6404 Take-home Exam 3: Wavelets, Categorical Data Analysis, Nonparametric Regression

Peter Williams, pwilliams60@gatech.edu

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1. Wavelets (50%)

Locate a one-dimensional data set that has sharp-changes like those presented in the recent lectures for applying the following wavelet procedures. It is best that the data size is larger than 512, and is in 2-factorial, e.g., 210 = 1024. Note that you need to locate proper R-package/codes to perform the tasks below.

For these exercises a dataset was located from the data science competition website, Kaggle (https://www.kaggle.com/datasets). The dataset consists of estimated energy consumption in Megawatts (MW) by hour provided by American Electric Power (AEP). AEP is a major investor-owned electric utility in the United States of America, which provides electricity to more than five million customers in 11 states including Texas and Ohio

To simplify subsequent wavelet analysis, a subset of the original dataset was utilized. It consists of n = 1024 observations of hourly Megawatt consumption estimates beginning from 9:00am June 21st, 2018 to 12:00am August 3rd, 2018. A preview of the original dataset is shown below:

Hourly (Datetime)	Energy Usage (MW)
6/21/18 9:00	15,286
6/21/18 10:00	15,958
6/21/18 11:00	16,495
6/21/18 12:00	17,001
6/21/18 13:00	17,398

6/21/18 14:00

Table 1: AEP Hourly Energy Consumption in Megawatts

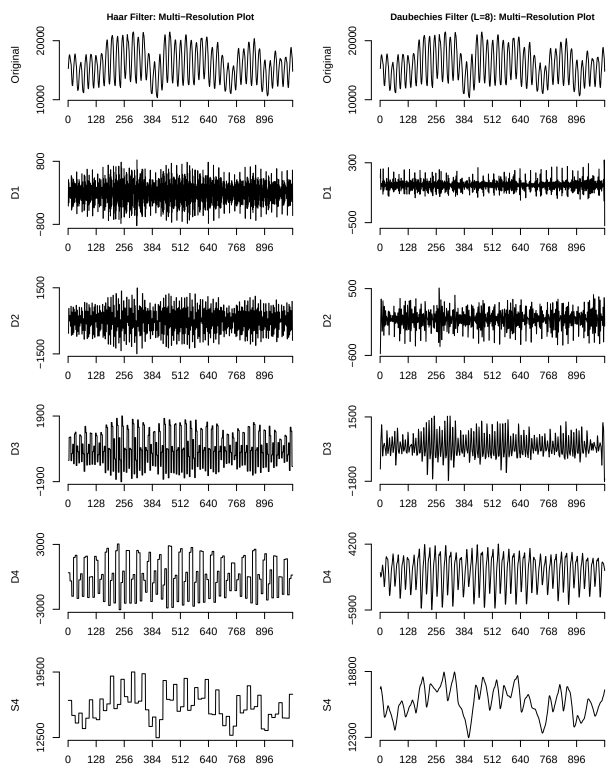
17,603

A multi-resolution plot, including the original time series of energy consumption data, provides a lot of context on the data. The plot of the original data shown below highlights significant shifts in energy consumption in the time series, and a persistent daily cyclical pattern.

To contrast the multi-resolution plots below, we have selected both the 'Haar' and 'Daubechies' filters for comparison in each column. For visual simplicity and interpretation both plots rely on 4 levels, or depth in terms of signal reconstruction. The MRA plot quickly highlights, for both families, that for the first scaling level *S*2, we have a good representation of the original function, that at further depth the signals are much noisier. Also worth noting is the contrast in smoothness of the 'Daubechies' first level decomposition vs. the 'Haar'. In fact the first level for the 'Daubechies' family captures more of the original signal, as the magnitude, or scale of signals at lower levels is tighter relative to the 'Haar' decomposition, as shown below:

i) Select two families of wavelets for completing the two tasks ii) and iii) below, and make comparisons for the results impacted from distinct wavelet families.

ii) Show a multi-resolution plot of mother and father discrete wavelet-coefficients (DWTs), and make comments about their values.



Note: To generate these plots, the 'waveslim' package was utilized in R, using the "mra" function with method "DWT" supplied as an argument which provides an additive decomposition of the original time series. Futher details are provided in the appendix.

iii) Apply two thresholding/shrinkage methods to reduce number of non-zero DWT coefficients. Apply the IDWT to reconstruct the original data signal by using the thresholded DWTs. Comment on the quality of the reconstructions from plotting the original and reconstructed signals (by overlaying them in one figure). Compare the two methods about their data-reduction ratios and the MSEs.

iv) Artificially alter the data in one "local-region" and one large-size "global region" for creating 3 distinct "fault-class" data sets. See lecture presentation about details of this task. Apply the best thresholding method (and the best wavelet-family) to model the data from all FOUR classes (one from the original data and the other 3 fault-class data). Use the multi-resolution plots of thresholded-DWTs to see how they are different in these FOUR classes of data signals.

v) Discuss the possibility and steps for developing a rigorous decision-making procedure to detect faulty-signals against the original data, and distinguish classes of faulty signals with reduced-size data presented by thresholded DWTs.

2. Categorical Data Analysis (25%)

Chapter 9 of the textbook on categorical data analysis includes 6 sections with various problems/data and methods. Locate three sets of problems/data matching three distinct methods taught in lectures. Apply proper statistical software (preferred to be in R-codes) to analyze the data. Provide in-depth comments about the findings in your statistical analyses.

1. Chi-square, goodness of fit

To discuss the Chi-square goodness of fit procedure, a dataset published by the *National Oceanic and Atmospheric Administration* was located that tracks the number of hurricanes that have made landfall on the continental United States by decade over the last ~ 60 years. Links to the actual source data is linked below. To get a sense of what the dataset looks like a brief preview is shown below, where the expected count of hurricanes is computed as the grand mean of observed landfalls across the entire dataset:

Table 2: NOAA: Continental United States: Hurricane Impacts/Landfalls

Count of Impacts	Expected Impacts
18	15.667
15	15.667
12	15.667
16	15.667
14	15.667
19	15.667
	18 15 12 16 14

Source: Continental United States Hurricane Impacts/Landfalls by decade as reported by the NOAA http://www.aoml.noaa.gov/hrd/hurdat/All_U.S._Hurricanes.html

Given the overall national interest in climate change, and its impact on weather patterns, it is of interest to many researchers if the frequency of hurricane landfalls in recent years in increasing. Given the dataset shown above, we can test this hypothesis in a crude way utilizing differences in observed vs. expected frequency of hurricane landfall. Where n_i refers to observed counts across i = 1, ..., k decades measured, and $N = \sum_{i=1}^{k} n_i$, is the total number of observed landfalls. We can then set up our test:

$$H_0: F_X(x) = F_0(x)$$

$$H_\alpha: F_X(x) \neq F_0(x)$$

And then take our test statistic T to be, $T = \sum_{i}^{r} \frac{(n_{i} - np_{i})^{2}}{np_{i}}$, which is approximately distributed $T \sim \chi^{2}_{r-1}$. Jumping into actual calculations, we have $T = \frac{(18 - np_{1})^{2}}{np_{1}} + \frac{(15 - np_{2})^{2}}{np_{2}} + ... + \frac{(19 - np_{6})^{2}}{np_{6}} = 2.128$, where $np_{i} = 15\frac{2}{3}$, for i = 1, ..., 6. Under our H_{0} , our p-value is 0.831, not sufficient evidence to reject our H_{0} and conclude

that hurricane landfalls are different across decades, but perhaps a contentious talking point in conversation. Obviously the scope and depth of the analysis here isn't sufficient to add meaningfully to the overall discussion about climate change.

2. Contingency tables, testing for homogeneity and independence

To demonstrate usage of contigency tables and associated tests using R, an article published by medical researchers that followed N=6272, Swedish men for 30 years to see whether there was any association between the amount of fish in their diet and prostate cancer was located. According to authors of the research, the original study actually used pairs of twins. This approach allowed researchers to share their findings with more confidence. In the table below, we show frequency counts among research respondents across their fish diet type, and whether or not they contracted prostate cancer across the 30 year study:

Table 3: Occurrence of Prostate Cancer, By Amount of Fish Consumption, N = 6272

	Large	Moderate	Small	Never
No	507	2769	2420	110
Yes	42	209	201	14

Source: Lancet, June 2001, "Fatty Fish Consumption and Risk of Prostate Cancer"

The table itself does not communicate readily any differences in cancer rates by diet type. To evaluate further how diet type may impact the rate of cancer, we utilize the chi-square test of independence, which is readily available via the function *chisq.est* in base R. Given a contingency table of consisting of m levels of a factor, along with an additional factor with k levels of observations, we can construct a matrix $N=(n_{ij}), i=1,...,m, j=1,...,k$, where each entry in N is an observed frequency, or count. The main gist of the chi-square test, is to study observed frequencies in N, n_{ij} , and their potential differences from the expected (under the hypothesis of independence) frequencies, denoted: $\hat{n}_{ij} = \frac{n_i \cdot n_j}{n_{...}}$, where $n_{i.} = \sum_{j=1}^k n_{ij}$, and $n_{j.} = \sum_{i=1}^m n_{ij}$, and compute our test statistic $T = \sum_{i=1}^m \sum_{j=1}^k \frac{(n_{ij} - n_{ij})^2}{n_{ij}}$, where $T \sim \chi^2$. The results of the test for our fish diet dataset are summarized in the table below:

Table 4: Chi-square Test for Independence Test, R (base), function: "chisq.test (df = 3)"

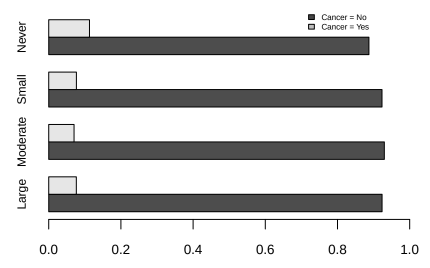
Result	Value
Test Statistic (Chi)	3.677
P-value	0.298

As shown, our resulting P-value 0.298 does not provide evidence for us to conclude that the observed frequency of prostate cancer by fish diet type, is unexpected by itself. While this finding may be interesting, this test alone isn't sufficient to rule out diet's role in respondent's contraction of prostate, as there are potentially many other factors which can contribute to prostate cancer in addition to diet alone.

3. Fisher exact test

Taking our analyis of prostate cancer, and fish diet further, visual inspection of the marginal proportion of prostate cancer by fish diet classification shows that those who never consumed fish in their diet, may have had slightly higher rates of prostate cancer, which was identified and estimated in the *Lancet* paper referenced above. Here is a basic barplot to visualize difference in rate of cancer by diet:

Marginal Proportion of Respondents With Prostate Cancer by Fish Diet



Since, as shown in the text, the Fisher exact test is based on the null hypothesis that two factors, each with two factor levels, are independent, conditional on fixing marginal frequencies for both factors. However, the fish diet dataset described above is 2×4 , therefore for the purposes of this question, we subset our data to just those with 'Never', and 'Large' fish diets for comparison.

As shown below, relying on the only the *fisher.test* function to compute *Fisher's Exact Test* in *R* (*base*) to detect any differences in counts across diet categories. The test shown below, relies on the odds ratio between the occurrence of cancer between groups ('Never' and 'Large'):

Table 5: Fisher's Exact Test: Fish Diet Comparison (Confidence = 0.95)

Result	Value
Odds Ratio	1.535
Upper Confidence Level	0.747
Lower Confidence Level	2.989
P-Value	0.207

Again, our resulting P-value from *Fisher's Exact Test* does not provide evidence for us to conclude that the observed frequency of prostate cancer by fish diet type, is unexpected. As highlighted before, there are potentially many other factors which can contribute to prostate cancer in addition to diet alone. The research referenced above in *Lancet* takes a deeper dive into this particular dataset and concludes that diet, in fact, did play a role describing respondents' rate of contracting prostate cancer, however, relying on more sophisticated statistical techniques.

3. Nonparametric Regression (25%)

Locate a data set suitable for both kernel and spline regressions. The data should include at least 3x-variables. It is okay to focus on additive-models discussed in lectures.

To consider procedures for both kernel and spline regression, we utilize the well-known automobile MPG dataset that is available in the UCI Machine Learning library. Link here: https://archive.ics.uci.edu/ml/machine-learning-databases/auto-mpg/auto-mpg.data. The dataset was used in the 1983 American Statistical Association Exposition, and is also well referenced in the following paper:

Quinlan,R. (1993). Combining Instance-Based and Model-Based Learning. In Proceedings on the Tenth International Conference of Machine Learning, 236-243, University of Massachusetts, Amherst.

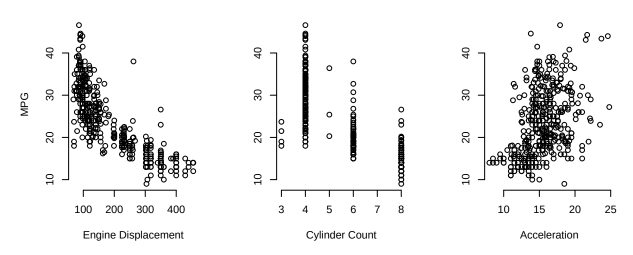
As highlighted by Quinlan in the paper above, this dataset is concerned with city-cycle fuel consumption in miles per gallon, to be predicted in terms of a number of attributes of each car. Each car in the dataset (N=392) was manufactured sometime in between 1970 and 1982. For convenience purposes we have removed observations with missing values. Based on the exercise specification we will rely on a car's *Mileage Per Gallon* (mpg), as our outcome of interest, and utilize 3 co-variates, namely the car's *Cylinder* count (cyl), *Engine Displacement* (disp), and *Acceleration* (accel). A preview of the data is provided below:

MPG	Displacement	Cylinder Count	Acceleration	Model
18	307	8	12.0	chevrolet chevelle malibu
15	350	8	11.5	buick skylark 320
18	318	8	11.0	plymouth satellite
16	304	8	12.0	amc rebel sst
17	302	8	10.5	ford torino
15	429	8	10.0	ford galaxie 500

Table 6: Preview of Automobile MPG Dataset (Published 1993)

One aspect of this dataset that makes it interestingly suite to kernel and spline regression, is that visually, there is evidence of non-linear relationship between the outcome of interest (MPG) and Engine Displacement. The relationship between MPG, Cylinder count and Acceleration is less clear, and shown below in a series of scatter plots:

MPG vs. Displacement, Cylinder Count and Acceleration



1) Go through one-variable-at-a-time kernel- and spline-regression fits to the data for all 3 x-variables. Compare the fitting results using the leave-one-out cross-validation procedure.

Iterating over the auto-mpg set, we fit both a spline, and kernel regression model, removing one target observation, and compared the prediction error generated out of sample for each model. Summary statistics resulting from the leave-one-out computation are presented below:

Table 7: Regression Model Error Comparisons: Leave-One-Out Procedure

Statistic	Kernel	Spline
Median Error	-0.30	-0.36

Statistic	Kernel	Spline
Error: 75th Percentile	2.09	2.03
Error: 25th Percentile	-2.50	-2.45
R-Squared	0.71	0.69
Mean Absolute Percentage Error	13.05	13.26

The summary statistics from this table clearly highlight that both models perform similarly across a number of simple error statistics. While the kernel regression approach may have a slight edge across a few statistics, in general, both models provided results with symmetric, generally unbiased out-of-sample prediction error for each leave-out observation. While the leave-one-out procedure doesn't highlight any significant differences in performance, further analysis and computation below does show some of the practical benefits of and limitations of each approach to solving the regression problem of predicting a car's *MPG* based on its characteristics.

Note that computational details for the leave-one-out procedure are detailed in the code appendix.

2) Select 2 sets of x-locations, e.g., (1st set: x1 = 3, x2 = 5, x3 = 2, where 3, 5, 2 are values within x-data range). These 2 sets of x-locations should be from a location close to the center of x-data-range and another location closer to the edge. Make predictions of Y at these x-data. Compare the predictions from kernel and spline methods, and also comment on the impact from x-data-locations.

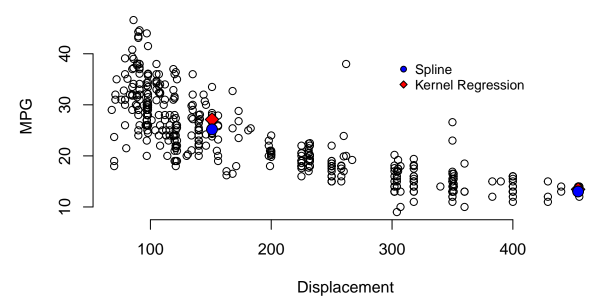
To select *X*-locations, we took the median across all *X* variables, i.e. *Displacement*, *Cylinder Count*, and *Acceleration* as one *X*-location, denoted as the "Center Point" in subsequence analysis. We then also created an "Outside Point" which is based on the maximum across all *X* variables, less 1. That is a point very close to the maximum across all co-variates. The table below summarizes the exact location of these points, and also provides associated predicted values from both kernel regression and spline models that were fit to the full auto-mpg dataset:

Table 8: Comparison of Predictions from Spline & Kernel Regression Methods

Displacement	Cylinder	Acceleration	X-Location	Kernel Prediction	Spline Prediction
151 454	4		center point	27.13	25.20
454	7	23.8	outside point	13.45	13.03

It is clear based on the table above that the spline provides a more conservative prediction of car MPG for both the center and outside points. To get a sense of how these predictions fare relative to our actual data, the scatter plot below shows our key predictor *Displacement* on the x-axis, against our outcome of interest *MPG* on the y-axis. The predicted MPG given each model is then shown in the colored points below:

Center & Outer X-Location Predictions vs. Data



The scatter plot again shows that the spline provides a slightly more conservative predictin vs. the kernal regression. However, both predictions do lie in a reasonable place relative to actual data points observed.

3) Construct the 90% Bias-Corrected Bootstrap CIs for the predictions at the selected 2-sets of x-locations. Show details of the bias-correction process. Compare the CIs at two x-locations, and also from two nonparametric regression methods.

Continuing our analysis of prediction at our two *X*-locations, we conduct a bootstrap sampling analysis. The procedure conducted consists of:

- 1. Taking a boostrap sample, or a uniform sample of observations (selected with replacement)
- 2. Fitting a spline or kernel regression model to the boostrap sample data, and
- 3. Predicting MPG at the given X-locations using the models fit in step 2

To create a more robust bootstrap estimate, we used bias-correction, or BCa, a technique similar to a percentile approach to boostrap sampling. However the bootstrap percentiles, but adjusted to account for bias and skewness. Underlying the the BCa approach is a notion of acceleration and skewness informing the parameter intervals, and also a connection to the normal distribution, as BCa relies on the CDF of a standard normal random variable to compute percentile estimates. The results of the bootstrap analysis are summarized in the table below:

Table 9: Bias-Corrected Bootstrap CIs - 90%, By Model Type and X-location

Point	Prediction	Lower	Upper	Method
Center Point	27.13	25.88	30.47	Kernel Regression
Outside Point	13.45	-0.22	16.08	Kernel Regression
Center Point	25.20	24.73	25.62	Spline
Outside Point	13.03	10.47	15.90	Spline

Note: Instability of the kernel regression fits from the np package, and instability of fits from smooth.spline in R (base) were both problematic due to sample size, and computation speed constraints further analysis would have likely yielded more stable results.

A key point to highlight from this table, is the instability of the kernel regression fits given our dataset.

Bootstrap replication actually created non-informative confidence intervals for our kernel regression model, given poor fits of in some boostrap replications. In contrast results from the spine model were more informative, and much narrower, and perhaps precise in comparison to kernel regression. The stability of the spline model in comparison does highlight its utility in this very specific case.

The bootstrap computational details were somewhat lengthy relative to other routines, and are provided in the *Code Appendix* at the end of this document.

Code Appendix

Multi-resolution Plot Generation (waveslim)

The code below demonstrates procedures to plot the additive components of a wavelet decomposition using the *waveslim* package in *R*. Note this code depends on usage of a time series exactly 1024 observations in length:

```
x <- as.numeric(aep$AEP_MW) #input megawatts time series
n.levels < -4
aep.haar <- mra(x = x, "haar", n.levels, "dwt")</pre>
nplots <- n.levels + 2</pre>
par(mfcol=c(nplots, 2), pty="m", mar=c(5-2.5, 4, 4-2, 1.5), cex.main = 0.9)
plot.ts(x, axes=F, main="Haar Filter: Multi-Resolution Plot", ylab = 'Original', bty='n')
axis(side=1, at=seq(0,1024,by=128),
     labels=seq(0,1024,by=128))
axis(side=2, at=c(round(min(x), digits = -4), round(max(x), digits = -4)),
     labels=c(round(min(x),digits = -4),round(max(x),digits = -4)))
for(i in 1:(n.levels +1)){
  plot.ts(aep.haar[[i]], axes=F, ylab=names(aep.haar)[i],bty='n')
  axis(side=1, at=seq(0,1024,by=128),
     labels=seq(0,1024,by=128))
  axis(side=2, at=c(round(min(aep.haar[[i]]),digits = -2),round(max(aep.haar[[i]]),digits = -2)),
       labels=c(round(min(aep.haar[[i]]),digits = -2),round(max(aep.haar[[i]]),digits = -2)))
}
#family daubechies
aep.la8 \leftarrow mra(x = x, "la8", n.levels, "dwt")
plot.ts(x, axes=F, main="Daubechies Filter (L=8): Multi-Resolution Plot", ylab = 'Original', bty='n')
axis(side=1, at=seq(0,1024,by=128),
     labels=seq(0,1024,by=128))
axis(side=2, at=c(round(min(x),digits = -4),round(max(x),digits = -4)),
     labels=c(round(min(x),digits = -4),round(max(x),digits = -4)))
for(i in 1:(n.levels +1)){
  plot.ts(aep.la8[[i]], axes=F, ylab=names(aep.la8)[i],bty='n')
  axis(side=1, at=seq(0,1024,by=128),
       labels=seq(0,1024,by=128))
  axis(side=2, at=c(round(min(aep.la8[[i]]),digits = -2),round(max(aep.la8[[i]]),digits = -2)),
       labels=c(round(min(aep.la8[[i]]),digits = -2),round(max(aep.la8[[i]]),digits = -2)))
```

Leave-One-Out Procedure, Kernel and Spline Regression

Below is exemplary code utilized in this report to generate additive model fits for both spline, and kernel regression models, along with the leave-one-out procedure. To speed up computation, which can be slow for kernel regression, the routine is distributed across CPUs utilizing the *parallel* library in *R*:

```
#leave one out procedure - spline and kernel regression
# CPU distributed (10 cores here)
spl_res <- rbindlist(mclapply(1:nrow(adat), function(i){
    train <- adat[-i]
    test <- adat[i]
    #additive spline fits
spl_fit_x1 <- smooth.spline(train$disp,train$mpg, cv= F,df = 3)</pre>
```

```
spl_fit_x2 <- smooth.spline(train$cyl,residuals(spl_fit_x1), cv= F,df = 3)</pre>
  spl_fit_x3 <- smooth.spline(train$accel,residuals(spl_fit_x2), cv= F,df = 3)</pre>
  test <- transform(test, prediction = predict(spl_fit_x1, data.frame(test$disp))$y +</pre>
                       predict(spl_fit_x2, data.frame(test$cyl))$y + predict(spl_fit_x3, data.frame(test$
  test$error <- test$mpg - test$prediction #actual vs. estimate</pre>
  test
}, mc.cores=10))
#this takes 1-2 minutes with 10 cores
kr_res <- rbindlist(mclapply(1:nrow(adat), function(i){</pre>
  train <- adat[-i]</pre>
  test <- adat[i]
  #additive kernel reg fits - simplified predict if data.frame is called
  kr_fit_x1 <- npreg(mpg~disp,gradients = TRUE, data = train)</pre>
  train$e1 <- residuals(kr_fit_x1)</pre>
  kr_fit_x2 <- npreg(e1~cyl,gradients = TRUE, data = train)</pre>
  train$e2 <- residuals(kr_fit_x2)</pre>
  kr_fit_x3 <- npreg(e2~accel,gradients = TRUE, data = train)</pre>
  test$prediction <- predict(kr_fit_x1, newdata = data.frame(disp = test$disp)) +</pre>
                       predict(kr_fit_x2, newdata = data.frame(cyl = test$cyl)) +
                       predict(kr_fit_x3, newdata = data.frame(accel = test$accel))
  test$error <- test$mpg - test$prediction #actual vs. estimate</pre>
}, mc.cores=10))
```

Bootstrap Confidence Intervals

Below is exemplary code utilized in this report to generate boostrap confidence intervals using the boot package, relying on the *boot* and *np* packages. Bootstrap sampling procedures were CPU distributed where possible to speed up computation:

```
B <- 1000 #bootstrap replication count
#estimator function
adat <- readRDS('data/auto_mpg.rds')</pre>
cpt <- data.table(disp = median(adat$disp),cyl = median(adat$cyl),</pre>
                   accel = median(adat$accel), type = 'center point')
opt <- data.table(disp = max(adat$disp) - 1,cyl = max(adat$cyl)-1,</pre>
                   accel = max(adat$accel) - 1, type = 'outside point')
spl_boot_cpt <- function(data, indices){</pre>
  #indices <- sample(1:nrow(adat), replace = T)</pre>
  bs_dat <- data[indices]</pre>
  sfit_x1 <- smooth.spline(bs_dat$disp,bs_dat$mpg, cv= F,df = 3)</pre>
  sfit_x2 <- smooth.spline(bs_dat$cyl,residuals(sfit_x1), cv= F,df = 3)</pre>
  sfit_x3 <- smooth.spline(bs_dat$accel,residuals(sfit_x2), cv= F,df = 3)</pre>
  prediction <- as.numeric(predict(sfit_x1, data.frame(cpt$disp))$y +</pre>
                               predict(sfit_x2, data.frame(cpt$cyl))$y +
                               predict(sfit_x3, data.frame(cpt$accel))$y)
  prediction
}
spl_boot_opt <- function(data, indices){</pre>
  #indices <- sample(1:nrow(adat),replace = T)</pre>
 bs_dat <- data[indices]</pre>
```

```
sfit_x1 <- smooth.spline(bs_dat$disp,bs_dat$mpg, cv= F,df = 3)</pre>
  sfit_x2 <- smooth.spline(bs_dat$cyl,residuals(sfit_x1), cv= F,df = 3)</pre>
  sfit_x3 <- smooth.spline(bs_dat$accel,residuals(sfit_x2), cv= F,df = 3)</pre>
  prediction <- as.numeric(predict(sfit_x1, data.frame(opt$disp))$y +</pre>
                               predict(sfit_x2, data.frame(opt$cyl))$y +
                               predict(sfit_x3, data.frame(opt$accel))$y)
 prediction
}
spl_cpt_res <- boot(adat, statistic = spl_boot_cpt,</pre>
                     R=B)
spl_opt_res <- boot(adat, statistic = spl_boot_opt,</pre>
                     R=B)
#bootstrap results - spline
cpt_results <- boot.ci(boot.out = spl_cpt_res,type = 'bca',conf = 0.9)</pre>
opt_results <- boot.ci(boot.out = spl_opt_res,type = 'bca',conf = 0.9)
boot_spl_bca <- data.table(</pre>
  Point = c('Center Point','Outside Point'),
  Prediction = c(cpt_results[[2]],opt_results[[2]]),
  Lower = c(cpt_results$bca[4],opt_results$bca[4]),
  Upper = c(cpt_results\bca[5],opt_results\bca[5]))
kr_boot_opt <- function(data, indices){</pre>
  #indices <- sample(1:nrow(adat), replace = T)</pre>
  kdat <- data[indices]</pre>
  kfit_x1 <- npreg(mpg~disp,gradients = TRUE, data = kdat)</pre>
  kdat$e1 <- residuals(kfit_x1)</pre>
  kfit_x2 <- npreg(e1~cyl,gradients = TRUE, data = kdat)</pre>
  kdat$e2 <- residuals(kfit_x2)</pre>
  kfit_x3 <- npreg(e2~accel,gradients = TRUE, data = kdat)</pre>
  prediction <- predict(kfit_x1, newdata = data.frame(disp = opt$disp)) +</pre>
    predict(kfit_x2, newdata = data.frame(cyl = opt$cyl)) +
    predict(kfit_x3, newdata = data.frame(accel = opt$accel))
  prediction
kr_boot_cpt <- function(data, indices){</pre>
  #indices <- sample(1:nrow(adat), replace = T)</pre>
  kdat <- data[indices]</pre>
  kfit_x1 <- npreg(mpg~disp,gradients = TRUE, data = kdat)</pre>
  kdat$e1 <- residuals(kfit_x1)</pre>
  kfit_x2 <- npreg(e1~cyl,gradients = TRUE, data = kdat)</pre>
  kdat$e2 <- residuals(kfit_x2)</pre>
  kfit_x3 <- npreg(e2~accel,gradients = TRUE, data = kdat)</pre>
  prediction <- predict(kfit_x1, newdata = data.frame(disp = cpt$disp)) +</pre>
    predict(kfit_x2, newdata = data.frame(cyl = cpt$cyl)) +
    predict(kfit_x3, newdata = data.frame(accel = cpt$accel))
  prediction
```

```
kr_cpt_res <- boot(adat, statistic = kr_boot_cpt,</pre>
                    R=B,parallel = 'multicore',
                    ncpus = 11)
kr_opt_res <- boot(adat, statistic = kr_boot_opt,</pre>
                    R=B,parallel = 'multicore',
                    ncpus = 11)
#bootstrap results - kernel regression
cpt_kresults <- boot.ci(boot.out = kr_cpt_res, type = 'bca', conf = 0.9)</pre>
opt_kresults <- boot.ci(boot.out = kr_opt_res,type = 'bca',conf = 0.9)</pre>
#results summary
boot_kr_bca <- data.table(</pre>
  Point = c('Center Point','Outside Point'),
  Prediction = c(cpt_kresults[[2]],opt_kresults[[2]]),
  Lower = c(cpt_kresults$bca[4],opt_kresults$bca[4]),
  Upper = c(cpt_kresults$bca[5],opt_kresults$bca[5]))
boot_kr_bca$Method <- 'Kernel Regression'</pre>
boot_spl_bca$Method <- 'Spline'</pre>
boot_sum <- rbindlist(list(boot_kr_bca,boot_spl_bca))</pre>
```