Package: TPACData (via r-universe)

October 18, 2024

Type Package
Title Human Protein Atlas Data for Tissue-Adjusted Pathway Analysis of Cancer (TPAC)
Version 0.1.0
Description Contains summary data on gene expression in normal human tissues from the Human Protein Atlas for use with the Tissue-Adjusted Pathway Analysis of cancer (TPAC) method. Frost, H. Robert (2023) ``Tissue-adjusted pathway analysis of cancer (TPAC)" <doi:10.1101 2022.03.17.484779="">.</doi:10.1101>
Depends R (>= 3.6.0)
Imports data.table
License GPL (>= 2)
Encoding UTF-8
LazyData true
LazyDataCompression bzip2
NeedsCompilation no
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Repository CRAN
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hpa.data

Gene expression data for normal human tissues from the Human Protein Atlas (HPA). This data was specially processed by the HPA group as FPKM values (file "HPA.normal.FPKM.GDCpipeline.csv") using a pipeline similar to that employed by GDC for the TCGA RNA-seq data.

Description

A data.table with 1,088,694 rows and 4 columns (Gene, Tissue, FPKM, TissueSpecificity):

• Gene: Ensembl gene ID

• Tissue: Human tissue type

- FPKM: Mean expression of that gene in units FPKM+1
- TissueSpecificity: The fold-change in expression between mean expression in that tissue and mean in all 18 tissues.

Usage

hpa.data

Format

A data.table with 1,088,694 rows and 4 columns (Gene, Tissue, FPKM, TissueSpecificity)

Source

Human Protein Atlas

References

• Ulhen et al. (2017). A pathology atlas of the human cancer transcriptome. Science. doi: 10.1126/science.aan2507

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