

Package: digitalPCR (via r-universe)

October 15, 2024

Type Package

Title Estimate Copy Number for Digital PCR

Version 1.1.0

Depends stats

Date 2016-03-07

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Description The assay sensitivity is the minimum number of copies that the digital PCR assay can detect. Users provide serial dilution results in the format of counts of positive and total reaction wells. The output is the estimated assay sensitivity and the copy number per well in the initial dilute.

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NeedsCompilation no

Date/Publication 2016-03-09 01:19:13

Repository <https://xutaodeng.r-universe.dev>

RemoteUrl <https://github.com/cran/digitalPCR>

RemoteRef HEAD

RemoteSha 3aa15672006597a3af6f91fb008acb9b97c3666b

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`digitalPCR`*Estimate copy number and assay sensitivity from dPCR serial dilution*

Description

The assay sensitivity is the minimum number of copies that the digital PCR assay can detect. Users provide serial dilution results in the format of counts of positive and total reaction wells. The output is the estimated assay sensitivity and the copy number per well in the initial dilute.

Usage

```
digitalPCR(pos, neg, dilution, Nboot, single.copy = c("FALSE", "TRUE"), upper.copy=100)
```

Arguments

<code>pos</code>	a numeric vector of number of positive reactions in the order of dilutions
<code>neg</code>	a numeric vector of number of negative reactions in the order of dilutions
<code>dilution</code>	a numeric vector of folds of dilutions relative to the initial dilution
<code>Nboot</code>	number of bootstrapping
<code>single.copy</code>	character tells whether the assay is assumed single-copy sensitive. default "FALSE"
<code>upper.copy</code>	the upper bound of copy number in highest concentration

Details

the `pos`, `neg` and `dilution` vector must be in the same order from high to low concentrations.

Value

The returned value is a list of four components:

<code>mean copy number</code>	mean copy number
<code>sd copy number</code>	standard deviation of copy number
<code>copy numbers</code>	bootstrapped copy number estimates
<code>thresholds</code>	bootstrapped assay sensitivity estimates

Author(s)

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Examples

```
#dilution at 1,2,4,8 fold
pos=c(221,97,39,14)
dilution=c(1,2,4,8)
neg=384-pos
#note in practice, set this 100 or larger
Nboot=10

result1=digitalPCR(pos, neg, dilution, Nboot)
print (paste(result1$"mean copy number", result1$"sd copy number"))
hist(result1$"thresholds",xlim=c(-2,10))

#the following example contains only 1 dilution
result2=digitalPCR(230, 355, 1, Nboot, "TRUE")
```

digitalPCR-internal *Internal functions*

Description

Internal functions

Author(s)

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