

# Package ‘BPM’

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**Type** Package

**Title** Bayesian Purity Model to Estimate Tumor Purity

**Version** 1.0.0

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**Description** Bayesian purity model to estimate tumor purity using methylation array data (DNA methylation Infinium 450K array data) without reference samples.

**Depends** R (>= 2.10)

**Imports** stats, limma

**License** GPL (>= 2)

**Encoding** UTF-8

**LazyData** true

**RoxygenNote** 6.1.0

**NeedsCompilation** no

**Repository** CRAN

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annotGeneNames	<i>gene names of probes in 450K array dat</i>
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**Description**

gene names of probes in 450K array dat

**Format**

A vector with length 480457

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ApiGetDMCs	<i>Get TOPK=500 DMCs and non-DMCs using moderated-t test</i>
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**Description**

Get TOPK=500 DMCs and non-DMCs using moderated-t test

**Usage**

```
ApiGetDMCs(betaValue, TOPK = 500, tumorNum = NULL,
  filterProbes = FALSE, userProbes = NULL)
```

**Arguments**

betaValue	A matrix from TCGA array data
TOPK	An integer number, default 500. Number of DMCs/non-DMCs.
tumorNum	A postive number, First tumorNum columns in betaValue are tumor samples. If tumorNum is NULL, first half of columns are considered as tumor samples,
filterProbes	Logistic. defalut is FALSE. The code use all probes in betaValue. If TRUE, you can use default good probes provided in our code. you can also provide your good probes in userProbes.
userProbes	A number list. The row numbers in betaValue. These rows are considered as good probes. return DMCs (TOPK DMCs and TOPK non-DMCs row index in betaValue)

**Note**

User can provide the good probes indexes (row number) to filter the probes. A global variable goodProbes are used in this function. goodProbes: probes with SNPs at the CpG or single base extension sites, and corss-reactive probes are removed. More details see the reference paper.

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BayPM *Bayesian Purity Model (BPM) Main functions.*

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### Description

Bayesian Purity Model (BPM) Main functions.

### Usage

```
BayPM(betaValue, TOPK = 500, tumorNum = NULL, filterProbes = FALSE,
      userProbes = NULL)
```

### Arguments

betaValue	A matrix,TCGA methylation array data. Each row: loci, Tumor1,Tumor2,...,Normal1,Nomral2,...
TOPK	A number. Number of DMCs/nonDMCs selected
tumorNum	The number of tumor samples. if NULL, the default number is half of column number of dataset.
filterProbes	Logistic. defalut is FALSE. The code use all probes in betaValue. If TRUE, you can use default good probes provided in our code. you can also provide your good probes in userProbes.
userProbes	A number list. The row numbers in betaValue. These rows are considered as good probes.

### Value

tumor purity estimation of tumor samples

### Examples

```
### need to install package "limma"
### source("https://bioconductor.org/biocLite.R");biocLite("limma");
BayPM(simUCEC,20,2);
```

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BPM *BPM software package*

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### Description

Bayesian model for purity estimation using DNA methylation data

### Details

The main function is [BayPM](#)

**Author(s)**

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**References**

Jianzhao Gao, Linghao Shen, and Xiaodan Fan, Bayesian model for purity estimation using DNA methylation data.(submitted)

**Examples**

```
### need to install package "limma"
### source("https://bioconductor.org/biocLite.R");biocLite("limma");
library(BPM);
BayPM(simUCEC,20,2);
```

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estimateNu	<i>Estimate noise intensity (nv) for non-DMCs, using maximum likelihood estimation.</i>
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**Description**

Estimate noise intensity (nv) for non-DMCs, using maximum likelihood estimation.

**Usage**

```
estimateNu(z, phi, maxit = 50, beginP = 20)
```

**Arguments**

z	A matrix. Observed mixed tumor samples.
phi	mode of beta-values of each row in pure normal samples y.
maxit	A positive integer. The iteration number used in maximum likelihood.
beginP	A number, where the method start to search from for root. return estimated nv (noise intensity)

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fullSampler	<i>Sampling xi and alpha (tumor purity)</i>
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**Description**

Sampling xi and alpha (tumor purity)

**Usage**

```
fullSampler(y, z, mstates, xprior = NULL, maxit = 1000,
  burnin = maxit, xpar = FALSE, n_ab0 = NULL, alp0 = NULL,
  xbar0 = NULL, trace = FALSE, verbose = FALSE)
```

**Arguments**

y	A matrix, observed pure normal samples
z	A matrix, observed mixed tumor samples
mstates	A matrix, hyper/hypo of dataset
xprior	A matrix, prior knowledge about purity
maxit	A number, maximum iteration
burnin	A number, "burn-in" sample
xpar	Logistic, default is FALSE
n_ab0	initial value of n_ab
alp0	initial value of alpha
xbar0	initial value of xbar
trace	Logisitic, check the values in code, default is FALSE
verbose	Logistic, output the message,default is FALSE

**Value**

x\_bar x\_mode, x\_last x2 x\_sample x\_sample xpar xprior2, nab n\_ab2, alp alp2

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goodProbes	<i>good probes in packages</i>
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**Description**

good probes removed Y chrome.

**Format**

A vector with length 425698

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simUCEC

*Simulated data to illustrate datasets in packages*

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**Description**

A dataset containing 100 gene and 4 samples, first two columns are tumor1 tumor2 last two columns are normal1 normal2

- x. the genes
- y. two tumor samples; two normal samples;

**Format**

A matrix with 100 rows and 4 columns

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