Package 'qPCRtools'

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Version 0.1.1
Title Tools for qPCR
Description A set of tools for qPCR data process.
URL https://github.com/lixiang117423/qPCRtools
BugReports https://github.com/lixiang117423/qPCRtools/issues
License MIT + file LICENSE
Imports broom, data.table, dplyr, ggplot2, ggpmisc, ggthemes, magrittr, multcomp, readxl, reshape2, rstatix, sjmisc, stringr, tibble, tidyr, xlsx
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CalCurve Standard Curve Calculation.

Description

The function can calculate the standard curve. At the same time, it can get the amplification efficiency of primer(s). Based on the amplification efficiency, we can know which method can be used to calculate the expression level.

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Arguments

cq.table The data frame of the position and Cq value.

concen.table The data frame of the position and concentration.

highest.concen The highest concentration for calculation. lowest.concen The lowest concentration for calculation.

dilution Dilution factor of cDNA template. The default value is 4.

by mean Cq value or not. The default value is TRUE.

Value

A list.

Author(s)

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Examples

```
df.1.path <- system.file("examples", "calsc.cq.txt", package = "qPCRtools")
df.2.path <- system.file("examples", "calsc.info.txt", package = "qPCRtools")
df.1 <- data.table::fread(df.1.path)
df.2 <- data.table::fread(df.2.path)
CalCurve(
    cq.table = df.1,
    concen.table = df.2,
    lowest.concen = 4,
    highest.concen = 4096,
    dilu = 4,
    by = "mean"
) -> p

p[["table"]]
p[["figure"]]
```

CalExp2ddCt

Calculate expression using standard curve.

Description

Calculate expression using standard curve.

Arguments

cq.table The data frame of the position and cq value.

design.table The data frame of the position and corresponding information.

correction Correct expression value by reference gene.

ref.gene The name of reference gene.
ref.group The name of reference group.

 ${\tt stat.method} \qquad {\tt Statistical\ method}.$

fig.type Calculation by mean cq value or not.

fig.ncol Number of columes of figure.

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Value

A list contain a table and a figure.

Author(s)

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Examples

 ${\tt CalExpCurve}$

Calculate expression using standard curve.

Description

Calculate expression using standard curve.

Arguments

cq.table The data frame of the position and Cq value.

 ${\tt design.table} \qquad {\tt The \ data \ frame \ of \ the \ position \ and \ corresponding \ information}.$

correction Correct expression value by reference gene.

ref.gene The name of reference gene.

stat.method Statistical method.

ref.group The name of reference group.

fig. type Calculation by mean Cq value or not.

fig.ncol Number of columes of figure.

Value

A list contain a table and a figure.

Author(s)

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Examples

```
df1.path = system.file("examples", "cal.exp.curve.cq.txt", package = "qPCRtools")
df2.path = system.file("examples", "cal.expre.curve.sdc.txt", package = "qPCRtools")
df3.path = system.file("examples", "cal.exp.curve.design.txt", package = "qPCRtools")
cq.table = data.table::fread(df1.path)
curve.table = data.table::fread(df2.path)
design.table = data.table::fread(df3.path)
CalExpCurve(
   cq.table,
   curve.table,
   design.table,
   correction = TRUE,
   ref.gene = "OsUBQ",
   stat.method = "t.test",
   ref.group = "CK",
   fig.type = "box",
   fig.ncol = NULL) -> res
res[["table"]]
res[["figure"]]
```

CalExpRqPCR

Calculate expression using standard curve.

Description

Calculate expression using standard curve.

Arguments

cq.table The data frame of the position and cq value.

design.table The data frame of the position and corresponding information.

correction Correct expression value by reference gene.

ref.gene The name of reference gene.
ref.group The name of reference group.

stat.method Statistical method.

fig.type Calculation by mean cq value or not.

fig.ncol Number of columes of figure.

Value

A list contain a table and a figure.

Author(s)

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Examples

CalRTable

Calculate RNA volume for reverse transcription.

Description

The first step of qPCR is usually the preparation of cDNA. We need to calculate the column of RNA for reverse transcription to cDNA. So, if we have the concentration of RNA, we can use the function 'CalRTable' to do that.

Arguments

data A data frame contained the sample names and the concentration value.

The default unit of concentration is ng/uL.

template A data.frame contained the information of reverse transcription.

RNA.weight RNA weight required for reverse transcription. Default is 1 ug.

Value

A list contain a table and a figure.

Author(s)

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Examples

```
df.1.path <- system.file("examples", "crtv.data.txt", package = "qPCRtools")
df.2.path <- system.file("examples", "crtv.template.txt", package = "qPCRtools")
df.1 <- data.table::fread(df.1.path)
df.2 <- data.table::fread(df.2.path)
result <- CalRTable(data = df.1, template = df.2, RNA.weight = 2)
head(result)</pre>
```

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