# Package 'FANTOM3and4CAGE'

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Title CAGE data from FANTOM3 and FANTOM4 projects
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<b>Depends</b> R (>= $2.15.0$ )
<b>Description</b> CAGE (Cap Analysis Gene Expression) data from FANTOM3 and FANTOM4 projects produced by RIKEN Omics Science Center.
License GPL-3
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FANTOMhumanSamples

List of FANTOM CAGE samples for human

## **Description**

A data.frame with information on all human CAGE samples from FANTOM3 and FANTOM4 projects available in this package.

#### Usage

data(FANTOMhumanSamples)

#### **Details**

FANTOMhumanSamples is a data. frame with 3 columns:

dataset: the name of the dataset that can be loaded using data() function group: the name of the group of samples that originate from the same tissue (*e.g.* blood) sample: the name of the specific sample

#### **Source**

The annotation of CAGE samples was obtained from the FANTOM web resource (http://fantom.gsc.riken.jp/4/download/

# References

Kawaji *et al.* (2010) Update of the FANTOM web resource: from mammalian transcriptional land-scape to its dynamic regulation, *Nucleic Acids Research* **39**:D856-D860.

# **Examples**

```
data(FANTOMhumanSamples)
head(FANTOMhumanSamples)
```

FANTOMmouseSamples

List of FANTOM CAGE samples for mouse

# **Description**

A data.frame with information on all mouse CAGE samples from FANTOM3 and FANTOM4 projects available in this package.

# Usage

data(FANTOMmouseSamples)

#### **Details**

FANTOMmouseSamples is a data.frame with 3 columns:

dataset: the name of the dataset that can be loaded using data() function

group: the name of the group of samples that originate from the same tissue (e.g. brain)

sample: the name of the specific sample

#### **Source**

The annotation of CAGE samples was obtained from the FANTOM web resource (http://fantom.gsc.riken.jp/4/download/

#### References

Kawaji *et al.* (2010) Update of the FANTOM web resource: from mammalian transcriptional land-scape to its dynamic regulation, *Nucleic Acids Research* **39**:D856-D860.

#### **Examples**

data(FANTOMmouseSamples)
head(FANTOMmouseSamples)

FANTOMtimecourseCAGEhuman

Human timecourse CAGE data produced in FANTOM4 project

# Description

This dataset contains all transcription start sites (TSSs) detected by CAGE and their frequency of usage in the human THP-1 cells induction timecourse from FANTOM4 project.

#### Usage

data(FANTOMtimecourseCAGEhuman)

# **Details**

FANTOMtimecourseCAGEhuman is a list with only one element (timecourse) named THP-1\_monocytic\_induction. This element is a data.frame with genomic positions of TSSs detected by CAGE and frequency of their usage in various timepoints during stimulation of human THP-1 myelomonocytic leukemia cells by phorbol myristate acetate (PMA). First three columns contain the chromosome (chr), the 1-based coordinate (pos) and the strand (strand) of the TSS, and the following columns contain the number of CAGE tags supporting that TSS in each specified timepoint.

#### Source

The CAGE detected TSSs (mapped to the hg18 human genome assembly) and numbers of CAGE tags for THP-1 monocytic induction were obtained from the FANTOM web resource (http://fantom.gsc.riken.jp/4/download The data were originally published by The FANTOM Consortium (Suzuki *et al.*, Nature Genetics 2009).

#### References

Kawaji *et al.* (2010) Update of the FANTOM web resource: from mammalian transcriptional land-scape to its dynamic regulation, *Nucleic Acids Research* **39**:D856-D860.

Suzuki *et al.* (2009) The transcriptional network that controls growth arrest and differentiation in a human myeloid leukemia cell line, *Nature Genetics* **41**:553-562.

#### **Examples**

data(FANTOMtimecourseCAGEhuman)
names(FANTOMtimecourseCAGEhuman)
head(FANTOMtimecourseCAGEhuman[["THP-1\_monocytic\_induction"]])

FANTOMtimecourseCAGEmouse

Mouse timecourse CAGE data produced in FANTOM projects

#### **Description**

This dataset contains all transcription start sites (TSSs) detected by CAGE and their frequency of usage in two mouse timecourses: adipogenic induction of DFAT-D1 preadipocytes and liver under constant darkness condition.

#### **Usage**

data(FANTOMtimecourseCAGEmouse)

#### **Details**

FANTOMtimecourseCAGEmouse is a list with two elements (timecourses) named adipogenic\_induction and liver\_under\_constant\_darkness. Each element is a data. frame with genomic positions of TSSs detected by CAGE and frequency of their usage in various timepoints during timecourse. First three columns contain the chromosome (chr), the 1-based coordinate (pos) and the strand (strand) of the TSS, and the following columns contain the number of CAGE tags supporting that TSS in each specified timepoint.

#### **Source**

The CAGE detected TSSs (mapped to the mm9 mouse genome assembly) and numbers of CAGE tags for both timecourses were obtained from the FANTOM web resource (http://fantom.gsc.riken.jp/4/download/). The data were originally published by The FANTOM Consortium (Carninci *et al.*, Science 2005).

#### References

Kawaji *et al.* (2010) Update of the FANTOM web resource: from mammalian transcriptional land-scape to its dynamic regulation, *Nucleic Acids Research* **39**:D856-D860.

Carninci *et al.* (2005) The Transcriptional Landscape of the Mammalian Genome, *Science* **309**(5740):1559-1563.

#### **Examples**

```
data(FANTOMtimecourseCAGEmouse)
names(FANTOMtimecourseCAGEmouse)
head(FANTOMtimecourseCAGEmouse[["adipogenic_induction"]])
```

 $\begin{tabular}{ll} FANTOM tissue CAGE data produced in FANTOM projects for various human tissues \\ and cell lines \\ \end{tabular}$ 

#### **Description**

This dataset contains all transcription start sites (TSSs) detected by CAGE and their frequency of usage in the various human tissues and cell lines generated in FANTOM3 and FANTOM4 projects.

# Usage

data(FANTOMtissueCAGEhuman)

#### **Details**

FANTOMtissueCAGEhuman is a list of groups of human samples, which are grouped by the tissue of origin. Each element is named according to the corresponding tissue and it is a data.frame with genomic positions of TSSs detected by CAGE and frequency of their usage in various samples originating from that tissue. First three columns in every data.frame contain the chromosome (chr), the 1-based coordinate (pos) and the strand (strand) of the TSS, and the following columns contain the number of CAGE tags supporting that TSS in each specific sample. These columns are named according to the corresponding sample.

#### Source

The CAGE detected TSSs (mapped to the hg18 human genome assembly) and numbers of CAGE tags were obtained from the FANTOM web resource (http://fantom.gsc.riken.jp/4/download/). The data were originally published by The FANTOM Consortium (Carninci *et al.*, Science 2005, and Carninci *et al.*, Nature Genetics 2006, and Faulkner *et al.*, Nature Genetics 2009).

#### References

Kawaji *et al.* (2010) Update of the FANTOM web resource: from mammalian transcriptional land-scape to its dynamic regulation, *Nucleic Acids Research* **39**:D856-D860.

Carninci et al. (2005) The transcriptional landscape of the mammalian genome, Science 309:1559-1563

Carninci *et al.* (2006) Genome-wide analysis of mammalian promoter architecture and evolution, *Nature Genetics* **6**:626-635.

Faulkner *et al.* (2009) The regulated retrotransposon transcriptome of mammalian cells, *Nature Genetics* **41**:563-571.

# **Examples**

data(FANTOMtissueCAGEhuman)
names(FANTOMtissueCAGEhuman)
head(FANTOMtissueCAGEhuman[["liver"]])

FANTOMtissueCAGEmouse CAGE data produced in FANTOM projects for various mouse tissues and cell lines

### **Description**

This dataset contains all transcription start sites (TSSs) detected by CAGE and their frequency of usage in the various mouse tissues and cell lines generated in FANTOM3 and FANTOM4 projects.

#### Usage

data(FANTOMtissueCAGEmouse)

#### **Details**

FANTOMtissueCAGEmouse is a list of groups of mouse samples, which are grouped by the tissue of origin. Each element is named according to the corresponding tissue and it is a data.frame with genomic positions of TSSs detected by CAGE and frequency of their usage in various samples originating from that tissue. First three columns in every data.frame contain the chromosome (chr), the 1-based coordinate (pos) and the strand (strand) of the TSS, and the following columns contain the number of CAGE tags supporting that TSS in each specific sample. These columns are named according to the corresponding sample.

#### Source

The CAGE detected TSSs (mapped to the mm9 mouse genome assembly) and numbers of CAGE tags were obtained from the FANTOM web resource (http://fantom.gsc.riken.jp/4/download/). The data were originally published by The FANTOM Consortium (Carninci *et al.*, Science 2005, and Faulkner *et al.*, Nature Genetics 2009).

#### References

Kawaji *et al.* (2010) Update of the FANTOM web resource: from mammalian transcriptional land-scape to its dynamic regulation, *Nucleic Acids Research* **39**:D856-D860.

Carninci *et al.* (2005) The transcriptional landscape of the mammalian genome, *Science* **309**:1559-1563.

Faulkner et al. (2009) The regulated retrotransposon transcriptome of mammalian cells, *Nature Genetics* **41**:563-571.

#### **Examples**

data(FANTOMtissueCAGEmouse)
names(FANTOMtissueCAGEmouse)
head(FANTOMtissueCAGEmouse[["lung"]])

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