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Biostatisti 2018 Autie Otantocen & Treatment

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ABOUT US

Our mission, vision & core values

Leadership

History

Equality, diversity & inclusion

Annual report

Give to MSK

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Bayesian Cell Mixture Model (CMM)

This is a zip file containing R code to implement CMM for analyzing tissue microarray data.

Download zip file

Reference: Shen, Taylor, and Ghosh (2008) Reconstructing tumor-wise protein expression in tissue microarray studies using a Bayesian cell mixture model. Bioinformatics. 24(24):2880-6.

Eigengene-based Linear Discriminant Analysis (ELDA)

This is a zip file containing R code to implement ELDA for tumor classification problems using gene expression microarray data. The README file describes how to use the code.

Download zip file

Reference: Shen et al. (2006) Eigengene-based linear discriminant model for tumor classification using gene expression microarray data. Bioinformatics. 22(21):2635-42.

- Download iCluster R package
- Download CMM R code
- Download ELDA R code

PREVIOUS

Hybrid Bayesian Laplacian Approach

NEXT

iClusterPlus

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