Computational & Mathematical Statistics - Fall 2023 Assignment 2

Information criteria; Forward-Backward; Penalized estimation

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Assignment 2 Part 1

 Perform residual diagnostics on the full second order model M3 from Assignment 1.

```
library(olsrr) # OLS diagnostics
library(L1pack) #LAD
library(caret)
data <- airquality
# Create the model with all the second and first-order terms
M3_{formula} \leftarrow Ozone \sim Solar.R + Wind + Temp + I(Solar.R^2) +
                        I(Wind^2) + I(Temp^2) + Solar.R:Wind+
                        Solar.R:Temp + Wind:Temp
set .seed(4)
# Remove rows with NA values
data <- na.omit(data)</pre>
# Shuffle the cleaned data
data_ <- data_[sample(nrow(data_)), ]</pre>
```

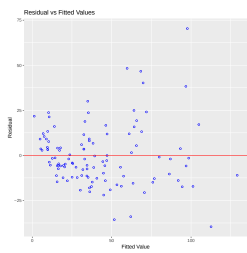




M3.ols <- lm(M3_formula, data = data_)
M3.lad <- lad(M3_formula, data = data_)

Residuals vs Fitted Plot

ols plot resid fit(M3.ols)

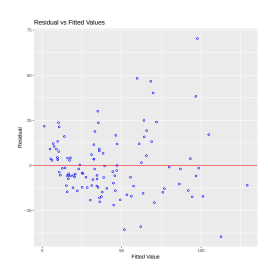


Model Diagnostics

- This plot helps assess whether the residuals have non-constant variance (heteroscedasticity).
- Scatter plot randomness around the zero line indicates the model's assumptions are likely correct, suggesting no non-linearity or misspecification.

AIC/BIC; Forward-Backward; Penalized est.

Residuals vs Fitted Plot



- In the provided plot, we would look for the absence of patterns and a roughly equal spread of residuals across the entire range of fitted values to satisfy the assumptions for the OLS.
- However, the plot seems to indicate a potential issue with heteroscedasticity, as indicated by a possible increasing spread in residuals as the fitted values increase.
- This would need to be further investigated.

Breusch Pagan Test for Heteroskedasticity

Data

Response: Ozone
Variables: fitted values of Ozone

Test Summary

DF = 1 Chi2 = 30.75916 Prob > Chi2 = 2.92122e-08

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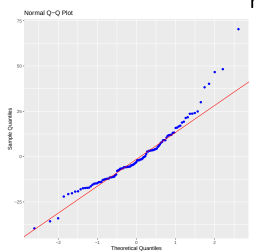
Breusch Pagan Test^{ab}

- One of the key assumptions of linear regression is that the residuals are distributed with equal variance at each level of the predictor variable. AKA homoscedasticity.
- When this assumption is violated, we say that heteroscedasticity is present in the residuals.

^aH₀: Homoscedasticity is present (the residuals are distributed with equal variance)

^bH_A: Heteroscedasticity is present (the residence are not distributed with equal variance)

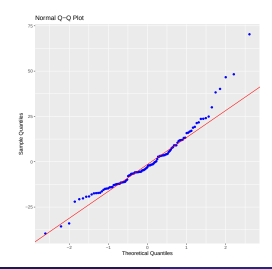
Normal Q-Q Plot



Normality Check

- This plot is used to determine if the residuals are approximately normally distributed.
- The residuals are plotted against a theoretical normal distribution in such a way that if the residuals are normally distributed, they will approximately lie on the diagonal line.

Normal Q-Q Plot

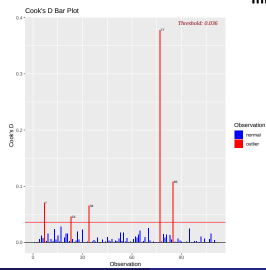


Normality Check

- The x-axis, labeled "Theoretical Quantiles", represents the quantiles from a standard normal distribution. These quantiles are the expected values if the residuals were normally distributed.
- The y-axis, labeled "Sample Quantiles", represents the quantiles of the residuals from our regression model. These are essentially the sorted values of the residuals, scaled to a normal distribution for comparison.

Cook's Distance Plot

ols plot cooksd bar(M3.ols)



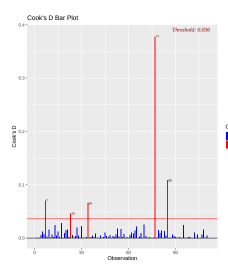
Influence Analysis

- The 'Cook's D Bar Plot' is utilized to detect influential observations in our regression model.
- Cook's D (Cook's Distance) measures the influence of individual data points on the regression coefficients.
- The threshold of 0.036 is used to identify points that have a potentially influential impact on the model.

outlier

Cook's Distance Plot

ols_plot_cooksd_bar(M3.ols)

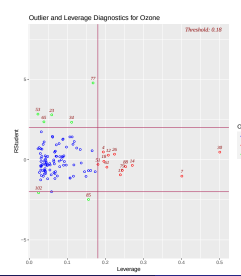


Influence Analysis

 Observation 77 is highly influential^aa, with a Cook's D value that greatly exceeds the threshold, suggesting a significant impact on the model.

^aThe term "highly influential" in the Observation Context of Cook's D refers to the impact that the removal of that observation would have on the fitted regression model. If an observation has a high Cook's D value (beyond the threshold), it means that the estimated regression coefficients change significantly when that observation excluded from the dataset.

Statistical Learning

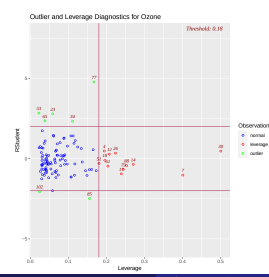


Outliers and Leverage^a

- The 'Residuals vs Leverage' plot is a diagnostic tool used to identify influential observations.
- o In general, the plot helps us understand which data points have the most influence on the calculation of the regression coefficients.

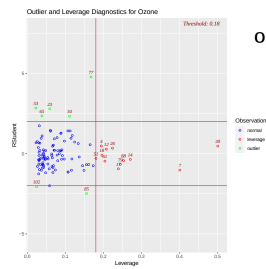
^aleverage is a measure of how far away the independent variable values of an observation are from those of the other observations. (Source)

ols_plot_resid_lev(M3.ols)



Outliers and Leverage

- The vertical red line represents a threshold for leverage, with points to the right deemed to have high leverage.
- The horizontal lines represent thresholds for standardized residuals, with points outside these lines considered to be outliers.
- The Rstudent axis represents studentized residuals, which are the residuals divided by their estimated standard deviation.



Outliers and Leverage

- The plot displays a few points that stand out:
 - Several points shown in blue are within the acceptable range for both leverage and studentized residuals, indicating they are neither high leverage points nor outliers.



Outlier and Leverage Diagnostics for Ozone Threshold: 0.18 Observation -5-0.1 0.2 0.4

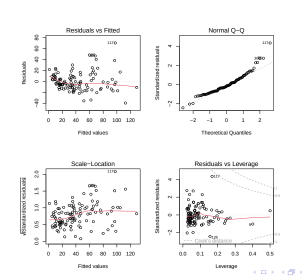
Leverage

Outliers and Leverage

- Point 7 is beyond the threshold for high leverage but has a low stud. residual, indicating that while it is a leverage point, it does not greatly influence the model's predictions.
- Point 53 is an outlier in terms of the Ozone, having a high stud. residual but is not a point of high leverage, suggesting it may not be influential in terms of the leverage but an outlier concernithe model's

Assignment 2 Part 1 - Diagnostics with plot() (OLS)

par(mfrow=c(2,2))
plot(M3.ols) # Alternative of olsrr

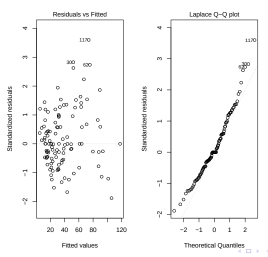






Assignment 2 Part 1 - Diagnostics with plot() (LAD)

par(mfrow=c(1,2))
plot(M3.lad)







Serial correlation

♦ Do not forget to evaluate whether your residuals are serially correlated ¹

independent across observations, but instead one residual is correlated with previous residuals.

Serial correlation

- Do not forget to evaluate whether your residuals are serially correlated.¹
- How can we address autocorrelation? One effective approach is to employ the **Durbin–Watson statistic**.

Durbin-Watson statistic

In statistics, the Durbin–Watson statistic is a test statistic used to detect the presence of autocorrelation. if e_t is the residual given by $e_t = \rho e_{t-1} + v_t$, the DW test statistic is

$$d = \frac{\sum_{t=2}^{T} (e_t - e_{t-1})^2}{\sum_{t=1}^{T} e_t^2} ,$$

where T is the number of observations. For large T, $d\approx 2(1-\hat{\rho})$, where $\hat{\rho}$ is the sample autocorrelation of the residuals. (Source)

independent across observations, but instead one residual is correlated with previous residuals and

Serial correlation (autocorrelation)

- A value of approximately 2 indicating no autocorrelation².
- A value of less than 2 suggesting positive autocorrelation.
- A value of more than 2 suggesting negative autocorrelation.
- 1.8 is close to 2. To obtain a more accurate estimation for the statistic, we can utilize **bootstrapping**.

relatively normal. (Source)

```
set.seed(42)
B <- 10000
bagging_preds <- matrix(NA, nrow=nrow(data_), ncol=B)</pre>
boot.results <- data.frame(dw_statistic=numeric(B),</pre>
                             p_value=numeric(B))
# Bagging loop
for (b in 1:B) {
# Draw a bootstrap sample
    bootstrap_indices <- sample(1:nrow(data_),</pre>
                                           size=nrow(data).
                                           replace=TRUE)
    bootstrap data <- data [bootstrap indices, ]
    bootstrap_model <- lm(M3_formula, data = bootstrap data)</pre>
    #Perform Durbin-Watson test for each bootstrap sample
    dw test result <- dwtest(bootstrap model)</pre>
    boot.results$dw_statistic[b] <- dw_test_result$statistic
    boot.results$p_value[b] <- dw_test_result$p.value
```

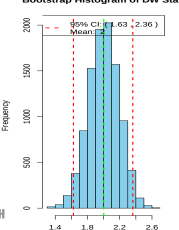


```
# Create histogram with 95% CI and mean value lines
boot.histogram <- function (data_vector, main_title, xlab_title) {</pre>
  # Histogram of the data
  hist(data vector, main = main title,
       xlab = xlab_title, col = "skyblue", border = "black")
  # Calculate the 95% confidence interval
  ci <- quantile(data_vector, probs = c(0.025, 0.975))</pre>
  mean_val <- mean(data_vector)</pre>
  # Add dotted vertical lines for the 95% confidence interval
  abline(v = ci[1], col = "red", lwd = 2, lty = 2) # lower quantile
  abline(v = ci[2], col = "red", lwd = 2, lty = 2) # upper quantile
  # Add a dashed green line for the mean
  abline(v = mean_val, col = "green", lwd = 2, lty = 2)
  # Add a legend with the confidence interval values and mean value
  legend("topright",
         legend = paste("95% CI: (", round(ci[1], 2), ',',
                    round(ci[2], 2), ')', "\nMean: ",
                        round(mean_val, 2)),
         lty = 2, lwd = 2, col = c("red", "green"))
}
```



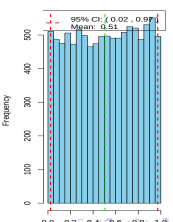
```
par(mfrow=c(1,2))
boot.histogram(boot.results$dw_statistic,
        'Bootstrap Histogram of DW Stat.', 'Durbin-Watson Statistic')
boot.histogram(boot.results$p_value, 'Histogram of DW Stat.',
               'Durbin-Watson Statistic')
```





1.8

Histogram of p-values.





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Observations:

- The 95% CI lies between $\underbrace{(1.5,2.5)}_{\supset (1.63,2.36)}$ (with a mean value of ≈ 2 .)
- Durbin—Watson Statistic approaches a normal distribution.
- Durbin—Watson p-values are uniformly distributed³.
- Hence, from our bootstrap analysis, we have substantial evidence to suggest that serial correlation does not present a concern in our dataset.

Methodological Alert

The above procedure is insignificant and should be avoided. We cannot use bootstrap to determine the presence of autocorrelation.



Observations:

- The 95% CI lies between $\underbrace{(1.5,2.5)}_{\supset (1.63,2.36)}$ (with a mean value of ≈ 2 .)
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How far is your analysis from the hypotheses of the Gauss-Markov Theorem?



$$\Pr(P < p) = \Pr(F(T) < p) = \Pr(T < F^{-1}(p)) = F(F^{-1}(p)) = p.$$
 (Source)

Gauss-Markov Theorem

Theorem

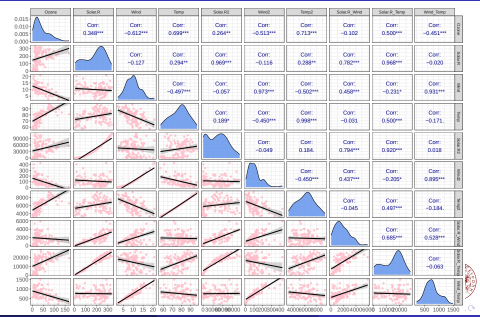
The Gauss-Markov Theorem is a fundamental theorem in linear regression that provides the conditions under which the OLS estimator is the Best Linear Unbiased Estimator (BLUE). In other words, under certain conditions, OLS provides the most precise (lowest variance) estimates of the regression coefficients without bias.

- For the OLS estimates to be BLUE, the following assumptions must be satisfied:
 - **1 Linearity**: The relationship between the dependent variable and the independent variables is linear in the parameters.
 - **2** Random Sampling: The data is a random sample from the population.
 - No Perfect Multicollinearity: The independent variables are not perfectly correlated with each other.
 - 4 Homoscedasticity: The variance of the residuals (errors) is constant across all levels of the independent variables.
 - No Autocorrelation: The residuals (errors) are not correlated with each other (no serial correlation).
 Zero Conditional Mean: the errors are uncorrelated with the features

Linearity - Pairplot via GGally, ggplot2

```
library(ggplot2)
library(GGally)
# Add the second order terms to data (frame)
data $Solar.R2 <- data $Solar.R^2
data $Wind2 <- data $Wind^2
data $Temp2 <- data $Temp^2
data_$Solar.R_Wind <- data_$Solar.R * data $Wind
data_$Solar.R_Temp <- data_$Solar.R * data_$Temp</pre>
data $Wind Temp <- data $Wind * data $Temp
ggpairs(data [c(-5,-6)],
  upper = list(continuous = wrap("cor", size=3.2, color="darkblue")),
  lower = list(continuous = wrap("smooth", colour="pink",
                                   fill="lightblue", alpha=0.9)),
  diag = list(continuous = wrap("densityDiag", fill="cornflowerblue",
                                  alpha=0.85)).
        axisLabels = "show") +
  theme bw() +
  theme(legend.position = "bottom", axis.text=element_text(size=12)
        plot.title = element_text(size=10, hjust=0.5))
 FORTH
```

Linearity - Pairplot via GGally, ggplot2



Gauss-Markov assumptions

- Linearity: We can observe some linearity from the pairplot.
- Random Sampling: We observed that in the scatter plots of the residuals versus predicted values, the residuals appear to be randomly scattered, suggesting random sampling.
- No Perfect Multicollinearity: Some of the features are highly (Pearson) correlated with each other, as we can observe from the pairplot. For instance, Solar.R:Temp with I(Solar.R)² and so forth. This might raise concerns regarding multicollinearity issues.
- Homoscedasticity: The Breusch-Pagan test yields a p-value that is very close to zero. Consequently, we can reject the null hypothesis, indicating the presence of heteroscedasticity.
- No Autocorrelation: We have substantial evidence to suggest that the residuals are not serially correlated. (DW statistic≈ 2)
- **Output Zero Cond. Mean** ($\mathbb{E}(\mu \mid \mathbf{X}) = 0$): We don't observe any patterns the Residuals-Fitted values plot, thus indicating that this assumption is satisfied.

Box-Cox transformation

♦ Evaluate the suggested Box-Cox transformation of the response





Box-Cox transformation

 Evaluate the suggested Box-Cox transformation of the response variable.

Definition

The **Box-Cox transformation** is defined as follows for a response variable Y and a transformation parameter λ :

$$Y(\lambda) = \begin{cases} \frac{Y^{\lambda} - 1}{\lambda} & \text{if } \lambda \neq 0, \\ \log(Y) & \text{if } \lambda = 0. \end{cases}$$

- Y is the response variable that we want to transform.
- λ is the transformation parameter. The value of λ is chosen to achieve the best approximation to normality for the transformed variable $Y(\lambda)$.
- When $\lambda = 1$, the transformation is the identity transformation (no change).
- When $\lambda = 0$, it becomes a logarithmic transformation.
- \bullet The choice of λ can significantly impact the shape of the distribution of the transformed variable.

Purpose of Box-Cox transformation

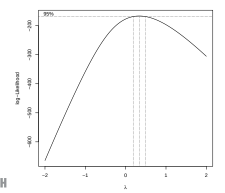
- Improving Normality: It's often used when residuals from a linear model do not appear to be normally distributed. Transforming the response variable can lead to residuals that are more normally distributed.
- Stabilizing Variance: It can be used when the variance of residuals is not constant across levels of an independent variable (heteroscedasticity). A Box-Cox transformation can help in stabilizing the variances.
- Enhancing Model Fit: By transforming the response variable, it can improve the linearity of the relationship between independent and dependent variables in a regression model.

The optimal value of λ is usually found empirically. Choose the one that maximizes the log-likelihood.⁴

The Box-Cox transformation is a powerful tool, particularly useful when dealing with non-normal data distributions and aiming to meet the assumptions of linear regressions a constant of the second s

Box-Cox transformation in R

```
library(MASS)
set.seed(4)
data_ <- na.omit(data) #redef the data
data_ <- data_[sample(nrow(data_)), ]
boxcox.result <- boxcox(M3.ols)
boxcox(M3.ols)</pre>
```



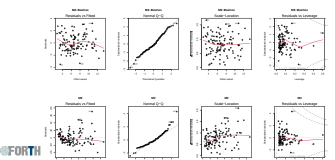




Box-Cox transformation in R

```
lambda.best <- boxcox.result$x[which.max(boxcox.result$y)]
y_boxcox <- ((data_$0zone^lambda.best) - 1)/lambda.best

M3_formula_ <- y_boxcox ~ Solar.R + Wind + Temp + I(Solar.R^2) +
    I(Wind^2) + I(Temp^2) + Solar.R:Wind +
    Solar.R:Temp + Wind:Temp
M3.ols_boxcox <- lm(M3_formula_, data = data_)
par(mfrow=c(2,4))
plot(M3.ols_boxcox, main="M3-BoxCox")
plot(M3.ols, main="M3")</pre>
```





Additional predictors

♦ Create 3 additional predictors Z1, Z2 and Z3. Z1 should be correlated





Additional predictors

 Create 3 additional predictors Z1, Z2 and Z3. Z1 should be correlated with Solar.R with rho approximately equal to 0.8, Z2 with Wind and Z3 with Temp (same rho for all noise predictors). Include the 3 noise predictors in your model (model M4).

Adjusted Approach for Creating Noisy Predictors:

- To achieve a specific correlation, we can use a linear combination of the original variable and independent random noise.
- ② The formula for creating a new variable Z with a desired correlation ρ with an original variable X is

$$Z =
ho imes X + \sqrt{1-
ho^2} imes exttt{Noise},$$

where Noise is randomly generated and independent of X.

Implementing the Revised Approach:

- Scale Solar.R, Wind, and Temp.
- Create Noise as a random normal variable.
- Construct Z1, Z2, and Z3 using the formula above.



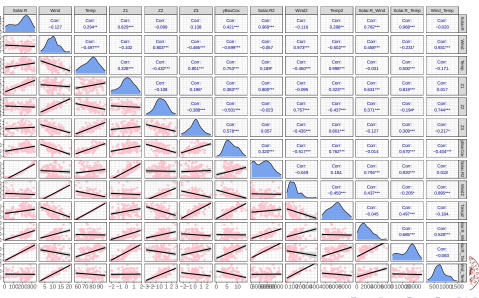
Create Z1, Z2, Z3 predictors

```
set.seed(3)
# Scaling func.
scale var <- function (x) {
  return((x - mean(x)) / sd(x))
}
# Function to create noisy predictor with desired correlation
create_noisy_predictor <- function (variable, rho) {</pre>
  noise <- rnorm(length(variable))</pre>
  return(rho * variable + sqrt(1 - rho^2) * noise)
data_$Z1 <- create_noisy_predictor(scale_var(data_$Solar.R), 0.8)
data_$Z2 <- create_noisy_predictor(scale_var(data_$Wind), 0.8)</pre>
data_$Z3 <- create_noisy_predictor(scale_var(data_$Temp), 0.8)</pre>
# Checking the correlations
cor(data_$Z1, data_$Solar.R) # Should be approximately 0.8
cor(data_$Z2, data_$Wind) # Should be approximately 0.8
cor(data_$Z3, data_$Temp) # Should be approximately 0.8
>> 0.8293
>> 0.8026
>> 0.8014
```



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Pairplot with the additional predictors



Tofallis 2015 paper

◆ Transform the response in Model 4 so that your model complies with Relative Error Minimization as discussed in the attached paper by Tofallis (2015). Is the transformation similar or different to the one suggested by Box-Cox? Compare your new model (M5) against M4 and perform residual diagnostics on M5; is there an improvement relative to M3?





Tofallis 2015 paper

◆ Transform the response in Model 4 so that your model complies with Relative Error Minimization as discussed in the attached paper by Tofallis (2015). Is the transformation similar or different to the one suggested by Box-Cox? Compare your new model (M5) against M4 and perform residual diagnostics on M5; is there an improvement relative to M3?

 $M4.ols \leftarrow lm(M4_formula, data = data_)$





Tofallis 2015 paper

Transform the response in Model 4 so that your model complies with Relative Error Minimization as discussed in the attached paper by Tofallis (2015). Is the transformation similar or different to the one suggested by Box-Cox? Compare your new model (M5) against M4 and perform residual diagnostics on M5; is there an improvement relative to M3?





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Toannis Maris

Breusch Pagan Test for Heteroskedasticity -----Ho: the variance is constant Ha: the variance is not constant Data

Data

Response : log_y

Variables: fitted values of log_y

Test Summary

DF = 1

Chi2 = 22.83046

Prob > Chi2 = 1.769392e-06

Durbin-Watson test

data: M5.ols

DW = 2.0301, p-value = 0.5549

alternative hypothesis: true autocorrelation is greater than ${\tt 0}$





```
summary(M5.ols)
>>
Coefficients:
              Estimate Std. Error t value Pr(>|t|)
             6.161e+00 5.671e+00 1.086
                                         0.2800
(Intercept)
Solar.R
           8.025e-03 6.183e-03 1.298 0.1974
Wind
          -4.039e-01 2.790e-01
                                 -1.448 0.1509
Temp
        -7.193e-02 1.195e-01
                                 -0.602 0.5487
I(Solar.R^2) -1.741e-05 7.196e-06
                                 -2.419 0.0174 *
I(Wind^2) 9.089e-03 4.219e-03 2.154
                                        0.0337 *
I(Temp^2) 5.612e-04 6.662e-04 0.842
                                         0.4016
Z1
            -1.469e-03 9.672e-02
                                 -0.015
                                         0.9879
Z2
             7.105e-02 7.305e-02 0.973
                                         0.3331
Z3
             2.386e-02 8.190e-02 0.291 0.7714
Solar.R:Wind -1.388e-04 1.802e-04
                                 -0.770 0.4429
Solar.R:Temp 2.621e-05 6.917e-05 0.379
                                         0.7056
Wind: Temp
             2.016e-03
                       2.723e-03 0.741
                                         0.4607
Residual standard error: 0.495 on 98 degrees of freedom
Multiple R-squared: 0.7089, Adjusted R-squared: 0.6732
F-statistic: 19.88 on 12 and 98 DF, p-value: < 2.2e-16
```



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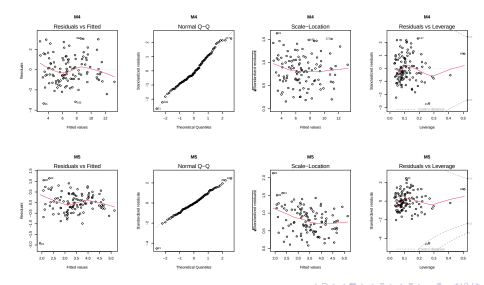
```
ols test breusch pagan(M4.ols)$p
ols_test_breusch_pagan(M5.ols)$p
>>
0.650352818060884 #M4 deals with heteroscedasticity
1.76939234875026e-06
```

We can not compare models with different response via AICc/BIC/R²

```
library(AICcmodavg)
AICc(M4.ols)
AICc(M5.ols)
BIC(M4.ols)
BIC(M5.ols)
summary(M4.ols)$r.squared
summary(M5.ols)$r.squared
>>
417.3133
177.4368
450.8717
210.9953
0.7528444
```

R-squared is the amount of variance in the response variable that is explained by the model. However, the R^2 of the first model describes the variance explained of y, which is something different from that in the transformed model $(\log(y))$. (Source)

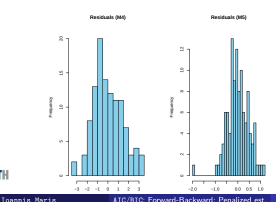
0.7088503



Box-Cox vs Tofalis log transformation:

- M4 deals with heteroscedasticity.
- M5 seems to have issues with heteroscedasticity.

```
par(mfrow = c(1, 2))
hist(residuals(M4.ols), main = "Residuals (M4)",
     xlab="Residuals", col="skyblue", breaks=35)
hist(residuals(M5.ols), main="Residuals (M5)",
    xlab="Residuals", col="skyblue", breaks=35)
```







Set up the metrics for cross-validation

```
LAR <- function (v. forcast) { #Log Acc. Ration
 n <- length(y)
  lar <- 0
 n <- 0 #counter
 for (i in 1:n) {
    if (v[i] != 0) {
      Q = as.numeric(forcast)[i]/v[i]
        if (0 > 0) {
          lar = lar + abs(log(Q))
          n_{-} = n_{-} + 1
        111
 lar <- lar/n_
  return(lar)
MAPE <- function (v, forcast) {
 n <- length(y)
 mape sum <- 0
 n <- 0 #counter
  for (i in 1:n) {
    if (v[i] != 0) {
      mape_sum = mape_sum + abs((y[i]-as.numeric(forcast)[i])/y[i])
      n_{-} = n_{-} + 1
    }
 mape \leftarrow (mape_sum/n_) * 100
 return (mape)
MAE <- function (actual, preds) mean(abs(actual-preds))
RMSE <- function (actual, preds) sqrt(mean((actual-preds)^2))
sMdAPE <- function (actual, preds) median(
                                      200*abs(actual-preds)/(actual+preds),
 FORTH
                                      na.rm = TRUE)
```



Cross-validation (with transformation of the response)

```
library(foreach)
KfoldCVPerf <- function (K = 10, data, model_type = 'lm',</pre>
                         formula, name = 'model name',
                    folds.seed = 42. inv.trans=function(x) return(x)) {
    ### Datatypes ###
    ## K: int #data: data.frame #model_type: 'lm' || 'lad'
    ## formula: as.formula() ## folds.seed: int: random folds
    #### inv.trans: inverse transformation of the response ####
    ### NOTE: The response y must be in the dataframe ###
    # Shuffle the data to get random folds:
    set.seed(folds.seed)
    data <- data[sample(nrow(data)), ]
    y <- all.vars(formula)[1] # get response name (str)
    rows per fold <- nrow(data)/K
    #Create a binning var
    binning variable <- cut(seq(1, nrow(data)), breaks = K, labels = FALSE)
    folds <- split(data, binning variable) # Split the folds
    folds list <- list() # Create a list of K folds
    for (i in 1:K) {
        folds list[[i]] = folds[[i]]
    perf_df <- data.frame(
        Fold = integer(0), RMSE = numeric(0), MAE = numeric(0),
        sMdAPE = numeric(0), LAR = numeric(0), MAPE = numeric(0))
    #Loop through the folds
    foreach (i = 1:K) %do% {
        val fold <- folds list[[i]]
        y_val <- val_fold[[y]]</pre>
        train_folds <- do.call(rbind, folds_list[-i])
        v train <- train folds[[v]]
 FORTH Change the format of the formula in order to convert response
        formula_ <- paste(all.vars(formula)[-1], collapse = ' + ')
```

Cross-validation (with transformation of the response)

```
if (model type == 'lm') {
         model <- lm(as.formula(paste("y_train~", formula_)),</pre>
                      data=train_folds)
     } else if (model type == 'lad') {
         model <- lad(as.formula(paste("v_train~", formula_)),</pre>
                       data=train_folds)
     } else {
         message("Unsupported model type.")
         break }
     preds <- predict(model, newdata = val_fold)
     y_val <- inv.trans(y_val)</pre>
     preds <- inv.trans(preds)</pre>
     rmse_val <- RMSE(y_val, preds)
     mae val <- MAE(v val, preds)
     smdape_val <- sMdAPE(y_val, preds)</pre>
     MAPE_val <- MAPE(v_val, preds)
     LAR val <- LAR(v val. preds)
     perf df <- rbind(perf df, data.frame(
                              Fold = i, RMSE = rmse_val,
                              MAE = mae val, sMdAPE = smdape val,
                              MAPE = MAPE val, LAR = LAR val))
 mean_rmse <- mean(perf_df$RMSE)
 mean mae <- mean(perf df$MAE)
 mean_smdape <- mean(perf_df$sMdAPE)</pre>
 mean_MAPE <- mean(perf_df$MAPE)</pre>
 mean LAR <- mean(perf df$LAR)
 # Add the means to the df
 perf df <- rbind(perf df, c("Mean", mean rmse, mean mae, mean smdape, mean MAPE, mean
 colnames(perf df)[-1] <- pasteO(name, ' (', colnames(perf df)[-1], ')')
return (perf df)
```

Cross-validation (with transformation of the response)

Example

Fold	M4 (RMSE)	M4 (MAE)	M4 (sMdAPE)	M4 (MAPE)	M4 (LAR)
1	13.4844639394214	11.722952456799	37.3501032481086	44.4310852974232	0.42197374564178
2	16.4394103204079	12.9306891690037	42.999233192946	49.2779504040391	0.404678422436324
3	35.1132950868163	19.9278180375665	27.6364194117723	39.7538616913861	0.441930070869637
4	23.4537311108301	15.1247076248887	29.354174271746	30.5885051836696	0.316859449685931
5	15.0432980699381	13.2118868133534	33.3835683169767	48.3219687016842	0.503947671845283
6	17.6450654467307	12.3756864535048	25.5236236495682	40.0221035385367	0.38466353961046
7	24.2386967241797	17.622639684578	41.8945739314198	53.6521920721395	0.536175225467513
8	14.1999279686414	10.8636915903395	40.3606155800208	50.2385711635589	0.419861171663608
9	11.2999901354722	10.0769371570312	23.3068862055817	51.1617299299498	0.367456908732039
10	21.2692243135157	14.9242266585308	48.6564201229144	130.38199296247	0.666525243883205
Mean	19.2187103115953	13.8781235645596	35.0465617931055	53.7829960944857	0.446407144983578

Repeated cross-validation

```
RepeatedCV <- function (repeats=500, K=10, data, formula,
                         model_name, inv.trans = function(y) y) {
    require(foreach)
    mean rows list <- list() # store the row means
    # Choose diff random.sds in order to get diff folds
    foreach (seed = 1:repeats) %do% {
        cv_results <- KfoldCVPerf(K = K, data=data, formula=formula,</pre>
                                    name=model name, folds.seed = seed.
                                    inv.trans = inv.trans)
        mean_row <- cv_results[cv_results[,1] == "Mean", -1]</pre>
        mean rows list[[seed]] <- as.numeric(mean row)</pre>
    }
    # Combine the mean rows into a df
    mean rows df <- do.call(rbind, mean rows list)
    # Average for each metric
    average_metrics <- colMeans(mean_rows_df, na.rm = TRUE)</pre>
    average_metrics_df <- as.data.frame(t(average_metrics))</pre>
    colnames(average_metrics_df) <- names(cv_results)[-1]</pre>
    rownames(average_metrics_df) <- "RCV Average"
    return(average metrics df)
```

Repeated cross-validation

Example

```
M3.RCV <- RepeatedCV(
    repeats = 500, K = 10, data = data_, formula = M3_formula, model_name = 'M3')

M4.RCV <- RepeatedCV(
    repeats = 500, K = 10, data = data_, formula = M4_formula, model_name = 'M4',
    inv.trans = function(y) (lambda.best * y + 1)^(1 / lambda.best))

M5.RCV <- RepeatedCV(
    repeats = 500, K = 10, data = data_, formula = M5_formula, model_name = 'M5',
    inv.trans = function(y) exp(y))

>>
```

M3.RCV	M3 (RMSE)	M3 (MAE)	M3 (sMdAPE)	M3 (MAPE)	M3 (LAR)
RCV Average	20.728841315252	16.1222941795992	33.8353230750363	67.7265253872971	0.438285472180775

M4.RCV	M4 (RMSE)	M4 (MAE)	M4 (sMdAPE)	M4 (MAPE)	M4 (LAR)
RCV Average	18.5803706748303	13.8479408576176	34.8100960781422	53.5017843693086	0.450559758646985

M5.RCV	M5 (RMSE)	M5 (MAE)	M5 (sMdAPE)	M5 (MAPE)	M5 (LAR)
RCV Average	18.731246022978	13.5511881738598	33.5722525817141	49.2004350320875	0.41236694504943

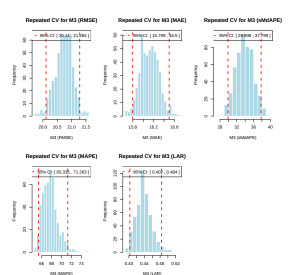
Repeated cross-validation: 95% CI

```
RCV.histograms <- function (repeats=100, K=10, data=data_,
                              formula, name, inv.trans=function(v) v) {
    require(foreach)
    metrics values list <- list()
    foreach (seed = 1:repeats) %do% {
        M5.cv <- KfoldCVPerf(K = K, data = data, formula = formula, name = name,
                             folds.seed = seed, inv.trans = inv.trans)
        # Extract the "Mean" row and store the metrics values
        mean_row <- M5.cv[M5.cv[,1] == "Mean", -1]
        for (metric in names(mean_row)) {
            metrics values list[[metric]] <- c(metrics values list[[metric]].
                                                as.numeric(mean row[metric]))
    # plot the histograms for each metric and add the CI lines
    par(mfrow=c(2, 3)) # Set up the plotting area to display multiple plots
    for (metric in names(metrics_values_list)) {
        # Calculate the 95% CI for the metric
        metric_values <- unlist(metrics_values_list[metric])</pre>
        ci <- quantile(metric_values, probs=c(0.025, 0.975))
        hist(metric values, main=paste("Repeated CV for", metric).
             xlab=metric. col='lightblue'. border='white'. break s=15)
        abline(v=ci[1], col="red", lwd=2, lty=2) # left CI line
        abline(v=ci[2], col="red", lwd=2, ltv=2) # right CI line
        legend("topright", legend=paste("95% CI: [", round(ci[1], 3),
                                     ',', round(ci[2], 3), "]"),
               col="red", lwd=2, lty=2)
    par(mfrow=c(1, 1))
```



Repeated cross-validation - M3

M3.RCVhists <- RCV.histograms(repeats=500, K=10, data=data_, formula=M3_formula, name='M3')

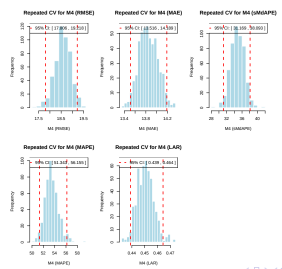






Repeated cross-validation - M4

M4.RCVhists <- RCV.histograms(repeats=500, K=10, data=data_, formula=M4_formula, name='M4', inv.trans=function(y) (lambda.best*y + 1)^(1/lambda.best))

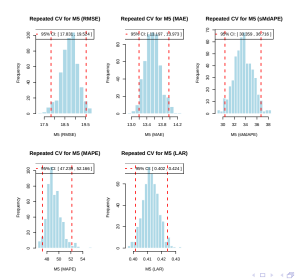






Repeated cross-validation - M5

M5.RCVhists <- RCV.histograms(repeats=500, K=10, data=data_, formula=M5_formula, name='M5', inv.trans=function(y) exp(y))



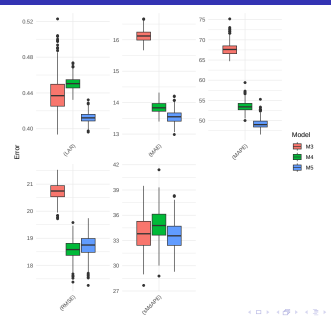




Repeated cross-validation - Box-Plots

```
RCV.boxplots combined <- function (repeats=500, K=10, data, M3 formula,
                                  M4_formula, M5_formula, inv.trans_M3,
                                  inv.trans_M4, inv.trans_M5) {
    require(ggplot2)
    combined_metrics <- data.frame(metric = character(),</pre>
                                   value = numeric(), model = factor(),
                                   stringsAsFactors = FALSE)
    # List of models and their respective formulas and inverse transformations
    models_list <- list(
        M3 = list(formula = M3 formula, trans = inv.trans M3),
        M4 = list(formula = M4 formula, trans = inv.trans M4),
        M5 = list(formula = M5_formula, trans = inv.trans_M5)
    for (model name in names(models list)) {
        foreach (seed = 1:repeats) %do% {
            cv_results <- KfoldCVPerf(K, data, 'lm',
                                      models list[[model name]] formula.
                                      model name, seed, models list[[model name]]$trans)
            mean_row <- cv_results[cv_results[, 1] == "Mean", -1]</pre>
            # Append results to the combined metrics data frame
            combined metrics <- rbind(combined metrics, data.frame(
                metric = names(mean_row), value = as.numeric(mean_row), model = model_name))
    # Extract only the metric name (after the space)
    combined_metrics$metric <- sapply(strsplit(combined_metrics$metric, " "), function(x) x[2])
    p <- ggplot(combined metrics, aes(x = metric, v = value, fill = model)) + geom boxplot() +
        facet wrap (~ metric, scales = 'free') + theme minimal() +
        theme(axis.text.x = element_text(angle = 45, hjust = 1),
              strip.text.x = element blank()) +
    ORTiabs(x = NULL, v = "Error", fill = "Model")
    print(p)
```

Repeated cross-validation - Box-Plots based on 500 repeats





Assignment 2 Part 2

Examine alternative tools for predictor selection on M5.

First use regsubsets to perform BIC-based forward and backward selection (F-BIC, B-BIC); which is the most parsimonious specification?





Statistical Learning

Assignment 2 Part 2

Examine alternative tools for predictor selection on M5.

First use regsubsets to perform BIC-based forward and backward selection (F-BIC, B-BIC); which is the most parsimonious specification?

```
M5.forward <- regsubsets(M5_formula, data=data_,
                                  method="forward". nvmax=12)
M5.backward <- regsubsets(M5_formula, data=data_,
                                  method="backward", nvmax=12)
M5.Fbic <- summary(M5.forward)$bic
M5.Bbic <- summary(M5.backward)$bic
M5.Fbic
M5.Bbic
>>
-81.616 -94.299 -105.724 -105.455 -103.401 -100.827 -97.223 -93.641 -89.421 -85.064
-80.45 -75.741
-81.616 \quad -94.137 \quad -103.77 \quad -105.762 \quad -106.207 \quad -102.545 \quad -98.415 \quad -93.857 \quad -89.612
-80.45 -75.741
FORTH
```

F-BIC

Examine alternative tools for predictor selection on M5.

summary(M5.forward)\$outmat

summary	Solar.R	Wind	Temp	I(Solar.R ²)	I(Wind ²)	I(Temp ²)	Z1	Z2	Z3	Solar.R:Wind	Solar.R:Temp	Wind:Temp
1(1)						*						
2 (1)						*					*	
3 (1)						*					*	*
4 (1)				*		*					*	*
5 (1)				*	*	*					*	*
6 (1)		*		*	*	*					*	*
7 (1)	*	*		*	*	*					*	*
8 (1)	*	*		*	*	*		*			*	*
9 (1)	*	*		*	*	*		*		*	*	*
10 (1)	*	*	*	*	*	*		*		*	*	*
11 (1)	*	*	*	*	*	*		*	*	*	*	
₽ ORTI	1.	*	*	*	*	*	*	*	*	*	*	
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B-BIC

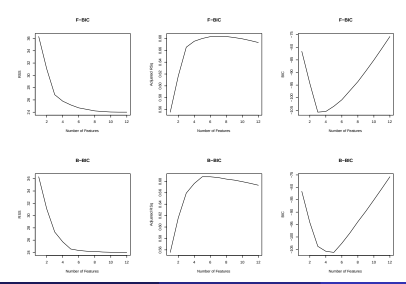
Examine alternative tools for predictor selection on M5.

summary(M5.backward)\$outmat

summary	Solar.R	Wind	Temp	I(Solar.R ²)	I(Wind ²)	I(Temp ²)	Z1	Z2	Z3	Solar.R:Wind	Solar.R:Temp	Wind:Temp
1(1)						*						
2 (1)	*					*						
3 (1)	*	*				*						
4 (1)	*	*			*	*						
5 (1)	*	*		*	*	*						
6 (1)	*	*		*	*	*		*				
7 (1)	*	*		*	*	*		*		*		
8 (1)	*	*		*	*	*		*		*		*
9 (1)	*	*	*	*	*	*		*		*		*
10 (1)	*	*	*	*	*	*		*		*	*	*
11 (1)	*	*	*	*	*	*		*	*	*	*	
FORT	H •	*	*	*	*	*	*	*	*	*	*	
									- ◆ □			₹ 1 00

F-BIC; B-BIC plots

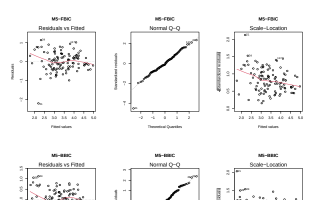
Examine alternative tools for predictor selection on M5.

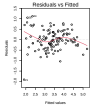


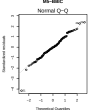


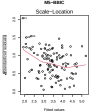
F-BIC; B-BIC residuals

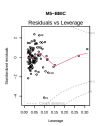
```
par(mfrow=c(2,4))
plot(M5.FBIC, main="M5-FBIC")
plot(M5.BBIC, main="M5-BBIC")
```











Leverage

0.02 0.04 0.06

M5-FBIC

Residuals vs Leverage



F-BIC: B-BIC est.

Examine alternative tools for predictor selection on M5.

```
coef(M5.forward, which.min(M5.Fbic))
coef(M5.backward, which.min(M5.Bbic))
>>
(Intercept): 2.279 Solar:R: 0.0075 Wind: -0.211 I(Solar.R^2): -1.49e-05 I(Wind^2):
     0.007 I(Temp^2) 0.00027
# Compare and choose the most parsimonious one
M5F lowest.bic <- M5.Fbic[which.min(M5.Fbic)]
M5B_lowest.bic <- M5.Bbic[which.min(M5.Bbic)]
M5F.len <- length(coef(M5.forward, which.min(M5.Fbic))[-1])
M5B.len <- length(coef(M5.backward, which.min(M5.Bbic))[-1])
# Parsimonious: if BICs are close choose the one with the < features
if (abs(M5F lowest.bic-M5B lowest.bic) < 3 && M5F.len < M5B.len) {
 cat("Most parsimonious model is from Forward Selection.".
      "BIC:", M5.Fbic[which.min(M5.Fbic)], ", features:", M5F.len)
} else {
 cat("Most parsimonious model is from Backward Selection.".
      "BIC:", M5.Bbic[which.min(M5.Fbic)], ", features:", M5B.len)
}
>>
Most parsimonious model is from Forward Selection, BIC: -105.7242 , features: 3
FORTH
```

F-BIC; B-BIC vs F-BIC; B-AIC

♦ Compare F-BIC, B-BIC to F-AIC (use ols_step_forward_aic from olsrr) and B-AIC (ols_step_backward_aic).





F-BIC; B-BIC vs F-BIC; B-AIC

♦ Compare F-BIC, B-BIC to F-AIC (use ols_step_forward_aic from olsrr) and B-AIC (ols_step_backward_aic).

```
M5.FAIC <- ols_step_forward_aic(M5.ols)
M5.BAIC <- ols_step_backward_aic(M5.ols)
M5.FAIC
M5.BAIC
>>
```





F-BIC; B-BIC vs F-BIC; B-AIC

>>

Selection Summary

Variable	AIC	Sum Sq	RSS	R-Sq	Adj. R-Sq
I(Temp^2)	196.992	46.152	36.317	0.55963	0.55559
Solar.R:Temp	181.599	51.420	31.050	0.62349	0.61652
Wind:Temp	167.464	55.620	26.850	0.67443	0.66530
I(Solar.R^2)	165.024	56.673	25.797	0.68720	0.67539
I(Wind^2)	164.369	57.283	25.187	0.69459	0.68005
Wind	164.233	57.763	24.707	0.70041	0.68313

>>

Backward Elimination Summary

Variable	AIC	RSS	Sum Sq	R-Sq	Adj. R-Sq
Full Model Z1 Z3 Solar.R:Temp	173.062	24.011	58.459	0.70885	0.67320
	171.062	24.011	58.459	0.70885	0.67650
	169.158	24.032	58.438	0.70860	0.67946
	167.320	24.067	58.403	0.70817	0.68217
Temp	165.784	24.168	58.302	0.70695	0.68397
Wind:Temp	163.936	24.201	58.269	0.70655	0.68661
Solar.R:Wind	162.515	24.328	58.142	0.70501	0.68799
Z2	161.562	24.558	57.912	0.70222	0.68804

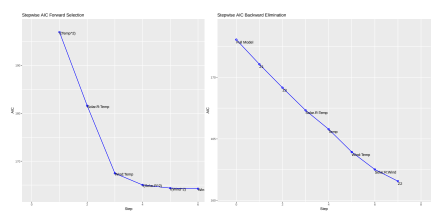




Statistical Learning

F-AIC; B-AIC plots

```
plot(M5.FAIC)
plot(M5.BAIC)
>>
```







Compare F-AIC; B-AIC with F-BIC; B-BIC

```
AIC_variables <- c(AIC(M5.FAIC$model), AIC(M5.BAIC$model), AIC(M5.FBIC), AIC(M5.FBIC))
AICc_variables <- c(AIC(M5.FAIC$model), AICc(M5.BAIC$model), AICc(M5.FBIC), AICc(M5.FBIC))
BIC_variables <- c(BIC(M5.FAIC$model), BIC(M5.BAIC$model), BIC(M5.FBIC))
metrics_matrix <- rbind(AIC = AIC_variables, AICc = AICc_variables, BIC = BIC_variables)
IC_df <- as.data.frame(metrics_matrix)
colnames(IC_df) <- c("F-AIC", "B-AIC", "F-BIC", "B-BIC")
IC_df
```

Criterion	F-AIC	B-AIC	F-BIC	B-BIC
AIC	164.232623164374	161.562457576751	167.464403165684	161.562457576751
AICc	165.644387870256	162.649836217528	168.035831737113	162.649836217528
віс	185.908864774872	180.529168985937	181.012054172246	180.529168985937





Statistical Learning

Cross-validation for stepwise models

We can modify the KfoldCVPerf.stepwise() function:

```
KfoldCVPerf.stepwise <- function (K=10, data, formula, name='model name',
                                  folds.seed = 42, inv.trans=function(y) return(y),
                                  method="forward", criterion="AIC", nvmax=12) {
 ### Same code as KfoldCVPerf() ###
 foreach (i = 1:K) %do% {
       #Keep the i-th fold as it is (testing)
        val fold <- folds list[[i]]
       y_val <- val_fold[[y]]</pre>
        #Combine the remaining folds (training)
        train folds <- do.call(rbind, folds list[-i])
        v_train <- train_folds[[v]]</pre>
        # Temporarily assign train folds to a global variable
        .GlobalEnv$train folds temp <- train folds #Important for F-BAIC
        #Choose model
        if (method == "forward" && criterion == "AIC") {
            # Use the global temporary variable for model fitting
            LM <- lm(formula, data = train_folds_temp)
            model.FAIC <- ols step forward aic(LM)
            model <- model.FATC$model
        } else if (method == "forward" && criterion == "BIC") {
            model.forward <- regsubsets(formula, data=train folds.
                                      method="forward", nymax=nymax)
            model.Fbic <- summary(model.forward)$bic</pre>
            FBIC vars <- names(coef(model.forward, which.min(model.Fbic)))
            FBIC_formula <- as.formula(paste("y_train ~",
                                 paste(FBIC_vars[-1], collapse=' + ')))
 FORTH
            model <- lm(FBIC formula, data=train folds)
```

Cross-validation for stepwise models

```
} else if (method == "backward" && criterion == "AIC") {
    # Use the global temporary variable for model fitting
    LM <- lm(formula, data = train_folds_temp)
    model.BAIC <- ols_step_backward_aic(LM)
    model <- model.BATC$model
} else if (method == "backward" && criterion == "BIC") {
    model.backward <- regsubsets(formula, data=train folds,
                             method="backward", nvmax=nvmax)
    model.Bbic <- summary(model.backward)$bic
    BBIC_vars <- names(coef(model.backward, which.min(model.Bbic)))
    BBIC_formula <- as.formula(paste("y_train ~",
                         paste(BBIC_vars[-1], collapse=' + ')))
    model <- lm(BBIC formula, data=train folds)
} else {
    stop("Unsupported method or criterion.")
rm(train folds temp, envir = .GlobalEnv) #remove globenv
preds <- predict(model, newdata = val fold)
v val <- inv.trans(v val)
preds <- inv.trans(preds)</pre>
### Same code as before ###
return (perf_df)
```



}



Repeated cross-validation for stepwise models

```
RepeatedCV.stepwise <- function (repeats=50, K=10, data, method, formula,
                         model_name, criterion, inv.trans = function(y) y) {
    require(foreach)
    mean rows list <- list() # store the row means
    foreach (seed = 1:repeats) %do% {
        cv_results <- KfoldCVPerf.stepwise(K=K, data=data, method=method,
                                   criterion=criterion, formula=formula,
                                   name=model name, folds.seed = seed,
                                   inv.trans=inv.trans)
        mean_row <- cv_results[cv_results[,1]=="Mean", -1]</pre>
        mean rows list[[seed]] <- as.numeric(mean row)
    mean_rows_df <- do.call(rbind, mean_rows_list)
    # Average for each metric
    average metrics <- colMeans(mean rows df, na.rm = TRUE)
    average_metrics_df <- as.data.frame(t(average_metrics))</pre>
    colnames(average_metrics_df) <- names(cv_results)[-1]</pre>
    rownames(average metrics df) <- "RCV Average"
    return(average_metrics_df)
M5 FAIC.RCV <- RepeatedCV.stepwise(
    repeats = 50.
    K = 10.
    data = data .
    method='forward'.
    criterion='AIC',
   formula = M5_formula,
    model_name = 'M5-FAIC',
    inv.trans = function(y) exp(y))
             ... define BAIC, FBIC, BBIC ...
                                                     ###
```



Repeated cross-validation - based on 50 repeats

M5_FAIC.RCV M5_FBIC.RCV M5_BAIC.RCV M5_BBIC.RCV >>

M5	M5-FAIC (RMSE)	M5-FAIC (MAE)	M5-FAIC (sMdAPE)	M5-FAIC (MAPE)	M5-FAIC (LAR)
RCV Average	20.7875055645847	14.5760054786763	33.7482568899758	50.21819388008	0.414177365614387
M5	M5-FBIC (RMSE)	M5-FBIC (MAE)	M5-FBIC (sMdAPE)	M5-FBIC (MAPE)	M5-FBIC (LAR)
RCV Average	20.1683579404873	14.4788907089285	33.713714195958	49.1706383379745	0.415243308699007
M5	M5-BAIC (RMSE)	M5-BAIC (MAE)	M5-BAIC (sMdAPE)	M5-BAIC (MAPE)	M5-BAIC (LAR)
RCV Average	20.5759123372757	14.2113507578813	32.8681337086692	50.023554489658	0.40249199693544



M5

RCV Average

M5-BBIC (MAPE)

48.2419010016839

M5-BBIC (sMdAPE)

33.8128440882392

M5-BBIC (MAE)

14.5744690252833

M5-BBIC (RMSE)

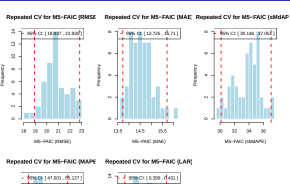
21.008066759466

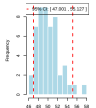
M5-BBIC (LAR)

0.4129241643574

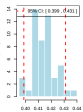
Repeated cross-validation - 95% CI for forward-AIC

based on 50 repeats





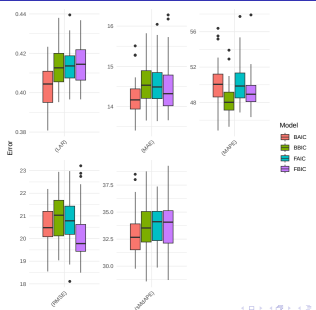
M5-FAIC (MAPE)







Repeated cross-validation - Box-Plots based on 50 repeats





 Regarding LAR, FBIC exhibits a lower median error compared to FAIC, while BAIC shows a lower median error than BBIC, indicating a better performance of the AIC criterion in the backward selection approach for this metric.





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- In terms of MAE, FBIC has a lower median error compared to FAIC, indicating better performance with forward selection using BIC. Conversely, BAIC has a lower median error than BBIC, suggesting better performance with backward selection using AIC.





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- In terms of MAE, FBIC has a lower median error compared to FAIC, indicating better performance with forward selection using BIC. Conversely, BAIC has a lower median error than BBIC, suggesting better performance with backward selection using AIC.
- With respect to MAPE, both FBIC and BBIC present lower median errors than FAIC and BAIC, respectively, which could imply that the BIC criterion is more effective in both forward and backward selection methods for minimizing MAPE.





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- In terms of MAE, FBIC has a lower median error compared to FAIC, indicating better performance with forward selection using BIC. Conversely, BAIC has a lower median error than BBIC, suggesting better performance with backward selection using AIC.
- With respect to MAPE, both FBIC and BBIC present lower median errors than FAIC and BAIC, respectively, which could imply that the BIC criterion is more effective in both forward and backward selection methods for minimizing MAPE.
- For RMSE, FBIC shows a lower median error than FAIC, while BAIC shows a lower median error than BBIC.



♦ Compare the previous models to F-P and B-P (use the p-value





♦ Compare the previous models to F-P and B-P (use the p-value threshold that was discussed in class).

We can just add the following code to the KfoldCVPerf.stepwise():

```
### ... ###
} else if (method == "forward" && criterion == "p_val") {
    # Use the global temporary variable for model fitting
    LM <- lm(formula, data = train_folds_temp)
    model.Fp <- ols_step_forward_p(LM, p_val=p) #def p as 0.005 (input)
    model <- model.Fp$model
} else if (method == "backward" && criterion == "p_val") {
    # Use the global temporary variable for model fitting
    LM <- lm(formula, data = train_folds_temp)
    model.Bp <- ols_step_backward_p(LM, p_val=p) #0.005
    model <- model.Bp$model
} ### ... same code ... ###</pre>
```





Statistical Learning

Repeated cross-validation - based on 50 repeats (Fp; Bp)

M5_Fp.RCV M5_Bp.RCV >>

M5	M5-Fp (RMSE)	M5-Fp (MAE)	M5-Fp (sMdAPE)	M5-Fp (MAPE)	M5-Fp (LAR)	
RCV Average	20.7095116438382	14.6475772954767	34.5658151361359	52.4466762438904	0.41670129690786	
M5	M5-Bn (RMSF)	M5-Rn (MAF)	M5-Rn (sMdAPF)	M5-Rn (MAPF)	M5-Rn (LAR)	
M5 RCV Average	M5-Bp (RMSE)	M5-Bp (MAE)	M5-Bp (sMdAPE)	M5-Bp (MAPE)	M5-Bp (LAR)	

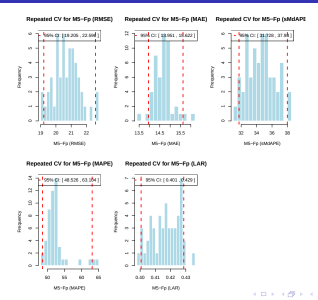




Statistical Learning

Repeated cross-validation - 95% CI for forward-p-value

based on 50 repeats

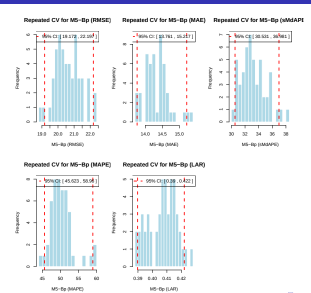






Repeated cross-validation - 95% CI for backward-p-value

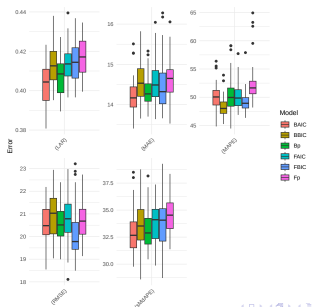
based on 50 repeats







Repeated cross-validation - Box-Plots based on 50 repeats





FORTH

LASSO and Ridge regression

♦ Finally estimate M5 using LASSO and ridge with lambda tuned via





LASSO and Ridge regression

 Finally estimate M5 using LASSO and ridge with lambda tuned via cross-validation. Compare against the previous models.

Again, we can modify the KfoldCVPerf.stepwise() function in order to get the performance for penalized models like Ridge and LASSO:

```
### ... same code ... ###
} else if (method == "ridge") {
    require(doMC)
    set_seed(42)
    registerDoMC(cores = 2) #parallel backend
    system.time(
      model <- cv.glmnet(
        x=model.matrix(formula, data).
        y=as.vector(data[y])[[1]],
        type.measure="mse", alpha=0, nfolds=10, parallel = TRUE))
} else if (method == "lasso") {
    require(doMC)
    set.seed(42)
    registerDoMC(cores = 2) #parallel backend
    system.time(
      model <- cv.glmnet(
        x=model.matrix(formula, data),
        y=as.vector(data[y])[[1]],
        type.measure="mse", alpha=1, nfolds=10, parallel = TRUE))
} else {
    stop("Unsupported method or criterion.")
```

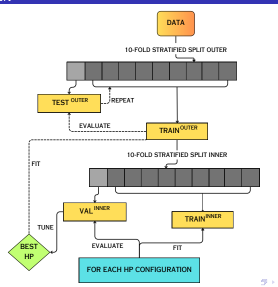
LASSO and Ridge regression

```
rm(train_folds_temp, envir = .GlobalEnv) #remove globenv
        # Check for nestedcy (ridge: lasso)
        if (method == "ridge" || method == "lasso") {
            preds <- as.vector(predict(model.
                              newx = model.matrix(formula, val fold).
                              s = "lambda.min"))
        } else {
            preds <- predict(model, newdata = val_fold)</pre>
        v_val <- inv.trans(v_val)</pre>
        preds <- inv.trans(preds)
        ### ... same code ... ###
# Repeated CV for Ridge and LASSO
M5_ridge.RCV <- RepeatedCV.stepwise(
    repeats = 50,
   K = 10
    data = data .
    method='ridge',
    formula = M5 formula,
    model name = 'M5-Ridge'.
    inv.trans = function(y) exp(y))
M5_lasso.RCV <- RepeatedCV.stepwise(
    repeats = 50.
   K = 10.
    data = data_,
    method='lasso',
    formula = M5 formula.
    model_name = 'M5-LASSO',
    invetrans = function(y) exp(y))
```



Nested cross-validation; tune lambda for Ridge, LASSO

$$\min_{(\beta_0,\beta)\in\mathbb{R}^{p+1}} \frac{1}{2N} \sum_{i=1}^{N} (y_i - \beta_0 - x_i^T \beta)^2 + \lambda \left[(1-a) \|\beta\|_2^2 / 2 + a \|\beta\|_1 \right], a = 0, 1.$$





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Repeated cross-validation for LASSO and Ridge

```
M5_ridge.RCV
M5_lasso.RCV
>>
```

M5	M5-Ridge (RMSE)	M5-Ridge (MAE)	M5-Ridge (sMdAPE)	M5-Ridge (MAPE)	M5-Ridge (LAR)
RCV Average	18.2001363127757	13.0288544659089	30.9505759289211	42.9486625659833	0.374310211087681

M5	M5-LASSO (RMSE)	M5-LASSO (MAE)	M5-LASSO (sMdAPE)	M5-LASSO (MAPE)	M5-LASSO (LAR)
RCV Average	18.5161292116031	13.2162243476291	31.2672848724556	44.2980742675429	0.378301429968909

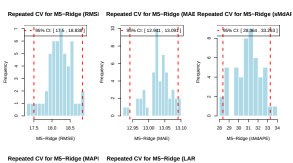
- Ridge Regression demonstrates superior performance compared to all previously evaluated models.
- Lasso Regression surpasses the earlier models as well, yet Ridge is preferred for its slightly lower errors.

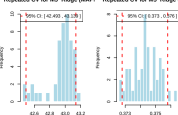


Toannis Maris

Repeated cross-validation - 95% CI for Ridge

based on 50 repeats





M5-Ridge (MAPE)





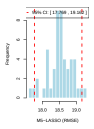
0.375

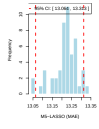
M5-Ridge (LAR)

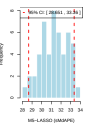
Repeated cross-validation - 95% CI for LASSO

based on 50 repeats

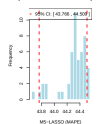
Repeated CV for M5-LASSO (RMS Repeated CV for M5-LASSO (MAIRepeated CV for M5-LASSO (sMdA

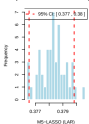






Repeated CV for M5-LASSO (MAP Repeated CV for M5-LASSO (LAF



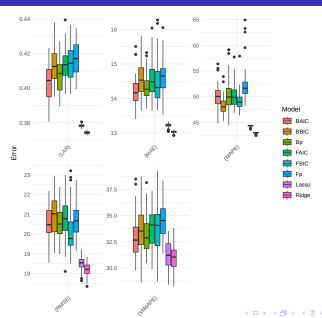






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Repeated cross-validation - Box-Plots based on 50 repeats





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Overall results based on repeated cross-validation

Model	RMSE	MAE	sMdAPE	MAPE	LAR
M3 RCV Average	20.7288	16.1223	33.8353	67.7265	0.4383
M4 RCV Average	18.5804	13.8479	34.8101	53.5018	0.4506
M5 RCV Average	18.7312	13.5512	33.5723	49.2004	0.4124
M5-FAIC RCV Average	20.7875	14.5760	33.7483	50.2182	0.4142
M5-FBIC RCV Average	20.1684	14.4789	33.7137	49.1706	0.4152
M5-BAIC RCV Average	20.5759	14.2114	32.8681	50.0236	0.4025
M5-BBIC RCV Average	21.0081	14.5745	33.8128	48.2419	0.4129
M5-Fp RCV Average	20.7095	14.6476	34.5658	52.4467	0.4167
M5-Bp RCV Average	20.6057	14.3282	33.2057	50.2604	0.4069
M5-Ridge RCV Average	18.2001	13.0289	30.9506	42.9487	0.3743
M5-LASSO RCV Average	18.5161	13.2162	31.2673	44.2981	0.3783

FORT

Table: Repeated Cross-validation average performance for different models



