

# Package ‘metaArray’

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**Title** Integration of Microarray Data for Meta-analysis

**Version** 1.58.0

**Author** Debasish Ghosh <ghoshd@psu.edu> Hyungwon Choi  
<hyung\_won\_choi@nuhs.edu.sg>

**Imports** Biobase, MergeMaid, graphics, stats

**Description** 1) Data transformation for meta-analysis of microarray  
Data: Transformation of gene expression data to signed  
probability scale (MCMC/EM methods) 2) Combined differential  
expression on raw scale: Weighted Z-score after stabilizing  
mean-variance relation within platform

**Maintainer** Hyungwon Choi <hyung\_won\_choi@nuhs.edu.sg>

**License** LGPL-2

**biocViews** Microarray, DifferentialExpression

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## R topics documented:

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mdata *metaArray sample dataset*

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### Description

Three datasets from liver, lung, and prostate cancer microarrays. Please refer to the bibliography in the vignette. Chen (30 primary, 9 metastatic), Garber (30 primary, 6 metastatic), Lapointe (30 primary, 9 metastatic)

### Usage

`data(mdata)`

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