

# Package: MRmediation (via r-universe)

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**Type** Package

**Title** A Causal Mediation Method with Methylated Region (MR) as the Mediator

**Version** 1.0.1

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**Description** A causal mediation approach under the counterfactual framework to test the significance of total, direct and indirect effects. In this approach, a group of methylated sites from a predefined region are utilized as the mediator, and the functional transformation is used to reduce the possible high dimension in the region-based methylated sites and account for their location information.

**License** GPL (>= 2)

**Encoding** UTF-8

**LazyData** true

**Depends** R (>= 3.5.0), fda

**Imports** MASS, stats

**RoxygenNote** 7.1.0

**Collate** 'MRmediation.R' 'mediation\_single.R' 'example\_data.R'

**NeedsCompilation** no

**Repository** CRAN

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## Contents

example_data . . . . .	2
mediation . . . . .	2
mediation_single . . . . .	3

<b>Index</b>	<b>5</b>
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example_data	<i>This is the data for examples</i>
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### Description

- data. phenotype file. 1st column is ID, 2nd column is continuous outcome, 3rd column is binary outcome, 4th column is exposure, 5th column is age, 6th column is gender, 7th-last columns are CpGs
- pos. CpG locations from the defined region and they are from the same chromosome.

### Usage

```
data(example_data)
```

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mediation	<i>A causal mediation method with methylated region as the mediator</i>
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### Description

A causal mediation method with methylated region as the mediator

### Usage

```
mediation(
  pheno,
  predictor,
  region,
  pos,
  order,
  gbasis,
  covariate,
  base = "bspline",
  family = "gaussian"
)
```

### Arguments

pheno	A vector of continuous or binary phenotypes (class: numeric).
predictor	A vector of values for the exposure variable (class: numeric).
region	A matrix of CpGs in a region. Each column is a CpG (class: data.frame).
pos	A vector of CpG locations from the defined region and they are from the same chromosome (class: integer).
order	A value for the order of bspline basis. 1: constant, 2: linear, 3: quadratic and 4: cubic.

gbasis	A value for the number of basis being used for functional transformation on CpGs.
covariate	A matrix of covariates. Each column is a covariate (class: data.frame).
base	"bspline" for B-spline basis or "fspline" for Fourier basis.
family	"gaussian" for continuous outcome or "binomial" for binary outcome.

### Value

1. pval\$TE: total effect (TE) p-value
2. pval\$DE: direct effect (DE) p-value
3. pval\$IE: indirect effect (IE) p-value
4. pval\_MX: p-value for the association between methylation and exposure

### Examples

```
#####
### Examples ###
#####
data("example_data")
predictor = data$exposure
region = data[,7:dim(data)[2]]
covariates = subset(data, select=c("age", "gender"))
# binary outcome
pheno_bin = data$pheno_bin
mediation(pheno_bin, predictor, region, pos, covariate=covariates, order=4,
gbasis=4, base="bspline", family="binomial")
# continuous outcome
pheno_con = data$pheno_con
mediation(pheno_con, predictor, region, pos, covariate=covariates, order=4,
gbasis=4, base="bspline", family="gaussian")
```

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mediation\_single      *A causal mediation method with a single CpG site as the mediator*

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### Description

A causal mediation method with a single CpG site as the mediator

### Usage

```
mediation_single(pheno, predictor, cpG, covariate, family = "gaussian")
```

**Arguments**

pheno	A vector of continuous or binary phenotypes (class: numeric).
predictor	A vector of values for the exposure variable (class: numeric).
cpg	A vector of a CpG (class: numeric).
covariate	A matrix of covariates. Each column is a covariate (class: data.frame).
family	"gaussian" for continuous outcome or "binomial" for binary outcome.

**Value**

1. pval\$TE: total effect (TE) p-value
2. pval\$DE: direct effect (DE) p-value
3. pval\$IE: indirect effect (IE) p-value
4. pval\_MX: p-value for the association between methylation and exposure

**Examples**

```
#####  
### Examples ###  
#####  
data("example_data")  
predictor = data$exposure  
cpg = data[,9] #any number in c(7:dim(data)[2])  
covariates = subset(data, select=c("age", "gender"))  
# binary outcome  
pheno_bin = data$pheno_bin  
mediation_single(pheno_bin, predictor, cpg, covariate=covariates, family="binomial")  
# continuous outcome  
pheno_con = data$pheno_con  
mediation_single(pheno_con, predictor, cpg, covariate=covariates, family="gaussian")
```

# Index

`data (example_data)`, [2](#)

`example_data`, [2](#)

`mediation`, [2](#)

`mediation_single`, [3](#)

`pos (example_data)`, [2](#)