Package 'popPCR'

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Title Classify Digital PCR Droplets by Fitting Fluorescence Populations

Version 0.1.1.1

Description Estimates DNA target concentration by classifying digital PCR (polymerase chain reaction) droplets as positive, negative, or rain, using Expectation-Maximization Clustering. The fitting is accomplished using the 'EMMIXskew' R package (v. 1.0.3) by Kui Wang, Angus Ng, and Geoff McLachlan (2018) as based on their paper ``Multivariate Skew t Mixture Models: Applications to Fluorescence-Activated Cell Sorting Data" <doi:10.1109/DICTA.2009.88>.

Imports mvtnorm

Depends graphics, stats, methods, grDevices, R (>= 2.10)

License GPL (>= 3)

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Author Joyce Emlyn Guiao [aut, cre]

Maintainer Joyce Emlyn Guiao <joyce_emlyn_guiao@dlsu.edu.ph>

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R topics documented:

calculateConc													•				•		2
popPCR																			3
printSummaryConc																			6
printSummaryFit .																			6
x_multiPop																			8
x_onePop																			8
x_twoPop																			9
_ 1																			

10

Index

calculateConc

Description

Mean target copies per partition (lambda) is derived using Poisson distribution as lambda = -ln(nneg / ntot). Target copies in sample is then calculated as conc = lambda * volSamp/(volDrp * 1000).

Usage

calculateConc(nneg, ntotal, volSamp, volDrp)

Arguments

nneg	numeric, negative droplet count
ntotal	numeric, total droplet count
volSamp	numeric, sample volume in microliter
volDrp	numeric, droplet (or partition) volume in nanoliter

Value

Returns a list with 2 named items lambda and conc

- lambda numeric, vector of mean target copies per partition (lambda) and its lower and upper 95% confidence interval
- conc numeric, vector of target copies in sample (based on the given sample volume (volSamp) and droplet volume (volDrp)) and its lower and upper 95% confidence interval

Examples

```
estimates <- calculateConc(5000, 20000, volSamp = 20, volDrp = 0.85)
estimates
    Output:
#
#
       $lambda
#
         lambda lower
                              upper
       1.386294 1.362289 1.410299
#
#
#
       $conc
          conc lower
#
                              upper
#
       32618.69 32053.87
                          33183.51
```

popPCR

Description

Estimates target concentration by counting positive droplets with Poisson correction. Positive, negative, and rain populations are fitted using EM. Droplets are then classified using Maximum A Posteriori rule

Usage

```
popPCR(
    x,
    dist,
    volSamp = 20,
    volDrp = 0.85,
    maxComponents = Inf,
    negProbThres = 1e-07,
    useOnlyNegProbThres = FALSE
)
```

Arguments

x	numeric, vector of droplet fluorescence amplitude					
dist	character, distribution of the mixture models ("normal", "skewed-normal", "t", "skewed-t")					
volSamp	numeric, sample volume in microliter					
volDrp	numeric, droplet (or partition) volume in nanoliter					
maxComponents	numeric, maximum number of components (e.g. populations)					
negProbThres	numeric, if only one population was detected, then its assumed as a negative population. Droplets will be classified as positive if its probability given the population < negProbThres.					
useOnlyNegProbThres						
	logical if TRUE then droplets will be classified as positive if its probability					

logical, if TRUE, then droplets will be classified as positive if its probability given the leftmost population < negProbThres. Default is FALSE, i.e. classification is done by Maximum A Posteriori rule.

Value

Returns a result.popPCR S4 class object with attributes

- · classification character, vector of droplet classification
- dist character, user-specified parameter for the mixture model
- dropletCount list, droplet classification count
- em list, returned value of EMMIXskew's EmSkew()

- estConc list, estimated target concentration as lambda and sample concentration (with 95% CI)
- G numeric, number of components fitted
- memberProb list, component membership probability of all droplets

Examples

```
library(popPCR)
```

```
# Plot histograms of available data
hist(x_onePop, breaks = 100)
hist(x_twoPop, breaks = 100)
hist(x_multiPop, breaks = 100)
# ---- Mixture model fitting ---- #
# Example 1. One population sample
result <- popPCR(x_onePop, dist = "t")</pre>
printSummaryConc(result)
#
    Output:
#
         Populations detected : 1
#
         Total droplets : 8000
         Positive : 1 (0.01%)
#
#
         Negative : 7999 (99.99%)
#
#
         Target copies in sample
                                         : 2.9414 ( 95% CI: [ -2.8237 , 8.7064 ] )
#
         Mean target copies per partition : 1e-04 ( 95% CI: [ -1e-04 , 4e-04 ] )
printSummaryFit(result)
     Output:
#
         Results of fitting a 1-component t mixture model
#
#
         Negative Population
#
         Mix prop. : 1
#
#
         Mu
             : 1024.1614
#
         Sigma
                   : 35253.1747
         Dof
                  : 2.005
#
# (Option) increase negProbThres to classify negative droplets more strictly
result <- popPCR(x_onePop, dist = "t", negProbThres = 1e-4)</pre>
printSummaryConc(result)
#
     Output:
         Populations detected : 1
#
#
         Total droplets : 8000
         Positive : 691 (8.64%)
#
         Negative : 7309 (91.36%)
#
#
                                       : 2125.5312 ( 95% CI: [ 1966.9936 , 2284.0688 ] )
#
        Target copies in sample
#
         Mean target copies per partition : 0.0903 ( 95% CI: [ 0.0836 , 0.0971 ] )
# Example 2. Two population sample
result <- popPCR(x_twoPop, dist = "t")</pre>
printSummaryConc(result)
#
     Output:
```

popPCR

```
#
         Populations detected : 2
#
         Total droplets : 10254
#
         Positive : 8693 (84.78%)
#
        Negative : 1561 (15.22%)
#
#
       Target copies in sample
                                     : 44290.3819 ( 95% CI: [ 43215.6408 , 45365.1231 ] )
#
        Mean target copies per partition : 1.8823 ( 95% CI: [ 1.8367 , 1.928 ] )
printSummaryFit(result)
#
     Output:
#
        Results of fitting a 2-component t mixture model
#
        Negative Population
#
#
        Mix prop. : 0.1522
#
        Mu
                  : 2136.7435
         Sigma
                   : 4126.8357
#
#
        Dof
                   : 12.3562
#
#
        Positive Population
#
        Mix prop. : 0.8478
#
        Mu
                  : 7580.1275
#
         Sigma
                   : 42621.1894
#
         Dof
                  : 2.415
# Example 3. Multiple population sample
result <- popPCR(x_multiPop, dist = "t", maxComponents = 4)</pre>
printSummaryConc(result)
     Output:
#
#
        Populations detected : 4
#
         Total droplets : 1814
#
        Positive : 44 (2.43%)
#
        Negative : 1252 (69.02%)
#
        Rain (1) : 258 (14.22%)
#
        Rain (2) : 260 (14.33%)
#
#
        Target copies in sample : 8724.5195 ( 95% CI: [ 7999.0578 , 9449.9812 ] )
#
        Mean target copies per partition : 0.3708 ( 95% CI: [ 0.34 , 0.4016 ] )
# In the output above, we see 2 rain populations! Let's examine its plot.
plot(stats::density(x_multiPop))
# We can see that Rain (1) is very close to the Negative population.
# Let's include droplets in Rain (1) in the negative droplet count.
nNegative <- result@dropletCount$neg + result@dropletCount$rain1</pre>
nTotal <- result@dropletCount$total</pre>
# Re-estimate concentration as follows
newEstimates <- calculateConc(nNegative, nTotal, volSamp = 20, volDrp = 0.85)</pre>
newEstimates
     Output:
#
#
        $lambda
           lambda
#
                      lower
                                upper
#
        0.1834247 0.1627763 0.2040731
#
#
        $conc
```

conc lower upper # 4315.875 3830.031 4801.719

printSummaryConc Print result summary of popPCR

Description

Summarizes the number of populations detected, total droplets, and number of classified positive, negative, and rain droplets. Also calculates the target copies in sample and the mean target copies per partition (lambda).

Usage

printSummaryConc(result.popPCR)

Arguments

result.popPCR returned value of popPCR()

Examples

```
result <- popPCR(x_twoPop, dist = "t")</pre>
printSummaryConc(result)
#
     Output:
#
         Populations detected : 2
         Total droplets : 10254
#
#
         Positive : 8693 (84.78%)
         Negative : 1561 (15.22%)
#
#
#
       Target copies in sample
                                       : 44290.3819 ( 95% CI: [ 43215.6408 , 45365.1231 ] )
#
         Mean target copies per partition : 1.8823 (95% CI: [ 1.8367 , 1.928 ] )
```

printSummaryFit Print fitted mixture model estimates from popPCR

Description

Summarizes the number of populations fitted and their estimate distribution parameters. If only 1 population was detected, then it is assumed and is identified to be a negative population. If 2 populations were detected, then the leftmost is identified as the Negative Population and the rightmost is the Positive Population. If 3 or more populations were detected, then the populations between the leftmost and the rightmost will be considered as Rain Populations; which are numbered to make it identifiable in case of multiple Rain Populations (i.e. Rain (1) and Rain (2)).

printSummaryFit

Usage

printSummaryFit(result.popPCR)

Arguments

result.popPCR returned value of popPCR()

Examples

```
result <- popPCR(x_twoPop, dist = "t")</pre>
printSummaryFit(result)
#
    Output:
#
        Results of fitting a 2-component t mixture model
#
#
        Negative Population
#
        Mix prop. : 0.1522
#
        Mu
                 : 2136.7435
#
        Sigma : 4126.8357
                  : 12.3562
#
        Dof
#
        Positive Population
#
#
        Mix prop. : 0.8478
#
        Mu
                 : 7580.1275
#
        Sigma
                : 42621.1894
#
        Dof
                  : 2.415
result <- popPCR(x_multiPop, dist = "t", maxComponents = 4)</pre>
printSummaryFit(result)
     Output:
#
#
        Results of fitting a 4-component t mixture model
#
        Negative Population
#
        Mix prop. : 0.6896
#
              : 1452.1416
#
        Mu
                  : 12526.8931
#
        Sigma
#
        Dof
                 : 21.3612
#
        Rain (1) Population
#
#
        Mix prop. : 0.142
                : 2142.1118
#
        Mu
#
        Sigma
               : 10762.5474
#
        Dof
                 : 186.2947
#
        Rain (2) Population
#
        Mix prop. : 0.1457
#
                 : 5119.0039
#
        Mu
#
                  : 334959.2499
        Sigma
                  : 2.3626
#
        Dof
#
#
        Positive Population
        Mix prop. : 0.0227
#
            : 8505.9682
#
        Mu
#
        Sigma : 192858.9044
```

Dof : 149.8677

x_multiPop dPCR sample w/ >=3 populations

Description

The reaction with ID 373 in the Dataset.zip provided in the repository from Lievens et. al. (2017)

Usage

x_multiPop

Format

A numeric vector with 1814 droplet fluorescence

Source

https://github.com/Gromgorgel/ddPCR/blob/master/Dataset.zip

x_onePop

dPCR sample w/ 1 population

Description

Simulated dataset with very high presence of rain and true mean copies per partition of 0.1.

Usage

x_onePop

Format

A numeric vector with 8000 droplet fluorescence

x_twoPop

Description

The reaction with ID 9 in the Dataset.zip provided in the repository from Lievens et. al. (2017)

Usage

x_twoPop

Format

A numeric vector with 10254 droplet fluorescence

Source

https://github.com/Gromgorgel/ddPCR/blob/master/Dataset.zip

Index

* datasets
 x_multiPop, 8
 x_onePop, 8
 x_twoPop, 9

calculateConc, 2

popPCR, 3
printSummaryConc, 6
printSummaryFit, 6

x_multiPop, 8
x_onePop, 8
x_twoPop, 9