

# Package ‘PLPE’

April 1, 2025

**Version** 1.66.0

**Date** 2009-07-22

**Title** Local Pooled Error Test for Differential Expression with Paired High-throughput Data

**Author** HyungJun Cho <hj4cho@korea.ac.kr> and  
Jae K. Lee <jaeklee@virginia.edu>

**Maintainer** Soo-heang Eo <hanansh@korea.ac.kr>

**Depends** R (>= 2.6.2), Biobase (>= 2.5.5), LPE, MASS, methods

**Description** This package performs tests for paired high-throughput data.

**biocViews** Proteomics, Microarray, DifferentialExpression

**LazyLoad** yes

**LazyData** yes

**License** GPL (>= 2)

**URL** <http://www.korea.ac.kr/~stat2242/>

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

**git\_branch** RELEASE\_3\_20

**git\_last\_commit** 62bff99

**git\_last\_commit\_date** 2024-10-29

**Repository** Bioconductor 3.20

**Date/Publication** 2025-03-31

## Contents

|                                  |    |
|----------------------------------|----|
| am.trans.paired . . . . .        | 2  |
| base.error.paired . . . . .      | 2  |
| generate.null . . . . .          | 3  |
| lpe.paired . . . . .             | 3  |
| lpe.paired.default . . . . .     | 4  |
| lpe.paired.fdr . . . . .         | 6  |
| lpe.paired.fdr.default . . . . . | 7  |
| plateletSet . . . . .            | 8  |
| print.lpe.paired . . . . .       | 8  |
| print.lpe.paired.fdr . . . . .   | 9  |
| summary.lpe.paired . . . . .     | 10 |
| summary.lpe.paired.fdr . . . . . | 11 |

|              |           |
|--------------|-----------|
| <b>Index</b> | <b>12</b> |
|--------------|-----------|

---

|                        |  |
|------------------------|--|
| <b>am.trans.paired</b> | <i>Local Pooled Error Test for Paired Data</i> |
|------------------------|--|

---

## Description

This is internal function in lpe.paired.

## Usage

```
am.trans.paired(y, design)
```

## Author(s)

HyungJun Cho and Jae K. Lee

## References

Cho H, Smalley DM, Ross MM, Theodorescu D, Ley K and Lee JK (2007). Statistical Identification of Differentially Labelled Peptides from Liquid Chromatography Tandem Mass Spectrometry, *Proteomics*, 7:3681-3692.

## See Also

[lpe.paired.default](#)

---

|                          |  |
|--------------------------|--|
| <b>base.error.paired</b> | <i>Local Pooled Error Test for Paired Data</i> |
|--------------------------|--|

---

## Description

This is internal function in lpe.paired.

## Usage

```
base.error.paired(x, design, est.A, estimator, q, data.type)
```

## Author(s)

HyungJun Cho and Jae K. Lee

## References

Cho H, Smalley DM, Ross MM, Theodorescu D, Ley K and Lee JK (2007). Statistical Identification of Differentially Labelled Peptides from Liquid Chromatography Tandem Mass Spectrometry, *Proteomics*, 7:3681-3692.

## See Also

[lpe.paired.default](#)

---

generate.null            *Local Pooled Error Test for Paired Data*

---

### Description

This is internal function in lpe.paired.

### Usage

```
generate.null(x, design, q)
```

### Arguments

|        |  |
|--------|--|
| x      | data matrix  |
| design | design matrix; condition index in the first column and pair index in the second column |
| q      | quantile for intervals of intensities  |

### Value

|        |  |
|--------|--|
| design | design matrix; condition index in the first column and pair index in the second column |
| q      | quantile for intervals of intensities  |

### Author(s)

HyungJun Cho and Jae K. Lee

### References

Cho H, Smalley DM, Ross MM, Theodorescu D, Ley K and Lee JK (2007). Statistical Identification of Differentially Labelled Peptides from Liquid Chromatography Tandem Mass Spectrometry, Proteomics, 7:3681-3692.

### See Also

[lpe.paired.default](#)

---

lpe.paired            *Local Pooled Error Test for Paired Data*

---

### Description

This invetigates differential expression for paired high-throughput data.

### Usage

```
lpe.paired(x,...)
```

**Arguments**

- x an object for which the extraction of model lpe.paired is meaningful.
- ... other arguments

**Value**

- x design matrix; condition index in the first column and pair index in the second column
- ... data type: 'ms' for mass spectrometry data, 'cdna' for cDNA microarray data

**Author(s)**

HyungJun Cho and Jae K. Lee

**References**

Cho H, Smalley DM, Ross MM, Theodorescu D, Ley K and Lee JK (2007). Statistical Identification of Differentially Labelled Peptides from Liquid Chromatography Tandem Mass Spectrometry, *Proteomics*, 7:3681-3692.

**See Also**

[lpe.paired.default](#)

**Examples**

```
#LC-MS/MS proteomic data for platelets MPs
library(PLPE)
data(plateletSet)
x <- exprs(plateletSet)
x <- log2(x)

cond <- c(1, 2, 1, 2, 1, 2)
pair <- c(1, 1, 2, 2, 3, 3)
design <- cbind(cond, pair)

out <- lpe.paired(x, design, q=0.1, data.type="ms")
out$test.out[1:10,]
```

**lpe.paired.default**      *Local Pooled Error Test for Paired Data*

**Description**

This investigates differential expression for paired high-throughput data.

**Usage**

```
## Default S3 method:
lpe.paired(x, design, data.type, q=0.01, probe.ID = NULL, estimator="median", w=0.5, w.estimator="f
```

**Arguments**

|             |  |
|-------------|--|
| x           | data matrix  |
| design      | design matrix; condition index in the first column and pair index in the second column                 |
| q           | quantile for intervals of intensities  |
| probe.ID    | probe set IDs; if NULL, row numbers are assigned.  |
| data.type   | data type: 'ms' for mass spectrometry data, 'cdna' for cDNA microarray data                            |
| estimator   | specification for the estimator: 'median', 'mean' and 'huber'  |
| w           | weight parameter between individual variance estimate and pooling variance estimate, $0 \leq w \leq 1$ |
| w.estimator | two approaches to estimate the weight: 'random' or 'fixed'   |
| iseed       | seed number  |
| ...         | other arguments  |

**Value**

|             |  |
|-------------|--|
| design      | design matrix; condition index in the first column and pair index in the second column                 |
| data.type   | data type: 'ms' for mass spectrometry data, 'cdna' for cDNA microarray data                            |
| q           | quantile for intervals of intensities  |
| estimator   | specification for the estimator: 'median', 'mean' and 'huber'  |
| w.estimator | two approaches to estimate the weight: 'random' or 'fixed'   |
| w           | weight parameter between individual variance estimate and pooling variance estimate, $0 \leq w \leq 1$ |
| test.out    | matrix for test results  |

**Author(s)**

HyungJun Cho and Jae K. Lee

**References**

Cho H, Smalley DM, Ross MM, Theodorescu D, Ley K and Lee JK (2007). Statistical Identification of Differentially Labelled Peptides from Liquid Chromatography Tandem Mass Spectrometry, Proteomics, 7:3681-3692.

**See Also**

[lpe.paired](#)

**Examples**

```
#LC-MS/MS proteomic data for platelets MPs
library(PLPE)
data(plateletSet)
x <- exprs(plateletSet)
x <- log2(x)

cond <- c(1, 2, 1, 2, 1, 2)
```

```

pair <- c(1, 1, 2, 2, 3, 3)
design <- cbind(cond, pair)

out <- lpe.paired(x, design, q=0.1, data.type="ms")
out$test.out[1:10,]
summary(out)

```

**lpe.paired.fdr***FDR for PLPE*

## Description

This computes FDR for PLPE.

## Usage

```
lpe.paired.fdr(x,...)
```

## Arguments

|     |                 |
|-----|-----------------|
| x   | data matrix     |
| ... | other arguments |

## Author(s)

HyungJun Cho and Jae K. Lee

## References

Cho H, Smalley DM, Ross MM, Theodorescu D, Ley K and Lee JK (2007). Statistical Identification of Differentially Labelled Peptides from Liquid Chromatography Tandem Mass Spectrometry, *Proteomics*, 7:3681-3692.

## See Also

[lpe.paired.fdr.default](#)

## Examples

```

#LC-MS/MS proteomic data for platelets MPs
library(PLPE)
data(plateletSet)
x <- exprs(plateletSet)
x <- log2(x)

cond <- c(1, 2, 1, 2, 1, 2)
pair <- c(1, 1, 2, 2, 3, 3)
design <- cbind(cond, pair)

out <- lpe.paired(x, design, q=0.1, data.type="ms")
out.fdr <- lpe.paired.fdr(x,obj=out)
out.fdr$FDR[1:10,]

```

---

lpe.paired.fdr.default  
*FDR for PLPE*

---

## Description

This computes FDR for PLPE.

## Usage

```
## Default S3 method:  
lpe.paired.fdr(x, obj, n.iter=5, lambda=0.9, ...)
```

## Arguments

|        |  |
|--------|--|
| x      | data matrix  |
| obj    | object created from lpe.paired                       |
| n.iter | number of iterations                                 |
| lambda | numeric vector of probabilities with values in [0,1] |
| ...    | other argument                                       |

## Value

|             |  |
|-------------|--|
| design      | design matrix; condition index in the first column and pair index in the second column                 |
| data.type   | data type: 'ms' for mass spectrometry data, 'cdna' for cDNA microarray data                            |
| estimator   | specification for the estimator: 'median', 'mean' and 'huber'  |
| w.estimator | two approaches to estimate the weight: 'random' or 'fixed'   |
| w           | weight parameter between individual variance estimate and pooling variance estimate, $0 \leq w \leq 1$ |
| pi0         | estimated proportion of non-null peptides  |
| FDR         | matrix for test results including FDRs   |
| ...         | other arguments  |

## Author(s)

HyungJun Cho and Jae K. Lee

## References

Cho H, Smalley DM, Ross MM, Theodorescu D, Ley K and Lee JK (2007). Statistical Identification of Differentially Labelled Peptides from Liquid Chromatography Tandem Mass Spectrometry, Proteomics, 7:3681-3692.

## See Also

[lpe.paired.fdr](#)

## Examples

```
#LC-MS/MS proteomic data for platelets MPs
library(PLPE)
data(plateletSet)
x <- exprs(plateletSet)
x <- log2(x)

cond <- c(1, 2, 1, 2, 1, 2)
pair <- c(1, 1, 2, 2, 3, 3)
design <- cbind(cond, pair)

out <- lpe.paired(x, design, q=0.1, data.type="ms")
out.fdr <- lpe.paired.fdr(x,obj=out)
out.fdr$FDR[1:10,]
```

**plateletSet**

*LCMS proteomic data for platele MPs*

## Description

This data set consists of LC-MS/MS data with three replicates of paired samples.

## Source

Garcia BA, Smalley DM, Cho H, Shabanowitz J, Ley K and Hunt DF (2005). The Platelet Microparticle Proteome, Journal of Proteome Research, 4:1516-1521.

**print.lpe.paired**

*Local Pooled Error Test for Paired Data*

## Description

This print lpe.paired.

## Usage

```
## S3 method for class 'lpe.paired'
print(x,...)
```

## Arguments

|     |                                   |
|-----|-----------------------------------|
| x   | an object created from lpe.paired |
| ... | other arguments                   |

## Author(s)

HyungJun Cho and Jae K. Lee

## References

Cho H, Smalley DM, Ross MM, Theodorescu D, Ley K and Lee JK (2007). Statistical Identification of Differentially Labelled Peptides from Liquid Chromatography Tandem Mass Spectrometry, *Proteomics*, 7:3681-3692.

## Examples

```
#LC-MS/MS proteomic data for platelets MPs
library(PLPE)
data(plateletSet)
x <- exprs(plateletSet)
x <- log2(x)

cond <- c(1, 2, 1, 2, 1, 2)
pair <- c(1, 1, 2, 2, 3, 3)
design <- cbind(cond, pair)

out <- lpe.paired(x, design, q=0.1, data.type="ms")
out
```

**print.lpe.paired.fdr**    *Local Pooled Error Test for Paired Data*

## Description

This prints lpe.paired.fdr

## Usage

```
## S3 method for class 'lpe.paired.fdr'
print(x,...)
```

## Arguments

|     |                                   |
|-----|-----------------------------------|
| x   | an object created from lpe.paired |
| ... | other arguments                   |

## Author(s)

HyungJun Cho and Jae K. Lee

## References

Cho H, Smalley DM, Ross MM, Theodorescu D, Ley K and Lee JK (2007). Statistical Identification of Differentially Labelled Peptides from Liquid Chromatography Tandem Mass Spectrometry, *Proteomics*, 7:3681-3692.

---

**summary.lpe.paired**      *Local Pooled Error Test for Paired Data*

---

**Description**

This summarize lpe.paired.

**Usage**

```
## S3 method for class 'lpe.paired'  
summary(object,...)
```

**Arguments**

|     |                                   |
|-----|-----------------------------------|
| x   | an object created from lpe.paired |
| ... | other arguments                   |

**Author(s)**

HyungJun Cho and Jae K. Lee

**References**

Cho H, Smalley DM, Ross MM, Theodorescu D, Ley K and Lee JK (2007). Statistical Identification of Differentially Labelled Peptides from Liquid Chromatography Tandem Mass Spectrometry, *Proteomics*, 7:3681-3692.

**Examples**

```
#LC-MS/MS proteomic data for platelets MPs  
library(PLPE)  
data(plateletSet)  
x <- exprs(plateletSet)  
x <- log2(x)  
  
cond <- c(1, 2, 1, 2, 1, 2)  
pair <- c(1, 1, 2, 2, 3, 3)  
design <- cbind(cond, pair)  
  
out <- lpe.paired(x, design, q=0.1, data.type="ms")  
summary(out)
```

---

**summary.lpe.paired.fdr**

*Local Pooled Error Test for Paired Data*

---

## Description

This summarize lpe.paired.

## Usage

```
## S3 method for class 'lpe.paired.fdr'  
summary(object,...)
```

## Arguments

|     |                                   |
|-----|-----------------------------------|
| x   | an object created from lpe.paired |
| ... | other arguments                   |

## Author(s)

HyungJun Cho and Jae K. Lee

## References

Cho H, Smalley DM, Ross MM, Theodorescu D, Ley K and Lee JK (2007). Statistical Identification of Differentially Labelled Peptides from Liquid Chromatography Tandem Mass Spectrometry, *Proteomics*, 7:3681-3692.

# Index

- \* **datasets**
  - plateletSet, 8
- \* **internal**
  - am.trans.paired, 2
  - base.error.paired, 2
  - generate.null, 3
  - print.lpe.paired, 8
  - print.lpe.paired.fdr, 9
  - summary.lpe.paired, 10
  - summary.lpe.paired.fdr, 11
- \* **models**
  - lpe.paired, 3
  - lpe.paired.default, 4
  - lpe.paired.fdr, 6
  - lpe.paired.fdr.default, 7
- am.trans.paired, 2
- base.error.paired, 2
- generate.null, 3
- lpe.paired, 3, 5
- lpe.paired.default, 2–4, 4
- lpe.paired.fdr, 6, 7
- lpe.paired.fdr.default, 6, 7
- plateletSet, 8
- print.lpe.paired, 8
- print.lpe.paired.fdr, 9
- summary.lpe.paired, 10
- summary.lpe.paired.fdr, 11