC applications.

To see complete information about the files, take a look into SamplesInfo.txt

#### 3 Session Info

```
sessionInfo()
## R version 3.2.0 (2015-04-16)
## Platform: x86_64-unknown-linux-gnu (64-bit)
## Running under: Ubuntu 14.04.2 LTS
##
## locale:
   [1] LC_CTYPE=en_US.UTF-8
                                   LC_NUMERIC=C
##
    [3] LC_TIME=en_US.UTF-8
                                   LC_COLLATE=C
    [5] LC_MONETARY=en_US.UTF-8
##
                                   LC_MESSAGES=en_US.UTF-8
   [7] LC_PAPER=en_US.UTF-8
                                   LC_NAME=C
   [9] LC_ADDRESS=C
                                   LC_TELEPHONE=C
## [11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
##
## attached base packages:
## [1] stats
                 graphics grDevices utils
                                                datasets methods
                                                                    base
##
## loaded via a namespace (and not attached):
## [1] formatR_1.1
                     tools_3.2.0
                                   highr_0.4.1
                                                  knitr_1.9
                                                                stringr_0.6.2
## [6] evaluate_0.6
```