

Regularization

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Load libraries, get data & set seed for reproducibility

```
set.seed(222)    # seed for reproducibility
library(glmnet)  # for ridge regression

## Loading required package: Matrix
## Loaded glmnet 4.0-2
library(dplyr)   # for data cleaning

##
## Attaching package: 'dplyr'
## The following objects are masked from 'package:stats':
##
##   filter, lag
## The following objects are masked from 'package:base':
##
##   intersect, setdiff, setequal, union
library(psych)   # for function tr() to compute trace of a matrix
data("mtcars")  #loading data
```

RIDGE REGRESSION

```
#1. RIDGE REGRESSION
# Center y, X will be standardized in the modelling function
#(ridge regression assumes the predictors are standardized and the response is centered)
y <- mtcars %>% select(mpg) %>% scale(center = TRUE, scale = FALSE) %>% as.matrix()
X <- mtcars %>% select(-mpg) %>% as.matrix()
```

Explanation: We scale the X matrix to ensure that the penalty term penalizes each coefficient equally. Instead of solving this heteroskedasticity problem by equalizing the variances of all predictors via scaling, we could just as well use them as weights in the estimation process! This is the idea behind the Differentially-weighted or Heteroskedastic Ridge Regression.

Perform 10-fold cross-validation to select lambda:

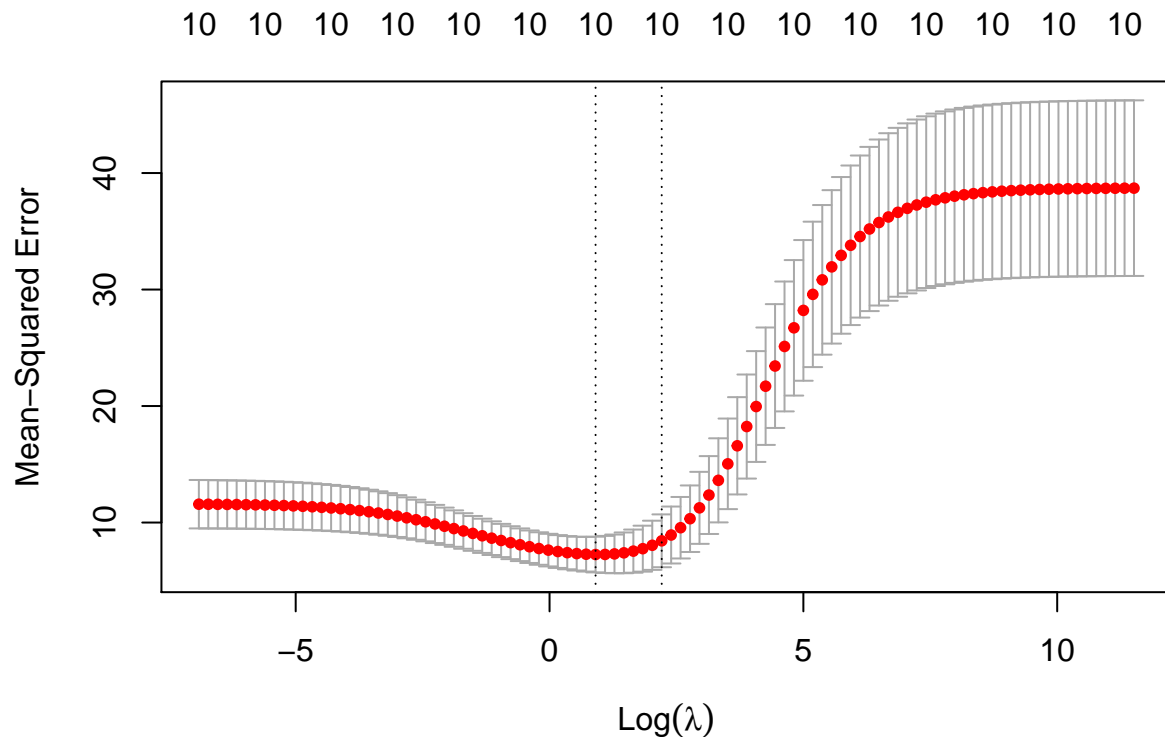
```
lambdas_to_try <- 10^seq(-3, 5, length.out = 100)
#10-fold cross-validation to select lambda
```

```

# Setting alpha = 0 implements ridge regression
ridge_cv <- cv.glmnet(X, y, alpha = 0, lambda = lambdas_to_try,
                     standardize = TRUE, nfolds = 10)
#function cv.glmnet does k-fold cross-validation for glmnet,
#produces a plot, and returns a value for lambda

# Plot cross-validation results
plot(ridge_cv)

```



```

# Best cross-validated lambda
lambda_cv <- ridge_cv$lambda.min
# Fit final model, get its sum of squared residuals and multiple R-squared
model_cv <- glmnet(X, y, alpha = 0, lambda = lambda_cv, standardize = TRUE)
y_hat_cv <- predict(model_cv, X)
ssr_cv <- t(y - y_hat_cv) %*% (y - y_hat_cv)
rsq_ridge_cv <- cor(y, y_hat_cv)^2
rsq_ridge_cv

```

```

##          s0
## mpg 0.8536968

```

Use information criteria to select lambda:

```

X_scaled <- scale(X)
aic <- c()
bic <- c()
for (lambda in seq(lambdas_to_try)) {
  # Run model

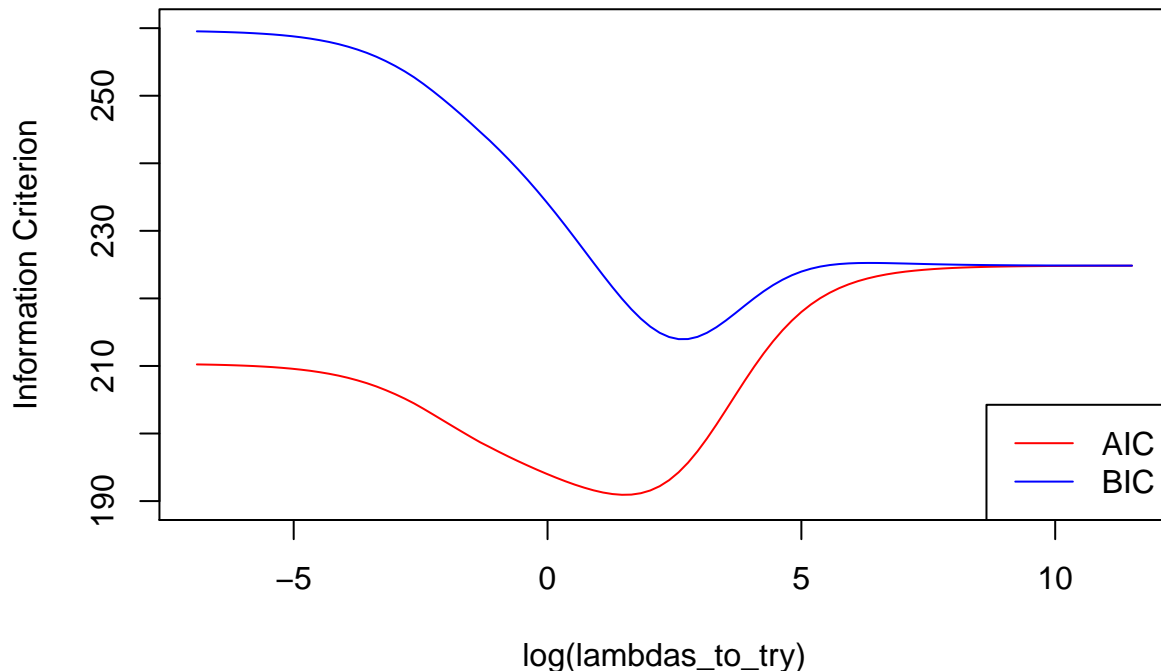
```

```

model <- glmnet(X, y, alpha = 0, lambda = lambdas_to_try[lambda], standardize = TRUE)
# Extract coefficients and residuals (remove first row for the intercept)
betas <- as.vector((as.matrix(coef(model))[-1, ]))
resid <- y - (X_scaled %*% betas)
# Compute hat-matrix and degrees of freedom
ld <- lambdas_to_try[lambda] * diag(ncol(X_scaled))
H <- X_scaled %*% solve(t(X_scaled) %*% X_scaled + ld) %*% t(X_scaled)
df <- tr(H)
# Compute information criteria
aic[lambda] <- nrow(X_scaled) * log(t(resid) %*% resid) + 2 * df
bic[lambda] <- nrow(X_scaled) * log(t(resid) %*% resid) + 2 * df * log(nrow(X_scaled))
}

# Plot information criteria against tried values of lambdas
plot(log(lambdas_to_try), aic, col = "red", type = "l",
     ylim = c(190, 260), ylab = "Information Criterion")
lines(log(lambdas_to_try), bic, col = "blue")
legend("bottomright", lwd = 1, col = c("red", "blue"), legend = c("AIC", "BIC"))

```



```

# Optimal lambdas according to both criteria
lambda_aic <- lambdas_to_try[which.min(aic)]
lambda_bic <- lambdas_to_try[which.min(bic)]

# Fit final models, get their sum of squared residuals and multiple R-squared
model_aic <- glmnet(X, y, alpha = 0, lambda = lambda_aic, standardize = TRUE)
y_hat_aic <- predict(model_aic, X)
ssr_aic <- t(y - y_hat_aic) %*% (y - y_hat_aic)
rsq_ridge_aic <- cor(y, y_hat_aic)^2

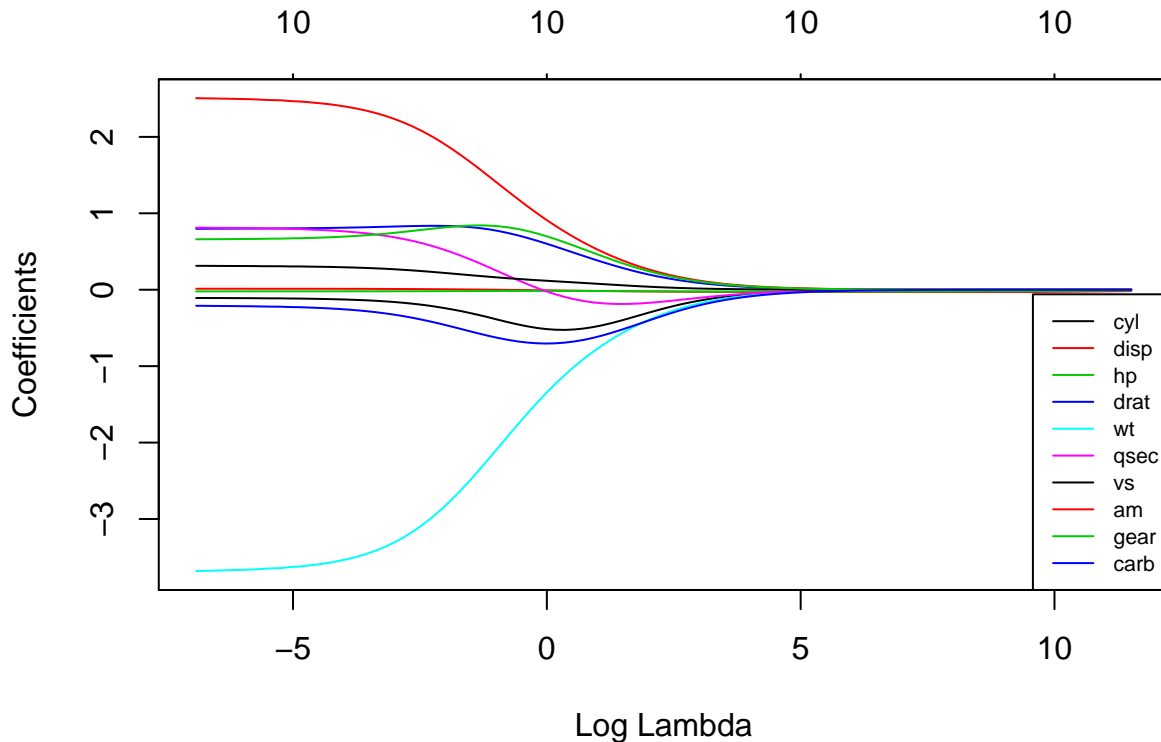
model_bic <- glmnet(X, y, alpha = 0, lambda = lambda_bic, standardize = TRUE)
y_hat_bic <- predict(model_bic, X)
ssr_bic <- t(y - y_hat_bic) %*% (y - y_hat_bic)

```

```
rsq_ridge_bic <- cor(y, y_hat_bic)^2
```

See how increasing lambda shrinks the coefficients :

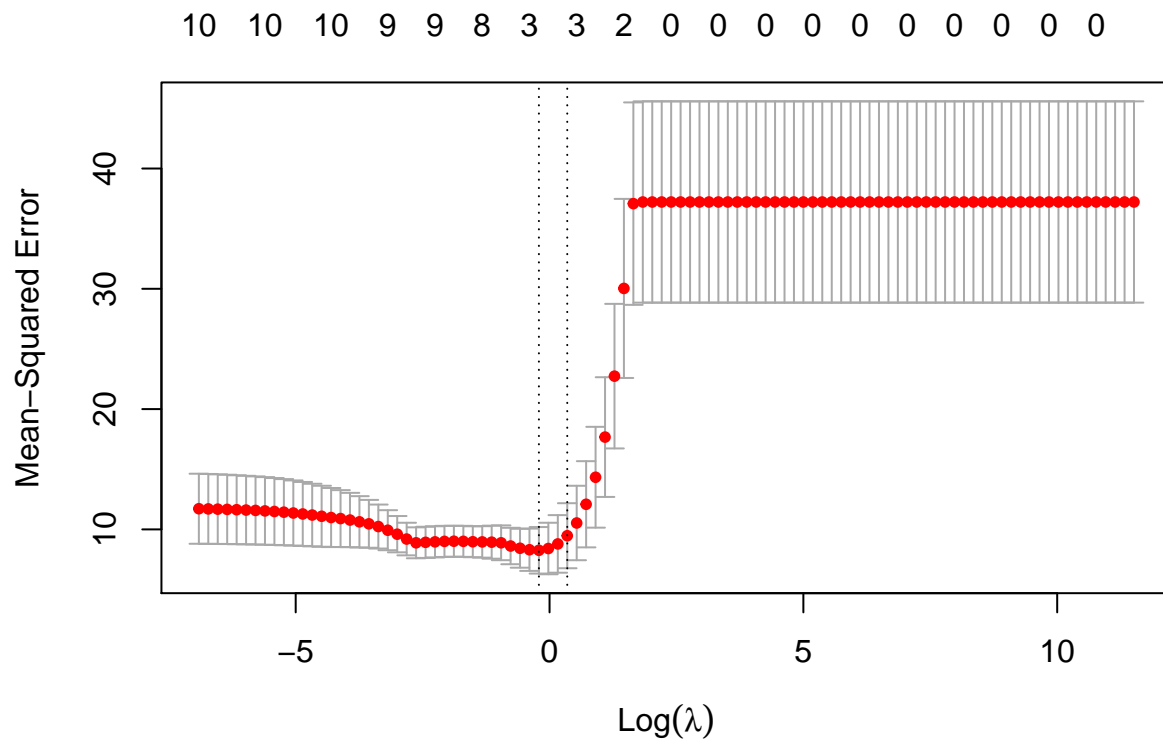
```
# Each line shows coefficients for one variables, for different lambdas.
# The higher the lambda, the more the coefficients are shrunk towards zero.
res <- glmnet(X, y, alpha = 0, lambda = lambdas_to_try, standardize = FALSE)
plot(res, xvar = "lambda")
legend("bottomright", lwd = 1, col = 1:6, legend = colnames(X), cex = .7)
```



LASSO REGRESSION

Lasso = Least Absolute Shrinkage and Selection Operator, is quite similar conceptually to ridge regression. It also adds a penalty for non-zero coefficients, but unlike ridge regression which penalizes sum of squared coefficients (the so-called L2 penalty), lasso penalizes the sum of their absolute values (L1 penalty). As a result, for high values of lambda, many coefficients are exactly zeroed under lasso, which is never the case in ridge regression.

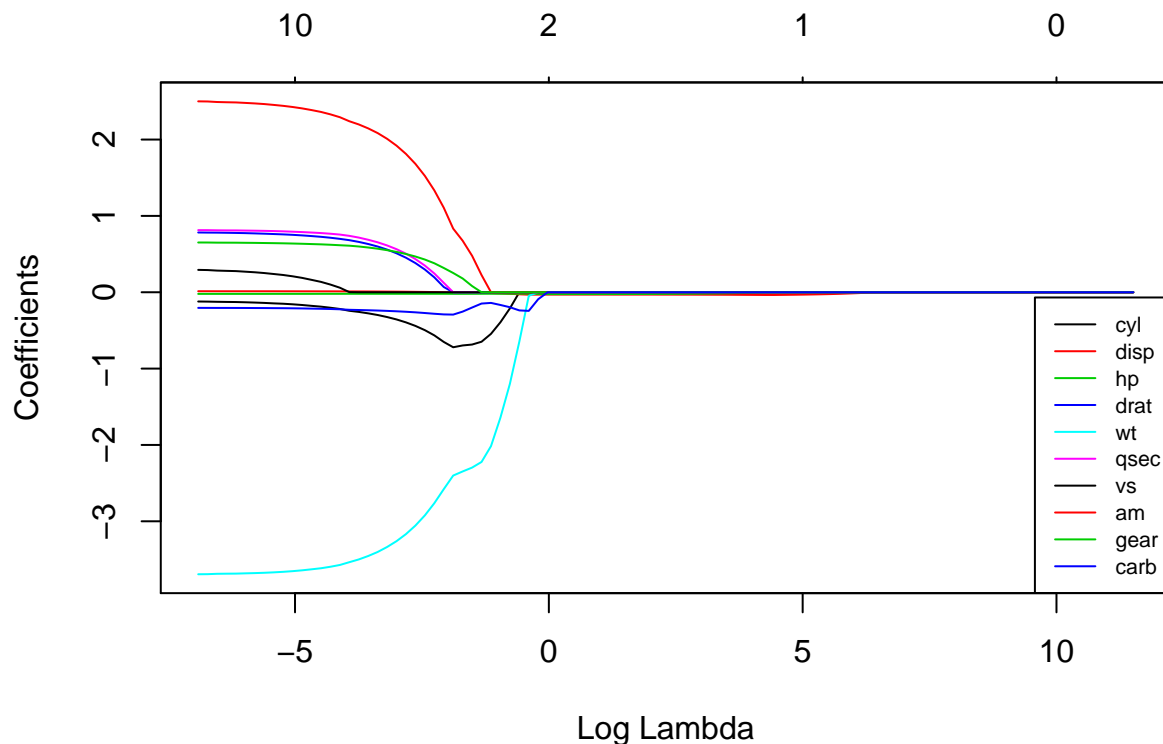
```
# Perform 10-fold cross-validation to select lambda
lambdas_to_try <- 10^seq(-3, 5, length.out = 100)
# Setting alpha = 1 implements lasso regression
lasso_cv <- cv.glmnet(X, y, alpha = 1, lambda = lambdas_to_try,
                      standardize = TRUE, nfolds = 10)
# Plot cross-validation results
plot(lasso_cv)
```



```
# Best cross-validated lambda
lambda_cv <- lasso_cv$lambda.min
# Fit final model, get its sum of squared residuals and multiple R-squared
model_cv <- glmnet(X, y, alpha = 1, lambda = lambda_cv, standardize = TRUE)
y_hat_cv <- predict(model_cv, X)
ssr_cv <- t(y - y_hat_cv) %*% (y - y_hat_cv)
rsq_lasso_cv <- cor(y, y_hat_cv)^2
```

See how increasing lambda shrinks the coefficients:

```
# Each line shows coefficients for one variables, for different lambdas.
# The higher the lambda, the more the coefficients are shrunk towards zero.
res <- glmnet(X, y, alpha = 1, lambda = lambdas_to_try, standardize = FALSE)
plot(res, xvar = "lambda")
legend("bottomright", lwd = 1, col = 1:6, legend = colnames(X), cex = .7)
```



Let us compare the multiple R-squared of various models we have estimated:

RIDGE VS. LASSO

```
rsq <- cbind("R-squared" = c(rsq_ridge_cv, rsq_ridge_aic, rsq_ridge_bic, rsq_lasso_cv))
rownames(rsq) <- c("ridge cross-validated", "ridge AIC", "ridge BIC", "lasso cross_validated")
print(rsq)
```

```
##                R-squared
## ridge cross-validated 0.8536968
## ridge AIC             0.8496310
## ridge BIC             0.8412011
## lasso cross_validated 0.8424217
```

Some more general considerations about how ridge and lasso compare:

- Often neither one is overall better.
- Lasso can set some coefficients to zero, thus performing variable selection, while ridge regression cannot.
- Both methods allow to use correlated predictors, but they solve multicollinearity issue differently:
- In ridge regression, the coefficients of correlated predictors are similar;
- In lasso, one of the correlated predictors has a larger coefficient, while the rest are (nearly) zeroed.
- Lasso tends to do well if there are a small number of significant parameters and the others are close to zero (ergo: when only a few predictors actually influence the response).
- Ridge works well if there are many large parameters of about the same value (ergo: when most predictors impact the response).

(However, in practice, we don't know the true parameter values, so the previous two points are somewhat theoretical. Just run cross-validation to select the more suited model for a specific case)

ELASTIC NET

(optional task) Elastic Net first emerged as a result of critique on lasso, whose variable selection can be too dependent on data and thus unstable. The solution is to combine the penalties of ridge regression and lasso to get the best of both worlds:

Check multiple R-squared:

```
y_hat_enet <- predict(elastic_net_model, X)
y_hat_enet
```

##	Mazda RX4	Mazda RX4 Wag	Datsun 710	Hornet 4 Drive
##	2.05551135	1.74280776	6.28216492	0.52249594
##	Hornet Sportabout	Valiant	Duster 360	Merc 240D
##	-3.27605721	0.15274583	-5.58554614	2.91074947
##	Merc 230	Merc 280	Merc 280C	Merc 450SE
##	3.20191054	-0.23145256	-0.15061543	-4.52522390
##	Merc 450SL	Merc 450SLC	Cadillac Fleetwood	Lincoln Continental
##	-3.98074275	-4.00295948	-8.12176404	-8.41163995
##	Chrysler Imperial	Fiat 128	Honda Civic	Toyota Corolla
##	-8.22108855	7.26534418	8.46202739	8.05969802
##	Toyota Corona	Dodge Challenger	AMC Javelin	Camaro Z28
##	3.86926293	-3.30051536	-2.67542264	-5.50883573
##	Pontiac Firebird	Fiat X1-9	Porsche 914-2	Lotus Europa
##	-4.13174939	7.59060420	6.05949380	6.85211961
##	Ford Pantera L	Ferrari Dino	Maserati Bora	Volvo 142E
##	-1.83268991	-0.04290647	-6.03188981	5.00416337

```
rsq_enet <- cor(y, y_hat_enet)^2
rsq_enet
```

```
##      [,1]
## mpg 0.8570287
```