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September 6, 2017

Contents

1	Overvi	iev	۷																					2
	1.1	Th	e m	ode	d																			2
2	Installation																3							
	2.1	Ins	tall	via	GitH	lub																		3
	2.2	Ins	tall	loca	ally .																			3
	2.3 I	Lo	ad t	he p	oack	age	9																	3
3	Analysis												3											
	3.1 I	Inp	ut .																					3
	3.2 I	Ru	n Ti	rend	dy																			4
	3.3	Visualize trends of the top dynamic genes																						
	3.4	Vis	sual	ize i	indiv	idu	al (ger	nes															7
	3.5	Ge	ne	spe	cific	est	im	ate	S															9
	3.6 I	Bre	eak	ooin	t dis	trib	uti	on	OV	er t	he	tir	ne	С	ou	rse	. 6							11
4	More a	adv	v an	cec	d ar	naly	/si	S.																11
	4.1	Tin	ne d	cour	se w	vith	no	n-ı	unif	orr	n s	sar	np	olir	ng									11
5	Extract genes with certain pattern													13										
6	Additio	Additional options													15									
7	Trendy	y s	hin	y a	pp .																			15
8	Sessio	onl	nfo	١																				18

1 Overview

Trendy is an R package fo analyzing high throughput expression data (e.g RNA-seq or microarray) with ordered conditions (e.g. time-course, spatial-course).

For each gene (or other features), Trendy fits a set of segmented (or breakpoint) regression models. The optimal model is chosen as the one with the lowest BIC. Each breakpoint represents a significant changes in the geneâĂŹs expression across the time-course.

The top dynamic genes are then identified as those that are well profiled by their gene-specific segmented regression model. Trendy also implements functions to visualize the dynamic genes and their trends, to order dynamic genes by their trends, and to compute breakpoint distribution at different time-points (e.g. detect time-points with a large number of expression changes).

1.1 The model

To illustrate Trendy, here we use time-course gene expression data as an example. Although, Trendy may also be applied to other types of features (e.g. isoform or exon expression) and/or other types of experiments with ordered conditions (e.g. spatial course).

Denote the normalized gene expression of gene g and sample/time t as $Y_{g,t}$. Denote the total number of genes as G and the total number of samples/times as N. For each gene, Trendy fits segmented regression models with varying numbers of breakpoints from 1 to K. K defaults to 3 but can also be specified by the user. The segmented R package is used to fit the segmented regression models.

For a given gene, among the models with varying k, Trendy selects the optimal number of breakpoints for this gene by comparing the BIC for each model.

To avoid overfitting, the optimal number of breakpoints will be set as $\vec{k_g} = \vec{k_g} - 1$ if at least one segment has less than c_{num} samples. The threshold c_{num} can be specified by the user; the default is 5.

Trendy reports the following for the optimal model:

- Gene specific adjusted R^2 (penalized for the chosen value of k)
- Segment slopes
- Segment trends (and associated p-values)
- Breakpoint estimates

Among all genes, the top dynamic genes are defined as those whose optimal model has high adjusted \mathbb{R}^2 s.

To compute the breakpoint distribution over the time-course, Trendy calculates the number of breakpoints for each time-point across all the genes.

The time-points with high D_t can be considered as those with global expression changes.

Trendy also summarizes the fitted trend or expression pattern of top genes. For samples between the i^{th} and $i+1^{th}$ breakpoint for a given gene, if the t-statistic of the segment slope has p-value greater than c_{pval} , the trend of this segment will be defined as no change. Otherwise the trend of this segment will be defined as up/down based on the slope coefficient. The default value of c_{pval} is 0.1, but may also be specified by the user.

2 Installation

2.1 Install via GitHub

The Trendy package can be installed using functions in the devtools package.

To install, type the following code into R:

```
install.packages("devtools")
library(devtools)
install_github("rhondabacher/Trendy/package/Trendy")
```

2.2 Install locally

Install packages segmented, parallel, and gplots:

```
install.packages(c("segmented","parallel","gplots"))
library("segmented")
library("parallel")
library("gplots")
```

Download the Trendy package from: https://github.com/rhonda/Trendy/tree/master/package And install the package locally.

2.3 Load the package

To load the Trendy package:

```
library(Trendy)
```

3 Analysis

3.1 Input

The input data should be a G-by-N matrix containing the expression values for each gene and each sample, where G is the number of genes and N is the number of samples. The samples should be sorted following the time course order. These values should exhibit expression data after normalization across samples. For example, for RNA-seq data, the raw counts may be normalized using MedianNorm and <code>GetNormalizedMat</code> function in <code>EBSeq</code>. More details can be found in the <code>EBSeq</code> vignette:

 $http://www.bioconductor.org/packages/devel/bioc/vignettes/EBSeq/inst/doc/EBSeq_Vignette.pdf$

The object TrendyExData is a simulated data matrix containing 50 rows of genes and 40 columns of samples.

```
data(TrendyExData)
str(TrendyExData)

## num [1:50, 1:40] 240 199 198 239 202 ...

## - attr(*, "dimnames")=List of 2

## ..$ : chr [1:50] "g1" "g2" "g3" "g4" ...

## ..$ : chr [1:40] "s1" "s2" "s3" "s4" ...
```

3.2 Run Trendy

The trendy function will fit multiple segmented regressions model for each gene (via the segmented R package) and select the the optimal model. Here we want to only consider a maximum of two breakpoints for each gene.

```
res <- trendy(Data = TrendyExData, Max.K = 2)
res.top <- topTrendy(res)</pre>
# default adjusted R square cutoff is 0.5
res.top$AdjustedR2
         g3
                   g1
                            g28
                                       g20
                                                 g15
                                                                      g2
                                                            g4
## 0.9787382 0.9775005 0.9751380 0.9739715 0.9729747 0.9711890 0.9710139
                  g23
        g10
                                        g8
                             g14
                                                  g5
                                                           g24
## 0.9705118 0.9701402 0.9694164 0.9691341 0.9689555 0.9656732 0.9652141
         g9
                   g12
                             g29
                                       g16
                                                 g22
                                                           g18
## 0.9649424 0.9644343 0.9632348 0.9630272 0.9627092 0.9626837 0.9619387
                                       g27
                            g30
                                                 g26
        g25
                   g11
                                                           g19
## 0.9611528 0.9600736 0.9597989 0.9592516 0.9572072 0.9536619 0.9529077
        g21
                   g13
## 0.9528854 0.9470928
```

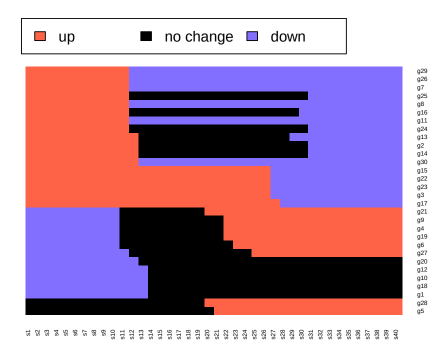
The topTrendy function may be used to extract top dynamic genes. By default, topTrendy will extract genes whose adjusted R^2 , \bar{R}^2 , is greater or equal to 0.5. To change this threshold, a user may specify the AdjR2.Cut parameter in the topTrendy function. The topTrendy function returns the Trendy output with genes sorted decereasingly by \bar{R}^2 .

By default the trendy function only considers genes whose mean expression is greater than 10. To use another threshold, the user may specify the parameter Mean.Cut.

3.3 Visualize trends of the top dynamic genes

contains the trend specification of the top genes. The function **trendHeatmap** can be used to display these trends:

```
res.trend <- trendHeatmap(res.top)
```



```
str(res.trend)

## List of 3

## $ firstup : Named num [1:17] 11.4 11.5 11.6 11.6 11.6 ...

## ..- attr(*, "names")= chr [1:17] "g29" "g26" "g7" "g25" ...

## $ firstdown : Named num [1:11] 10.7 10.9 10.9 10.9 11 ...

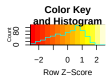
## ..- attr(*, "names")= chr [1:11] "g21" "g9" "g4" "g19" ...

## $ firstnochange: Named num [1:2] 19 20.4

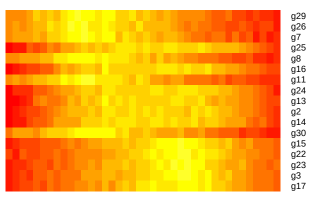
## ..- attr(*, "names")= chr [1:2] "g28" "g5"
```

The trendHeatmap function classifies the top dynamic genes into three groups: start with up, start with down and start with no change. Within each group, genes are sorted by the position of the first breakpoint.

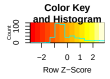
To generate an expression heatmap of the first group of genes (first go up):



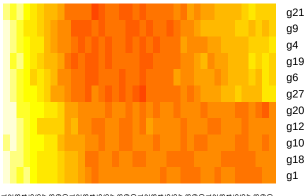
top genes (first go up)



Similarly, to generate an expression heatmap of the second group of genes (first go down):

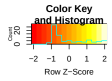


top genes (first go down)

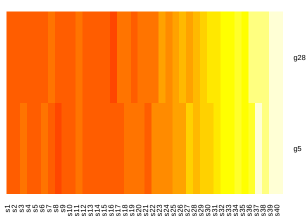


To generate an expression heatmap of the second group of genes (first no change):

```
heatmap.2(TrendyExData[names(res.trend$firstnochange),],
   trace="none", Rowv=F,Colv=F,
        scale="row", main="top genes (first no change)",
        cexRow=.8)
```



top genes (first no change)

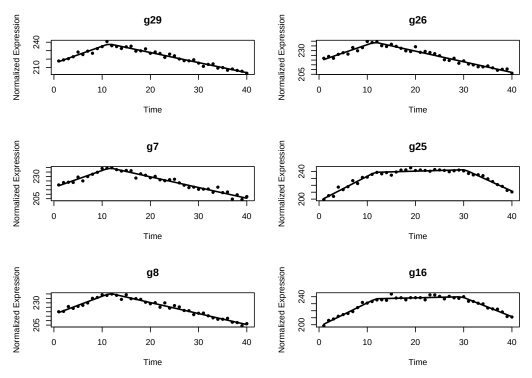


3.4 Visualize individual genes

The plotFeature function may be used to plot expression of individual features/genes and the fitted lines.

For example, to plot the top six genes in the first group of genes (first go up):

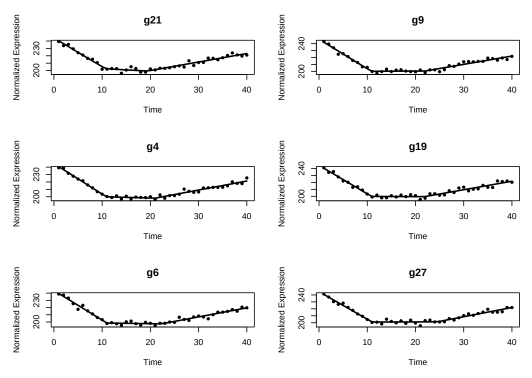
Trendy: Segmented regression analysis of expression dynamics for high-throughput ordered profiling experiments



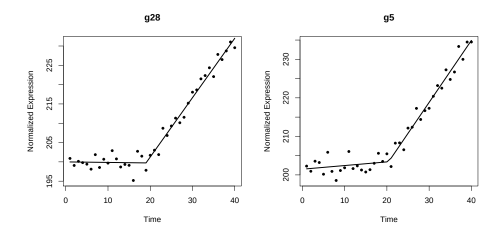
The input of function plotFeature requires the expression data and a list of genes of interest. The parameter Trendy.Out are the results from the trendy function. If it is not specified, then plotFeature will run trendy on the genes of interest before plotting. Specifying the output obtained from previous steps will save time by avoiding fitting the models again.

Similarly, to plot the top six genes in the second group of genes (first go down):

Trendy: Segmented regression analysis of expression dynamics for high-throughput ordered profiling experiments

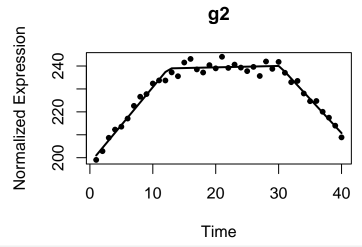


To plot the two genes in the third group of genes (first no change):



3.5 Gene specific estimates

For a given gene of interest, its estimated parameters can be obtained by (using g2 as an example):



The above printouts show that for gene g2 the optimal number of breakpoints is 2. Two estimated breakpoints are around time-points s12 and s30. The fitted slopes for the 3 segments are 3.31, 0.06 and -2.97, which indicate the trend is up-same-down.

These estimates can also be automatically formatted using the function formatResults which can be saved as a .txt. or .csv file. The output currently includes the estimated slope, p-value, and trend of each segment, the estimated breakpoints, the trend for each sample, and the adjusted \mathbb{R}^2 .

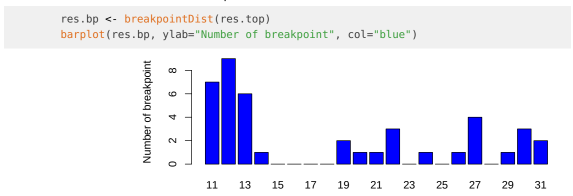
```
trendy.summary <- formatResults(res.top)

# write.table(trendy.summary, file="trendy_summary.txt")</pre>
```

The NA indicates that g3 does not have a segment 3 slope since it only has one breakpoint (i.e two segments).

3.6 Breakpoint distribution over the time course

To calculate number of breakpoints over the time course:



The bar plot indicates that a number of genes have breakpoints around s12 and s13.

4 More advanced analysis

4.1 Time course with non-uniform sampling

If the samples were collected with different time intervals then it is highly suggested to use the original time (instead of a vector of consecutive numbers). To do so, the user may specify the order/times via the T.Vect parameter in the trendy function.

For example, suppose for the example data, the first 30 samples were collected every hour and the other 10 samples were collected every 5 hours. We may define the time vector as:

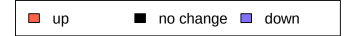
```
t.v <- c(1:30, seq(31, 80, 5))
names(t.v) <- colnames(TrendyExData)
print(t.v)

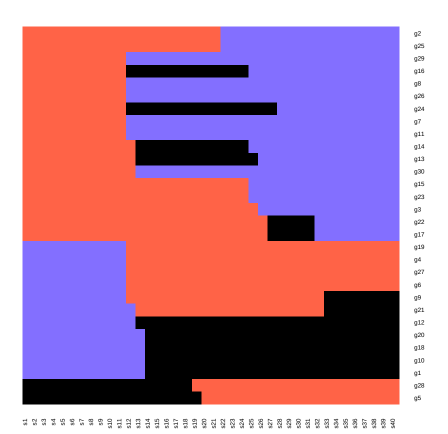
## s1 s2 s3 s4 s5 s6 s7 s8 s9 s10 s11 s12 s13 s14 s15 s16 s17 s18 s19
## 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19
## s20 s21 s22 s23 s24 s25 s26 s27 s28 s29 s30 s31 s32 s33 s34 s35 s36 s37 s38
## 20 21 22 23 24 25 26 27 28 29 30 31 36 41 46 51 56 61 66
## s39 s40
## 71 76</pre>
```

To run Trendy model using the empirical collecting time instead of sample ID (1-40):

```
res2 <- trendy(Data = TrendyExData, T.Vect=t.v, Max.K=2)
res.top2 <- topTrendy(res2)
res.trend2 <- trendHeatmap(res.top2)</pre>
```

Trendy: Segmented regression analysis of expression dynamics for high-throughput ordered profiling experiments





```
str(res.trend2)

## List of 3

## $ firstup : Named num [1:17] 10.2 10.6 11.4 11.4 11.4 ...

## ..- attr(*, "names")= chr [1:17] "g2" "g25" "g29" "g16" ...

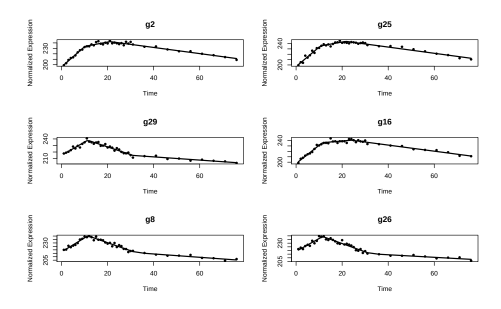
## $ firstdown : Named num [1:11] 11.2 11.4 11.4 11.5 11.7 ...

## ..- attr(*, "names")= chr [1:11] "g19" "g4" "g27" "g6" ...

## $ firstnochange: Named num [1:2] 19 19.5

## ..- attr(*, "names")= chr [1:2] "g28" "g5"
```

To plot the first six genes that have up-regulated pattern at the beginning of the time course:

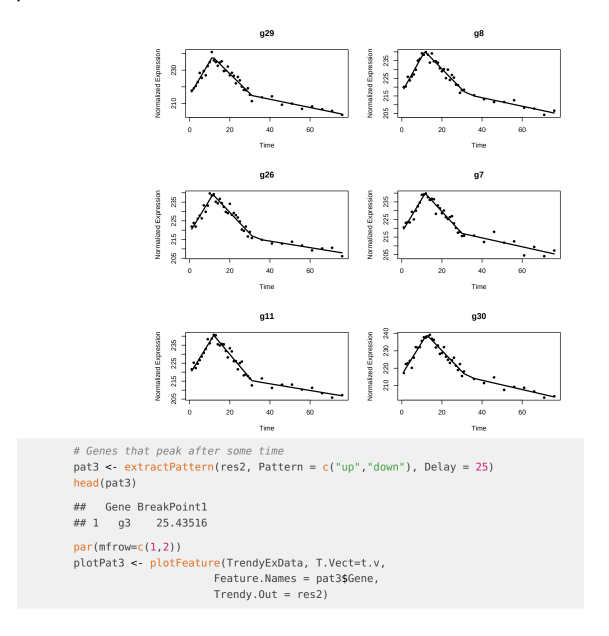


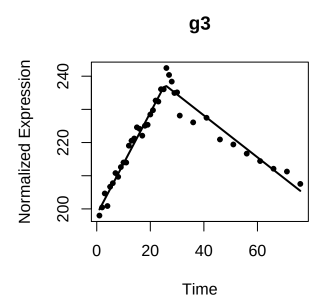
5 Extract genes with certain pattern

Genes that have a peak along the time-course will have fitted trend somewhere as "up-down". Genes that are oscillating may have the fitted trend "up-down". To extract a list of such genes we can use the extractPattern:

```
# Genes that peak
pat1 <- extractPattern(res2, Pattern = c("up", "down"))</pre>
head(pat1)
##
      Gene BreakPoint1
## 5
       g29
               11.35849
##
   2
        g8
               11.43482
## 8
               11.56147
       g26
               11.65663
## 10
        g7
## 4
       g11
               11.99780
## 6
       g30
               12.38463
par(mfrow=c(3,2))
plotPat1 <- plotFeature(TrendyExData, T.Vect=t.v,</pre>
                       Feature.Names = pat1$Gene[1:6],
                       Trendy.Out = res2)
```

Trendy: Segmented regression analysis of expression dynamics for high-throughput ordered profiling experiments





6 Additional options

In the trendy function, the thresholds c_{num} and c_{pval} can be specified via parameters Min.Num.In.Seg and Pval.Cut, respectively.

7 Trendy shiny app

The Trendy shiny app requires an .RData object output from the trendy function, which can be obtained by setting Save.Object=TRUE.

```
res <- trendy(TrendyExData, Max.K=2, Save.Object = TRUE, File.Name="exampleObject")
Then in R run:
library(shiny)
runGitHub('rhondabacher/Trendy')</pre>
```

Trendy Upload .RData object first, then obtain list of genes according to some pattern or visualize genes one by one. Input .Rdata from trendy() run:

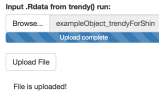


Figure 1: Upload shiny object

Trendy Upload .RData object first, then obtain list of genes according to some pattern or visualize genes one by one. Input .Rdata from trendy() run: Browse... exampleObject_trendyForSh Upload File File is uploaded! Obtain gene patterns Visualize genes Please select a folder for output : Select Output Folder Enter pattern (separate by comma, no spaces): up,down Only consider genes with adjusted R squared greater than: Only consider genes with pattern after time-point: 0 Output a plot of patterned genes? Output file name (will default to pattern) Submit for processing

Figure 2: Find all genes with a given pattern

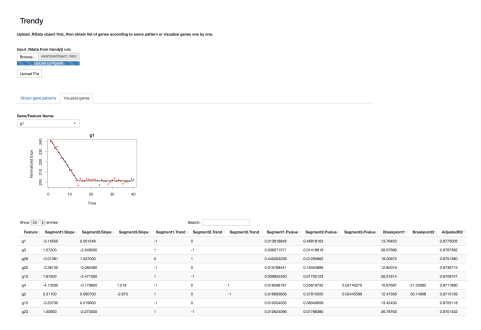


Figure 3: Search genes individually

8 SessionInfo

```
print(sessionInfo())
## R version 3.4.1 (2017-06-30)
## Platform: x86_64-apple-darwin15.6.0 (64-bit)
## Running under: OS X El Capitan 10.11.6
## Matrix products: default
## BLAS: /Library/Frameworks/R.framework/Versions/3.4/Resources/lib/libRblas.0.dylib
## LAPACK: /Library/Frameworks/R.framework/Versions/3.4/Resources/lib/libRlapack.dylib
## locale:
## [1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/c/en_US.UTF-8
## attached base packages:
## [1] stats graphics grDevices utils datasets methods base
##
## other attached packages:
## [1] gplots_3.0.1 Trendy_0.99.0 knitr_1.17
## loaded via a namespace (and not attached):
## [1] Rcpp_0.12.12 gtools_3.5.0 digest_0.6.12
## [4] rprojroot_1.2 bitops_1.0-6 backports_1.1.0
## [7] formatR_1.5 magrittr_1.5 evaluate_0.10.1
## [10] KernSmooth_2.23-15 highr_0.6 stringi_1.1.5
## [13] gdata_2.18.0 rmarkdown_1.6 BiocStyle_2.5.31
## [16] tools_3.4.1 stringr_1.2.0 parallel_3.4.1
                                                           BiocStyle_2.5.31
## [16] tools_3.4.1 stringr_1.2.0 ## [19] yaml_2.1.14 compiler_3.4.1
                                                           segmented_0.5-2.1
## [22] caTools_1.17.1 htmltools_0.3.6
```