## Package 'ObMiTi'

December 19, 2024

Type Package

Title Ob/ob Mice Data on Normal and High Fat Diet

**Version** 1.15.0

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 devel

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Author Omar Elashkar [aut, cre] (ORCID: <https://orcid.org/0000-0002-5505-778X>), Mahmoud Ahmed [aut] (ORCID: <https://orcid.org/0000-0002-4377-6541>)

Maintainer Omar Elashkar <omar.i.elashkar@gmail.com>

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ObMiTi package

#### 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")</pre>

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

```
# print object
ob_counts
```

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