

Package: bdots (via r-universe)

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Type Package

Title Bootstrapped Differences of Time Series

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BugReports <https://github.com/collinn/bdots/issues>

Depends R (>= 4.0.0), data.table

Imports nlme, mvtnorm, parallel, stats, graphics, utils, ggplot2,
gridExtra

LazyData TRUE

Description Analyze differences among time series curves with p-value
adjustment for multiple comparisons introduced in Oleson et al
(2015) <[DOI:10.1177/0962280215607411](https://doi.org/10.1177/0962280215607411)>.

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URL <https://github.com/collinn/bdots>

NeedsCompilation no

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Suggests knitr, rmarkdown, tinytest

VignetteBuilder knitr, rmarkdown

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ar1Solver	<i>Compute AR1 correlation coefficient</i>
-----------	--

Description

Computes value for AR1 correlation coefficient for use in `p_adjust`

Usage

```
ar1Solver(t)
```

Arguments

`t` A numeric vector of t-statistics

Value

Estimated AR1 correlation coefficient

See Also

[p_adjust](#)

Examples

```
t <- rt(1e3, df = 1)
rho <- ar1Solver(t)
```

bdotsBoot	<i>Create bootstrapped curves from bdotsObj</i>
-----------	---

Description

Creates bootstrapped curves and performs alpha adjustment. Can perform "difference of difference" for nested comparisons

Usage

```
bdotsBoot(
  formula,
  bdObj,
  Niter = 1000,
  alpha = 0.05,
  padj = "oleson",
  permutation = FALSE,
  cores = 0,
  ...
)
```

Arguments

formula	See details.
bdObj	An object of class 'bdotsObj'
Niter	Number of iterations of bootstrap to draw
alpha	Significance level
padj	Adjustment to make to pvalues for significance. Will be able to use anything from p.adjust function, but for now, just "oleson"
permutation	Boolean indicating whether to use permutation testing rather than adjusting alpha to control FWER. WARNING: This option is very much in beta testing and not recommended for general use. Also not available for paired tests or difference of difference
cores	Number of cores to use in parallel. Default is zero, which uses half of what is available.
...	not used

Details

The formula is the only tricky part of this. There will be a minor update to how it works in the future. The three parts we will examine here are Groups, the LHS, and the RHS. For all variable names, special characters should be included with backticks, i.e., ``my-var``

Groups

The Groups are the values input in `group` in the `bdotsFit` function, which are columns of the dataset used. These will be denoted G_i . Within each group, we will designate the unique values within each group as v_j , ..., whereby $G_i(v_1, v_2)$ will designate unique two unique values within G_i . The possible values of v_i will be implied by the group with which they are associated.

For example, if we have groups `vehicle` and `color`, we could specify that we are interested in all red cars and trucks with the expression `vehicle(car, truck) + color(red)`.

Formula**### Bootstrapped difference of curves**

This illustrates the case in which we are taking a simple bootstrapped difference between two curves within a single group

If only one group was provided in `bdotsFit`, we can take the bootstrapped difference between two values within the group with

```
y ~ Group1(val1, val2)
```

If more than two groups were provided, we must specify within which values of the other groups we would like to compare the differences from `Group1` in order to uniquely identify the observations. This would be

```
y ~ Group1(val1, val2) + Group2(val1)
```

For example, bootstrapping the differences between cars and trucks when `color` was provided as a second group, we would need `y ~ vehicle(car, truck) + color(red)`.

Bootstrapped difference of difference curves

This next portion illustrates the case in which we are interested in studying the difference between the differences between two groups, which we will call the `innerGroup` and the `outerGroup` following a nested container metaphor. Here, we must use caution as the order of these differences matter. Using again the vehicle example, we can describe this in two ways:

1. We may be interested in comparing the difference between red trucks and cars (`d_red`) with the difference between blue trucks and cars (`d_blue`). In this case, we will be finding the difference between cars and trucks twice (one for blue, one for red). The vehicle type is the `innerGroup`, nested within the `outerGroup`, in this case, `color`.
2. We may also be interested in comparing the difference between red trucks and blue trucks (`d_truck`) with the difference between red and blue cars (`d_car`). Here, `innerGroup` is the `color` and `outerGroup` is the `vehicle`

As our primary object of interest here is not the difference in outcome itself, but the difference of the outcome within two groups, the LHS of the formula is written `diffs(y, Group1(val1, val2))`, where `Group1` is the `innerGroup`. The RHS is then used to specify the groups of which we want to take the outer difference of. The syntax here is the same as above. Together, then, the formula looks like

```
diffs(y, Group1(val1, val2)) ~ Group2(val1, val2)
```

in the case in which only two grouping variables were provided to `bdotsFit` and

```
diffs(y, Group1(val1, val2)) ~ Group2(val1, val2) + Group3(val1) + ...
```

is used to uniquely identify the sets of differences when three or more groups were provided.

Value

Object of class 'bdotsBootObj'

Examples

```
## Not run:
```

```
## fit <- bdotsFit(cohort_unrelated, ...)
```

```
boot1 <- bdotsBoot(formula = diffs(Fixations, LookType(Cohort, Unrelated_Cohort)) ~ Group(50, 65),
                  bdObj = fit,
                  N.iter = 1000,
```

```

      alpha = 0.05,
      p.adj = "oleson",
      cores = 4)

boot2 <- bdotsBoot(formula = Fixations ~ Group(50, 65) + LookType(Cohort),
  bdObj = fit,
  N.iter = 1000,
  alpha = 0.05,
  p.adj = "oleson",
  cores = 4)

## End(Not run)

```

bdotsCorr	<i>Correlation with fixed value in bdots</i>
-----------	--

Description

Find the correlation of a fixed value with the bdots fitted curves at each time point

Usage

```
bdotsCorr(bdObj, val, ciBands = FALSE, method = "pearson")
```

Arguments

bdObj	Object of class 'bdotsObj'
val	Character string of fixed value for correlation in dataset from 'bdotsFit'
ciBands	Boolean for including confidence intervals
method	Arguments for 'cor' or 'cor.test'. The default option us 'method = "pearson"'

bdotsFit	<i>Fit nlme curves to grouped observations</i>
----------	--

Description

Creates observation level curves to use in bdotsBoot

Usage

```
bdotsFit(
  data,
  subject,
  time,
  y,
  group,
  curveType = doubleGauss(concave = TRUE),
  cor = TRUE,
  numRefits = 0,
  cores = 0,
  verbose = FALSE,
  ...
)
```

Arguments

data	Dataset used
subject	Column name of dataset containing subject identifiers
time	Column name containing time variable
y	Column name containing outcome of interest
group	Character vector containing column names of groups. Can be greater than one
curveType	See details/vignette
cor	Boolean. Autocorrelation?
numRefits	Integer indicating number of attempts to fit an observation if the first attempt fails
cores	number of cores. Default is 0, indicating half cores available
verbose	currently not used
...	Secret

Details

This is step one of the three step bdots process. Things should be more or less straight forward. The only tricky part involves curveType. For now know that one can use doubleGauss(concave = TRUE/FALSE) or logistic(). Should be passed in as a call. See the vignette on customizing this

Value

Object of class 'bdotsObj', inherits from data.table

Examples

```
## Not run:
res <- bdotsFit(data = cohort_unrelated,
  subject = "Subject",
  time = "Time",
```

```

y = "Fixations",
group = c("Group", "LookType"),
curveType = doubleGauss(concave = TRUE),
cor = TRUE,
numRefits = 2,
cores = 0,
verbose = FALSE)

## End(Not run)

```

bdotsFitter

Fits Individual Subject Curve

Description

The one subject version of bdotsFit

Usage

```

bdotsFitter(
  dat,
  curveType,
  rho,
  numRefits = 0,
  verbose,
  getCovOnly = NULL,
  params = NULL,
  splitVars = NULL,
  datVarNames = NULL,
  ...
)

```

Arguments

dat	data for single subject/group combo
curveType	this is actually a function. Should rename
rho	correlation coefficient
numRefits	number of refit attempts
verbose	not used
getCovOnly	only find covariance matrix from starting parameter values
params	starting parameters, if wanting to add manually
splitVars	variables used to identify group. Might combine with datVarNames
datVarNames	character vector indicating reponse and time values from parent call
...	not used

bdotsRefit

*Refit Observations Returned from bdotsFit***Description**

Refit Observations Returned from bdotsFit

Usage

```
bdotsRefit(
  bdObj,
  fitCode = 1L,
  quickRefit = FALSE,
  numRefits = 2L,
  paramDT = NULL,
  ...
)
```

Arguments

<code>bdObj</code>	An object of class 'bdotsObj' returned from <code>bdotsFit</code>
<code>fitCode</code>	A length one integer indicating observations to refit. See Details
<code>quickRefit</code>	Boolean indicating if a quick refit should be used. If TRUE, rather than prompting the user for adjustments for each observation, <code>bdotsRefit</code> will jitter the parameters of all observations indicated by <code>fitCode</code> and attempt to refit. Between the original and the refitted curve, this will place priority on the higher <code>fitCode</code> . If these are equal, R2 will take precedence. Otherwise, the original fit will be kept.
<code>numRefits</code>	Integer indicating the number of refit attempts after jittering parameters, either with <code>quickRefit</code> or when done individually
<code>paramDT</code>	A <code>data.table</code> or <code>data.frame</code> that matches the what is returned by <code>coefWriteout(bdObj)</code> . That is, it should have columns uniquely identifying observations with subjects and groups, as well as named columns for the parameters. NA parameters are OK. Can also be a subset of the original rows. Note, if this argument is not NULL, the remaining arguments will be ignored.
<code>...</code>	not used

Details

`fitCode` indicates lower bound on observations to refit. For example, if `fitCode = 4`, `bdotsRefit` will prompt user to refit all observations with `fitCode = 4, 5, 6`. The `quickRefit` option will attempt to jitter and refit all observations selected by `fitCode`. Otherwise, the user will be prompted through a menu to individually refit observations

ValueReturns `bdObj` with updated fits

bdRemove	<i>bdots Remove Function</i>
----------	------------------------------

Description

Remove observations with a specified fitCode and optionally all pairs

Usage

```
bdRemove(bdObj, fitCode = 6L, removePairs = TRUE)
```

Arguments

bdObj	bdots object
fitCode	min fitCode to remove. Default is 6, which removes all subjects with NULL fits (fitCode = 5 would remove 5 and 6)
removePairs	Boolean. Remove subject pairs is one of pair is removed. Default is TRUE to retain paired t-test

Details

This function is used to remove all bdots observations with a fit code equal to or larger than the argument passed to fitCode without refitting. If removePairs = TRUE, all entries for a subject will be removed if their fit failed in any of the groups in which they were a member

ci	<i>ci dataset</i>
----	-------------------

Description

ci dataset - need to include details

Usage

```
ci
```

Format

An object of class data.frame with 108216 rows and 5 columns.

coef.bdotsObj	<i>Extract bdotsFit Moedel Coefficients</i>
---------------	---

Description

Returns coefficient matrix for bdotsFit object

Usage

```
## S3 method for class 'bdotsObj'
coef(object, ...)
```

Arguments

object	A bdotsObj
...	not used

Value

Returns matrix of model coefficients for observations in object

coefWriteout	<i>Create data.table with bdotsObj parameters</i>
--------------	---

Description

Creates an object of class data.table that matches parameter values for each observation. This can then be passed to the bdotsRefit function

Usage

```
coefWriteout(bdObj)
```

Arguments

bdObj	An object returned from bdotsFit or bdotsRefit
-------	--

Value

A data.table matching parameter values to observations

Examples

```
## Not run:
fit <- bdotsFit(data = cohort_unrelated,
               subject = "Subject",
               time = "Time",
               y = "Fixations",
               group = c("Group", "LookType"),
               curveType = doubleGauss(concave = TRUE),
               cor = TRUE,
               numRefits = 2,
               cores = 0,
               verbose = FALSE)
parDT <- coefWriteout(fit)

## End(Not run)
```

cohort_unrelated	<i>cohort_unrelated dataset</i>
------------------	---------------------------------

Description

cohort_unrelated dataset - need to include details

Usage

```
cohort_unrelated
```

Format

An object of class `data.frame` with 50100 rows and 6 columns.

createCurveList	<i>Create old bdots curve list</i>
-----------------	------------------------------------

Description

Create old bdots curve list

Usage

```
createCurveList(x, prs, splitGroups)
```

Arguments

x	list of group distributions
prs	named list of parsed call
splitGroups	splitGroups

 curveFitter

Curve Fitter

Description

Used in bdotsFit

Usage

```
curveFitter(dat, ff, params, rho, numRefits = 0, getCovOnly = NULL, ...)
```

Arguments

dat	data used in building curve
ff	formula used in building curve
params	starting parameters
rho	correlation coefficient
numRefits	number of refit attempts
getCovOnly	only find covariance matrix from starting parameter values
...	don't know that this is used, can maybe get rid of it

 df_cohort_unrelated

df_cohort_unrelated dataset

Description

df_cohort_unrelated dataset - need to include details

Usage

```
df_cohort_unrelated
```

Format

An object of class `data.frame` with 78156 rows and 5 columns.

df_target	<i>df_target dataset</i>
-----------	--------------------------

Description

df_target dataset - need to include details

Usage

df_target

Format

An object of class data.frame with 37575 rows and 4 columns.

doubleGauss	<i>Double Gauss curve function for nlme</i>
-------------	---

Description

Double Gauss function used in fitting nlme curve for observations

Usage

doubleGauss(dat, y, time, params = NULL, concave = TRUE, ...)

Arguments

dat	subject data to be used
y	outcome variable, character vector
time	time variable, character vector
params	NULL unless user wants to specify starting parameters for gnls
concave	Boolean
...	just in case

Details

User should only have to worry about setting concavity of this function

$$y \sim (\text{time} < \mu) * (\exp(-1 * (\text{time} - \mu) ^ 2 / (2 * \text{sig1} ^ 2)) * (\text{ht} - \text{base1}) + \text{base1}) + (\mu <= \text{time}) * (\exp(-1 * (\text{time} - \mu) ^ 2 / (2 * \text{sig2} ^ 2)) * (\text{ht} - \text{base2}) + \text{base2})$$

doubleGauss2	<i>DoubleGauss2 curve function for nlme</i>
--------------	---

Description

DoubleGauss2 function used in fitting nlme curve for observations

Usage

```
doubleGauss2(dat, y, time, params = NULL, concave = TRUE, ...)
```

Arguments

dat	subject data to be used
y	outcome variable, character vector
time	time variable, character vector
params	NULL unless user wants to specify starting parameters for gnls
concave	Boolean
...	just in case

Details

User should only have to worry about setting concavity of this function. Presently only work for time series scaled out to 2000ms

$$y \sim (\text{time} < \mu) * (\exp(-1 * (\text{time} - \mu) ^ 2 / (2 * \text{sig1} ^ 2)) * (\text{ht} - \text{base1}) + \text{base1}) + (\mu <= \text{time}) * (\exp(-1 * (\text{time} - \mu) ^ 2 / (2 * \text{sig2} ^ 2)) * (\text{ht} - \text{base2}) + \text{base2})$$

effectiveAlpha_f	<i>Effective Alpha Functional</i>
------------------	-----------------------------------

Description

Functional that returns function for computing effective alpha for given parameters and distribution

Usage

```
effectiveAlpha_f(rho, n = 10, df = NULL, method = "norm")
```

Arguments

rho	Correlation coefficient
n	Number of observations
df	Degrees of freedom if method = "t"
method	Character string. Determines distribution for adjusted alpha can be either "norm" for normal distribution or "t" for t-dist

expCurve *Exponential curve function*

Description

Exponential function used in fitting nlme curve for observations

Usage

```
expCurve(dat, y, time, params = NULL, ...)
```

Arguments

dat	subject data to be used
y	outcome variable
time	time variable
params	NULL unless user wants to specify starting parameters for gnls
...	just in case

Details

Remove any values of zero, or jitter, before using with bdotsFit

$y \sim x_0 \exp(k \beta)$

findModifiedAlpha *Find modified alpha*

Description

find modified alpha

Usage

```
findModifiedAlpha(
  rho,
  n,
  df,
  alpha = 0.05,
  errorAcc = 0.001,
  gradDiff = ifelse(cores > 3, 0.5, 0.1),
  cores = 0,
  verbose = FALSE,
  method = "t"
)
```


Arguments

rho	correlation coefficient
n	number of observations
df	degrees of freedom if method == "t"
alpha	starting alpha from which to adjust
errorAcc	acceptable error for alphastar
gradDiff	gradient steps in algorithm
cores	number of cores. Default is zero, or half of what's available
verbose	will probably remove this
method	either "t" or "norm"

fwerAlpha	<i>fwerAlpha</i>
-----------	------------------

Description

Family wise alpha calculation

Usage

```
fwerAlpha(rho, k, n = 10)
```

Arguments

rho	Correlation coefficient
k	Bounds of non-critical region
n	Number of observations

Details

Returns effective alpha, given number of tests and the correlation coefficient. This isn't explicitly checked, but there is no reason this function should take any non-scalar values. Derivation of this can be found on pg 12 of Jake's 'Detecting time-specific differences'. This function performs the expression

$$1 - P(I_t)P(I_t | I_{t-1})^{N-1}$$

getBootDist	<i>Create Distribution for Groups</i>
-------------	---------------------------------------

Description

Create Distribution for Groups

Usage

```
getBootDist(x, b = 1000)
```

Arguments

x	A subset object from bdObj
b	Number of bootstraps

getFitCorforGroups	<i>Get Fit Correlations</i>
--------------------	-----------------------------

Description

Helper function for finding correlation of fixed value and fitted values within group

Usage

```
getFitCorforGroups(x, val, ciBands = FALSE, method = "pearson")
```

Arguments

x	A split object of class 'bdObj' split by identifiers
val	Fixed value from dataset
ciBands	boolean for including cibands
method	method for correlation function

getSubCurveValues *Return fitted values*

Description

Returns fitted values at observed times

Usage

```
getSubCurveValues(bd, origNames = TRUE, origTime = TRUE)
```

Arguments

bd	Single row of bdObj
origNames	use original names for y and time, or use "y" and "time"
origTime	Boolean. Do I actually want fitted values at observed times for that subject, or data.table with fitted values at the union of times

Details

Given a single row of bdObj, this returns fitted values at the observed times to use in conjunction with whatever else

getT *Generate t-distribution for permutation*

Description

Creates vector of t-statistics for given permutation returning either entire vector or max values

Usage

```
getT(x, idx, group, whole = FALSE)
```

Arguments

x	bdObj
idx	permutation to use
group	group that we are permuting against
whole	return vector of T stats or just the max

linear	<i>Linear curve function</i>
--------	------------------------------

Description

Linear function used in fitting nlme curve for observations

Usage

```
linear(dat, y, time, params = NULL, ...)
```

Arguments

dat	subject data to be used
y	outcome variable
time	time variable
params	NULL unless user wants to specify starting parameters for gnls
...	just in case

Details

Don't use this function please
 $y \sim \text{slope} * \text{time} + \text{intercept}$

logistic	<i>Logistic curve function for nlme</i>
----------	---

Description

Logistic function used in fitting nlme curve for observations

Usage

```
logistic(dat, y, time, params = NULL, ...)
```

Arguments

dat	subject data to be used
y	outcome variable
time	time variable
params	NULL unless user wants to specify starting parameters for gnls
...	just in case

Details

$y \sim \text{mini} + (\text{peak} - \text{mini}) / (1 + \exp(4 * \text{slope} * (\text{cross} - (\text{time})) / (\text{peak} - \text{mini})))$

parTest2	<i>Parameter t-test</i>
----------	-------------------------

Description

Perform t-test on curve parameters of `bdotsFit` object

Usage

```
parTest2(bdObj, group, vals = NULL)
```

Arguments

<code>bdObj</code>	Object of class <code>bdObj</code>
<code>group</code>	Length one character of grouping column in which to perform t-test
<code>vals</code>	Character vector of values within grouping column in which to perform the test. If NULL, it will do all pairwise tests

Details

Performs pairwise t-test. Currently only tests at $\alpha = 0.95$. Also currently only allows t-test within single grouping column. Ability to test across grouping columns to come later

Value

List of t-test results of class `bdotsPars_ttest`

Examples

```
## Not run:
res <- bdotsFit(data = cohort_unrelated,
  subject = "Subject",
  time = "Time",
  y = "Fixations",
  group = c("Group", "LookType"),
  curveType = doubleGauss(concave = TRUE),
  cor = TRUE,
  numRefits = 2,
  cores = 0,
  verbose = FALSE)
tstats <- parTest(res, group = "LookType", vals = c("Cohort", "Unrelated_Cohort"))

## End(Not run)
```

permTest	<i>Permutation testing for bboot</i>
----------	--------------------------------------

Description

Provides alternative for controlling FWER

Usage

```
permTest(x, prs, alpha, P)
```

Arguments

x	this is splitGroups (each item of list bdotsObj)
prs	from parser, indicates inner/outer groups
alpha	alpha for fwer
P	number of permutations

plot.bdotsBootObj	<i>Plot for object of class bdotsBootObj</i>
-------------------	--

Description

Allows a number of different but also unstable option for plotting an object of class bdotsBoot

Usage

```
## S3 method for class 'bdotsBootObj'
plot(x, alpha = NULL, ciBands = TRUE, plotDiffs = TRUE, group = NULL, ...)
```

Arguments

x	An object of class bdotsBootObj
alpha	Significance level for plotting confidence intervals.
ciBands	Boolean indicating whether or not to include confidence intervals around fitted curves
plotDiffs	Boolean to plot difference curve
group	Specify group to plot if difference of difference was used. The user can also subset the bdotsBootObj prior to plotting. Currently not used
...	ignore for now, but will eventually allow plot parameters

Details

This plot function is also a bit unstable and is expected to change

Value

List of ggplot objects, which may be helpful if the margins are weird

plot.bdotsCorrObj *Plots for bdotsCorr*

Description

Plots correlation of fixed value with fitted curves over time

Usage

```
## S3 method for class 'bdotsCorrObj'
plot(x, ciBands = FALSE, window = NULL, ...)
```

Arguments

x	object of class 'bdotsCorrObj'
ciBands	boolean. Whether or not to include confidence intervals in plots. Must have been selected in 'bdotsCorr'
window	A length 2 numeric vector with start and end points for the plotting window
...	super secret, don't use

plot.bdotsObj *Plot a bdotsFit object*

Description

Plot individual fits or model fit parameters from an object of class 'bdotsObj'. These functions are not very stable

Usage

```
## S3 method for class 'bdotsObj'
plot(x, fitCode = NULL, gridSize = NULL, plotfun = "fits", ...)
```

Arguments

x	An object of class 'bdotsObj' returned from bdotsFit
fitCode	Currently not used
gridSize	Length one numeric indicating size of plot grid. Default is 2x2. For right now, they are square
plotfun	Plot either subject fits or model parameters with "fits" or "pars"
...	ignore for now (other args to plot.generic)

Details

Right now, these functions are a bit unstable and expected to change. The largest current issue is with the placement of the legend, which cannot be adjusted. If you are running into issues with seeing things correctly, try making the "Plots" window in RStudio larger before running this function

Value

This will return a list of all of the plots rendered.

polynomial	<i>Polynomial curve function for nlme</i>
------------	---

Description

Polynomial function used in fitting nlme curve for observations

Usage

```
polynomial(dat, y, time, degree, raw = TRUE, params = NULL, ...)
```

Arguments

dat	subject data to be used
y	outcome variable
time	time variable
degree	degree of polynomial
raw	Boolean, use raw polynomials?
params	NULL unless user wants to specify starting parameters for gnls
...	just in case

Details

It's recommended that one uses raw polynomials for this function for numerical stability. As inference is not performed on the parameters themselves, this should have minimal consequences

$$y \sim \text{mini} + (\text{peak} - \text{mini}) / (1 + \exp(4 * \text{slope} * (\text{cross} - (\text{time})) / (\text{peak} - \text{mini})))$$

`print.bdotsBootObj` *Print 'bdotsBootObj'*

Description

Prints argument. Really, just the summary function

Usage

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

Arguments

`x` An object of class 'bdotsBootObj'
`...` Top secret alpha one code red

Details

Generic for printing 'bdotsBootObj'

`print.bdotsBootSummary`
Print bdotsBoot Summary

Description

That's pretty much it. This is a print method, so there is likely not much need to call it directly

Usage

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

Arguments

`x` generic name, but this will be an object of bdotsBootSummary
`...` ignored for now

```
print.bdotsPars_ttest2
```

Print Parameter Test Summary

Description

Print Parameter Test Summary

Usage

```
## S3 method for class 'bdotsPars_ttest2'  
print(x, ...)
```

Arguments

x	object to be printed
...	not used

Details

That's pretty much it. This is a print method, so there is likely not much need to call it directly

```
print.bdotsSummary     Print bdotsObj Summary
```

Description

Print bdotsObj Summary

Usage

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

Arguments

x	object to be printed
...	not used

Details

That's pretty much it. This is a print method, so there is likely not much need to call it directly

p_adjust

Adjust P-values for Multiple Comparisons

Description

Identical to `stats::p.adjust`, but includes `method = "oleson"`

Usage

```
p_adjust(p, method = "oleson", n = length(p), alpha = 0.05, df, rho, cores = 0)
```

Arguments

p	numeric vector of p-values (possibly with NAs).
method	correction method, a character string. Can be any of the methods in <code>p.adjust.methods</code> , with the additional value <code>method = "oleson"</code>
n	number of comparisons, must be at least <code>length(p)</code> ; only set this (to non-default) when you know what you are doing!
alpha	adjustment to be made with method <code>oleson</code>
df	degrees of freedom, if using <code>method = "oleson"</code>
rho	AR1 correlation coefficient, if using <code>method = "oleson"</code>
cores	number of cores for use in parallel, only valid for <code>method = "oleson"</code> . Default is zero, using half of the available cores

Details

This function works identically to the function `p.adjust`, with the additional option to use `method = "oleson"`. For this option, user must include a value for `df`, `alpha`. If `method = "oleson"` and no value is given for `rho`, 0.9 will be used. To compute a value for `rho` from t-statistics, use `ar1Solver`.

Value

Returns a vector of adjusted p-values just as in `p.adjust`, but with additional attributes for `alphastar` and `rho`.

See Also

[ar1Solver](#)

```
rbindlist.bdObjList    rbindlist for bdotsObjects
```

Description

Similar to data.table::rbindlist, but preserves bdotsObjects attributes

Usage

```
## S3 method for class 'bdObjList'
rbindlist(x, ...)
```

Arguments

x	bdotsObject
...	for compatability with data.table

```
split.bdObj            Split object of class bdotsObj
```

Description

Analogous to other splitting functions, but retains necessary attributes across the split object. As of now, it can only be unsplit with bdots::rbindlist

Usage

```
## S3 method for class 'bdotsObj'
split(x, f, drop = FALSE, by, ...)
```

Arguments

x	Object of class bdotsObj
f	For consistency with generic, but is not used
drop	logical. Default FALSE will not drop empty list elements caused by factor levels not referred by that factor. Analagous to data.table::split
by	Character vector of column names on which to split. Usually will be Subject or one of the fitted groups
...	not used

subset.bdotsBootObj *Subset a nested group bdotsBoot objects*

Description

Subset a nested group bdotsBoot objects

Usage

```
## S3 method for class 'bdotsBootObj'
subset(x, group, adjustAlpha = NULL, ...)
```

Arguments

x	An object returned from bdotsBoot
group	A group to subset. Must be an outer group
adjustAlpha	currently not used. Will give option to recompute adjusted alpha
...	Not used

Details

This function is used to subset a bdotsBootObject that was fit to compute the difference of differences. This allows the user to subset out the outer group in the comparison for plotting and investigation

summary.bdotsBootObj *Summary for bdotsBootObj*

Description

Provides summary information for bdotsBootObj

Usage

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

Arguments

object	An object of class bdotsObj
...	Ignored for now

Value

Returns an object of class "bdotsBootSummary". There is some summarized information included if assigned to an object, i.e., 'summ <- summary(bdotsBootObj)' then 'str(summ)'

summary.bdotsObj	<i>Summary for bdotsObj</i>
------------------	-----------------------------

Description

Provides summary information for bdotsObj

Usage

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

Arguments

object	An object of class bdotsObj
...	not used

Value

Returns an object of class "bdotsSummary". There is some summarized information included if assigned to an object, i.e., 'summ <- summary(bdObj)' then 'str(summ)'

target	<i>target dataset</i>
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Description

target dataset - need to include details

Usage

```
target
```

Format

An object of class data.frame with 25050 rows and 4 columns.

writeCSV	<i>Write fits from bdotsBoot to csv file</i>
----------	--

Description

The function is used to write out columns for each group for which a curve was bootstrapped

Usage

```
writeCSV(bootObj, file, alpha = 0.05, ...)
```

Arguments

bootObj	An object of class bdotsBootObj
file	file name to write out csv
alpha	alpha level for upper/lower CI
...	Other arguments passed to <code>data.table::fread</code>

Details

This is potentially useful for constructing plots in a separate application. There is an additional column, `Significant` indicating if a particular time point was considered significant between the difference curves. For difference of difference objects, this only indicates significance for the outer difference.

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