

# Package ‘did’

June 21, 2019

**Title** Treatment Effects with Multiple Periods and Groups

**Version** 1.2.2

**Description** The standard Difference-in-Differences (DID) setup involves two periods and two groups -- a treated group and untreated group. Many applications of DID methods involve more than two periods and have individuals that are treated at different points in time. This package contains tools for computing average treatment effect parameters in Difference in Differences models with more than two periods and with variation in treatment timing using the methods developed in Callaway and Sant'Anna (2018) <<https://ssrn.com/abstract=3148250>>. The main parameters are group-time average treatment effects which are the average treatment effect for a particular group at a particular time. These can be aggregated into a fewer number of treatment effect parameters, and the package deals with the cases where there is selective treatment timing, dynamic treatment effects, calendar time effects, or combinations of these. There are also functions for testing the Difference in Differences assumption, and plotting group-time average treatment effects.

**Depends** R (>= 2.10)

**License** GPL-2

**Encoding** UTF-8

**LazyData** true

**Imports** BMisc (>= 1.3.1), MASS, pbapply, stats, ggplot2, knitr, utils, gridExtra

**RoxygenNote** 6.1.1

**VignetteBuilder** knitr

**Suggests** rmarkdown

**NeedsCompilation** no

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## R topics documented:

AGGTE . . . . .	2
did . . . . .	3
expf . . . . .	4
gdid . . . . .	4
indicator . . . . .	5
MP . . . . .	6
mp.spatt . . . . .	7
mp.spatt.test . . . . .	9
MP.TEST . . . . .	10
mpdta . . . . .	11
onefun . . . . .	11
summary.AGGTE . . . . .	12
summary.MP . . . . .	12
summary.MP.TEST . . . . .	13
<b>Index</b>	<b>14</b>

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AGGTE

*AGGTE*

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### Description

AGGTE class for aggregate treatment effects

### Usage

```
AGGTE(simple.att = NULL, simple.se = NULL, selective.att = NULL,
      selective.se = NULL, selective.att.g = NULL, selective.se.g = NULL,
      dynamic.att = NULL, dynamic.se = NULL, dynamic.att.e = NULL,
      dynamic.se.e = NULL, calendar.att = NULL, calendar.se = NULL,
      calendar.att.t = NULL, calendar.se.t = NULL, dynsel.att.e1 = NULL,
      dynsel.se.e1 = NULL, dynsel.att.ee1 = NULL, dynsel.se.ee1 = NULL,
      groups = NULL, times = NULL)
```

### Arguments

simple.att	simple weighted average of group-time average treatment effects
simple.se	the standard error for simple.att
selective.att	aggregated group-time average treatment effects when there is selective treatment timing
selective.se	the standard error for selective.att
selective.att.g	aggregated group-time average treatment effects when there is selective treatment timing for each particular group
selective.se.g	the standard error for selective.att.g

dynamic.att	aggregated group-time average treatment effects when there are dynamic treatment effects
dynamic.se	the standard error for dynamic.att
dynamic.att.e	aggregated group-time average treatment effects when there are dynamic treatment effects for each length of exposure to treatment
dynamic.se.e	the standard error for dynamic.att.e
calendar.att	the aggregated group-time average treatment effects when there are calendar time effects
calendar.se	the standard error for calendar.att
calendar.att.t	the aggregated group-time average treatment effects when there are calendar time effects for each time period
calendar.se.t	the standard error for calendar.att.t
dynsel.att.e1	aggregated group-time average treatment effects when there are dynamic treatment effects and selective treatment timing. Here, e1 is the number of periods that a group is required to be treated in order to be included in the results.
dynsel.se.e1	the standard error for dynsel.att.e1
dynsel.att.ee1	aggregated group-time average treatment effects when there are dynamic treatment effects and selective treatment timing. Here, e1 is the number of periods that a group is required to be treated in order to be included in the results and for each length of exposure to treatment
dynsel.se.ee1	the standard error for dynsel.att.ee1
groups	vector of all groups
times	vector of all times

---

did *did: Difference in Differences with multiple periods and variation in treatment timing*

---

### Description

did: Difference in Differences with multiple periods and variation in treatment timing

expf

*expf*

---

**Description**

exponential weighting function

**Usage**

```
expf(X, u)
```

**Arguments**

X                    matrix of X's from the data  
u                    a particular value to multiply times the X's

**Value**

numeric vector

**Examples**

```
data(mpdata)  
dta <- subset(mpdata, year==2007)  
X <- model.matrix(~lpop, data=dta)  
X <- expf(X, X[1,])
```

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ggdid*ggdid*

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**Description**

Function to plot MP objects

**Usage**

```
ggdid(mpobj, type = c("attgt", "dynamic", "selective", "calendar",  
"dynsel"), ylim = NULL, xlab = NULL, ylab = NULL,  
title = "Group", xgap = 1, ncol = 1, e1 = 1)
```

**Arguments**

<code>mpobj</code>	an MP object
<code>type</code>	the type of plot, should be one of "attgt", "dynamic", "selective", "calendar", "dynsel". "attgt" is the default and plots all group-time average treatment effects separately by group (including pre-treatment time periods); "dynamic" plots dynamic treatment effects – these are the same as event studies; "selective" plots average effects of the treatment separately by group (which allows for selective treatment timing); "calendar" plots average treatment effects by time period; and "dynsel" plots dynamic effects allowing for selective treatment timing (this also requires setting the additional parameter <code>e1</code> )
<code>ylim</code>	optional y limits for the plot; setting here makes the y limits the same across different plots
<code>xlab</code>	optional x-axis label
<code>ylab</code>	optional y-axis label
<code>title</code>	optional plot title
<code>xgap</code>	optional gap between the labels on the x-axis. For example, <code>xgap=3</code> indicates that the labels should show up for every third value on the x-axis. The default is 1.
<code>ncol</code>	The number of columns to include in the resulting plot. The default is 1.
<code>e1</code>	only used when plot type is "dynsel", this specifies the number of post-treatment periods that need to be available for particular groups to be included in the resulting plot when there are dynamic treatment effects and selective treatment timing

**Examples**

```
## Not run:
data(mpdt)
out <- mp.spatt(lmp ~ treat, xformula=~lpop, data=mpdt,
               panel=TRUE, first.treat.name="first.treat",
               idname="countyreal", tname="year",
               bstrap=FALSE, se=TRUE, cband=FALSE)
ggdid(out)

## End(Not run)
```

---

indicator

*indicator*

---

**Description**

indicator weighting function

**Usage**

```
indicator(X, u)
```

**Arguments**

X                    matrix of X's from the data  
u                    a particular value to compare X's to

**Value**

numeric vector

**Examples**

```
data(mpdata)
dta <- subset(mpdata, year==2007)
X <- model.matrix(~lpop, data=dta)
X <- indicator(X, X[1,])
```

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MP

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*MP*


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**Description**

multi-period object

**Usage**

```
MP(group, t, att, V, c, inffunc, n = NULL, W = NULL, Wpval = NULL,
    aggte = NULL)
```

**Arguments**

group                which group (defined by period first treated) an group-time average treatment effect is for  
t                    which time period a group-time average treatment effect is for  
att                  the group-average treatment effect for group group and time period t  
V                    the variance matrix for group-time average treatment effects  
c                    critical value if one is obtaining uniform confidence bands  
inffunc              the influence function for estimating group-time average treatment effects  
n                    the number of observations  
W                    the Wald statistic for pre-testing the common trends assumption  
Wpval                the p-value of the Wald statistic for pre-testing the common trends assumption  
aggte                an aggregate treatment effects object

**Value**

MP object

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mp.spatt	<i>mp.spatt</i>
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**Description**

mp.spatt computes the ATT in the case where there are more than two periods of data and allowing for treatment to occur at different points in time extending the method of Abadie (2005). This method relies on once individuals are treated they remain in the treated state for the duration.

**Usage**

```
mp.spatt(formla, xformla = NULL, data, tname, aggte = TRUE, w = NULL,
panel = FALSE, idname = NULL, first.treat.name, alp = 0.05,
method = "logit", se = TRUE, bstrap = FALSE, biters = 100,
clustervars = NULL, cband = FALSE, citers = 100, seedvec = NULL,
pl = FALSE, cores = 2, printdetails = TRUE)
```

**Arguments**

formla	The formula $y \sim d$ where $y$ is the outcome and $d$ is the treatment indicator ( $d$ should be binary)
xformla	A optional one sided formula for additional covariates that will be adjusted for. E.g $\sim \text{age} + \text{education}$ . Additional covariates can also be passed by name using the $x$ paramater.
data	The name of the data.frame that contains the data
tname	The name of the column containing the time periods
aggte	boolean for whether or not to compute aggregate treatment effect parameters, default TRUE
w	A vector of weights for each observation (not implemented)
panel	Boolean indicating whether the data is panel or repeated cross sections
idname	The individual (cross-sectional unit) id name
first.treat.name	The name of the variable in data that contains the first period when a particular observation is treated. This should be a positive number for all observations in treated groups. It should be 0 for observations in the untreated group.
alp	the significance level, default is 0.05
method	The method for estimating the propensity score when covariates are included
se	Boolean whether or not to compute standard errors
bstrap	Boolean for whether or not to compute standard errors using the multiplier bootstrap. If standard errors are clustered, then one must set <code>bstrap=TRUE</code> .

<code>biters</code>	The number of bootstrap iterations to use. The default is 100, and this is only applicable if <code>bstrap=TRUE</code> .
<code>clustervars</code>	A vector of variables to cluster on. At most, there can be two variables (otherwise will throw an error) and one of these must be the same as <code>idname</code> which allows for clustering at the individual level.
<code>cband</code>	Boolean for whether or not to compute a uniform confidence band that covers all of the group-time average treatment effects with fixed probability $1-\alpha p$ . The default is <code>FALSE</code> and the resulting standard errors will be pointwise.
<code>citers</code>	Computing uniform confidence bands requires the bootstrap, if <code>cband = TRUE</code> , then this is the number of bootstrap iterations to compute the confidence band. The default is 100.
<code>seedvec</code>	Optional value to set random seed; can possibly be used in conjunction with bootstrapping standard errors# (not implemented)
<code>p1</code>	Boolean for whether or not to use parallel processing
<code>cores</code>	The number of cores to use for parallel processing
<code>printdetails</code>	Boolean for showing detailed results or not

**Value**

MP object

**References**

Callaway, Brantly and Sant'Anna, Pedro. "Difference-in-Differences with Multiple Time Periods and an Application on the Minimum Wage and Employment." Working Paper <<https://ssrn.com/abstract=3148250>> (2018).

**Examples**

```
data(mpdtA)

## with covariates
out1 <- mp.spatt(lemp ~ treat, xformula=~lpop, data=mpdtA,
  panel=TRUE, first.treat.name="first.treat",
  idname="countyreal", tname="year",
  bstrap=FALSE, se=TRUE, cband=FALSE)
## summarize the group-time average treatment effects
summary(out1)
## summarize the aggregated treatment effect parameters
summary(out1$aggte)

## without any covariates
out2 <- mp.spatt(lemp ~ treat, xformula=NULL, data=mpdtA,
  panel=TRUE, first.treat.name="first.treat",
  idname="countyreal", tname="year",
  bstrap=FALSE, se=TRUE, cband=FALSE)
summary(out2)
```



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mp.spatt.test	<i>mp.spatt.test</i>
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### Description

integrated moments test for conditional common trends holding in all pre-treatment time periods across all groups

### Usage

```
mp.spatt.test(formla, xformlalist = NULL, data, tname,
  weightfun = NULL, w = NULL, panel = FALSE, idname = NULL,
  first.treat.name, alp = 0.05, method = "logit", biters = 100,
  clustervarlist = NULL, pl = FALSE, cores = 2)
```

### Arguments

formla	The formula $y \sim d$ where $y$ is the outcome and $d$ is the treatment indicator ( $d$ should be binary)
xformlalist	A list of formulas for the X variables. This allows to test using different specifications for X, if desired
data	The name of the data.frame that contains the data
tname	The name of the column containing the time periods
weightfun	A function that takes in two arguments, X and u, to compute the weighting function for the test. The default is $1*(X \leq u)$
w	A vector of weights for each observation (not implemented)
panel	Boolean indicating whether the data is panel or repeated cross sections
idname	The individual (cross-sectional unit) id name
first.treat.name	The name of the variable in data that contains the first period when a particular observation is treated. This should be a positive number for all observations in treated groups. It should be 0 for observations in the untreated group.
alp	the significance level, default is 0.05
method	The method for estimating the propensity score when covariates are included
biters	The number of bootstrap iterations to use. The default is 100, and this is only applicable if bstrap=TRUE.
clustervarlist	A list of cluster variables. This allows to conduct the test using different levels of clustering, if desired.
pl	Boolean for whether or not to use parallel processing
cores	The number of cores to use for parallel processing

### Value

list containing test results

## References

Callaway, Brantly and Sant'Anna, Pedro. "Difference-in-Differences with Multiple Time Periods and an Application on the Minimum Wage and Employment." Working Paper <<https://ssrn.com/abstract=3148250>> (2018).

## Examples

```
## Not run:
data(mpdta)
mptest <- mp.spatt.test(lemp ~ treat, xformlalist=list(~lpop), data=mpdta,
  panel=TRUE, first.treat.name="first.treat",
  idname="countyreal", tname="year", clustervarlist=list(NULL))
summary(mptest[[1]])

## End(Not run)

data(mpdta)
mptest <- mp.spatt.test(lemp ~ treat, xformlalist=list(NULL), data=mpdta,
  panel=TRUE, first.treat.name="first.treat",
  idname="countyreal", tname="year", clustervarlist=list(NULL))
summary(mptest[[1]])
```

---

MP.TEST

*MP.TEST*

---

## Description

MP.TEST objects

## Usage

```
MP.TEST(CvM, CvMb, CvMcval, CvMpval, KS, KSb, KScval, KSpval, clustervars,
  xformla)
```

## Arguments

CvM	Cramer von Mises test statistic
CvMb	a vector of bootstrapped Cramer von Mises test statistics
CvMcval	CvM critical value
CvMpval	p-value for CvM test
KS	Kolmogorov-Smirnov test statistic
KSb	a vector of bootstrapped KS test statistics
KScval	KS critical value
KSpval	p-value for KS test
clustervars	vector of which variables were clustered on for the test
xformla	formla for the X variables used in the test

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mpdta	<i>County teen employment</i>
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**Description**

A dataset containing (the log of) teen employment in 500 counties in the U.S. from 2004 to 2007. This is a subset of the dataset used in Callaway and Sant'Anna (2018). See that paper for additional descriptions.

**Usage**

mpdta

**Format**

A data frame with 2000 rows and 5 variables:

**year** the year of the observation

**countyreal** a unique identifier for a particular county

**lpop** the log of 1000s of population for the county

**lemp** the log of teen employment in the county

**first.treat** the year that the state where the county is located raised its minimum wage, it is set equal to 0 for counties that have minimum wages equal to the federal minimum wage over the entire period.

**treat** whether or not a particular county is treated in that year

**Source**

Callaway and Sant'Anna (2018)

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onefun	<i>onefun</i>
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---

**Description**

just return the value 1

**Usage**

onefun(X, u)

**Arguments**

X matrix of X's from the data

u a particular value to compare X's to

**Value**

numeric vector

**Examples**

```
data(mpdt)
dta <- subset(mpdt, year==2007)
X <- model.matrix(~lpop, data=dta)
X <- onefun(X, X[1,])
```

---

summary.AGGTE

*summary.AGGTE*

---

**Description**

print a summary of an AGGTE object

**Usage**

```
## S3 method for class 'AGGTE'
summary(object, type = c("dynamic", "selective",
  "calendar", "dynsel"), e1 = 1, ...)
```

**Arguments**

object	an AGGTE object
type	which type of summary to print, options are "dynamic", "selective", "calendar", and "dynsel"
e1	if the type is "dynsel", this is the number of post-treatment periods required in order for a group to be used to construct aggregated parameters with selective treatment timing and dynamic effects; otherwise not used
...	other variables

---

summary.MP

*summary.MP*

---

**Description**

prints a summary of a MP object

**Usage**

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

**Arguments**

object	an MP object
...	extra arguments

---

<code>summary.MP.TEST</code>	<i>summary.MP.TEST</i>
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**Description**

print a summary of test results

**Usage**

```
## S3 method for class 'MP.TEST'  
summary(object, ...)
```

**Arguments**

object	an MP.TEST object
...	other variables

# Index

## \*Topic **datasets**

mpdta, [11](#)

AGGTE, [2](#)

did, [3](#)

did-package (did), [3](#)

expf, [4](#)

ggdid, [4](#)

indicator, [5](#)

MP, [6](#)

mp.spatt, [7](#)

mp.spatt.test, [9](#)

MP.TEST, [10](#)

mpdta, [11](#)

onefun, [11](#)

summary.AGGTE, [12](#)

summary.MP, [12](#)

summary.MP.TEST, [13](#)