Package ‘l1kdeconv’

July 8, 2017

Type Package
Title Deconvolution for LINCS L1000 Data
Version 1.2.0
Date 2017-07-05
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Description LINCS L1000 is a high-throughput technology that allows the gene expression measurement in a large number of assays. However, to fit the measurements of ~1000 genes in the ~500 color channels of LINCS L1000, every two landmark genes are designed to share a single channel. Thus, a deconvolution step is required to infer the expression values of each gene. Any errors in this step can be propagated adversely to the downstream analyses. We present a LINCS L1000 data peak calling R package l1kdeconv based on a new outlier detection method and an aggregate Gaussian mixture model. Upon the remove of outliers and the borrowing information among similar samples, l1kdeconv shows more stable and better performance than methods commonly used in LINCS L1000 data deconvolution.

Imports stats, mixtools, ggplot2
License GPL (>= 2)
Depends R (>= 3.2.0)
RoxygenNote 6.0.1
NeedsCompilation no
Repository CRAN
Date/Publication 2017-07-08 04:41:45 UTC

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getclusterranges

*Get the Cluster Ranges in a Vector of 1D Coordinates*

**Description**

Get the Cluster Ranges in a Vector of 1D Coordinates

**Usage**

```r
getclusterranges(x, gap)
```

**Arguments**

- `x`: a numeric vector
- `gap`: the size for the recognition of data free gaps

**Examples**

```r
x = c(1:3, 11:13)
getclusterranges(x, 3)
```

---

gmmplot

*Plot the Fit Results of 2-Component Gaussian Mixture Model*

**Description**

Plot the Fit Results of 2-Component Gaussian Mixture Model

**Usage**

```r
gmmplot(x, mu1, mu2, sigma, lambda, nbins = 15, xlim)
```

**Arguments**

- `x`: a numeric vector
- `mu1`: the mean of the 1st cluster
- `mu2`: the mean of the 2nd cluster
- `sigma`: the common variance of both clusters
- `lambda`: the proportion parameter
- `nbins`: the number of bins per cluster (6*sigma)
- `xlim`: the limitation of x scale
multigmmmanydata

Examples

```r
set.seed(0)
x=list(c(
  rnorm(150, mean=0)
, rnorm(50, mean=10)
))
fit_res=multigmmsamedistribu(x)

with(
  as.list(fit_res$par_conv)
  , gmmplot(x[[1]]
    , mu1=mu1
    , mu2=mu2
    , sigma=sigma
    , lambda=lambda
    , xlim=range(unlist(x))
  )
)
```

---

**multigmmmanydata**  
*Split the input dataset into several sub list to deconvolution.*

---

**Description**

Due to the limitation of optimization that too many data would dramatically slow down the speed.

**Usage**

```r
multigmmmmanydata(x, grp_size = 3, lambda_lower = 0.1, lambda_upper = 1 - lambda_lower, sigma_lower = 0.01, debug = F)
```

**Arguments**

- `x`: a list of numeric vector
- `grp_size`: the normal group size for each group
- `lambda_lower`: the lower bound of \( \lambda \)
- `lambda_upper`: the upper bound of \( \lambda \)
- `sigma_lower`: the lower bound of \( \sigma \)
- `debug`: enable the debug mode to show par and fn

**Examples**

```r
set.seed(0)
x1=c(rnorm(150, mean=0), rnorm(50, mean=10))
x2=c(rnorm(150, mean=20), rnorm(50, mean=40))
x3=c(rnorm(150, mean=30), rnorm(50, mean=60))
x4=c(rnorm(150, mean=30), rnorm(50, mean=60))
```
multigmmplot

Plot the Fit Results of aggregate 2-Component Gaussian Mixture Model

Usage

```r
multigmmplot(x, fit_res, nbins = 15)
```

Arguments

- `x`: a list of a numeric vector
- `fit_res`: the result of AGMM
- `nbins`: the number of bins per cluster

Examples

```r
params=list(
  c(mu1=0, mu2=10, sd = 1),
  c(mu1=10, mu2=20, sd = 1)
)
set.seed(0)
x=lapply(
  params ,
  function(v) {
    c(
      rnorm(100, mean=v[['mu1']], sd = v[['sd']]),
      rnorm(50, mean=v[['mu2']], sd = v[['sd']])
    )
  }
)
multigmmplot(x, multigmmsamedistribu(x))
```
multigmmsamedistribu

*Description*

Fit Multi 2-Component Gaussian Mixture Model in same distribution with a Fixed Proportion

*Usage*

```r
multigmmsamedistribu(x, lambda_lower = 0.1, lambda_upper = 1 - lambda_lower, sigma_lower = 0.01, debug = F)
```

*Arguments*

- `x`: a list of numeric vector
- `lambda_lower`: the lower bound of \( \lambda \)
- `lambda_upper`: the upper bound of \( \lambda \)
- `sigma_lower`: the lower bound of \( \sigma \)
- `debug`: enable the debug mode to show par and fn

*Examples*

```r
set.seed(0)
x1=c(rnorm(150, mean=0), rnorm(50, mean=10))
x2=c(rnorm(150, mean=20), rnorm(50, mean=40))
x3=c(rnorm(150, mean=30), rnorm(50, mean=60))
x=list(x1, x2, x3)
multigmmsamedistribu(x)
```

multigmmsamedistribulik

*Description*

The sum of Log-Likelihoods of 1D Multi Same Distribution Gaussian Mixture Model

*Usage*

```r
multigmmsamedistribulik(x)
```

*Arguments*

- `x`: a list of numeric vectors
Examples

```r
set.seed(0)
x1 = c(
    rnorm(100, mean=0),
    rnorm(100, mean=1)
)
x = list(x1)
multigmmsamedistribulik(x)(c(0.5, 1, 0.5, 1))
```

---

**multigmmsamedistribumulti**

*Split the input dataset into several sub list to deconvolution.*

---

**Description**

Due to the limitation of optimization that too many data would dramatically slow down the speed.

**Usage**

```r
multigmmsamedistribumulti(x, lambda_lower = 0.1, lambda_upper = 1 - lambda_lower, sigma_lower = 0.01, debug = F)
```

**Arguments**

- **x**: a list of numeric vector
- **lambda_lower**: the lower bound of \( \lambda \)
- **lambda_upper**: the upper bound of \( \lambda \)
- **sigma_lower**: the lower bound of \( \sigma \)
- **debug**: enable the debug mode to show par and fn

**Examples**

```r
set.seed(0)
x1 = c(rnorm(150, mean=0), rnorm(50, mean=10))
x2 = c(rnorm(150, mean=20), rnorm(50, mean=40))
x3 = c(rnorm(150, mean=30), rnorm(50, mean=60))
x4 = c(rnorm(150, mean=30), rnorm(50, mean=60))
x5 = c(rnorm(150, mean=30), rnorm(50, mean=60))
x6 = c(rnorm(150, mean=30), rnorm(50, mean=60))
x = list(x1, x2, x3, x4, x5, x6)
multigmmmmanydata(x)
```
**rmoutlier1d**

Remove the Outliers in a Vector of 1D Coordinates

**Usage**

```
rmoutlier1d(x, dy_thr = dnorm(4), clustersize_thr = 3, gapsize = 10)
```

**Arguments**

- `x`: a numeric vector
- `dy_thr`: the threshold for dy
- `clustersize_thr`: the threshold for cluster size
- `gapsize`: the threshold of points in recognizing data free gap

**Examples**

```r
x <- c(1, 10:30, 50)
par(mfrow=c(2, 1))
plot(density(x))
plot(density(rmoutlier1d(x)))
```

---

**splitgrp**

Split a list with size n into groups with at least m elements

**Description**

Split a list with size n into groups with at least m elements

**Usage**

```
splitgrp(n, m)
```

**Arguments**

- `n`: an integer indicating the total length
- `m`: the min group size

**Examples**

```
splitgrp(1, 2)
splitgrp(2, 2)
splitgrp(3, 2)
```
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