cummeRbund: Visualization and Exploration of Cufflinks High-throughput Sequencing Data

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Contents

1	Requirements				
2	Introduction	2			
3	CummeRbund Classes				
	3.1 CuffSet Class	2			
	3.2 CuffData Class	2			
	3.3 CuffFeatureSet Class	3			
	3.4 CuffFeature Class	3			
4	Reading cuffdiff output	4			
	4.1 Adding additional feature annotation	5			
5	Global statistics 5				
6	Accessing Data	10			
	6.1 Writing your own SQL accessors	12			
7		13			
	7.1 Geneset level plots	13			
8	Individual Genes	19			
	8.1 Gene-level plots	20			
9	Data Exploration	23			
	9.1 Finding similar genes	23			
10	Miscellaneous	25			
11	Session info	26			

1 Requirements

- Cufflinks \geq v1.1.0
- $R \ge v2.7.0$
- Packages:
 - RSQLite
 - ggplot2
 - reshape
 - plyr

2 Introduction

cummeRbund is a visualization package for Cufflinks high-throughput sequencing data. The base class, *cuffSet* is a 'pointer' to cufflinks data that are stored out-of-memory in a sqlite database.

3 CummeRbund Classes

3.1 CuffSet Class

A pointer class to control access to the sqlite tables holding the Cufflinks data. The primary slot is @DB which contains the RSQLite connection object. The additional slots (genes, isoforms, TSS, and CDS) are each instances of the *Cuff-Data* class and are pointers to sets of tables for each data subtype. This is the default class created by *readCufflinks*. By default, *CuffData* accessor methods applied to a *CuffSet* class will operate on the 'genes' slot.

3.2 CuffData Class

The CuffData class is also a pointer class to the SQL backend, but each instance is specific for a data subtype (genes, isoforms, TSS, CDS). Again, there is an @DB slot that contains the RSQLite connection object. There are several accessor, setter, and plotting methods that allow for global analysis of all features within a CuffData class. Subsetting is currently being re-written, however, it is primarily done through the 'gene_id' field. Available slots for the CuffData class are:

- DB: RSQLite connection object
- tables: A *list* of tables in the SQLite DB that contain the cufflinks data.
- filters: A *list* of filters for subsetting (not implemented yet).
- type: A character field describing the data (ie. 'genes', 'isoforms', 'TSS', 'CDS', 'other')

• idField: The name of the identifying index field for this object (eg. 'gene_id' for type='gene', or 'isoform_id' for type='isoform')

Making the best use of either the CuffSet or CuffData classes will enable you to keep the entire dataset out of memory and significantly improve performance for large cufflinks datasets.

3.3 CuffFeatureSet Class

The CuffFeatureSet class is a data-storage container that holds all available data for a pre-determined list of features. Slots for FPKM data, differential regulation data, and feature-level annotation are all available. Unlike the previous classes, this class contains no connection information to the SQL database, but rather contains several slots with data.frame objects storing multiple-features worth of information. There are available accessors, and plotting methods that are designed to present multiple-features worth of information (eg. heatmaps, scatterplots, etc) Available slots for a CuffFeatureSet object include:

- annotation: Holds all feature-level annotation information for all features in object.
- fpkm: A data frame of FPKM data across all samples, for all features in object.
- diff: A data frame of differential expression/regulation data for all features in object.

A specialized sub-class of *CuffFeatureSet* is the *CuffGeneSet* class. This subclass adds additional slots to contain all isoforms, TSS, and CDS information for a given set of gene_ids. The *CuffGeneSet* class is designed to aggregate all relevant information for a set of genes into one object for easy analysis and/or manipulation. The *CuffGeneSet* object adds the following slots:

- ids: A 'character' list of all gene_ids used in object.
- isoforms: A CuffFeatureSet object for all isoforms of genes in object.
- TSS: A CuffFeatureSet object for all TSS of genes in object.
- CDS: A CuffFeatureSet object for all CDS of genes in object.

3.4 CuffFeature Class

The CuffFeature class is designed for single-feature-level data analysis and plotting. The methods available for this object are designed to analyze or visualize information about a specific feature. This is a 'data' object, as opposed to a 'pointer' object to the database backend. There is a validity requirement that a CuffFeature object only point to data from a single feature. Available slots for a CuffFeature object include:

- annotation: Holds feature-level annotation information for a given feature.
- fpkm: A data frame of FPKM data across all samples for a given feature.
- diff: A data frame of differential expression/regulation data for a given feature.

A specialized sub-class of *CuffFeature* is the *CuffGene* class. This subclass adds additional slots to contain all isoform, TSS, and CDS information for a given gene. The *CuffGene* object adds the following slots:

- $\bullet\,$ id: The common 'gene_id' for all data in object
- isoforms: A CuffFeature object for all isoforms of a given gene.
- TSS: A CuffFeature object for all TSS of a given gene.
- CDS: A CuffFeature object for all CDS of a given gene.

Note: Future versions of cummeRbund may try to collapse the redundant functionality of the CuffFeature and CuffFeatureSet classes.

4 Reading cuffdiff output

One of the principle benefits of using cummeRbund is that data are stored in a SQLite database. This allows for out-of-memory analysis of data, quick retrieval, and only a one-time cost to setup the tables. By default, cummeRbund assumes that all output files from cuffdiff are in the current working directory. To read these files, populate the 'cuffData.db' database backend, and return the CuffSet pointer object, you can do the following.

Again, by default *dir* is assumed to be the current working directory and cuff<-readCufflinks() should work if all appropriate files are in the current working directory. Should you need to rebuild the SQLite backend for any reason, you

can add the option rebuild=T to readCufflinks. Once the database is created, readCufflinks will default to using the SQL backend and should not need to rebuild this database. Each R session should begin with a call to readCufflinks so as to initialize the database connection and create an object with the appropriate RSQLite connection information.

4.1 Adding additional feature annotation

Gene- or feature-level annotation can be permanently added to the database tables for future querying. If you have a data.frame where the first column contains the 'tracking_id' (eg. 'gene_id' for genes, 'isoform_id' for isoforms, etc). You can easily add feature level annotation using the addFeatures() function:

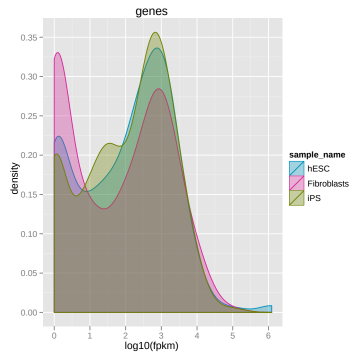
By default, features added to a *CuffSet* object are assumed to be gene-level annotations, but the level can selected using the argument *level*. Features added to a *CuffData* object are assumed to be of the same type as the object@type value (e.g. gene-level features for 'genes', isoform-level features for isoforms, etc.)

5 Global statistics

Several plotting methods are available that allow for quality-control or global analysis of cufflinks data. For example, to assess the distributions of FPKM scores across samples, you can use the *csDensity* plot (Figure 1).

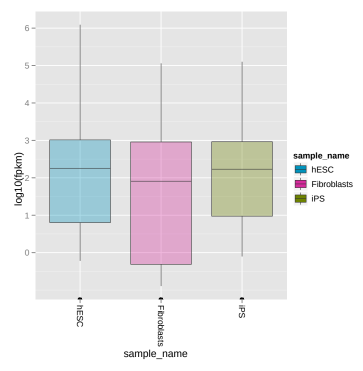
```
> dens <- csDensity(cuff@genes)</pre>
```

> dens

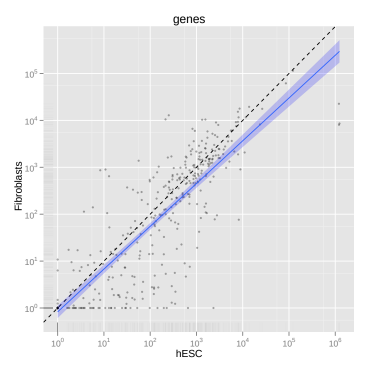


Boxplots can be visualized using the csBoxplot method (Figure 2).

> b <- csBoxplot(cuff@genes)
> b

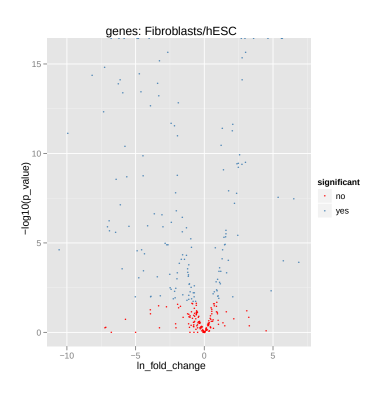


Pairwise comparisons can be made by using csScatter. You must specify the sample names to use for the x and y axes:



Volcano plots are also available for the ${\it CuffData}$ objects. Again, you must specify the comparisons by sample name.

> v <- csVolcano(cuff@genes, "hESC", "Fibroblasts")
> v



6 Accessing Data

Feature-level information can be accessed directly from a *CuffData* object using the *fpkm*, *diffData*, or *features* methods:

```
> gene.features <- features(cuff@genes)
> head(gene.features)
      gene_id class_code nearest_ref_id gene_short_name
1 XLOC_000001
                     <NA>
                                                     <NA>
                                    <NA>
2 XLOC_000002
                     <NA>
                                     <NA>
                                                    OR4F5
3 XLOC_000003
                     <NA>
                                    <NA>
                                                     <NA>
4 XLOC_000004
                     <NA>
                                     <NA>
                                                      <NA>
5 XLOC_000005
                     <NA>
                                     <NA>
                                                      <NA>
6 XLOC_000006
                     <NA>
                                     <NA>
                                                   OR4F16
               locus length coverage gene_id
    chr1:11873-29961
                          NA
1
                                   NA
                                          <NA>
2
    chr1:69090-70008
                          NA
                                   NA
                                          <NA>
3 chr1:321083-321114
                          NA
                                   NA
                                          <NA>
4 chr1:321145-321223
                                          <NA>
                          NA
                                   NA
5 chr1:322036-328580
                          NA
                                   NA
                                          <NA>
6 chr1:367658-368595
                                          <NA>
                          NA
                                   NA
> gene.fpkm <- fpkm(cuff@genes)</pre>
> head(gene.fpkm)
      gene_id sample_name
                                fpkm
                                         conf_hi conf_lo
1 XLOC_000001 Fibroblasts 16.401100
                                       428.14700
                                                        0
2 XLOC_000001
                     hESC 0.723836
                                         3.01108
                                                        0
                       iPS 54.067200 1402.31000
3 XLOC_000001
                                                        0
4 XLOC_000002 Fibroblasts
                            0.000000
                                         0.00000
                                                        0
                     hESC
                                         0.00000
5 XLOC_000002
                            0.000000
                                                        0
6 XLOC_000002
                      iPS 0.000000
                                         0.00000
  quant_status
1
       LOWDATA
2
            OK
3
       LOWDATA
4
            OK
5
            OK
6
            OK
> isoform.fpkm <- fpkm(cuff@isoforms)</pre>
> head(isoform.fpkm)
      isoform_id sample_name
                                   fpkm conf_hi conf_lo
1 TCONS_00000001 Fibroblasts 11.910700 19.96650
                                                   3.85498
2 TCONS_00000001
                         hESC 0.000000 0.00000 0.00000
```

```
3 TCONS_00000001
                         iPS 9.563700 23.68410 0.00000
4 TCONS_00000002 Fibroblasts 0.000000 8.55378 0.00000
5 TCONS_00000002
                  hESC 0.723836 3.01108 0.00000
6 TCONS_00000002
                        iPS 32.934400 47.93760 17.93130
  quant_status
1
            OK
2
            OK
3
       LOWDATA
4
            OK
5
            OK
6
            OK
> gene.diff <- diffData(cuff@genes)</pre>
> head(gene.diff)
      gene_id sample_1
                          sample_2 status
                                              value_1
                                                          value_2
1 XLOC_000001
                  hESC Fibroblasts
                                       OK 7.23836e-01
                                                          16.4011
2 XLOC_000002
                  hESC Fibroblasts NOTEST 0.00000e+00
                                                           0.0000
3 XLOC_000003
                  hESC Fibroblasts NOTEST 0.00000e+00
                                                           0.0000
4 XLOC_000004
                  hESC Fibroblasts
                                       OK 1.20000e+06 22616.4000
5 XLOC_000005
                  hESC Fibroblasts
                                       OK 1.13903e+03
                                                          41.1644
                  hESC Fibroblasts NOTEST 0.00000e+00
6 XLOC_000006
                                                           0.0000
  ln_fold_change test_stat p_value q_value significant
         4.50198 -0.246654 0.805176 0.893616
1
2
         0.00000 0.000000 1.000000 1.000000
                                                      no
3
         0.00000 0.000000 1.000000 1.000000
                                                      nο
4
        -5.72952 1.310270 0.190105 0.300329
                                                      no
5
        -4.79027 10.857600 0.000000 0.000000
                                                     yes
         0.00000 0.000000 1.000000 1.000000
6
```

Vectors of sample names and feature names are available by using the *samples* and *featureNames* methods:

To facilitate Bioconductor-like operations, an 'FPKM-matrix' can be returned easily using the fpkmMatrix method:

```
> gene.matrix <- fpkmMatrix(cuff@genes)
> head(gene.matrix)
```

	hESC	${\tt Fibroblasts}$	iPS
XLOC_000001	7.23836e-01	16.4011	54.06720
XLOC_000002	0.00000e+00	0.0000	0.00000
XLOC_000003	0.00000e+00	0.0000	0.00000
XLOC_000004	1.20000e+06	22616.4000	0.00000
XLOC_000005	1.13903e+03	41.1644	944.30800
XLOC_000006	0.00000e+00	0.0000	9.00455

6.1 Writing your own SQL accessors

Since the cufflinks is a SQLite database backend, if you are familiar with SQL and/or RSQLite query construction, you can simply design your own SQL queries to access the data that you are after. PUT DATABASE SCHEMA HERE...

7 Creating Gene Sets

Gene Sets (stored in a *CuffGeneSet* object) can be created using the *getGenes* method on a CuffSet object. You must first create a vector of 'gene_id' or 'gene_short_name' values to identify the genes you wish to select:

```
> data(sampleData)
> myGeneIds <- sampleIDs
> myGeneIds

[1] "XLOC_001363" "XLOC_001297" "XLOC_001339" "XLOC_000132"
[5] "XLOC_001265" "XLOC_000151" "XLOC_001359" "XLOC_000069"
[9] "XLOC_000170" "XLOC_000105" "XLOC_001262" "XLOC_001348"
[13] "XLOC_001411" "XLOC_001369" "XLOC_000158" "XLOC_001370"
[17] "XLOC_001263" "XLOC_000115" "XLOC_000089" "XLOC_001240"
> myGenes <- getGenes(cuff, myGeneIds)
> myGenes
CuffGeneSet instance for genes c("XLOC_000069", "XLOC_000089")
```

CuffGeneSet instance for genes c("XLOC_000069", "XLOC_000089", "XLOC_000105", "XLOC_000115", Short name: ESPN PGD MFN2 PRAMEF1 EFHD2 PADI1 NA FAM43B UBE2J2 Clorf86 SLC2A7 SPATA2 Slots:

annotation

fpkm

diff

isoforms

CuffFeatureSet instance of size 45

TSS

CuffFeatureSet instance of size 18

CDS

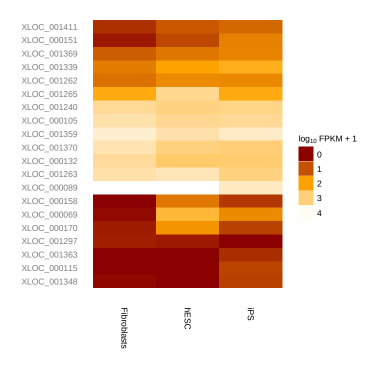
CuffFeatureSet instance of size 31

The same fpkm, fpkmMatrix, features, diffData, samples, and featureNames are available for instances of the CuffGeneSet class.

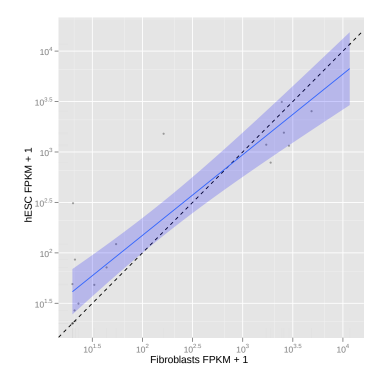
7.1 Geneset level plots

There are several plotting functions available for gene-set-level visualization:

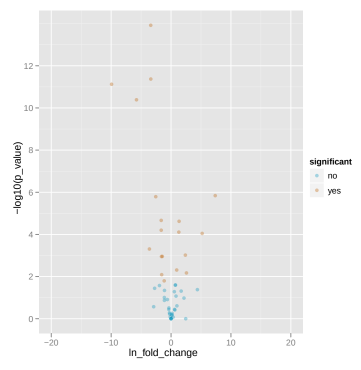
```
> h <- csHeatmap(myGenes, cluster = "both")
> h
```



> s <- csScatter(myGenes, "Fibroblasts", "hESC", smooth = T) > s



> v <- csVolcano(myGenes, cluster = "both")
> v

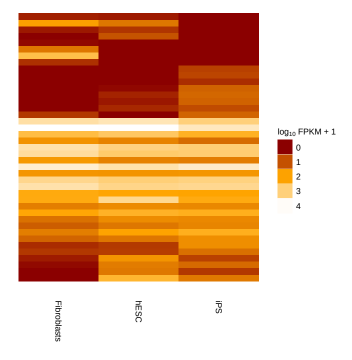


Similar plots can be made for all sub-level features of a CuffGeneSet class by specifying which slot you would like to plot (eg. @isoforms, @TSS, @CDS).

```
> ih <- csHeatmap(myGenes@isoforms, cluster = "both",</pre>
```

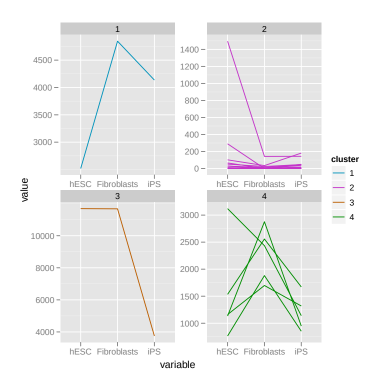
⁺ labRow = F)

> ih



Rudimentary k-means clustering is implemented as well.

- > ic <- csCluster(myGenes, k = 4)
 > ic



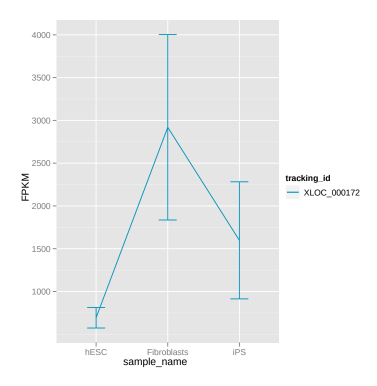
8 Individual Genes

An individual CuffGene object can be created by using the getGene function for a given 'gene_id'.

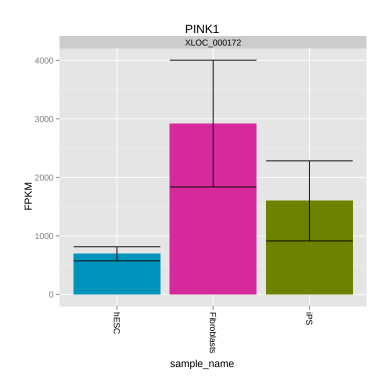
```
> myGeneId <- "PINK1"
> myGene <- getGene(cuff, myGeneId)
> myGene
CuffGene instance for gene PINK1
Short name:
                    PINK1
Slots:
         annotation
         fpkm
         diff
         isoforms
                          CuffFeature instance of size 2
         TSS
                             CuffFeature instance of size 2
         CDS
                             CuffFeature instance of size 2
> head(fpkm(myGene))
      gene_id sample_name
                              fpkm conf_hi conf_lo
1 XLOC_000172 Fibroblasts 2919.340 4002.960 1835.730
                     hESC 693.465 813.869 573.062
2 XLOC_000172
3 XLOC_000172
                      iPS 1598.040 2282.380 913.710
  quant_status
1
            OK
2
            OK
3
            OK
> head(fpkm(myGene@isoforms))
      isoform_id sample_name
                                 fpkm conf_hi
                                                  conf_lo
1 TCONS_00000480 Fibroblasts 2101.640 3111.330 1091.9400
2 TCONS_00000481 Fibroblasts 817.704 1391.700
                                                243.7120
3 TCONS_00000480
                        hESC 573.512 668.688
                                                478.3370
                        hESC 119.953 152.675
4 TCONS_00000481
                                                 87.2311
5 TCONS_00000480
                         iPS 1598.040 2282.380
                                                913.7100
6 TCONS_00000481
                         iPS
                                0.000
                                         0.000
                                                  0.0000
  quant_status
1
            OK
2
            OK
3
            OK
4
            OK
5
            OK
6
            OK
```

8.1 Gene-level plots

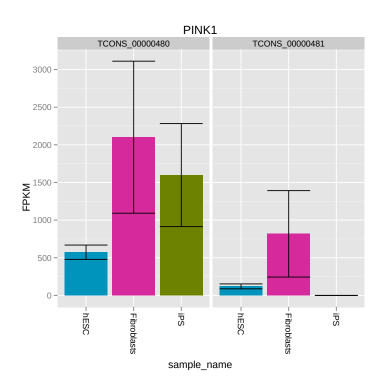
> gl <- expressionPlot(myGene)
> gl



- > gb <- expressionBarplot(myGene)
 > gb



- > igb <- expressionBarplot(myGene@isoforms)
 > igb



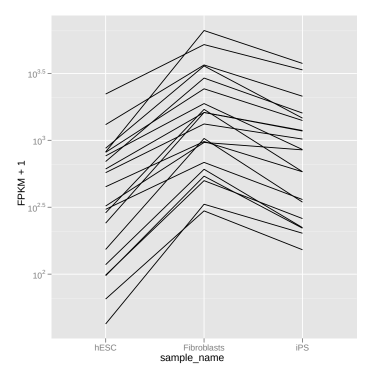
9 Data Exploration

The cummeRbund package is more than just a visualization tool as well. We are working to implement several different means of data exploration from gene and condition clustering, finding features with similar expression profiles, as well as incorporating Gene Ontology analysis.

9.1 Finding similar genes

One common question in large-scale gene expression analyses is 'How can I find genes with similar expression profiles to gene x?'. We have implemented a method, findSimilar to allow you to identify a fixed number of the most similar genes to a given gene of interest. For example, if you wanted to find the 20 genes most similar to "PINK1", you could do the following:

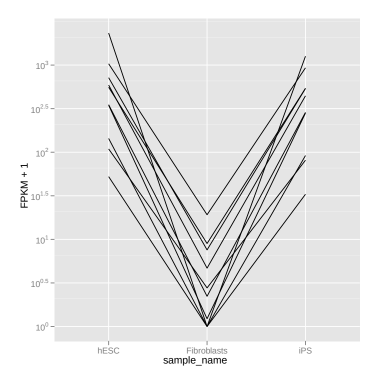
```
> mySimilar <- findSimilar(cuff, "PINK1", n = 20)
> mySimilar.expression <- expressionPlot(mySimilar,
+ logMode = T, showErrorbars = F)</pre>
```



You are also able to provide your own expression profile in lieu of a 'gene_id'. The vector provided must match the order and length of samples().

```
> myProfile <- c(500, 0, 400)
> mySimilar2 <- findSimilar(cuff, myProfile, n = 10)
```

> mySimilar2.expression <- expressionPlot(mySimilar2,
+ logMode = T, showErrorbars = F)</pre>



findSimilar() uses the Jensen-Shannon distance between the probability distributions of each gene across conditions to determine the similarity. We have found this to be a more robust way to determine distance between genes using the high dynamic range of FPKM data. Future versions may allow for other dissimilarity measures to be used instead.

10 Miscellaneous

- All plotting functions return ggplot objects and the resulting objects can be manipulated/faceted/altered using standard ggplot2 methods.
- There are occasional DB connectivity issues that arise. Not entirely sure why yet. If necessary, just readCufflinks again and this should solve connectivity issues with a new RSQLite connection object. If connectivity continues to be a problem, try cuff<-readCufflinks(rebuild=T)
- I am still working on fully documenting each of the methods. There are a good number of arguments that exist, but might be hard to find without looking at the source.

11 Session info

```
[1] TRUE
> sessionInfo()
R version 2.13.1 (2011-07-08)
Platform: x86_64-apple-darwin9.8.0/x86_64 (64-bit)
locale:
[1] en_US.UTF-8/en_US.UTF-8/C/C/en_US.UTF-8/en_US.UTF-8
attached base packages:
[1] grid
            stats
                        graphics grDevices utils
                                                      datasets
[7] methods base
other attached packages:
[1] cummeRbund_0.1.3 ggplot2_0.8.9
                                      proto_0.3-9.2
[4] reshape_0.8.4
                    plyr_1.6
                                      RSQLite_0.9-4
[7] DBI_0.2-5
loaded via a namespace (and not attached):
[1] digest_0.5.0 tools_2.13.1
```