

bdots

Abstract

The Bootstrapped Differences of Timeseries (bdots) was first introduced by Oleson (and others) as a method for controlling type I error in a composite of serially correlated tests of differences between two time series curves in the context of eye tracking data. This methodology was originally implemented in R by Seedorff 2018. Here, we revisit the original package, both expanding the underlying theoretical components and creating a more robust implementation.

1 Introduction

In 2017, Oleson et al. introduced a method for detecting time-specific differences in densely sampled time series. This largely centered around bootstrapping and computing a series of highly correlated t -statistics and using estimates of the autocorrelation as an adjustment for the family-wise error rate. This paper was presented in the context of the Visual World Paradigm (VWP), an experimental paradigm combining eyetracking with an interactive environment to measure dynamics in language processing. In 2018, R software was introduced on CRAN to perform a limited version of the analysis proposed in Oleson. In this paper, we introduce `bdots` v2, an update to the CRAN package that broadly expands the capabilities of the original.

This paper is not intended to serve as a complete use guide to updates in the `bdots` package. Rather, the purpose is to showcase major changes and improvements to the package, with those seeking a more comprehensive treatment directed to the package vignettes. Updates to the `bdots` package have been such that there is little resemblance to the original. Rather than taking a “compare and contrast” approach, we will first enumerate the major changes, followed by a general demonstration of the package use:

1. Simplified user interface
2. User defined curves
3. Permit fitting for arbitrary number of groups
4. Updates to bootstrapping algorithm and introduction of permutation test

5. Automatic detection of paired tests based on subject identifier
6. Allows for non-homogenous sampling of data across subjects and groups
7. Introduce formula syntax for bootstrapping difference function
8. The fitted `bdots` object inherits from `data.table` class
9. `bdots` is now stylized “`bdots`”

Bootstrapped differences in time series The high level motivation, abstracted from the particulars, is more or less as follows: we are often interested in generally comparing time series between two or more groups and in particular, identifying a window of time in which they differ significantly without a priori specification of any regions. A full(er) review of previous methods can be found in Seedorff, though here we limit the scope of interest to specifying that we are interested in “developing a statistical tool to (1) detect differences in two time series (such as the VWP) and (2) to offer a precise characterization of the time window in which a difference occurs [?].” This is subsequently done in two steps. First, we use a bootstrapping procedure to estimate the group distributions of two time series; and second, we use either a FWER correction or permutation testing to identify time windows in which differences occur.

A typical instantiation of this problem occurs when we have two groups (or experimental conditions, etc.,) in which subjects have an associated time series. For example, we may be interested in comparing the growth of a particular tumor in mice over time between a control group and several candidate treatment groups. It’s assumed that each group has some distribution of associated functions in time, and we are interested in identifying windows in time in which these distributions of functions are significantly different.

FOUND SOURCE FOR THIS: *McMurray, Klein-Packard, Tomblin 2019, real time mechanics..pg 7*

Need to read that context and then weave it in here

[Not sure if we include this here]There are also situations in which we are interested in the difference of differences. I don’t remember where or why this is done, some justification for doing this instead of some complicated F test, but I can find that later. Instead, let me offer an example. I’m already not going to like this example, so let me just put it anyways knowing that it will be deleted. Suppose we are interested in understanding how the color of a vehicle differentially impacts performance based on the vehicle type. We know that there is some difference between cars and trucks. Suppose then that we look at the difference between red cars and red trucks and then the difference between blue cars and blue trucks. If color does not mediate this difference, the difference between red cars and trucks should be the same as the difference

between blue cars and trucks. This information can be determined if we look then at the difference between the differences.

The original `bdots` package was predicated on comparing differences between dense, highly correlated time series by first specifying functional forms and then performing statistical tests on each of the observed time points. With version 2.0, capabilities have drastically improved, and `bdots` is able to fit parametric functions to any type of data observed in time. Along with methodological improvements, we have included more options in determining statistical significance in the differences of curves, utilizing a **robust** permutation testing framework when the assumptions of autocorrelation do not hold. In addition to methodology, a number of quality-of-life improvements have also been made, greatly simplifying syntax, creating more **robust** functions, and including a collection of useful methods for handling returned objects.

In summary, `bdots` has transitioned from a package focused exclusively on densely sampled timeseries assuming a limited number of functional forms to a **robust** framework for identifying time windows of significant difference across a wide breadth of timeseries-adjacent data.

2 Methodology and Overview

There are a few minor changes and additions in the underlying methodology used in the `bdots` package which we will briefly review here. For those interested in a more detailed description, see chapter 3 of my dissertation.

Actually need to nail down how I want to organize this. There is basically:

1. Creating group distributions
 - (a) Getting fits
 - (b) Bootstrapping for dist
2. Finding areas of significance
 - (a) Permutation testing
 - (b) FWER/oleson adjustment

2.1 Creating Group Distributions

Broadly, there are two steps to performing an analysis with the `bdots` package: fitting the curves to observed data and bootstrapping differences between groups. The first step involves specifying an underlying curve f ,

which is assumed to be parametric¹. Along with the observed data y for each i th subject, `bdots`, via fitting with `gnls`, returns a set of parameters along with an estimate of their covariance.

$$F : f \times y_i \rightarrow N(\hat{\theta}_i, V_i), \quad (1)$$

where θ is a length- p vector representing the parameters of the function.

Once fits have been made, we are ready for testing the bootstrapped difference between curves. After specifying the groups of interest for analysis, two algorithms are implemented: a bootstrapping algorithm is used to determine the distribution of each group of curves, and either permutation testing or bootstrapping is used to specify regions of statistically significant differences, depending on the underlying assumptions made. The algorithm for bootstrapping for each group is as follows:

1. For a group of size n , select n subjects from the group, *with replacement*. This controls for the between subject variability
2. For each selected subject, draw a set of parameters from the distribution $\theta_i^* \sim N(\hat{\theta}_i, V_i)$. This permits us to account for within subject variability
3. For each of the resampled θ_i^* , find the b th bootstrap estimate for the group $\theta_b = \frac{1}{n} \sum_{i=1}^n \theta_i^*$
4. Perform this sequence B times

The end results is a $B \times p$ matrix containing a bootstrapped sample of the group distribution for θ . Each row of this matrix is used to create a $1 \times T$ vector representing f_θ evaluated at T time points. This results in a $B \times T$ matrix representing a collection of bootstrapped curves evaluated at each time point, in total representing a bootstrapped distribution of the curves.

Next we attend to identifying regions in which a statistically significant difference between curves is present, choosing from one of the two methods currently available.

2.2 Evaluating significance differences

see maybe here introduce both and the point, along with basic assumptions

2.2.1 Permutation Testing

The simplest method implemented for identifying time-specific differences is permutation testing, ideal when minimal assumptions can be made on the observed data.

¹the option to include non-parametric functions is anticipated in the future work of this package. The process will be similar, however, with θ then representing the number and location of the knots for splines

We begin by computing a t -statistic of the difference at each time point,

$$T(t) = \frac{|\bar{f}_1(t) - \bar{f}_2(t)|}{\sqrt{\frac{1}{n_2} \text{Var}(f_1(t)) + \frac{1}{n_2} \text{Var}(f_2(t))}}, \quad \Leftarrow \text{check these (specifically } n \text{ in denom)} \quad (2)$$

or, in the case of paired groups,

$$T(t) = \frac{\bar{f}_D(t)}{\sqrt{\frac{1}{n} \text{Var}(f_D(t))}}. \quad (3)$$

Next, we go about creating a null distribution against which to test our hypothesis that there is no difference between each group at each time point. We do this with permutations, the algorithm being as follows: for two groups, with n_1 and n_2 subjects in each:

1. Assign to each subject a label indicating group membership
2. Randomly shuffle the labels indicating membership, creating two new groups with n_1 and n_2 subjects in each
3. Recalculate the t -statistic, $T(t)$ and record the maximum of each permutation

The collection of maximum values for $T(t)$ will serve as the null distribution against which to compare our observed $T(t)$. Regions in which the observed t statistic are beyond the specified α in the null distribution are then considered significant.

2.2.2 FWER Adjustment

In addition to permutation testing, there are also adjustments that can be made to control for the family-wise error rate. As was done with permutation testing, we begin by computing a t -statistic at each time point for the observed data,

$$T(t) = \frac{|\bar{f}_1(t) - \bar{f}_2(t)|}{\sqrt{\frac{1}{n_2} \text{Var}(f_1(t)) + \frac{1}{n_2} \text{Var}(f_2(t))}}, \quad \Leftarrow \text{check these (specifically } n \text{ in denom)} \quad (4)$$

or, again in the case of paired groups,

$$T(t) = \frac{\bar{f}_D(t)}{\sqrt{\frac{1}{n} \text{Var}(f_D(t))}}. \quad (5)$$

Unlike the case with the permuted data, we have no need for a null distribution, instead determining significance by considering the observed statistics against a modified α . Adjustments that can be made

include all of the adjustments found in `stats::p.adjust`, and adjustment made on the false discovery rate (FDR), and finally the adjustment “oleson” used in the original `bdots` package. In short, the “oleson” adjustment makes use of an autocorrelation parameter to adjust for the highly correlated series of t -statistics. A full treatment of this methodology is included in Oleson 2017.

3 Example Data

We will illustrate use of the updated `bdots` package with a worked example, using an artificial dataset to help detail some of the newer aspects of the package. The dataset will consist of outcomes for a collection of vehicles, consisting of eight distinct groups. These groups will be nested in order of vehicle origin (foreign or domestic), vehicle class (car or truck), and vehicle color (red or blue). Further, vehicles of different color but within the same origin and class groups will be considered paired observations. A table detailing the relationship of the groups is shown here:

Origin	Class	Color
foreign	car	red
		blue
	truck	red
		blue
domestic	car	red
		blue
	truck	red
		blue

The outcome here is simply y due to a lack of creativity, but the functional form assumed (and used in data generation) follows the four parameter logistic,

$$f_{\theta}(t) = b + \frac{p - b}{1 + \exp\left(\frac{4s}{p-b}(c - t)\right)}, \quad (6)$$

where b , p , s , and c represent the baseline, peak, slope, and crossover points, respectively

4 Fitting Curves

The curve fitting process is performed with the `bfit` function (previously `bdotsFit`), taking the following arguments: (removing ‘cor’ and numRefits)

```
bfit(data, subject, time, y, group, curveType, cores, ...)
```

Curve functions Each of `subject`, `time`, and `y`, are length one character vectors representing columns of the dataset used in `data`, while `group` is a character vector (of varying length), also column(s) found in `data`. New here is `curveType`, taking as an argument an R call to a particular curve, for example the four parameter logistic, `bdots::logistic()`. This is done to self-contain any additional arguments associated with the fitting curve, for example the concavity of the double-Gaussian (`curveType = doubleGauss(concave = TRUE)`) or the number of degrees in a polynomial (`curveType = polynomial(degree = 5)`). A number of curves are included with the `bdots` package, including those for the four-parameter logistic, the double-Gaussian, an exponential curve, and polynomials of arbitrary degree. A detailed vignette on writing custom curves can be found with `vignette("bdots")`

Notably, `bdots` can now fit curves to an arbitrary number of groups at once, so long as all have the same parametric specification. Fitting a collection of curves to our vehicle data on all of the groups with the logistic function would look like

```
# Need to change these once I get real data
fit <- bfit(data = Vehicle, subject = "vehicle",
  time = "Time", y = "out", group = c("origin", "class", "color"),
  curveType = logistic(), cores, ...).
```

Return object and generics The function `bfit` returns an object of class `bdotsObj`, inheriting from class `data.table`. As such, each row uniquely identifies one permutation of subject and group values. Included in this row are the subject identifier, group classification, summary statistics regarding the curves, and a nested `gnls` object.

Several methods exist for this object, including `plot`, `summary`, and `coef`, returning a matrix of fitted coefficients returned from `gnls`. One consequence of inheriting from `data.table`, we are able to utilize `data.table` syntax. Note, for example, the differences between `coef(fit)`, `coef(fit[group == "A",])`, and `coef(fit[group == "B",])`. This is especially helpful when looking to plot only a subset of the fitted curves, i.e., `plot(fit[group == "A",])`

Fit Codes need to decide what i'm going to do about this. would be nice if the metric used here was module, and it can be, just not right now because that would be a lot of extra work for nothing. I could just ignore this issue all togettrher, indicate that this is for correlation and R2, and made absolutely no mention of how this possibly conflicts with other types of data

5 Bootstrapping

Once fits have been made, we are ready to begin estimating the group distributions.

Like the fitting function, the bootstrapping process has been consolidated to a single function, `bboot` (previously `bdotsBoot`). The number of options included in the `bboot` function have expanded to include a new formula syntax for specifying the analysis of interest as well as to include options for permutation testing. A call to `bboot` takes the following form

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

By default, significance is determined with an adjustment to the family-wise error rate, as was done in the original implementation of `bdots`, using method `padj = "oleson"`. When setting the argument `permutation = TRUE`, `padj` is ignored and permutation testing for regions of difference is used instead. Bootstrapping is still used for determining the group distribution, however.

A key component of the bootstrapping function is specifying which groups in our dataset we are wishing to analyze and how. This is done with a formula syntax unique to `bdots` explained in the next section.

5.1 Bootstrapping Formula

The formula argument serves two functions in `bboot`: first, it specifies the collection of curves we wish to investigate the difference between, and second, it determines if we are interested in directly comparing the differences or the difference of differences between curves.

To begin, let's reintroduce the structure of the groups we have in our dataset. Recall that we have foreign and domestic cars and trucks, and each of these vehicles comes in red and blue. Recall also that the different colors of each vehicle are considered paired observations.

Origin	Class	Color
foreign	car	red
		blue
	truck	red
		blue
domestic	car	red
		blue
	truck	red
		blue

Let's begin with a simple case: supposing we want to investigate the difference in outcome between foreign and domestic cars. Notionally, we would write

$$y \sim \text{Origin}(\text{foreign}, \text{domestic}).$$

Specifically, note that this is of the form

$$\text{outcome} \sim \text{GroupName}(\text{value1}, \text{value2})[\text{<- include this?}]$$

Note that this involves the grouping variable, `Origin`, with the two values we are interested in comparing, `domestic` and `foreign`. With this specification, the distribution of functions considered in `domestic` include all red and blue domestic cars and trucks.

If we wanted to limit our investigation to only foreign and domestic cars, we would do this by including an extra term specifying the group and the desired value. In this case,

$$y \sim \text{Origin}(\text{foreign}, \text{domestic}) + \text{Class}(\text{car}).$$

To compare only foreign and domestic *red* cars, we would add an additional term for color:

$$y \sim \text{Origin}(\text{foreign}, \text{domestic}) + \text{Class}(\text{car}) + \text{Color}(\text{red}).$$

[this section needs rewritten still its not correct] In each of these cases, we have specified particular groups or nesting of groups whose outcomes we wish to compare. Alternatively, we may be interested in comparing the *difference of differences*. For example, suppose we suspect that there may be some difference between red and blue vehicles, and that this difference may be different for cars compared to trucks. Whereas previously we were interested in comparing the differences in y between origin, we are now interested in comparing the differences of y between colors. Consequently, we will include this difference on the left hand side of the formula as our new outcome. This is done using `diffs` syntax as such:

$$\text{diffs}(y, \text{Color}(\text{red}, \text{blue})) \sim \text{Class}(\text{car}, \text{truck})$$

Similar as to the case before, if we wanted to limit this difference of differences investigation to only include domestic vehicles, we can do so by including an additional term:

$$\text{diffs}(y, \text{Color}(\text{red}, \text{blue})) \sim \text{Class}(\text{car}, \text{truck}) + \text{Origin}(\text{domestic}).$$

The formula syntax was originally contrived to make comparisons within groups or within nested groups. Conceivably, however, one could be interested in making the comparison between domestic red trucks and foreign blue cars. Doing so requires a bit of a work around. Examples detailing how one might go about doing this are included in the appendix.

First, there would be some function of sorts, something like `makeUniqueGroups` which would create a new group column with each permutation of previous groups being given a unique identifier. Doing this on the vehicle example would look something like `fit <- makeuniquewhatever` resulting in the following grouping structure (for example) (and maybe you could specify group name and values who knows, kinda like factor this is just a working thought example)

Origin	Class	Color	bgroup
foreign	car	red	A
		blue	B
	truck	red	C
		blue	D
domestic	car	red	E
		blue	F
	truck	red	G
		blue	H

To then investigate differences in outcome between a foreign red car and a domestic blue truck would simply then be

$$y \sim \text{bgroup}(A, H)$$

yeah not like sexy or anything but whatever it would work.

5.2 Bootstrapping and Permutation

I guess just quickly recap what was said previously and how it might be used here.

Once we have determined the groups we are interested in comparing, we are ready to call the `bboot` function. As detailed earlier, `bboot` is now able to perform permutation testing on curve differences in addition to bootstrapping. We will discuss each of these briefly in more detail.

Bootstrapping As in the original iteration of the package, `bdots` seeks to identify differences in curves without a priori specification of a time window. This is done as in (other paper), where curves are bootstrapped to create a sampling distribution, and a collection of t -tests is performed at each observed timepoint. The FWER is controlled against this series of tests by making an adjustment to the nominal α – the novel

contribution of the original **bdots** package was implementing a correction based on an estimate of the autocorrelation of tests (oleson paper). In addition to this adjustment, **bdots** also allows corrections to be made using a number of other methods, including `fdr`, `bonferonni`, and I think maybe another. The type of correction made is specified with the `padj` argument in `bboot`.

Permutation In addition to the bootstrapping algorithm just described, **bdots** now also includes the option for running a permutation test to establish a null distribution on the differences between curves. This is done by setting the argument in `bboot(..., permutation = TRUE)`. `Niter` now refers to the number of permutations performed rather than the number of bootstraps. Additionally, when `permutation = TRUE` is set, the argument to `padj` is ignored. Permutation testing may have less power in some cases (explored in other paper), but is maximally robust and ideal for situations in which the number of assumptions made is limited.

5.3 Summary and Analysis

Let's begin first by running `bboot` using bootstrapping to compare the outcome between domestic cars and trucks

```
boot <- bboot(y ~ Vehicle(car, truck) + Origin(domestic), fit)
```

This returns an object of class

Also need to include in this section

1. plots
2. summary output example
3. maybe more plots

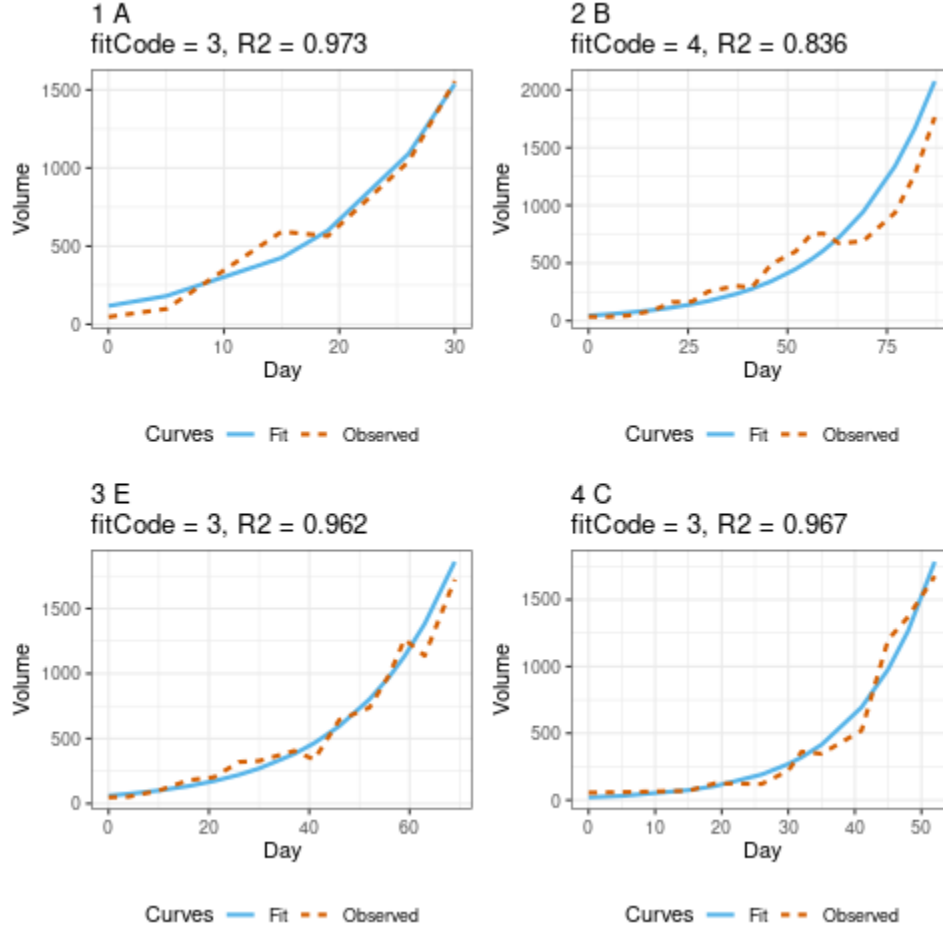
6 Extensions? Plots? I don't know!

Let's do a brief tour of some of the other additions to **bdots** that probably doesn't warrant its own section for use

6.1 Non-homogenous sampling

The **bdots** package now has support for data with non-homogenous time sampling across subjects or trials. For example, here is data collected comparing tumor growth for 451LuBr cell line in mice with repeated measures and five treatment groups

```
with(tumrdata[subjects 1-4, ], plot(observations as points))
```



It is not a problem to fit these groups and perform our bootstrapping analysis either on the union of observed time, or some custom range in between

```
<bfit, bboot, summary, plot> but for rats
```

`bdots` also allows for repeated observations, as is the case with saccade data from the VWP. Here, an individual subject has 30 trials with saccades taken at the trial level. That is, rather than taking a sequence of observations for each subject, `bfit` allows for an unordered set with observations and associated time, $\mathcal{S}_i = \{(y_j, t_j)\}$ across j observations. As this relates to the VWP, you can read more about his development in my dope ass other paper called chapter 2.

6.2 Refitting

There are sometimes situations in which the fitted function returned by `bfit` is a poor fit for some of the subjects. This can be evidenced by the `fitCode` or via a visual inspection of the fitted functions against the

observations for each subject. When this occurs, there are several options available to the user, all of which are provided through the function `brefit` (previously `bdotsRefit`). `brefit` takes the following arguments:

```
Do I need an image for the side by side plots? I assume no brefit(bdObj, fitCode = 1L, quickRefit =  
FALSE, numRefits = 2L, paramDT = NULL, ...)
```

The first of these arguments, outside of the object itself, is `fitCode`, indicating the minimum fit code to be included in the refitting process. This is a convenient way to limit the refitting process to those of a particular quality. The `quickRefit` option allows the fitter to run automatically, jittering the previous set of parameters for each refitted subject and comparing the new fit to the previous, keeping the better of the two. `numRefits` indicates how many attempts the fitter should make in doing this. Finally, `paramDT` allows for a `data.table` with columns for subject, group identifiers, and parameters to be passed in as a new set of starting parameters. This `data.table` requires the same format as that returned by `bdots::coefWriteout`.

When `quickRefit = FALSE`, the user is put through a series of prompts whereby for each subject to be refit, in addition to being given a series of diagnostics, they have the option to:

1. Keep the original fit
2. Jitter starting parameters
3. Adjust starting parameters manually
4. Remove AR1 assumptions (come back to this)
5. See original fit metrics again
6. Delete subject
7. Save and exit the refitter

```
idk show comparison of plots for refitting i guess
```

As the menu item suggests, users have the ability to end the manually refitting process early and save where they had left off. This looks like this

```
refit <- brefit(fit, ...)  
refit <- brefit(refit, ...) # pass that shit back in
```

A final note should be said regarding the option to delete a subject. As `bdots` now automatically determines if subjects are paired based on subject identifiers (necessary for calculations in the significance

testing step), it is critical that if a subject has a poor fit in on group and must be removed that he or she is also removed from all additional groups in order to retain paired status. This can be overwritten with a final prompt in the `brefit` function before they are removed.

Additionally, this can be done independent of the refitter with the new function `bdRemove`

6.3 User created curves

I know I mentioned this elsewhere, but I might erase it there and move a fuller discussion of it here

6.4 Correlations

There are sometimes cases in which we are interested in determining the correlation of a fixed attribute with group outcome responses across time (what such a case may be, I have no idea). This can be done with the `bcorr` function (previously `bdotsCorr`), which takes as an argument an object of class `bdotsObj` as well as a character vector representing a column from the original dataset used in `bfit`

```
bcorr(fit, "value", ciBands, method = "pearson")
```

This returns a thing that can be plotted. Idk, it really doesn't seem that important to me

6.5 α Adjustment

This probably last section that needs anything special, and that is an update to the `p.adjust` function, `p.adjust`, identical to `p.adjust` except that it accepts method "oleson" and takes additional arguments `rho`, `df`, and `cores`. `rho` determines the autocorrelation estimate for the oleson adjustment while `df` returns the degrees of freedom used to compute the original vector of t-statistics. If an estimate of `rho` isn't available, one can be computed on a vector of t-statistics using the `ar1Solver` function:

```
t <- diffinv(rnorm(999))
rho <- ar1Solver(t)
```

7 Discussion

need to redo this section pretty much from scratch

I'm not really sure what to include in the discussion. We don't need to compare it to other approaches for analyzing this data, as that's already been done. I can point to full methodology paper to see improvements in CI coverage and difference detection/power. Reporting should be fairly simple, an α is given which is

used to set the threshold for permutation tests – nothing else needs to be done. The previous `bdots` package suggested reporting quality of individual fits that made up the bootstrapped curves, though that was based on R^2 and $AR(1)$ status. The former of these is problem specific, the latter now irrelevant.

raff raff raff raff raff

Improvements made to the `bdots` package have drastically improved the ease of use of the package and the scope of the types of problems it is able to address. The consolidation of major components into two functions has also streamlined use. Quality of life improvements include multiple group fitting, formula syntax for bootstraps, tractable return objects, and others. Generics have also been good. The package is also now statistically correct. It has extended the types of data that can be accommodated, including heterogenous observations across time and the ability to construct user-specified curves. it really is a pretty neat package.