FANTOM3and4CAGE: an R data package with CAGE data from FANTOM3 and FANTOM4 projects

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1 Introduction

This document briefly describes the content of the FANTOM3and4CAGE data package. FANTOM3and4CAGE is a Bioconductor-compliant R package that contains Cap Analysis of Gene Expression (CAGE) sequencing data produced by FANTOM consortium (http://fantom.gsc.riken.jp/). CAGE (Kodzius et al. (2006)) is a high-throughput method for transcriptome analysis that utilizes "cap-trapping" (Carninci et al. (1996)), a technique based on the biotinylation of the 7-methylguanosine cap of Pol II transcripts, to pulldown the 5'-complete cDNAs reversely transcribed from the captured transcripts. This enables the sequencing of short fragments from 5' ends, which can be mapped back to the referent genome to infer the exact position of the transcription start sites (TSSs) used for transcription of captured RNAs. Number of CAGE tags supporting each TSS gives the information on relative frequency of its usage and can be used as a measure of expression from that specific TSS. Thus, CAGE provides information on two aspects of capped transcriptome: genome-wide 1bp-resolution map of transcription start sites and

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transcript expression levels. This information can be used for various analyses, from 5' centered expression profiling (Takahashi et al. (2012)) to studying promoter architecture (Carninci et al. (2006)).

This data package contains genomic coordinates of TSSs and number of CAGE tags supporting each TSS in various mouse and human samples analysed by CAGE in FAN-TOM3 and FANTOM4 projects. The data was originally published in main FANTOM publications (Carninci et al. (2005), Carninci et al. (2006), Suzuki et al. (2009), Faulkner et al. (2009)), and represents a valuable resource of genome-wide TSSs for mouse and human. All of the data was downloaded from the FANTOM web resource (Kawaji et al. (2010), http://fantom.gsc.riken.jp/4/download/) and was organized into datasets by organism and tissue of origin. All human data is mapped to hg18 assembly, and mouse data to mm9 assembly of the genome. Figure 1 schematically describes the organization and the structure of the data in the FANTOM3and4CAGE package.

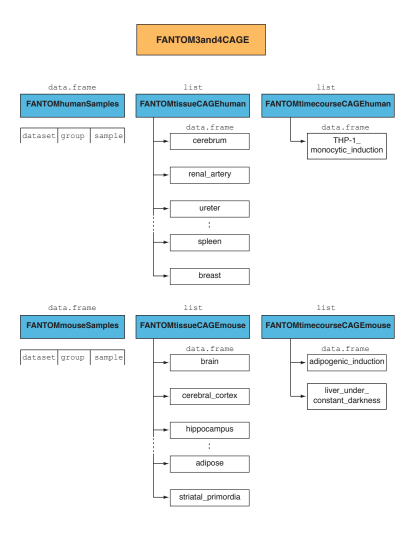


Figure 1: Content and structure of data in FANTOM3and4CAGE data package

2 Getting started

To load the FANTOM3and4CAGE package into your R environment type:

> library(FANTOM3and4CAGE)

2.1 Listing available CAGE samples

As shown in Figure 1, there are six datasets (shaded in blue) that can be loaded via call to data() function. Two of them are data.frames that describe the content of the remaining four datasets. These are FANTOMhumanSamples and FANTOMmouseSamples, for human and mouse, respectively.

To load the list of human samples type:

- > data(FANTOMhumanSamples)
- > head(FANTOMhumanSamples, 10)

	dataset	group	sample
1	${\tt FANTOMtissueCAGEhuman}$	cerebrum	cerebrum
2	${\tt FANTOMtissueCAGEhuman}$	renal_artery	renal_artery
3	${\tt FANTOMtissueCAGEhuman}$	ureter	ureter
4	${\tt FANTOMtissueCAGEhuman}$	urinary_bladder	urinary_bladder
5	${\tt FANTOMtissueCAGEhuman}$	kidney	malignancy
6	${\tt FANTOMtissueCAGEhuman}$	kidney	kidney
7	${\tt FANTOMtissueCAGEhuman}$	${\tt small_intestine}$	small_intestine
8	${\tt FANTOMtissueCAGEhuman}$	rectum	malignancy
9	${\tt FANTOMtissueCAGEhuman}$	rectum	rectum
10	${\tt FANTOMtissueCAGEhuman}$	cecum	malignancy

The information is organized into three 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

2.2 CAGE datasets for various tissues

The FANTOMtissueCAGEhuman and FANTOMtissueCAGEmouse datasets contain CAGE data organized by tissue of origin:

- > data(FANTOMtissueCAGEhuman)
- > names(FANTOMtissueCAGEhuman)

```
[1] "cerebrum"
                        "renal_artery"
                                            "ureter"
[4] "urinary_bladder" "kidney"
                                            "small_intestine"
 [7] "rectum"
                                            "liver"
                        "cecum"
[10] "large_intestine" "prostate_gland"
                                           "mammary_gland"
[13] "epididymis"
                        "skin"
                                            "adipose"
[16] "pancreas"
                        "thymus"
                                            "undefined"
                                            "adrenal_gland"
[19] "blood"
                        "lung"
[22] "colon"
                        "brain"
                                           "cerebellum"
[25] "testis"
                        "embryo"
                                            "bone marrow"
[28] "heart"
                        "muscle"
                                           "frontal_lobe"
                                            "spleen"
[31] "occipital_lobe"
                        "parietal_lobe"
[34] "breast"
```

It is a named list, where names correspond to entries in the group column (in the data.frame listing all the samples) and indicate tissue of origin. Each element of the list is a data.frame with genomic coordinates of TSSs detected in that group of samples followed by columns with numbers of CAGE tags supporting each TSS in every individual sample. The names of columns correspond to entries in the sample column (in the data.frame listing all the samples) and describe individual samples.

```
> lung_group <- FANTOMtissueCAGEhuman[["lung"]]
> head(lung_group)
```

	chr	pos	strand	R	RCB-0702_WI-38	RCB-0098_A549
1	chr1	558799	+		0	1
2	chr1	559777	+		0	1
3	chr1	703878	_		0	1
4	chr1	703904	_		0	0
5	chr1	752741	_		0	0
6	chr1	752754	_		0	0
	RCB-0)465_Lu-	-130 lui	ng	5	
1			0	0)	
2			0	0)	
3			0	0)	
4			1	0)	
5			1	0)	
6			1	0)	

2.3 CAGE timecourse datasets

In addition to CAGE data for various tissue types, there are timecourse datasets available in *FANTOM3and4CAGE* package. These are FANTOMtimecourseCAGEhuman and FANTOMtimecourseCAGEmouse, for human and mouse, respectively.

- > data(FANTOMtimecourseCAGEmouse)
- > names(FANTOMtimecourseCAGEmouse)
- [1] "adipogenic_induction"
- [2] "liver_under_constant_darkness"
- > head(FANTOMtimecourseCAGEmouse[["adipogenic_induction"]])

	chr	pos	strand	DFAT-D1	_preadipocytes_Odays
1	chr1	3091394	+		0
2	chr1	3641339	_		1
3	chr1	3661787	_		0
4	chr1	3936146	+		0
5	chr1	3968879	_		0
6	chr1	4569858	+		0
	DFAT-	-D1_pread	dipocyte	es_2days	DFAT-D1_preadipocytes_4days
1				0	1
2				0	0
3				0	0
4				0	1
5				0	0
6				0	0
	DFAT-	-D1_pread	dipocyte	es_6days	DFAT-D1_preadipocytes_8days
1				0	0
2				0	0
3				0	1
4				0	0
5				0	1
6				0	1

They are organized in the same way as tissue datasets described above, *i.e.* each element of the list is a data.frame with CAGE detected TSSs for one timecourse.

3 Session Info

[3] LC_TIME=C LC_COLLATE=C

[5] LC_MONETARY=C LC_MESSAGES=en_US.UTF-8

[7] LC_PAPER=C LC_NAME=C

[9] LC_ADDRESS=C LC_TELEPHONE=C

[11] LC_MEASUREMENT=no_NO.UTF-8 LC_IDENTIFICATION=C

attached base packages:

[1] stats graphics grDevices utils datasets

[6] methods base

other attached packages:

[1] FANTOM3and4CAGE_0.99.0

loaded via a namespace (and not attached):

[1] tools_3.0.0

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