Package 'ChIPXpressData'

March 25, 2013

Type Package

Title ChIPXpress Pre-built Databases

Version 0.99.1

Date 2012-07-24

Author George Wu

Maintainer George Wu <gewu@jhsph.edu>

Description Contains pre-built mouse (GPL1261) and human (GPL570) database of gene expression profiles to be used for ChIPXpress ranking.

License GPL (>=2)

Depends bigmemory

biocViews AffymetrixChIP, Homo_sapiens, Mus_musculus

R topics documented:

| ChIPXpressData-package | 1 |
|------------------------|---|
| DB_GPL1261.bigmemory | 2 |
| DB_GPL570.bigmemory | 3 |
| | |

5

Index

ChIPX pressData-package

ChIPXpress Gene Expression Databases

Description

Pre-built databases of gene expression profiles for ChIPXpress analysis in big.matrix format. DB_GPL1261 contains mouse data. DB_GPL570 contains human data.

Details

| ChIPXpressData |
|----------------|
| Package |
| 0.99.0 |
| 2012-07-24 |
| GPL 2.0 |
| |

Author(s)

George Wu Maintainer: George Wu <gewu@jhsph.edu>

References

McCall M.N., Bolstad B.M., and Irizarry R.A. (2010) Frozen robust multiarray analy-sis (fRMA). Biostatistics 11, 242-253.

Barrett T., et al. (2007) NCBI GEO: mining tens of millions of expression profiles - database and tools update. Nucl. Acids Res. 35, D760-D765.

DB GPL1261.bigmemory

Database of gene expression profiles from the Affymetrix Mouse 430 2.0 Array (GPL1261) in big.matrix format

Description

The data set contains 9634 mouse profiles downloaded from NCBI GEO, processed using fRMA, and normalized. It is in big.matrix format.

Format

The format is: Formal class 'big.matrix' [package "bigmemory"] with 1 slots ..@ address:<externalptr>

Details

The database is formatted as a big.matrix for more efficient loading into memory. It is stored in DB_GPL1261.bigmemory and the corresponding description file is DB_GPL1261.bigmemory.desc. To utilize the big.matrix format, it requires the package bigmemory to be loaded. See the bigmemory package for more information.

The database contains 20757 rows and 9643 columns. Each row represents the expression vector for each gene and each column represents the gene expression measurements for a sample from NCBI GEO obtained using the GPL1261 platform. Each gene will match uniquely to a single probe ID; only the probe with the highest variance in the compendium apriori to normalization are retained as the representative measurement for each gene.

Source

www.ncbi.nlm.nih.gov/geo/

References

McCall M.N., Bolstad B.M., and Irizarry R.A. (2010) Frozen robust multiarray analysis (fRMA). Biostatistics 11, 242-253.

Barrett T., et al. (2007) NCBI GEO: mining tens of millions of expression profiles - database and tools update. Nucl. Acids Res. 35, D760-D765.

Examples

Load the GPL1261 database library(bigmemory) path <- system.file("extdata",package="ChIPXpressData") DB_GPL1261 <- attach.big.matrix("DB_GPL1261.bigmemory.desc",path=path) ## DB_GPL1261 is then ready for input into the ChIPXpress function.

To see info about the database matrix describe(DB GPL1261)

DB GPL570.bigmemory

Database of gene expression profiles from the Affymetrix Human U133 Plus 2.0 array (GPL570) in big.matrix format

Description

The data set contains 18257 human profiles downloaded from NCBI GEO, processed using fRMA, and normalized. It is in big.matrix format.

Format

The format is: Formal class 'big.matrix' [package "bigmemory"] with 1 slots ..@ address:<externalptr>

Details

The database is formatted as a big.matrix for more efficient loading into memory. It is stored in DB_GPL570.bigmemory and the corresponding description file is DB_GPL570.bigmemory.desc. To utilize the big.matrix format, it requires the package bigmemory to be loaded. See the bigmemory package for more information.

The database contains 19798 rows and 18257 columns. Each row represents the expression vector for each gene and each column represents the gene expression measurements for a sample from NCBI GEO obtained using the GPL570 platform. Each gene will match uniquely to a single probe ID; only the probe with the highest variance in the compendium apriori to normalization are retained as the representative measurement for each gene.

Source

www.ncbi.nlm.nih.gov/geo/

References

McCall M.N., Bolstad B.M., and Irizarry R.A. (2010) Frozen robust multiarray analysis (fRMA). Biostatistics 11, 242-253.

Barrett T., et al. (2007) NCBI GEO: mining tens of millions of expression profiles - database and tools update. Nucl. Acids Res. 35, D760-D765.

Examples

Load the GPL570 database library(bigmemory) path <- system.file("extdata",package="ChIPXpressData") DB_GPL570 <- attach.big.matrix("DB_GPL570.bigmemory.desc",path=path) ## DB_GPL570 is then ready for input into the ChIPXpress function.

To see info about the database matrix describe (DB_GPL570)

Index

*Topic datasets,GPL1261,database DB_GPL1261.bigmemory, 2 *Topic datasets,GPL570,database DB_GPL570.bigmemory, 3 *Topic package, database, ChIPXpress ChIPXpressData-package, 1

ChIPXpressData (ChIPXpressData-package), 1 ChIPXpressData-package, 1

DB_GPL1261.bigmemory, 2 DB_GPL570.bigmemory, 3