Package 'ObMiTi'

October 31, 2024

Type Package Title Ob/ob Mice Data on Normal and High Fat Diet Version 1.14.0 Year 2021 Description The package provide RNA-seq count for 2 strains of mus musclus; Wild type and Ob/Ob. Each strain was divided into two groups, and each group received either chow diet or high fat diet. RNA expression was measured after 20 weeks in 7 tissues. License GPL-3 URL https://github.com/OmarElAshkar/ObMiTi BugReports https://github.com/OmarElAshkar/ObMiTi/issues **Encoding** UTF-8 RoxygenNote 7.1.1 Depends R (>= 4.1), SummarizedExperiment, ExperimentHub Suggests knitr, rmarkdown, BiocManager, GenomicFeatures, S4Vectors, devtools, testthat VignetteBuilder knitr biocViews ExperimentHub, GEO, RNASeqData git_url https://git.bioconductor.org/packages/ObMiTi git_branch RELEASE_3_20 git_last_commit 654990d git_last_commit_date 2024-10-29 Repository Bioconductor 3.20 Date/Publication 2024-10-31 Author Omar Elashkar [aut, cre] (<https://orcid.org/0000-0002-5505-778X>), Mahmoud Ahmed [aut] (<https://orcid.org/0000-0002-4377-6541>) Maintainer Omar Elashkar <omar.i.elashkar@gmail.com> **Contents**

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ObMiTi

Description

Title: Ob/ob Mice Data on Normal and High Fat Diet The package provide RNA-seq count for 2 strains of mus musclus; Wild type and Ob/Ob. Each strain was divided into two groups, and each group received either chow diet or high fat diet. RNA expression was measured after 12 weeks in 7 tissues.

Details

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

assay The read counts matrix.

colData The phenotype data of the samples

rowRanges The feature data and annotation of the peaks.

metadata extra details about the sample and associated phenotype studies. This is a data.frame of bibliography information of the studies from which the samples were collected for.

Examples

```
# load the data object
library(ExperimentHub)
# query package resources on ExperimentHub
eh <- ExperimentHub()
query(eh, "ObMiTi")
```

```
# load data from ExperimentHub
ob_counts <- query(eh, "ObMiTi")[[1]]</pre>
```

print object
ob_counts

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