# Package 'lumiBarnes'

June 5, 2025

00110 0, 2020
Type Package
Title Barnes Benchmark Illumina Tissues Titration Data
Version 1.49.0
Date 2010-10-12
Author Pan Du
Maintainer Pan Du <dupan@northwestern.edu></dupan@northwestern.edu>
<b>Description</b> The Barnes benchmark dataset can be used to evaluate the algorithms for Illumina microarrays. It measured a titration series of two human tissues, blood and placenta, and includes six samples with the titration ratio of blood and placenta as 100:0, 95:5, 75:25, 50:50, 25:75 and 0:100. The samples were hybridized on Human-Ref-8 BeadChip (Illumina, Inc) in duplicate. The data is loaded as an LumiBatch Object (see documents in the lumi package).
License LGPL
<b>Depends</b> R (>= 2.0), Biobase (>= 2.5.5), lumi (>= 1.1.0)
biocViews ExperimentData, Tissue, MicroarrayData, ChipOnChipData
git_url https://git.bioconductor.org/packages/lumiBarnes
git_branch devel
git_last_commit 6725a43
git_last_commit_date 2025-04-15
Repository Bioconductor 3.22
Date/Publication 2025-06-05
Contents
lumiBarnes
Index

2 lumiBarnes

lumiBarnes

Barnes Benchmark Illumina Tissues Titration Data

### **Description**

The Barnes data set measured a titration series of two human tissues, blood and placenta. It includes six samples with the titration ratio of blood and placenta as 100:0, 95:5, 75:25, 50:50, 25:75 and 0:100. The samples were hybridized on HumanRef-8 BeadChip (Illumina, Inc) in duplicate. See (Barnes, et al., 2005) for details. The data is saved as a LumiBatch object and should be use together with lumi package.

Because the Barnes data utilized the pre-released version of HumanRef-8 version 1 BeadChip, some probes on the chip do not exist in the public released HumanRef-8 version 1 BeadChip. For annotation consistence, these probes was removed in the lumiBarnes package. For the interested users, the raw data can be downloaded from the paper companion website: http://www.bioinformatics.ubc.ca/pavlidis/lab/platform0

### Usage

data(lumiBarnes)

#### **Format**

lumiBarnes is a LumiBatch-class object.

#### Source

Barnes, M., Freudenberg, J., Thompson, S., Aronow, B. and Pavlidis, P. (2005) Ex-perimental comparison and cross-validation of the Affymetrix and Illumina gene expression analysis platforms, Nucleic Acids Res, 33, 5914-5923.

## **Examples**

data(lumiBarnes)
lumiBarnes

# Index

 $*\ datasets$ 

lumiBarnes, 2

 ${\tt lumiBarnes}, {\tt 2}$