## Package 'curatedAdipoChIP'

April 13, 2022

Type Package

```
Title A Curated ChIP-Seq Dataset of MDI-induced Differentiated
     Adipocytes (3T3-L1)
Version 1.10.0
Year 2019
Description A curated dataset of publicly available ChIP-sequencing of
     transcription factors, chromatin remodelers and histone modifications in the
     3T3-L1 pre-adipocyte cell line. The package document the data collection,
     pre-processing and processing of the data. In addition to the documentation,
     the package contains the scripts that was used to generated the data.
License GPL-3
URL https://github.com/MahShaaban/curatedAdipoChIP
BugReports https://github.com/MahShaaban/curatedAdipoChIP/issues
Encoding UTF-8
RoxygenNote 6.1.1
Depends R (>= 3.6), SummarizedExperiment, ExperimentHub
Suggests knitr, rmarkdown, GenomicFeatures, ChIPseeker, AnnotationDbi,
     S4Vectors, DESeq2, fastqcr, devtools, testthat, readr, dplyr,
     tidyr, ggplot2
VignetteBuilder knitr
biocViews ExperimentData, ExperimentHub, GEO, ChIPSeqData,
     SequencingData
git_url https://git.bioconductor.org/packages/curatedAdipoChIP
git_branch RELEASE_3_14
git_last_commit 9a6f121
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#### **Description**

A Curated ChIP-Seq Dataset of MDI-induced Differentiated Adipocytes (3T3-L1)

#### **Details**

A curated dataset of publicly available ChIP-sequencing of transcription factors, chromatin remodelers and histone modifications in the 3T3-L1 pre-adipocyte cell line. The package document the data collection, pre-processing and processing of the data. In addition to the documentation, the package contains the scripts that was used to generated the data.

The dataset can be accessed through the ExperimentHub as a RangedSummarizedExperiment object contains:

assay The read counts matrix.

**colData** The phenotype data and quality control data of the samples.

rowRanges The feature data and annotation of the peaks.

**metadata** The study level metadata which contains one object called studies. This is a data. frame of bibliography information of the studies from which the samples were collected.

#### **Examples**

```
## Not run:
# load the data object
library(ExperimentHub)

# query package resources on ExperimentHub
eh <- ExperimentHub()
query(eh, "curatedAdipoChIP")

# load data from ExperimentHub
peak_counts <- query(eh, "curatedAdipoChIP")[[1]]

# print object
peak_counts
## End(Not run)</pre>
```

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