Statistical Computing with R: Masters in Data Sciences 503 (S22) Third Batch, SMS, TU, 2024

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Review Preview

- Simple Linear Regression
 - Gradient Decent fit
 - Model Accuracy
 - Model Prediction
 - Model Validation
 - Validation set, Leave one out cross validation, k-folds cross validation, repeated k-folds cross-validation etc.)

- Multiple Linear Regression
 - Simple linear regression +
 - Multicollinearity, its assessment and solutions
 - Regularization
 - Ridge
 - Lasso
 - Elastic Net (Ridge+Lasso)

Simple Linear Regression: Gradient Decent

https://towardsdatascience.com/linear-regression-using-gradient-descent-97a6c8700931

 Linear Regression can also be fitted with Gradient Decent algorithm instead of OLS

$$egin{split} D_m &= rac{1}{n} \sum_{i=0}^n 2(y_i - (mx_i + c))(-x_i) \ D_m &= rac{-2}{n} \sum_{i=0}^n x_i (y_i - ar{y}_i) \end{split}$$

 Here we minimize the loss function (E) to find m (slope b) and c (constant a)

$$E = rac{1}{n} \sum_{i=0}^{n} (y_i - (mx_i + c))^2$$

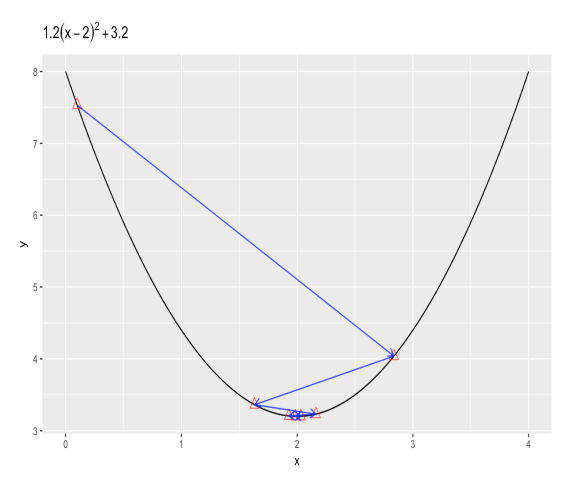
E = MSE = Mean Sum of Square = Mean Squared Error = Variation

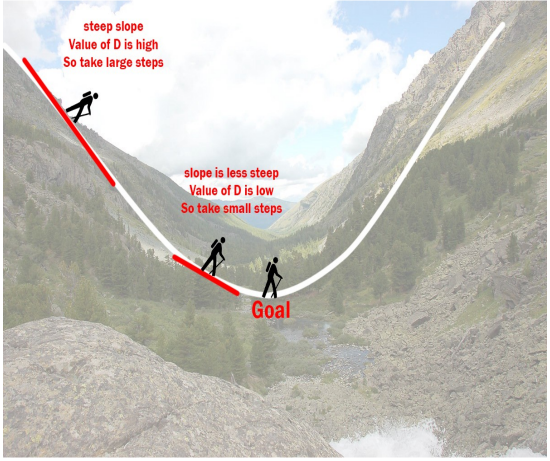
$$D_c = rac{-2}{n} \sum_{i=0}^n (y_i - ar{y}_i)$$

We can solve these two equations to get the value of m and c. This is an **optimization problem** as we need to MINIMIZE the loss function **MSE** using multiple iterations!

Illustration:

http://ethen8181.github.io/machine-learning/linear_regression/linear_regession.html





How to get MSE of Simple Linear Model?

- Using the residuals of linear model:
 lm1 <- lm(mpg ~ wt, data=mtcars)
 (mse <- mean(lm1\$residuals^2))
- Saving predicted values:
 data <- data.frame(pred = predict(lm1), actual = mtcars\$mpg)
 head(data)
 mean((data\$actual data\$pred)^2)

- > (mse <- mean(lm1\$residuals^2))
- [1] 8.697561
- > mean((data\$actual data\$pred)^2)
- [1] 8.697561

Model Accuracy of Linear Model:

- R-square Explained variance (higher is better!)
- RMSE Root of MSE (lower is better)
- MAE Mean Absolute Error (lower is better)
- MAPE Mean Absolute Percentage Error (lower is better)

$$MAPE = \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{1000\%}{n} \sum_{\text{Each residual is scaled against the actual value}} \frac{100\%}{n} \sum_{\text{Each resid$$

#Better to use "caret" package

Install.packages("caret")

library(caret)

R2 <- R2(data\$pred, data\$actual)

• 0.7528328

RMSE <- RMSE(data\$pred, data\$actual)

• 2.949163

MAE <- MAE(data\$pred, data\$actual)

• 2.340642

Get MAPE in R as:

• 12.60733

If LINE is valid after BLUE then we can predict: (So, we will use "Im1<-Im(mpg~wt, data=mtcars)" model to do it!)

- We need to save independent variable value/values in a new data:
 new.wt <- data.frame(wt = 6)
- We can then use this data to predict the value of the dependent variable based on the fitted model as:

```
predict(lm1, newdata = new.wt)
```

Result = 5.218297

• Interpretation: Cars with 6000 lbs weight will (only) give 5.22 miles per gallon as per the linear regression algorithm!)

Validation & Cross-validation for Predictive Modelling including Linear Model:

- In statistics, we normally use the "full" data to do predictions
- In machine learning, we use validation/cross-validation sets to do the predictions
- Validation/Cross-validation can be done with:
 - Validation set with split data
 - Leave one out cross-validation (LOOCV) with full data
 - K-fold cross-validation with full data
 - Repeated k-fold validation with full data

Validation: Validation set (widely used!)

- Here the full data is "randomly" divided into two sets:
 - Training set
 - Testing set (validation set)
- Then model is fitted in the training set
- The model fit is then validated in the testing set using prediction
- This is the most widely used cross-validation method in the supervised machine learning

Lets do it for "mtcars" data:

test.data <- data[ind==2,]

```
#Define the mtcars data as "data":
data <- mtcars
#Use random seed to replicate the result
set.seed(1234)
#Do random sampling to divide the cases into two independent samples
ind <- sample(2, nrow(mtcars), replace = T, prob = c(0.7, 0.3))
#Data partition
train.data <- data[ind==1,]
```

Model Fit, Prediction and Cross-Validation: Validation set approach

```
lm4 <- lm(mpg~wt, data = train.data)</pre>
library(dplyr)
library(caret)
predictions <- Im4 %>%
predict(test.data)
data.frame(R2 = R2(predictions,
test.data$mpg),
      RMSE = RMSE(predictions,
      test.data$mpg),
      MAE = MAE(predictions,
      test.data$mpg))
```

Model Accuracy of Training dataset: summary(lm4)

- Multiple R-squared: 0.7013
- MSE = 9.526359 (How?)
- RMSE = SQRT(MSE) = 3.08648

Model Accuracy of Testing dataset:

R2 RMSE MAE

0.90310852.2793031.698583

Model Fit, Prediction and Cross-Validation: Leave-One-Out Cross-Validation approach:

```
#Leave one out CV
library(caret)
# Define training control
train.control <- trainControl(method = "LOOCV")</pre>
# Train the model
model1 <- train(mpg ~wt, data = mtcars, method = "lm", trControl = train.control)
#Why mtcars data, why not train.data?
# Summarize the results
print(model1)
```

Linear Regression

32 samples

1 predictor

No pre-processing

Resampling: Leave-One-Out Cross-Validation

Summary of sample sizes: 31, 31, 31, 31, 31, 31, ...

Resampling results:

RMSE Rsquared MAE

3.201673 0.7104641 2.517436

Tuning parameter 'intercept' was held constant at a value of TRUE

Prediction with LOOCV:

```
predictions1 <- model1 %>%
predict(test.data)

data.frame(R2 = R2(predictions1,
test.data$mpg),

    RMSE = RMSE(predictions1,
test.data$mpg),

    MAE = MAE(predictions1,
test.data$mpg))
```

R2 RMSE MAE

0.9031085 2.244232 1.714515

Model Fit, Prediction and Cross-Validation: K-folds Cross-Validation approach

```
#k-fold cross validation
library(caret)
# Define training control
set.seed(123)
train.control <- trainControl(method = "cv", number
= 10)
# Train the model
model2 <- train(mpg ~ wt, data = mtcars, method =
"lm",
        trControl = train.control)
# Summarize the results
print(model2)
```

```
Linear Regression 32 samples
```

1 predictor

No pre-processing

Resampling: Cross-Validated (10 fold)

Summary of sample sizes: 28, 28, 29, 29, 29, 30, ...

Resampling results:

RMSE Rsquared MAE

2.85133 0.7346939 2.375068

Tuning parameter 'intercept' was held constant at a value of TRUE

Predictions with k-folds CV:

```
predictions2 <- model2 %>%
predict(test.data)

data.frame(R2 = R2(predictions2,
test.data$mpg),

    RMSE = RMSE(predictions2,
test.data$mpg),

    MAE = MAE(predictions2,
test.data$mpg))
```

R2 RMSE MAE

• 0.9031085 2.244232 1.714515

Model Fit, Prediction and Cross-Validation: Repeated K-folds Cross-Validation approach

```
#repeated k-fold cross validation
library(caret)
# Define training control
set.seed(123)
train.control <- trainControl(method = "repeatedcv",
                 number = 10, repeats = 3)
# Train the model
model <- train(mpg ~wt, data = mtcars, method =
"lm",
        trControl = train.control)
# Summarize the results
print(model)
```

```
Linear Regression
32 samples
1 predictor
No pre-processing
Resampling: Cross-Validated (10 fold, repeated 3 times)
Summary of sample sizes: 28, 28, 29, 29, 29, 30, ...
Resampling results:
 RMSE
                     Rsquared
                                           MAE
 2.975392
                     0.8351572
                                           2.539797
Tuning parameter 'intercept' was held constant at a value of
TRUE
```

Prediction with repeated k-folds CV:

```
predictions3 <- model3 %>%
predict(test.data)
data.frame(R2 = R2(predictions3,
test.data$mpg),
      RMSE = RMSE(predictions3,
test.data$mpg),
      MAE = MAE(predictions3,
test.data$mpg))
```

R2 RMSE MAE

0.9031085 2.244232 1.714515

Summary: Which one should be used based on R-squared values of "Im" model?

• R-square for training set: 0.7013

• R-square for testing set: 0.9031085

- R-square for training with LOOCV:
 0.7104641
- R-square for testing with LOOCV: 0.9031085

- R-square for training with k-folds CV: 0.7346939
- R-square for testing with k-folds CV: 0.9031085

- R-square for training with repeated k-folds CV: 0.8351572
- R-square for testing with repeated k-folds CV: 0.9031085

Summary: Which one should be used based on RMSE value?

• RMSE for training set: 3.08648

• RMSE for testing test: 2.279303

• RMSE for training with LOOCV: 3.201673

RMSE for testing with LOOCV:
 2.244232

RMSE for training with k-folds CV:
 2.85133

• RMSE for testing with k-folds CV: 2.244232

 RMSE for training with repeated kfolds CV: 2.975392 RMSE for testing with repeated kfolds CV: 2.244232

Quick Think!

 Which model must be selected: Based on R-square or based on RMSE? If BLUE and LINE test is a must then which training model should be checked?

- Do we need to check the BLUE and LINE assumptions for the fit done with the training data?
- Validation set model?
- LOOCV set model?
- K-fold CV set model?
- Repeated K-fold CV set model?

Question/queries so far?

Multiple linear regression:

- It is an extension of the simple linear regression
- Multiple linear regression have more than one (two or more) independent variables
- Multiple linear regression has one

 (1) continuous dependent variable
 so it is a supervised learning

- All the assumptions of the simple linear regression are also applicable here
- There is one more condition:

 Multicollinearity must not be present i.e. correlations between independent variables must not be "high"

Multiple Linear Regression

A multiple linear regression model of Y on X_1 , X_{2_1} ... X_n in stochastic form in statistics is written as:

$$Y = \alpha + \beta_1 X_1 + \beta_2 X_2 + ... + \beta_n X_n + u$$

where α is called y-intercept and and β_1 , β_2 , ..., β_n are called slopes of X_1 , X_2 , ..., X_n respectively, and u is called error or disturbance term, which is <u>erratic or</u> random in nature.

Multiple Linear Regression Estimation

$$\hat{y} = a + b_1 x_1 + b_2 x_2 + ... + b_n x_n$$

• where a, b_1 , b_2 , ..., b_n are <u>least square estimates</u> of α and β_1 , β_2 , ..., β_n respectively.

• We need to calculate best solutions of n equations each containing unknown parameters α , β 1, β_2 , ..., β_n using Ordinary Least Square (OLS) method using statistical software R

Multiple linear regression process:

 We must fit bi-variate linear regression for all the predictors and can take ONLY the statistically significant predictors in the final model

 However, must must assess the MULTICOLLINEARITY of the predictor variables before taking them into the final model

 Multicollinearity can be assessed using Pearson's correlation coefficient but it is better to use Variance Inflation Factor (VIF)

Fit simple linear regression models of mpg with all the predictors of "mtcars" data

• What did you find?

Which predictors can be taken to the final model?

What to do now?

What will be your conclusion?

Assessing multicollinearity:

- Pearson correlation coefficients can be used
- Variance Inflation Factor (VIF) is most commonly used to assess multicollinearity
- We need to get a correlation matrix and flag the correlations with more than 0.75
- We can get the VIF for each independent variable

 These pair/s of independent variables influences the linear model coefficients Multicollinearity will be confirmed for an independent variable with VIF > 10 for linear models

Fitting multiple linear regression model using "mtcars" data: AFTER BIVARIATE MODELS!

- mlr <- lm(mpg ~., data = mtcars)
- summary(mlr)
- library(car)
- vif(mlr)
- We need to drop the independent variable with highest VIF and run the model again until all the VIF <10!

- None of the variables used in the model are statistically significant!
- > vif(mlr)

7.908747

cyl	disp	hp
15.373833	21.620241	9.832037
drat	wt	qsec
3.374620	15.164887	7.527958
VS	am	gear
4.965873	4.648487	5.357452
carb		

Fitting multiple linear regression using "mtcars" data:

#Removing "disp" variable:

• The "wt" variable is significant

mlr1 <- lm(mpg ~ cyl+hp+drat+wt+qsec+vs+am+gear+c arb, data = mtcars) summary(mlr1) vif(mlr1)

 We need to drop the independent variable with highest VIF and run the model again until all the VIF <10! • > vif(mlr1) hp drat cyl 14.284737 7.123361 3.329298 qsec wt VS 6.189050 6.914423 4.916053 carb gear am 4.645108 5.324402 4.310597

Fitting multiple linear regression using "mtcars" data:

#Removing "cyl" variable:

```
mlr2 <- lm(mpg ~
hp+drat+wt+qsec+vs+am+gear+carb,
data = mtcars)
summary(mlr1)
vif(mlr1)
```

- We need to drop the independent variable with highest VIF and run the model again until VIF <10!
- If all the VIF < 10 then we can interpret the model and do the predictions

• The "wt" variable is significant, b = -2.60968 (1 unit increase in wt reduces the mpg by 2.61 unit controlling for other independent variables)

• > vif(mlr2)

hp	drat	wt
6.015788	3.111501	6.051127
qsec	VS	am
5.918682	4.270956	4.285815
gear carb		
4.690187	4.290468	

Assignment: Self-Learning!

- Use the validation and crossvalidation methods for the multiple linear regression (mlr2) model
- Which model is the best model?
- Why?
- Predict the weight of the cars based on the best model identified using the test.data

- Change all the variables (except mpg) as standardized variable using "scale" command in R/R Studio
- Fit the multiple linear regression model with these standardized variables
- Does it solve the multicollinearity issue?
- Why? Write conclusions.

Alternative way to deal with multicollinearity in data science/machine learning:

 We can use the "regularization" methods

- The most common ones are:
 - Ridge regression
 - Lasso regression
 - Elastic net regression

 Once the "multicollinearity" problem is fixed then we can do the predictions and use the validation indices to select the best model for our data!

• I will post a YouTube video link and you can fit these models for the "mtcars" data and learn from them!

Regularized Regression resources:

• https://www.datacamp.com/tutorial/tutorial-ridge-lasso-elastic-net

https://daviddalpiaz.github.io/r4sl/regularization.html

• https://www.r-bloggers.com/2017/07/machine-learning-explained-regularization/

• https://medium.com/edureka/regularization-in-machine-learning-4e041bbbdae

Question/queries?

Next class

 Other regression models used in the supervised learning: polynomial regression, KNN algorithm etc.

Neural Network, Perceptron

MLP, neural net regression etc.

Thank you!

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