Package 'OryzaProbe'

October 17, 2022

Title Rice Microarray Probe ID Conversion, from Probe ID to RAP-DB ID

Type Package

```
Version 0.1.0
Description
      Microarray probe ID is not convenient for further enrichment analysis and target gene selection.
      The package is created for the rice microarray probe ID conversion.
      This package can convert microarray probe ID from GPL6864 <a href="https:">https:</a>
      //www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GPL6864>, GPL8852 <https:</pre>
      //www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GPL8852>, and GPL2025 < https:
      //www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GPL2025> platforms to RAP-
      DB ID. RAP-DB "The Rice Annota-
      tion Project Database" <a href="https://rapdb.dna.affrc.go.jp">https://rapdb.dna.affrc.go.jp</a> is a well-
      known database for rice Oryza sativa, and the gene ID in this database is widely used in many ar-
      eas related to rice research.
      For multiple probes representing a single gene, This package can merge them by tak-
      ing the mean, max, or min value of these probes.
      Or we can keep multiple probes by appending sequence numbers to duplicate the RAP-DB ID.
License GPL-3
Encoding UTF-8
Imports stats
Depends R (>= 2.10)
RoxygenNote 7.2.1
Suggests knitr, rmarkdown, testthat (>= 3.0.0)
Config/testthat/edition 3
VignetteBuilder knitr
NeedsCompilation no
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Repository CRAN
```

Date/Publication 2022-10-17 11:55:27 UTC

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Description

Convert rice microarray probe ID to RAP-DB ID and merge multiple probes for one gene

Usage

```
probeConvert(exprMatrix, probeMerge = FALSE, mergeBy = "mean")
```

Arguments

exprMatrix A expression matrix is required. Row names of the matrix should be the mi-

croarray probe ID.

probeMerge A instruction is needed. If probeMerge = F, the multiple probes for one gene

will not be merged. If probeMerge = T, the multiple probes for one gene will be

merged. The default value is F.

mergeBy A method for merging multiple probes for one gene if probeMerge = T. Mean,

max and min value is available.

Value

A matrix.

Examples

```
expr <- system.file("test_file", header = TRUE, sep="\t", header=TRUE, stringsAsFactors = FALSE)
formatted_expr <- probeConvert(exprMatrix = expr)
formatted_expr1 <- probeConvert(exprMatrix = expr, probeMerge = TRUE)</pre>
```

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