## Package 'curatedAdipoRNA'

March 28, 2024

Title A Curated RNA-Seq Dataset of MDI-induced Differentiated

Type Package

```
Adipocytes (3T3-L1)
Version 1.18.0
Year 2019
Description A curated dataset of RNA-Seq samples. The samples are MDI-induced
     pre-phagocytes (3T3-L1) at different time points/stage of differentiation.
     The package document the data collection, pre-processing and processing. 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/curatedAdipoRNA
BugReports https://github.com/MahShaaban/curatedAdipoRNA/issues
Encoding UTF-8
RoxygenNote 6.1.1
LazyData TRUE
Depends R (>= 3.6), SummarizedExperiment
Suggests knitr, rmarkdown, DESeq2, fastqcr, devtools, testthat, readr,
     dplyr, tidyr, ggplot2, S4Vectors
VignetteBuilder knitr
biocViews ExperimentData, GEO, RNASeqData, SequencingData
git_url https://git.bioconductor.org/packages/curatedAdipoRNA
git_branch RELEASE_3_18
git_last_commit a9918d6
git_last_commit_date 2023-10-24
Repository Bioconductor 3.18
Date/Publication 2024-03-28
Author Mahmoud Ahmed [aut, cre] (<a href="https://orcid.org/0000-0002-4377-6541">https://orcid.org/0000-0002-4377-6541</a>)
Maintainer Mahmoud Ahmed <mahmoud.s.fahmy@students.kasralainy.edu.eg>
```

2 adipo\_counts

### **R** topics documented:

	adipo_cour curatedAdi																
Index																	4
adipo	o_counts	Gene	cour	its in	ı diff	fere	ntic	ıtin	g a	dipe	осуі	tes					

#### Description

Gene counts in differentiating adipocytes

#### Usage

adipo\_counts

#### **Format**

A RangedSummarizedExperiment object contains:

```
assay The gene counts matrix.
```

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

rowRanges The feature data at gene level.

**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**

```
# load the data object
data('adipo_counts')
# print the object
adipo_counts
```

curatedAdipoRNA 3

curatedAdipoRNA	curatedAdipoRNA package	

#### Description

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

#### **Details**

A curated dataset of RNA-Seq samples. The samples are MDI-induced pre-phagocytes (3T3-L1) at different time points/stage of differentiation. The package document the data collection, pre-processing and processing. In addition to the documentation, the package contains the scripts that was used to generated the data. The datasets and the pipeline used to process it are documented in adipo\_counts and the package vignette.

# **Index**