Package 'mammaPrintData'

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Title RGLists from the Glas and Buyse breast cancer studies
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Date 2013-05-13
Description Gene expression data for the two breast cancer cohorts published by Glas and Buyse in 2006
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Author Luigi Marchionni <marchion@jhu.edu></marchion@jhu.edu>
Maintainer Luigi Marchionni <marchion@jhu.edu></marchion@jhu.edu>
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License Artistic-2.0
URL http://luigimarchionni.org/breastTSP.html
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R topics documented: mammaPrintData-package buyseRGcy3 buyseRGcy5 glasRGcy3 glasRGcy5
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mammaPrintData-package

Gene expression, annotations and clinical information for the Glas and Buse cohorts

Description

This package contains the two RGList-class instances corresponding to the breast cancer patients' cohorts published by Glas and colleagues in BMC Genomics (2006), and by Buyse and colleagues in JNCI (2006). Since in both studies a two-colors dye swap design was applied, this package contains two distinct RGList-class instances for each data set, one for each dye-swap set of hybridizations. This package contains unprocessed data, as obtained from the original raw data files available from the ArrayExpress repository.

Usage

data(glasRG)
data(buyseRGa)

Format

The RGList-class instances contained in this package (glasRGcy5, glasRGcy5, buyseRGcy5, and buyseRGcy5, accounts for 1900 microarray features and 162 and 307 samples respectively.

Details

Package: mammaPrintData

Type: Package
Version: 0.99.5.
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Author(s)

Luigi Marchionni <marchion@gmail.com>

Source

http://www.ncbi.nlm.nih.gov/pubmed/?term=17074082 http://www.ncbi.nlm.nih.gov/pubmed/ ?term=16954471

References

Annuska M Glas et al., "Converting a breast cancer microarray signature into a high-throughput diagnostic test", *BMC Genomics*, 2006, **7**:278-288

Marc Buyse et al., "Validation and clinical utility of a 70-gene prognostic signature for women with node-negative breast cancer", *JNCI*, 2006, **98**(17):1183-1192

See Also

See RGList-class

Examples

```
### load limma package
library(limma)
### load the glasRG dataset: two different dye-swap sets:
###glasRGcy5: information provided for RNA labeled with Cy5
###glasRGcy3: information provided for RNA labeled with Cy3
data(glasRG)
### load the buyseRG dataset: two different dye-swap sets:
###buyseRGcy5: reference RNA was labeled with Cy5
###buyseRGcy3: reference RNA was labeled with Cy3
data(buyseRG)
### show the class of the glasRGcy3 dataset
class(glasRGcy5)
### show the class of the buyseRGcy3 dataset
class(buyseRGcy3)
### show the dimentions of the glasRGcy3 dataset
dim(glasRGcy5)
### show the dimentions of the buyseRGcy3 dataset
dim(buyseRGcy3)
### show the first 10 rows of the glasRGcy3 phenotype data
head(glasRGcy3$targets, n=10)
### show the first 10 rows of the buyseRGcy5 phenotype data
head(buyseRGcy5$targets, n=10)
### show the first 10 features of the glasRGcy5 phenotype data
head(glasRGcy5$genes, n=10)
### show the first 10 features of the buyseRGcy3 phenotype data
head(buyseRGcy3$genes, n=10)
```

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buyseRGcy3	Gene expression, annotations and clinical information for the Buyse cohort: set of dye-swap hybridizations in which the reference RNA was labeled with Cy3

Description

This package contains the RGList-class instance (see limma) for the gene expression data set published by Buyse and colleagues. This object contains the set of dye-swap hybridizations in which the reference RNA was labeled with Cy3.

Usage

data(buyseRG)

Format

The buyseRGcy3 RGList-class contains 1900 microarray features and 307 samples. The following components were included:

- buyseRGcy3\$targets: a data.frame for the breast cancer patients clinical information;
- buyseRGcy3\$genes: a data. frame containing the microarray annotations;
- buyseRGcy3\$R: the raw median gene expression foreground intensities from the Red channel;
- buyseRGcy3\$Rb: the raw median gene expression background intensities from the Red channel;
- buyseRGcy3\$G: the raw median gene expression foreground intensities from the Green channel:
- buyseRGcy3\$Gb: the raw median gene expression background intensities from the Green channel;
- buyseRGcy3\$logRatio: the Log ratio between Red and Green channel, as available from the raw data files;
- buyseRGcy3\$logRatioError: the Log ratio Error associated with the Log Ratio between Red and Green channel, as available from the raw data files;
- buyseRGcy3\$ID: the microarray features identifiers, as available from the raw data files;

Details

This dataset corresponds to the breast cancer patients' cohort published by Buyse and collleagues in JNCI (2006).

Source

http://www.ncbi.nlm.nih.gov/pubmed/?term=16954471

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References

Marc Buyse et al., "Validation and clinical utility of a 70-gene prognostic signature for women with node-negative breast cancer", *JNCI*, 2006, **98**(17):1183-1192

See Also

```
See RGList-class
```

Examples

```
### load limma package
library(limma)

### load the buyseRGcy3 dataset: dye-swap set in which the reference
###RNA was labeled with Cy3 (from the Glas cohort)
data(buyseRG)

### show the class of the buyseRGcy3 dataset
class(buyseRGcy3)

### show the dimentions of the buyseRGcy3 dataset
dim(buyseRGcy3)

### show the first 10 rows of the buyseRGcy3 phenotype data
head(buyseRGcy3$targets, n=10)

### show the first 10 features of the buyseRGcy3 phenotype data
head(buyseRGcy3$genes, n=10)
```

buyseRGcy5

Gene expression, annotations and clinical information for the Buyse cohort: set of dye-swap hybridizations in which the reference RNA was labeled with Cy5

Description

This package contains the RGList-class instance (see limma) for the gene expression data set published by Buyse and colleagues. This object contains the set of dye-swap hybridizations in which the reference RNA was labeled with Cy5.

Usage

```
data(buyseRG)
```

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Format

The buyseRGcy5 RGList-class contains 1900 microarray features and 307 samples. The following components were included:

- buyseRGcy5\$targets: a data.frame for the breast cancer patients clinical information;
- buyseRGcy5\$genes: a data. frame containing the microarray annotations;
- buyseRGcy5\$R: the raw median gene expression foreground intensities from the Red channel;
- buyseRGcy5\$Rb: the raw median gene expression background intensities from the Red channel;
- buyseRGcy5\$G: the raw median gene expression foreground intensities from the Green channel:
- buyseRGcy5\$Gb: the raw median gene expression background intensities from the Green channel;
- buyseRGcy5\$logRatio: the Log ratio between Red and Green channel, as available from the raw data files;
- buyseRGcy5\$logRatioError: the Log ratio Error associated with the Log Ratio between Red and Green channel, as available from the raw data files;
- buyseRGcy5\$ID: the microarray features identifiers, as available from the raw data files;

Details

This dataset corresponds to the breast cancer patients' cohort published by Buyse and collleagues in JNCI (2006).

Source

```
http://www.ncbi.nlm.nih.gov/pubmed/?term=16954471
```

References

Marc Buyse et al., "Validation and clinical utility of a 70-gene prognostic signature for women with node-negative breast cancer", *JNCI*, 2006, **98**(17):1183-1192

See Also

```
See RGList-class
```

Examples

```
### load limma package
library(limma)

### load the buyseRGcy5 dataset: dye-swap set in which the reference
###RNA was labeled with Cy3 (from the Glas cohort)
data(buyseRG)

### show the class of the buyseRGcy5 dataset
class(buyseRGcy5)
```

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```
### show the dimentions of the buyseRGcy5 dataset
dim(buyseRGcy5)

### show the first 10 rows of the buyseRGcy5 phenotype data
head(buyseRGcy5$targets, n=10)

### show the first 10 features of the buyseRGcy5 phenotype data
head(buyseRGcy5$genes, n=10)
```

glasRGcy3

Gene expression, annotations and clinical information for the Glas cohort: set of dye-swap hybridizations in which the information was associated with RNA samples labeled with Cy3

Description

This package contains the RGList-class instance (see limma) for the gene expression data set published by Glas and colleagues. This object contains the set of dye-swap hybridizations in which the information provided in the ArrayExpress SDRF table was associated with the RNA samples labeled with Cy3.

Usage

data(glasRG)

Format

The glasRGcy3 RGList-class contains 1900 microarray features and 307 samples. The following components were included:

- glasRGcy3\$targets: a data. frame for the breast cancer patients clinical information;
- glasRGcy3\$genes: a data.frame containing the microarray annotations;
- glasRGcy3\$R: the raw median gene expression foreground intensities from the Red channel;
- glasRGcy3\$Rb: the raw median gene expression background intensities from the Red channel;
- glasRGcy3\$G: the raw median gene expression foreground intensities from the Green channel;
- glasRGcy3\$Gb: the raw median gene expression background intensities from the Green channel:
- glasRGcy3\$logRatio: the Log ratio between Red and Green channel, as available from the raw data files;
- glasRGcy3\$logRatioError: the Log ratio Error associated with the Log Ratio between Red and Green channel, as available from the raw data files;
- glasRGcy3\$ID: the microarray features identifiers, as available from the raw data files;

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Details

This dataset corresponds to the breast cancer patients' cohort published by Glas and collleagues in BMC Genomics (2006).

Source

```
http://www.ncbi.nlm.nih.gov/pubmed/?term=17074082
```

References

Annuska M Glas et al., "Converting a breast cancer microarray signature into a high-throughput diagnostic test", *BMC Genomics*, 2006, **7**:278-288

See Also

```
See RGList-class
```

Examples

```
### load limma package
library(limma)

### load the glasRGcy3 dataset: dye-swap set in which the information provided
###inthe SDRF table was associated with the samples labeled with Cy3
data(glasRG)

### show the class of the glasRGcy3 dataset
class(glasRGcy3)

### show the dimentions of the glasRGcy3 dataset
dim(glasRGcy3)

### show the first 10 rows of the glasRGcy3 phenotype data
head(glasRGcy3$targets, n=10)

### show the first 10 features of the glasRGcy3 phenotype data
head(glasRGcy3$genes, n=10)
```

glasRGcy5

Gene expression, annotations and clinical information for the Glas cohort: set of dye-swap hybridizations in which the information was associated with RNA samples labeled with Cy5

Description

This package contains the RGList-class instance (see limma) for the gene expression data set published by Glas and colleagues. This object contains the set of dye-swap hybridizations in which the information provided in the ArrayExpress SDRF table was associated with the RNA samples labeled with Cy5.

glasRGcy5

Usage

```
data(glasRG)
```

Format

The glasRGcy5 RGList-class contains 1900 microarray features and 307 samples. The following components were included:

- glasRGcy5\$targets: a data. frame for the breast cancer patients clinical information;
- glasRGcy5\$genes: a data. frame containing the microarray annotations;
- glasRGcy5\$R: the raw median gene expression foreground intensities from the Red channel;
- glasRGcy5\$Rb: the raw median gene expression background intensities from the Red channel;
- glasRGcy5\$G: the raw median gene expression foreground intensities from the Green channel;
- glasRGcy5\$Gb: the raw median gene expression background intensities from the Green channel:
- glasRGcy5\$logRatio: the Log ratio between Red and Green channel, as available from the raw data files;
- glasRGcy5\$logRatioError: the Log ratio Error associated with the Log Ratio between Red and Green channel, as available from the raw data files;
- glasRGcy5\$ID: the microarray features identifiers, as available from the raw data files;

Details

This dataset corresponds to the breast cancer patients' cohort published by Glas and collleagues in BMC Genomics (2006).

Source

```
http://www.ncbi.nlm.nih.gov/pubmed/?term=17074082
```

References

Annuska M Glas et al., "Converting a breast cancer microarray signature into a high-throughput diagnostic test", *BMC Genomics*, 2006, 7:278-288

See Also

```
See RGList-class
```

Examples

```
### load limma package
library(limma)

### load the glasRGcy3 dataset: dye-swap set in which the information provided
###inthe SDRF table was associated with the samples labeled with Cy5
data(glasRG)
```

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show the class of the glasRGcy5 dataset
class(glasRGcy5)

show the dimentions of the glasRGcy5 dataset
dim(glasRGcy5)

show the first 10 rows of the glasRGcy5 phenotype data
head(glasRGcy5\$targets, n=10)

show the first 10 features of the glasRGcy5 phenotype data head(glasRGcy5\$genes, n=10)

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