

# Package ‘MiPP’

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**Title** Misclassification Penalized Posterior Classification

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**Depends** R (>= 2.4)

**Imports** Biobase, e1071, MASS, stats

**Description** This package finds optimal sets of genes that separate samples into two or more classes.

**License** GPL (>= 2)

**URL** <http://www.healthsystem.virginia.edu/internet/hes/biostat/bioinformatics/>

**biocViews** Microarray, Classification

**git\_url** <https://git.bioconductor.org/packages/MiPP>

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|       |                                              |
|-------|----------------------------------------------|
| colon | <i>Gene expression data for colon cancer</i> |
|-------|----------------------------------------------|

---

**Description**

This data set consists of gene expression of colon cancer study.

**Usage**

data(colon)

**Format**

A matrix containing 2000 probe sets and 2 classes (T, F)

**Source**

Alon, U., Barkai, N., Notterman, D.A., Gish, K., Ybarra, S., Mack, D., Levine, A.J. (1999). Broad Patterns of Gene Expression Revealed by Clustering Analysis of Tumor and Normal Colon Tissues probed by Oligonucleotide Arrays, PNAS, 96(12), 6745–6750.

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|              |                                      |
|--------------|--------------------------------------|
| cv.mipp.rule | <i>Fitting cross-validation MiPP</i> |
|--------------|--------------------------------------|

---

**Description**

Fits cross-validation MiPP

---

|          |                        |
|----------|------------------------|
| get.mipp | <i>Choosing a rule</i> |
|----------|------------------------|

---

**Description**

Choose a rule to compute MiPP

---

|              |                                    |
|--------------|------------------------------------|
| get.mipp.lda | <i>Fitting LDA to compute MiPP</i> |
|--------------|------------------------------------|

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**Description**

Fits LDA to compute MiPP

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|                   |                                               |
|-------------------|-----------------------------------------------|
| get.mipp.logistic | <i>Fitting logistic model to compute MiPP</i> |
|-------------------|-----------------------------------------------|

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**Description**

Fits logistic model to compute MiPP

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|              |                                    |
|--------------|------------------------------------|
| get.mipp.qda | <i>Fitting QDA to compute MiPP</i> |
|--------------|------------------------------------|

---

**Description**

Fits QDA to compute MiPP

---

|                     |                                             |
|---------------------|---------------------------------------------|
| get.mipp.svm.linear | <i>Fitting SVM (linear) to compute MiPP</i> |
|---------------------|---------------------------------------------|

---

**Description**

Fits SVM (linear) to compute MiPP

---

|                               |                                          |
|-------------------------------|------------------------------------------|
| <code>get.mipp.svm.rbf</code> | <i>Fitting SVM (RBF) to compute MiPP</i> |
|-------------------------------|------------------------------------------|

---

**Description**

Fits SVM (RBF) to compute MiPP

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|                    |                                          |
|--------------------|------------------------------------------|
| <code>leuk1</code> | <i>Gene expression data for leukemia</i> |
|--------------------|------------------------------------------|

---

**Description**

This data set consists of gene expression of leukemia study.

**Usage**

```
data(leukemia)
```

**Format**

A matrix containing 6817 probe sets and 38 samples (2 classes: AML, ALL)

**Source**

Golub, T.R., Slonim, D.K., Tamayo, P., Huard, C., Gaasenbeek, M., Mesirov, P., Coller, H., Loh, M.L., Downing, J.R., Caliguri, M.A., Bloomfield, C.D., and Lander, E.S. (1999) Molecular classification of cancer: class discovery and class prediction by gene expression monitoring. *Science*, 286, 531-537.

---

|                    |                                          |
|--------------------|------------------------------------------|
| <code>leuk2</code> | <i>Gene expression data for leukemia</i> |
|--------------------|------------------------------------------|

---

**Description**

This data set consists of gene expression of leukemia study.

**Usage**

```
data(leukemia)
```

**Format**

A matrix containing 6817 probe sets and 34 samples (2 classes: AML, ALL)

**Source**

Golub, T.R., Slonim, D.K., Tamayo, P., Huard, C., Gaasenbeek, M., Mesirov, P., Coller, H., Loh, M.L., Downing, J.R., Caliguri, M.A., Bloomfield, C.D., and Lander, E.S. (1999) Molecular classification of cancer: class discovery and class prediction by gene expression monitoring. *Science*, 286, 531-537.

---

leukemia

*Gene expression data for leukemia*

---

**Description**

This data set consists of gene expression of leukemia study.

**Usage**

```
data(leukemia)
```

**Format**

A matrix containing 6817 probe sets and 2 classes (AML, ALL)

**Source**

Golub, T.R., Slonim, D.K., Tamayo, P., Huard, C., Gaasenbeek, M., Mesirov, P., Coller, H., Loh, M.L., Downing, J.R., Caliguri, M.A., Bloomfield, C.D., and Lander, E.S. (1999) Molecular classification of cancer: class discovery and class prediction by gene expression monitoring. *Science*, 286, 531-537.

---

```
linearkernel.decision.function
```

*SVM (linear) kernel to compute MiPP*

---

**Description**

SVM (linear) kernel to compute MiPP

mipp

*MiPP-based Classification***Description**

Finds optimal sets of genes for classification

**Usage**

```
mipp(x, y, x.test = NULL, y.test = NULL, probe.ID = NULL,
     rule = "lda", method.cut = "t.test", percent.cut = 0.01,
     model.sMiPP.margin = 0.01, min.sMiPP = 0.85, n.drops = 2,
     n.fold = 5, p.test = 1/3, n.split = 20,
     n.split.eval = 100)
```

**Arguments**

|                    |                                                                                                                   |
|--------------------|-------------------------------------------------------------------------------------------------------------------|
| x                  | data matrix                                                                                                       |
| y                  | class vector                                                                                                      |
| x.test             | test data matrix if available                                                                                     |
| y.test             | test class vector if available                                                                                    |
| probe.ID           | probe set IDs; if NULL, row numbers are assigned.                                                                 |
| rule               | classification rule: "lda", "qda", "logistic", "svmlin", "svmrbf"; the default is "lda".                          |
| method.cut         | method for pre-selection; t-test is available.                                                                    |
| percent.cut        | proportion of pre-selected genes; the default is 0.01.                                                            |
| model.sMiPP.margin | smallest set of genes s.t. sMiPP <= (max sMiPP-model.sMiPP.margin); the default is 0.01.                          |
| min.sMiPP          | Adding genes stops if max sMiPP is at least min.sMiPP; the default is 0.85.                                       |
| n.drops            | Adding genes stops if sMiPP decreases (n.drops) times, in addition to min.sMiPP criterion.; the default is 2.     |
| n.fold             | number of folds; default is 5.                                                                                    |
| p.test             | partition percent of train and test samples when test samples are not available; the default is 1/3 for test set. |
| n.split            | number of splits; the default is 20.                                                                              |
| n.split.eval       | numbr of splits for evalutation; the default is 100.                                                              |

**Value**

|            |                                                                     |
|------------|---------------------------------------------------------------------|
| model      | candiadate genes (for each split if no indep set is available       |
| model.eval | Optimal sets of genes for each split when no indep set is available |

```
#####
#Example 1: When an independent test set is available

data(leukemia)

#Normalize combined data
leukemia <- cbind(leuk1, leuk2)
leukemia <- mipp.preproc(leukemia, data.type="MAS4")

#Train set
x.train <- leukemia[,1:38]
y.train <- factor(c(rep("ALL",27),rep("AML",11)))

#Test set
x.test <- leukemia[,39:72]
y.test <- factor(c(rep("ALL",20),rep("AML",14)))

#Compute MiPP
out <- mipp(x=x.train, y=y.train, x.test=x.test, y.test=y.test, probe.ID = 1:nrow(x.train), n.fold=5, percent.cut=0.5)

#Print candidate models
out$model

#####
#Example 2: When an independent test set is not available

data(colon)

#Normalize data
x <- mipp.preproc(colon)
y <- factor(c("T", "N", "T", "N", "T", "N", "T", "N", "T", "N",
  "T", "N", "T", "N", "T", "N", "T", "N", "T", "N",
  "T", "N", "T", "N", "T", "T", "T", "T", "T", "T",
  "T", "T", "T", "T", "T", "T", "T", "N", "T",
  "T", "N", "N", "T", "T", "T", "T", "N", "T", "N",
  "N", "T", "T", "N", "N", "T", "T", "T", "T", "N",
```

```

      "T", "N"))

#Deleting contaminated chips
x <- x[,-c(51,55,45,49,56)]
y <- y[ -c(51,55,45,49,56)]

#Compute MiPP
out <- mipp(x=x, y=y, probe.ID = 1:nrow(x), n.fold=5, p.test=1/3, n.split=5, n.split.eval=100,
percent.cut= 0.1, rule="lda")

#Print candidate models for each split
out$model

#Print optimal models and independent evaluation for each split
out$model.eval

```

---

mipp.preproc

*Preprocessing*


---

## Description

Performs IQR normalization, thesholding, and log2-transformation

## Usage

```
mipp.preproc(x, data.type = "MAS5")
```

## Arguments

|           |                                   |
|-----------|-----------------------------------|
| x         | data                              |
| data.type | data type is MAS5, MAS4, or dChip |

## See Also

[mipp](#)

## Examples

```

library(MiPP)

data(colon)
colon.nor <- mipp.preproc(colon)

```

mipp.rule

*Computing MiPP***Description**

Computes MiPP

mipp.seq

*MiPP-based Classification***Description**

sequentially finds optimal sets of genes for classification

**Usage**

```
mipp.seq(x, y, x.test = NULL, y.test = NULL, probe.ID = NULL,
  rule = "lda", method.cut = "t.test", percent.cut = 0.01,
  model.sMiPP.margin = 0.01, min.sMiPP = 0.85, n.drops = 2,
  n.fold = 5, p.test = 1/3, n.split = 20, n.split.eval = 100,
  n.seq=3, cutoff.sMiPP=0.7, remove.gene.each.model="all")
```

**Arguments**

|                    |                                                                                                                   |
|--------------------|-------------------------------------------------------------------------------------------------------------------|
| x                  | data matrix                                                                                                       |
| y                  | class vector                                                                                                      |
| x.test             | test data matrix if available                                                                                     |
| y.test             | test class vector if available                                                                                    |
| probe.ID           | probe set IDs; if NULL, row numbers are assigned.                                                                 |
| rule               | classification rule: "lda", "qda", "logistic", "svmlin", "svmrbf"; the default is "lda".                          |
| method.cut         | method for pre-selection; t-test is available.                                                                    |
| percent.cut        | proportion of pre-selected genes; the default is 0.01.                                                            |
| model.sMiPP.margin | smallest set of genes s.t. sMiPP <= (max sMiPP-model.sMiPP.margin); the default is 0.01.                          |
| min.sMiPP          | Adding genes stops if max sMiPP is at least min.sMiPP; the default is 0.85.                                       |
| n.drops            | Adding genes stops if sMiPP decreases (n.drops) times, in addition to min.sMiPP criterion.; the default is 2.     |
| n.fold             | number of folds; default is 5.                                                                                    |
| p.test             | partition percent of train and test samples when test samples are not available; the default is 1/3 for test set. |

`n.split`            number of splits; the default is 20.  
`n.split.eval`      numbr of splits for evaluation; the default is 100.  
`n.seq`              Number of sequential gene model selection; the default is 3.  
`cutoff.sMiPP`      Cutoff point of 5 percent sMiPP to select gene models  
`remove.gene.each.model`      Re-run after removing all genes in the selected models if "all" and the first gene for each of the selected models if "first"

### Value

`model`              candiadate genes (for each split if no indep set is available)  
`model.eval`        Optimal sets of genes for each split when no indep set is available  
`genes.selected`    a list of genes selected by sequential selection

### Author(s)

Soukup M, Cho H, and Lee JK

### References

Soukup M, Cho H, and Lee JK (2005). Robust classification modeling on microarray data using misclassification penalized posterior, *Bioinformatics*, 21 (Suppl): i423-i430.  
 Soukup M and Lee JK (2004). Developing optimal prediction models for cancer classification using gene expression data, *Journal of Bioinformatics and Computational Biology*, 1(4) 681-694

### Examples

```
#####
#Example 1: When an independent test set is available

data(leukemia)

#Normalize combined data
leukemia <- cbind(leuk1, leuk2)
leukemia <- mipp.preproc(leukemia, data.type="MAS4")

#Train set
x.train <- leukemia[,1:38]
y.train <- factor(c(rep("ALL",27),rep("AML",11)))

#Test set
x.test <- leukemia[,39:72]
y.test <- factor(c(rep("ALL",20),rep("AML",14)))

#Compute MiPP
out <- mipp.seq(x=x.train, y=y.train, x.test=x.test, y.test=y.test, n.fold=5, percent.cut=0.01, rule="lda", n.seq=3)

#Print candidate models
```

```

out$model

#Print the genes selected
out$genes.selected

#####
#Example 2: When an independent test set is not available

data(colon)

#Normalize data
x <- mipp.preproc(colon)
y <- factor(c("T", "N", "T", "N", "T", "N", "T", "N", "T", "N",
  "T", "N", "T", "N", "T", "N", "T", "N", "T", "N",
  "T", "N", "T", "N", "T", "T", "T", "T", "T", "T",
  "T", "T", "T", "T", "T", "T", "T", "T", "N", "T",
  "T", "N", "N", "T", "T", "T", "T", "N", "T", "N",
  "N", "T", "T", "N", "N", "T", "T", "T", "T", "N",
  "T", "N"))

#Deleting contaminated chips
x <- x[,-c(51,55,45,49,56)]
y <- y[ -c(51,55,45,49,56)]

#Compute MiPP
out <- mipp.seq(x=x, y=y, n.fold=5, p.test=1/3, n.split=5, n.split.eval=100,
percent.cut= 0.05, rule="lda", n.seq=2)

#Print candidate models for each split
out$model

#Print optimal models and independent evaluation for each split
out$model.eval

#Print the genes selected
out$genes.selected

```

---

pre.select

---

Pre-selection

---

## Description

Pre-select genes

---

|                           |                               |
|---------------------------|-------------------------------|
| <code>quant.normal</code> | <i>Quantile normalization</i> |
|---------------------------|-------------------------------|

---

**Description**

Performs quantile normalization

---

|                            |                               |
|----------------------------|-------------------------------|
| <code>quant.normal2</code> | <i>Quantile normalization</i> |
|----------------------------|-------------------------------|

---

**Description**

Performs quantile normalization

---

|                                          |                                         |
|------------------------------------------|-----------------------------------------|
| <code>rbfkernel.decision.function</code> | <i>SVM (RBF) kernel to compute MiPP</i> |
|------------------------------------------|-----------------------------------------|

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**Description**

SVM (RBF) kernel to compute MiPP

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