# ${\bf Package\ 'breast Cancer MAINZ'}$

October 17, 2019

Type Package
<b>Title</b> Gene expression dataset published by Schmidt et al. [2008] (MAINZ).
Version 1.22.0
<b>Date</b> 2011-02-10
<b>Description</b> Gene expression data from the breast cancer study published by Schmidt et al. in 2008, provided as an eSet.
biocViews ExperimentData, CancerData, BreastCancerData, MicroarrayData, GEO
<b>Author</b> Markus Schroeder, Benjamin Haibe- Kains, Aedin Culhane, Christos Sotiriou, Gianluca Bontempi, John Quackenbush
Maintainer Markus Schroeder <mschroed@jimmy.harvard.edu>, Benjamin Haibe-Kains    </mschroed@jimmy.harvard.edu>
<b>Depends</b> R (>= 2.5.0)
Suggests survcomp, genefu, Biobase
LazyLoad yes
License Artistic-2.0
<pre>URL http://compbio.dfci.harvard.edu/</pre>
git_url https://git.bioconductor.org/packages/breastCancerMAINZ
git_branch RELEASE_3_9
git_last_commit 41b8b64
git_last_commit_date 2019-05-02
Date/Publication 2019-10-17
R topics documented:
mainz
Index

2 mainz

mainz

Gene expression, annotations and clinical data from Schmidt et al. 2008

#### **Description**

This dataset contains the gene expression, annotations and clinical data as published in Schmidt et al. 2008.

### Usage

data(mainz)

#### **Format**

ExpressionSet with 22283 features and 200 samples, containing:

- exprs(mainz): Matrix containing gene expressions as measured by Affymetrix hgu133a technology (single-channel, oligonucleotides).
- fData(mainz): AnnotatedDataFrame containing annotations of Affy microarray platform hgu133a.
- pData(mainz): AnnotatedDataFrame containing Clinical information of the breast cancer patients whose tumors were hybridized.
- experimentalData(mainz): MIAME object containing information about the dataset.
- annotation(mainz): Name of the affy chip.

### Details

This dataset represents the study published by Schmidt et al. 2008.

• Abstract: Estrogen receptor (ER) expression and proliferative activity are established prognostic factors in breast cancer. In a search for additional prognostic motifs, we analyzed the gene expression patterns of 200 tumors of patients who were not treated by systemic therapy after surgery using a discovery approach. After performing hierarchical cluster analysis, we identified coregulated genes related to the biological process of proliferation, steroid hormone receptor expression, as well as B-cell and T-cell infiltration. We calculated metagenes as a surrogate for all genes contained within a particular cluster and visualized the relative expression in relation to time to metastasis with principal component analysis. Distinct patterns led to the hypothesis of a prognostic role of the immune system in tumors with high expression of proliferation-associated genes. In multivariate Cox regression analysis, the proliferation metagene showed a significant association with metastasis-free survival of the whole discovery cohort [hazard ratio (HR), 2.20; 95% confidence interval (95% CI), 1.40-3.46]. The B-cell metagene showed additional independent prognostic information in carcinomas with high proliferative activity (HR, 0.66; 95% CI, 0.46-0.97). A prognostic influence of the B-cell metagene was independently confirmed by multivariate analysis in a first validation cohort enriched for high-grade tumors (n = 286; HR, 0.78; 95% CI, 0.62-0.98) and a second validation cohort enriched for younger patients (n = 302; HR, 0.83; 95% CI, 0.7-0.97). Thus, we could show in three cohorts of untreated, node-negative breast cancer patients that the humoral immune system plays a pivotal role in metastasis-free survival of carcinomas of the breast.

mainz 3

#### **Source**

http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE11121

#### References

Marcus Schmidt and Daniel Boehm and Christian von Toerne and Eric Steiner and Alexander Puhl and Heryk Pilch and Hans-Anton Lehr and Jan G. Hengstler and Hainz Koelbl and Mathias Gehrmann (2008)"The Humoral Immune System Has a Key Prognostic Impact in Node-Negative Breast Cancer", *Cancer Research*, **68**(13):5405-5413

## **Examples**

```
## load Biobase package
library(Biobase)
## load the dataset
data(mainz)
## show the first 5 rows and columns of the expression data
exprs(mainz)[1:5,1:5]
## show the first 6 rows of the phenotype data
head(pData(mainz))
## show first 20 feature names
featureNames(mainz)[1:20]
## show the experiment data summary
experimentData(mainz)
## show the used platform
annotation(mainz)
## show the abstract for this dataset
abstract(mainz)
```

# Index

\*Topic datasets mainz, 2

mainz, 2