# **Titanic Survival Prediction**

```
library(data.table)
library(tidyverse)
library(dplyr)
library(stringr)
library(caret)
library(randomForest)
library(randomForest)
library(e1071)
```

```
train <- fread("train.csv")%>% data.table()
test <- fread("test.csv") %>% data.table()
test$Survived <- NA
combi = rbind(train, test)
ntrain <- nrow(train)</pre>
```

```
str(train)
```

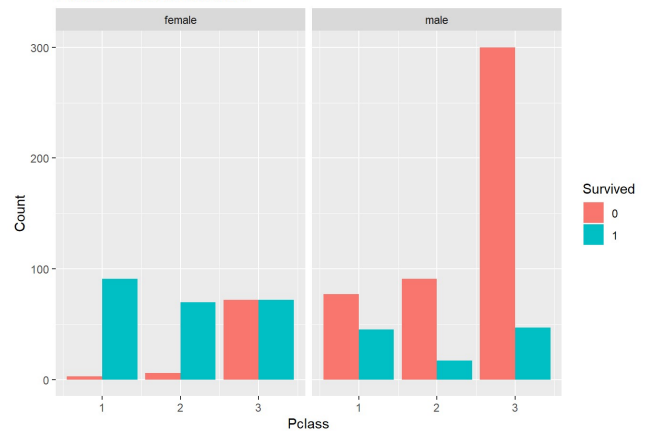
```
## Classes 'data.table' and 'data.frame': 891 obs. of 12 variables:
## $ PassengerId: int 1 2 3 4 5 6 7 8 9 10 ...
## $ Survived : int 0 1 1 1 0 0 0 0 1 1 ...
## $ Pclass
               : int 3 1 3 1 3 3 1 3 3 2 ...
               : chr "Braund, Mr. Owen Harris" "Cumings, Mrs. John Bradley (Florenc
## $ Name
e Briggs Thayer)" "Heikkinen, Miss. Laina" "Futrelle, Mrs. Jacques Heath (Lily May Pee
1)" ...
## $ Sex
               : chr "male" "female" "female" ...
## $ Age
               : num 22 38 26 35 35 NA 54 2 27 14 ...
## $ SibSp
              : int 1101000301...
## $ Parch
              : int 0000000120...
               : chr "A/5 21171" "PC 17599" "STON/02. 3101282" "113803" ...
## $ Ticket
## $ Fare
               : num 7.25 71.28 7.92 53.1 8.05 ...
              : chr "" "C85" "" "C123" ...
## $ Cabin
## $ Embarked : chr "S" "C" "S" "S" ...
## - attr(*, ".internal.selfref")=<externalptr>
```

```
str(test)
```

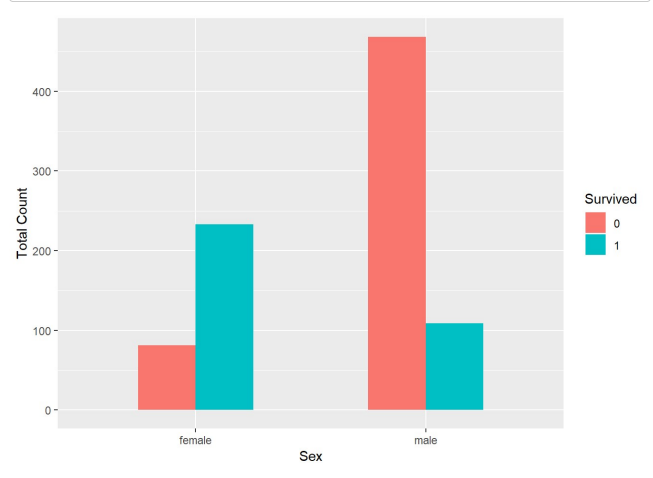
```
## Classes 'data.table' and 'data.frame': 418 obs. of 12 variables:
  $ PassengerId: int 892 893 894 895 896 897 898 899 900 901 ...
   $ Pclass
                : int 3 3 2 3 3 3 3 2 3 3 ...
                     "Kelly, Mr. James" "Wilkes, Mrs. James (Ellen Needs)" "Myles,
   $ Name
                : chr
Mr. Thomas Francis" "Wirz, Mr. Albert" ...
                : chr "male" "female" "male" "male" ...
   $ Sex
##
   $ Age
                : num 34.5 47 62 27 22 14 30 26 18 21 ...
##
   $ SibSp
                : int 0100100102...
##
   $ Parch
                : int 0000100100...
                : chr "330911" "363272" "240276" "315154" ...
##
   $ Ticket
   $ Fare
                : num 7.83 7 9.69 8.66 12.29 ...
                      ... ... ...
   $ Cabin
                : chr
##
                : chr "Q" "S" "Q" "S" ...
##
  $ Embarked
   $ Survived : logi NA NA NA NA NA NA ...
   - attr(*, ".internal.selfref")=<externalptr>
```

```
ggplot(combi[1:ntrain,], aes(x = factor(Pclass), fill = factor(Survived))) +
  geom_bar(width = 0.5, position="dodge") +
  xlab("Pclass") +
  ylab("Total Count") +
  labs(fill = "Survived")
```

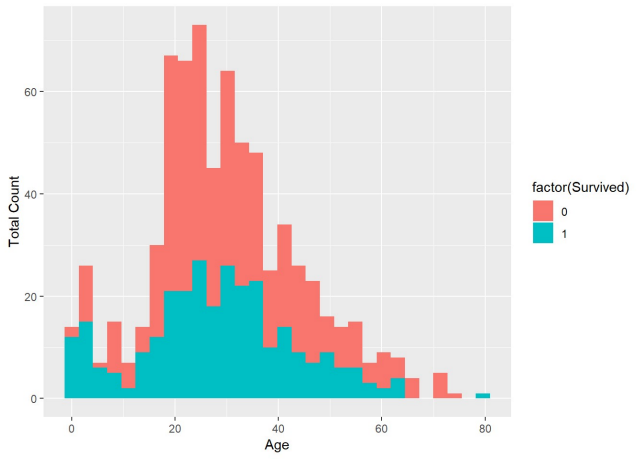
#### Pclass vs Sex vs Survived



```
ggplot(combi[1:ntrain,], aes(x = factor(Sex), fill = factor(Survived))) +
  geom_bar(width = 0.5, position="dodge") +
  xlab("Sex") +
  ylab("Total Count") +
  labs(fill = "Survived")
```

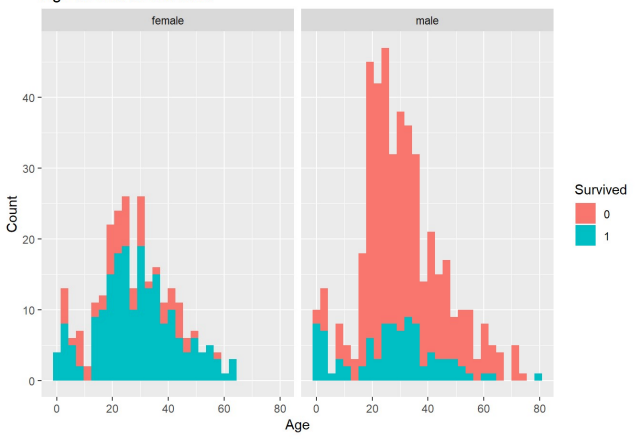


```
ggplot(subset(combi[1:ntrain,],!is.na(Age)), aes(x = Age, fill = factor(Survived))) +
  geom_histogram(bins = 30) +
  xlab("Age") +
  ylab("Total Count")
```



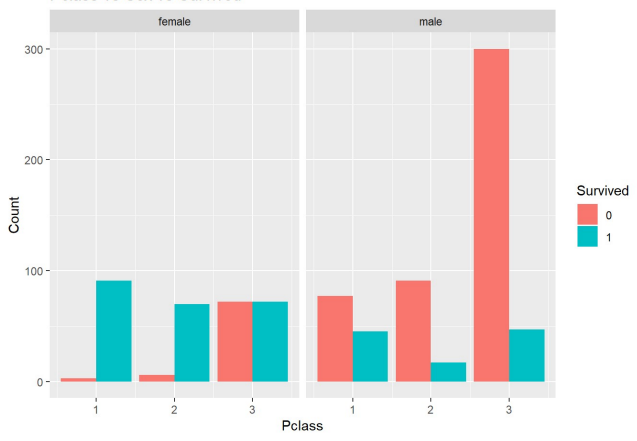
```
ggplot(subset(combi[1:ntrain,],!is.na(Age)), aes(Age, fill = factor(Survived))) +
  geom_histogram(bins=30) +
  xlab("Age") +
  ylab("Count") +
  facet_grid(.~Sex)+
  scale_fill_discrete(name = "Survived") +
  ggtitle("Age vs Sex vs Survived")
```

# Age vs Sex vs Survived



```
ggplot(combi[1:ntrain,], aes(Pclass, fill = factor(Survived))) +
  geom_bar(stat = "count",position = "dodge")+
  xlab("Pclass") +
  facet_grid(.~Sex)+
  ylab("Count") +
  scale_fill_discrete(name = "Survived") +
  ggtitle("Pclass vs Sex vs Survived")
```

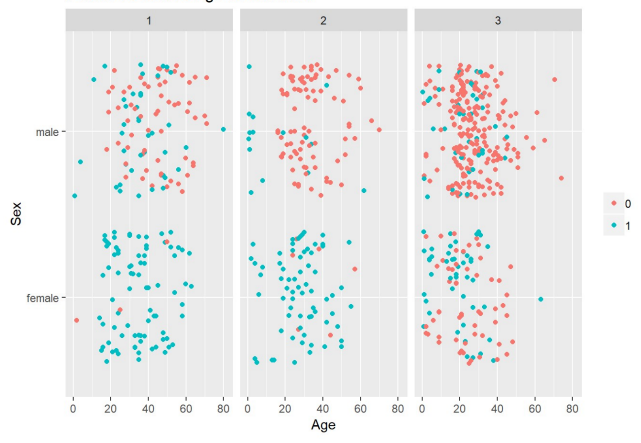
#### Pclass vs Sex vs Survived



```
ggplot(combi[1:ntrain,], aes(x = Age, y = Sex)) +
  geom_jitter(aes(colour = factor(Survived))) +
  theme(legend.title = element_blank())+
  facet_wrap(~Pclass) +
  labs(x = "Age", y = "Sex", title = "Pclass vs Sex vs Age vs Survived")+
  scale_fill_discrete(name = "Survived") +
  scale_x_continuous(name="Age",limits=c(0, 81))
```

## Warning: Removed 177 rows containing missing values (geom\_point).

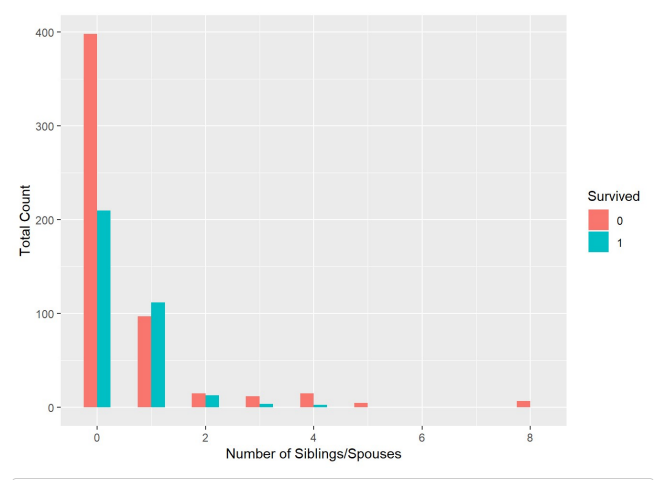
# Pclass vs Sex vs Age vs Survived



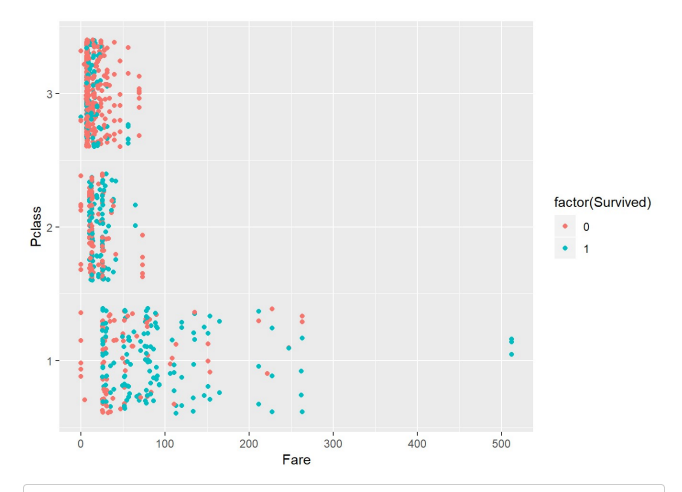
```
table(combi[1:ntrain,]$SibSp)
```

```
##
## 0 1 2 3 4 5 8
## 608 209 28 16 18 5 7
```

```
ggplot(combi[1:ntrain,], aes(x = SibSp, fill = factor(Survived))) +
  geom_histogram(binwidth=0.5, position="dodge") +
  xlab("Number of Siblings/Spouses") +
  ylab("Total Count") +
  labs(fill = "Survived")
```



```
ggplot(combi[1:ntrain,], aes(x=Fare , y = Pclass)) +
  geom_jitter(aes(color = factor(Survived)))
```

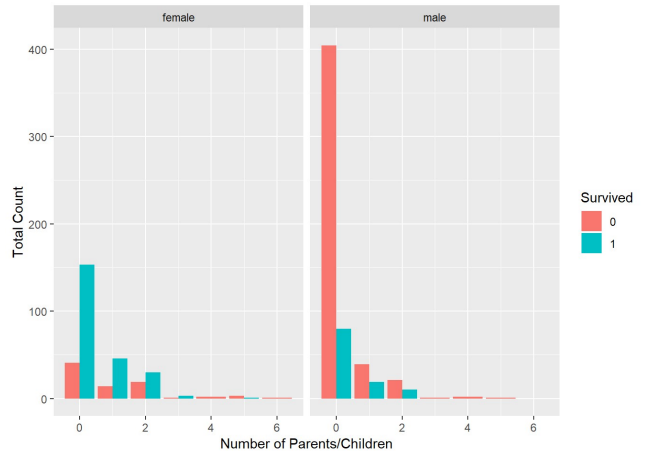


### table(combi[1:ntrain,]\$Parch)

```
##
## 0 1 2 3 4 5 6
## 678 118 80 5 4 5 1
```

```
ggplot(combi[1:ntrain,], aes(x = Parch, fill = factor(Survived))) +
  geom_histogram(binwidth=0.5, position="dodge", stat = "count") +
  facet_grid(.~Sex)+
  xlab("Number of Parents/Children") +
  ylab("Total Count") +
  labs(fill = "Survived")
```

## Warning: Ignoring unknown parameters: binwidth, bins, pad



finding percentage of missing values in each column

```
sapply(combi, function(x) {ifelse(sum(is.na(x))!=0 , round(sum(is.na(x))*100/nrow(combi),2) , round(sum(x=="")*100/nrow(combi),2))})
```

#	## Pas	ssengerId	Survived	Pclass	Name	Sex	Age
#	##	0.00	31.93	0.00	0.00	0.00	20.09
#	##	SibSp	Parch	Ticket	Fare	Cabin	Embarked
4	##	0.00	0.00	0.00	0.08	77.46	0.15

Ignoring attribute Cabin since 77% of data is missing, replacing NA in Embarked to S

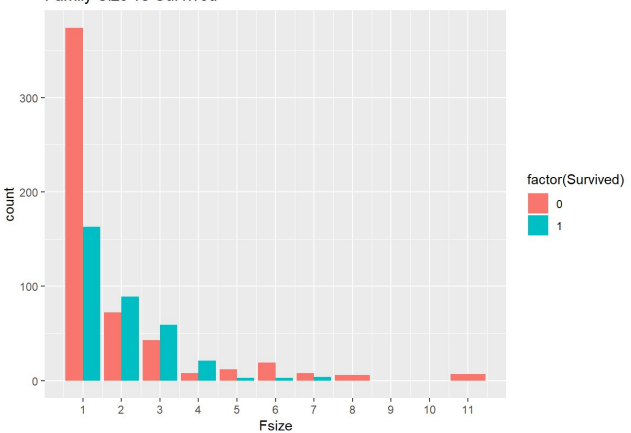
```
table(combi$Embarked)
```

```
##
## C Q S
## 2 270 123 914
```

```
combi$Embarked[combi$Embarked ==""] <- 'S'
combi$Embarked[combi$Embarked ==""] <- 'S'
combi$Fsize <- combi$SibSp + combi$Parch + 1</pre>
```

```
ggplot(combi[1:ntrain,] , aes(x=Fsize , fill = factor(Survived)))+geom_bar(stat = "cou
nt" , position = "dodge")+
    scale_x_continuous(breaks = seq(0,max(combi[1:ntrain,]$Fsize),1)) +
    ggtitle("Family Size vs Survived")
```

## Family Size vs Survived

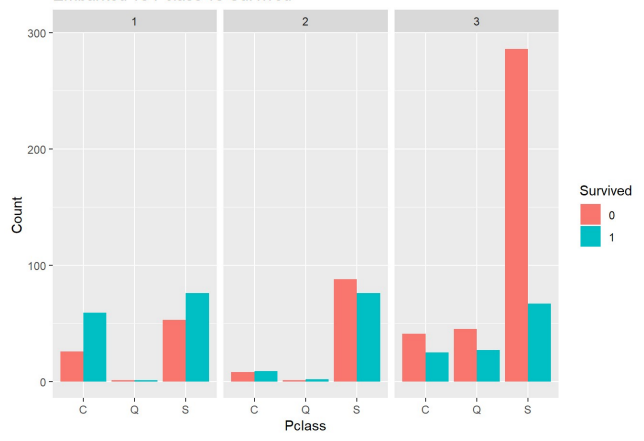


#### table(combi\$Pclass)

```
##
## 1 2 3
## 323 277 709
```

```
ggplot(combi[1:ntrain,], aes(Embarked, fill = factor(Survived))) +
  geom_bar(stat = "count",position = "dodge")+
  xlab("Pclass") +
  ylab("Count") +
  facet_grid(.~Pclass) +
  scale_fill_discrete(name = "Survived") +
  ggtitle("Embarked vs Pclass vs Survived")
```

#### Embarked vs Pclass vs Survived



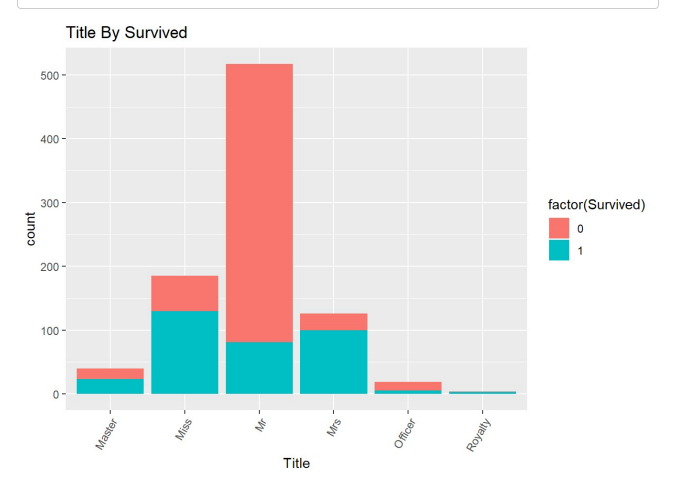
```
combi$Title <- NA
combi$Title <- sapply(combi$Name , function(x) str_trim(str_split(x,"[,.]")[[1]][2],si
de ="both"))
unique(combi$Title)</pre>
```

```
"Mrs"
                                        "Miss"
##
   [1] "Mr"
                                                        "Master"
                                        "Dr"
                        "Rev"
                                                        "Mme"
##
   [5] "Don"
   [9] "Ms"
                                                        "Sir"
                        "Major"
                                        "Lady"
                        "Col"
                                        "Capt"
                                                        "the Countess"
## [13] "Mlle"
## [17] "Jonkheer"
                        "Dona"
```

```
combi$Title[combi$Title%in%c("Mme")] <- "Mrs"
combi$Title[combi$Title%in%c("Mlle","Ms")] <- "Miss"
officer <- c('Capt', 'Col', 'Don', 'Dr', 'Major', 'Rev')
royalty <- c('Dona', 'Lady', 'the Countess', 'Sir', 'Jonkheer')
combi$Title[combi$Title %in% royalty] <- 'Royalty'
combi$Title[combi$Title %in% officer] <- 'Officer'</pre>
```

```
ggplot(combi[1:ntrain,] , aes(x=Title , fill = factor(Survived))) +
  geom_histogram(bins = 6 , stat = "count") +
  ggtitle("Title By Survived")+
  theme(axis.text.x=element_text(angle=60, hjust=1))
```

```
## Warning: Ignoring unknown parameters: binwidth, bins, pad
```



imputing missing age by predicting the age based on variables Pclass, Sex, title, SibSp, Parch, fare, Title

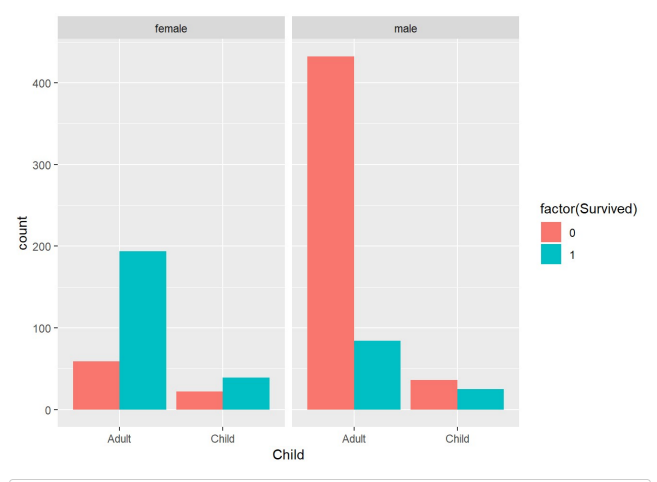
```
## Call:
## rpart(formula = Age ~ Pclass + Sex + Embarked + SibSp + Parch +
       Fsize + Fare + Title, data = combi[!is.na(combi$Age), ],
       method = "anova")
##
##
     n = 1046
##
##
             CP nsplit rel error
                                     xerror
                                                  xstd
                     0 1.0000000 1.0021794 0.04531621
## 1 0.21028409
## 2 0.10512853
                     1 0.7897159 0.7921058 0.03520690
## 3 0.09537135
                     2 0.6845874 0.7502329 0.03593644
## 4 0.01436395
                     3 0.5892160 0.5953120 0.03049945
## 5 0.01266967
                     4 0.5748521 0.6028027 0.03090432
## 6 0.01056208
                     5 0.5621824 0.5919662 0.03086995
## 7 0.01000000
                     6 0.5516203 0.5884412 0.03089030
##
## Variable importance
##
      Title
                                                                Sex Embarked
                Fare
                       Pclass
                                  Parch
                                           Fsize
                                                     SibSp
                                                                  8
##
         28
                  16
                            16
                                     11
                                              11
                                                         8
                                                                           3
##
## Node number 1: 1046 observations,
                                         complexity param=0.2102841
     mean=29.88114, MSE=207.5502
##
##
     left son=2 (266 obs) right son=3 (780 obs)
##
     Primary splits:
##
         Title splits as LLRRRR,
                                          improve=0.21028410, (0 missing)
                            to the right, improve=0.15460490, (0 missing)
##
         Pclass < 1.5
                            to the right, improve=0.07107333, (0 missing)
##
         SibSp < 2.5
##
                < 49.5021 to the left, improve=0.05839866, (1 missing)
         Fare
                            to the right, improve=0.05804572, (0 missing)
##
         Fsize < 2.5
##
     Surrogate splits:
                                            agree=0.782, adj=0.143, (0 split)
##
         Sex
                  splits as LR,
##
         SibSp
                  < 2.5
                              to the right, agree=0.773, adj=0.109, (0 split)
                              to the right, agree=0.762, adj=0.064, (0 split)
##
         Fsize
                  < 4.5
                  < 1.5
                              to the right, agree=0.751, adj=0.023, (0 split)
##
         Parch
##
         Embarked splits as
                             RLR,
                                            agree=0.748, adj=0.008, (0 split)
##
## Node number 2: 266 observations,
                                        complexity param=0.09537135
##
     mean=18.56831, MSE=164.0627
##
     left son=4 (128 obs) right son=5 (138 obs)
     Primary splits:
##
##
         Parch < 0.5
                           to the right, improve=0.4744399, (0 missing)
##
         Fsize < 2.5
                           to the right, improve=0.3884753, (0 missing)
##
         Sex
               splits as
                          RL,
                                         improve=0.2597037, (0 missing)
                                         improve=0.2597037, (0 missing)
         Title splits as
##
                          LR----,
         SibSp < 0.5
                          to the right, improve=0.2127207, (0 missing)
##
##
     Surrogate splits:
         Fsize < 1.5
##
                          to the right, agree=0.932, adj=0.859, (0 split)
##
         SibSp < 0.5
                          to the right, agree=0.786, adj=0.555, (0 split)
##
         Fare < 13.20835 to the right, agree=0.744, adj=0.469, (0 split)
```

```
##
         Sex
               splits as RL,
                                        agree=0.711, adj=0.398, (0 split)
##
         Title splits as LR----,
                                        agree=0.711, adj=0.398, (0 split)
##
## Node number 3: 780 observations,
                                        complexity param=0.1051285
##
     mean=33.7391, MSE=163.8521
##
     left son=6 (562 obs) right son=7 (218 obs)
##
     Primary splits:
         Pclass < 1.5
                           to the right, improve=0.17857830, (0 missing)
##
##
                < 24.86875 to the left,
                                         improve=0.13383420, (1 missing)
         Fare
##
         Title splits as
                           --LRRR,
                                          improve=0.03939711, (0 missing)
                splits as RL,
                                          improve=0.01923511, (0 missing)
##
         Sex
##
         Parch < 0.5
                           to the left, improve=0.01239742, (0 missing)
##
     Surrogate splits:
                  < 26.26875 to the left, agree=0.908, adj=0.670, (0 split)
##
         Fare
##
         Embarked splits as RLL,
                                            agree=0.765, adj=0.161, (0 split)
##
         Title
                  splits as --LLRR,
                                            agree=0.731, adj=0.037, (0 split)
##
## Node number 4: 128 observations,
                                       complexity param=0.01266967
##
     mean=9.407578, MSE=73.70615
##
     left son=8 (103 obs) right son=9 (25 obs)
##
     Primary splits:
##
         Fare
               < 48.2
                           to the left, improve=0.2915456, (0 missing)
##
         Pclass < 1.5
                           to the right, improve=0.2562854, (0 missing)
                                          improve=0.1522839, (0 missing)
##
         Sex
                splits as RL,
                                          improve=0.1522839, (0 missing)
##
         Title splits as
                           LR----,
         SibSp < 0.5
                           to the right, improve=0.0349171, (0 missing)
##
##
     Surrogate splits:
                           to the right, agree=0.961, adj=0.8, (0 split)
##
         Pclass < 1.5
##
## Node number 5: 138 observations,
                                       complexity param=0.01436395
     mean=27.06522, MSE=97.83633
##
##
     left son=10 (69 obs) right son=11 (69 obs)
     Primary splits:
##
##
         Pclass
                  < 2.5
                             to the right, improve=0.2309667000, (0 missing)
         Fare
                  < 26.275
                             to the left, improve=0.1953148000, (0 missing)
##
##
         Embarked splits as
                             RLL,
                                            improve=0.0374038300, (0 missing)
                  < 0.5
                             to the right, improve=0.0008847728, (0 missing)
##
         SibSp
##
         Fsize
                             to the right, improve=0.0008847728, (0 missing)
                  < 1.5
##
     Surrogate splits:
##
         Fare
                  < 10.17085 to the left, agree=0.935, adj=0.870, (0 split)
##
         Embarked splits as RLL,
                                            agree=0.645, adj=0.290, (0 split)
                             to the right, agree=0.514, adj=0.029, (0 split)
##
         SibSp
                  < 0.5
         Fsize
                             to the right, agree=0.514, adj=0.029, (0 split)
##
                  < 1.5
##
         Sex
                  splits as RL,
                                            agree=0.507, adj=0.014, (0 split)
##
## Node number 6: 562 observations,
                                        complexity param=0.01056208
     mean=30.37011, MSE=116.7829
##
##
     left son=12 (361 obs) right son=13 (201 obs)
##
     Primary splits:
```

```
##
         Pclass < 2.5
                             to the right, improve=0.03493722, (0 missing)
##
         Fare
                  < 9.54375 to the left, improve=0.03140571, (1 missing)
                             --LRR-,
                                           improve=0.02300209, (0 missing)
##
         Title
                  splits as
##
         Embarked splits as LRL,
                                           improve=0.01586441, (0 missing)
##
         Parch
                  < 3.5
                             to the left, improve=0.01382681, (0 missing)
##
     Surrogate splits:
         Fare < 10.48125 to the left, agree=0.835, adj=0.537, (0 split)
##
         Title splits as --LRR-,
                                        agree=0.669, adj=0.075, (0 split)
##
##
               splits as RL,
                                        agree=0.651, adj=0.025, (0 split)
         Sex
##
## Node number 7: 218 observations
     mean=42.42431, MSE=180.5023
##
##
## Node number 8: 103 observations
##
     mean=7.123786, MSE=43.27704
##
## Node number 9: 25 observations
##
     mean=18.8168, MSE=89.05189
##
## Node number 10: 69 observations
    mean=22.31159, MSE=42.00074
##
##
## Node number 11: 69 observations
    mean=31.81884, MSE=108.4781
##
##
## Node number 12: 361 observations
##
    mean=28.86288, MSE=100.2727
##
## Node number 13: 201 observations
    mean=33.07711, MSE=135.0276
##
```

```
combi$Age[is.na(combi$Age)] <- predict(agefit , combi[is.na(combi$Age),])
## child or adult based on age
combi$Child[combi$Age < 18] <- 'Child'
combi$Child[combi$Age >= 18] <- 'Adult'</pre>
```

```
ggplot(data = combi[1:ntrain,] , aes(x=Child , fill = factor(Survived))) +
  geom_bar(stat = "count", position = "dodge") + facet_grid(.~Sex)
```



```
combi$Pclass <- factor(combi$Pclass)
combi$Sex <- as.integer(combi$Sex=="male")
combi$Child <- as.integer(combi$Child=="Child")
combi$Embarked <- factor(combi$Embarked)
combi$Title <- as.factor(combi$Title)

sapply(combi, function(x) {ifelse(sum(is.na(x))!=0 , round(sum(is.na(x))*100/nrow(combi),2) , round(sum(x=="")*100/nrow(combi),2))})</pre>
```

##	PassengerId	Survived	Pclass	Name	Sex	Age
##	0.00	31.93	0.00	0.00	0.00	0.00
##	SibSp	Parch	Ticket	Fare	Cabin	Embarked
##	0.00	0.00	0.00	0.08	77.46	0.00
##	Fsize	Title	Child			
##	0.00	0.00	0.00			

```
#creating indices
trainIndex <- createDataPartition(combi[1:ntrain,]$Survived,p=1,list=FALSE)

#splitting data into training/testing data using the trainIndex object
train_titanic <- combi[trainIndex,]
test_titanic <- combi[-trainIndex,]

#creating indices to split train into train and validation
Index2 <- createDataPartition(train_titanic$Survived,p=0.8,list=FALSE)
train <- train_titanic[Index2,]
validation <- train_titanic[-Index2,]</pre>
```

## Logistic Regression

```
model_glm <- glm(Survived ~ Pclass+Sex+Fsize+Child+Fare+Embarked+Title ,data = train,
family = binomial(link = "logit") )
summary(model_glm)</pre>
```

```
##
## Call:
## glm(formula = Survived ~ Pclass + Sex + Fsize + Child + Fare +
      Embarked + Title, family = binomial(link = "logit"), data = train)
##
##
## Deviance Residuals:
      Min
               10
                   Median
                               3Q
                                      Max
## -2.1463 -0.5656 -0.3809 0.5329
                                  2.6420
##
## Coefficients:
##
                Estimate Std. Error z value Pr(>|z|)
## (Intercept)
               19.153678 500.065168 0.038
                                           0.9694
## Pclass2
               -1.100031
                         0.350589 -3.138
                                           0.0017 **
## Pclass3
               -2.145112
                         0.344665 -6.224 4.85e-10 ***
## Sex
              -14.758669 500.064477 -0.030
                                          0.9765
## Fsize
              ## Child
               0.488847
                         0.432319 1.131
                                          0.2582
               0.003918 0.003286 1.192
## Fare
                                          0.2332
## EmbarkedQ
              -0.212344 0.478981 -0.443 0.6575
## EmbarkedS
              -0.358904
                         0.292326 -1.228
                                          0.2195
## TitleMiss -15.530429 500.064856 -0.031
                                           0.9752
## TitleMr
              ## TitleMrs
              -15.032857 500.064975 -0.030
                                           0.9760
## TitleOfficer -4.292622 0.923427 -4.649 3.34e-06 ***
## TitleRoyalty -3.517950 1.590117 -2.212
                                          0.0269 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 939.87 on 712 degrees of freedom
## Residual deviance: 568.60 on 699 degrees of freedom
## AIC: 596.6
## Number of Fisher Scoring iterations: 13
```

```
# validation
predglm <- predict(model_glm, validation , type ="response" )
logit_survived = as.numeric(predglm >= 0.5)
table(logit_survived)
```

```
## logit_survived
## 0 1
## 105 73
```

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction 0 1
            0 86 14
##
            1 19 59
##
##
##
                  Accuracy : 0.8146
                    95% CI: (0.7496, 0.8688)
##
##
       No Information Rate: 0.5899
       P-Value [Acc > NIR] : 1.274e-10
##
##
##
                     Kappa: 0.6208
   Mcnemar's Test P-Value: 0.4862
##
##
##
               Sensitivity: 0.8190
               Specificity: 0.8082
##
            Pos Pred Value: 0.8600
##
##
            Neg Pred Value: 0.7564
##
                Prevalence: 0.5899
            Detection Rate: 0.4831
##
##
      Detection Prevalence : 0.5618
##
         Balanced Accuracy: 0.8136
##
##
          'Positive' Class : 0
```

```
# predicting Test
test_glm <- predict(model_glm, test_titanic , type ="response" )
print(RMSE(validation$Survived,logit_survived))</pre>
```

```
## [1] 0.4305732
```

#### Random Forest

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction 0 1
##
            0 91 9
            1 24 54
##
##
##
                  Accuracy : 0.8146
##
                    95% CI: (0.7496, 0.8688)
##
       No Information Rate: 0.6461
##
       P-Value [Acc > NIR] : 6.012e-07
##
                     Kappa : 0.6153
##
   Mcnemar's Test P-Value: 0.01481
##
##
##
               Sensitivity: 0.7913
               Specificity: 0.8571
##
            Pos Pred Value: 0.9100
##
##
            Neg Pred Value: 0.6923
                Prevalence: 0.6461
##
##
            Detection Rate : 0.5112
      Detection Prevalence: 0.5618
##
##
         Balanced Accuracy: 0.8242
##
##
          'Positive' Class: 0
##
```

```
print(RMSE(validation$Survived,rf_pred))
```

#### SVM

```
## SVM
# Set up the 5-fold CV
fitControl <- caret::trainControl(method = "repeatedcv",</pre>
                                   number = 5,
                                   repeats = 5)
# Define ranges for the two parameters
              sapply(seq(-1,3,0.0125), function(x) 10^x)
sigma_range = sapply(seq(-3,1,0.0125), function(x) 10^x)
# Create the grid of parameters
fitGrid <- expand.grid(C= C_range,</pre>
                       sigma = sigma_range)
Rsvm <- caret::train(Survived ~ Pclass+Sex+Fsize+Child+Fare+Embarked+Title, data=trai
n,
                     method = "svmLinear",
                     trControl = fitControl,
                      preProcess = c("center", "scale"))
```

```
## Warning in train.default(x, y, weights = w, ...): You are trying to do
## regression and your outcome only has two possible values Are you trying to
## do classification? If so, use a 2 level factor as your outcome column.
```

```
svm_pred <- predict(Rsvm , validation )
svm_pred = as.numeric(svm_pred >= 0.5)
confusionMatrix(as.factor(validation$Survived) ,as.factor(svm_pred))
```

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction 0 1
##
            0 80 20
##
            1 17 61
##
##
                  Accuracy : 0.7921
##
                    95% CI : (0.7251, 0.8492)
       No Information Rate: 0.5449
##
       P-Value [Acc > NIR] : 5.144e-12
##
##
##
                     Kappa: 0.5796
   Mcnemar's Test P-Value : 0.7423
##
##
##
               Sensitivity: 0.8247
               Specificity: 0.7531
##
            Pos Pred Value : 0.8000
##
##
            Neg Pred Value : 0.7821
##
                Prevalence: 0.5449
##
            Detection Rate: 0.4494
##
      Detection Prevalence: 0.5618
##
         Balanced Accuracy: 0.7889
##
          'Positive' Class : 0
##
##
```

```
print(RMSE(validation$Survived,svm_pred))
```

```
## [1] 0.4559223
```

## **Gradient Boosting**

```
#gradient boosting
fitControl <- trainControl(method = 'repeatedcv',</pre>
                            number = 5,
                            repeats = 5)
# for caret, there are only four tuning parameters below.
# tune n.trees
newGrid <- expand.grid(n.trees = c(50, 100, 200, 300),</pre>
                        interaction.depth = c(6),
                        shrinkage = 0.01,
                        n.minobsinnode = 10
)
fit_gbm <- train(Survived ~ Pclass+Sex+Fsize+Child+Fare+Embarked+Title, data=train,</pre>
                 method = 'gbm',
                 trControl = fitControl,
                 tuneGrid = newGrid,
                 bag.fraction = 0.5,
                 verbose = FALSE,
                 preProcess = c("center", "scale"))
fit_gbm$bestTune
```

```
## n.trees interaction.depth shrinkage n.minobsinnode
## 4 300 6 0.01 10
```

```
gbm_pred <- predict(fit_gbm , validation )
gbm_pred = as.numeric(gbm_pred >= 0.5)
confusionMatrix(as.factor(validation$Survived) ,as.factor(gbm_pred))
```

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction 0 1
##
            0 95 5
            1 25 53
##
##
##
                  Accuracy: 0.8315
##
                    95% CI : (0.7682, 0.8833)
       No Information Rate : 0.6742
##
       P-Value [Acc > NIR] : 1.724e-06
##
##
##
                     Kappa : 0.6478
   Mcnemar's Test P-Value : 0.0005226
##
##
##
               Sensitivity: 0.7917
               Specificity: 0.9138
##
            Pos Pred Value: 0.9500
##
##
            Neg Pred Value : 0.6795
##
                Prevalence: 0.6742
            Detection Rate: 0.5337
##
##
      Detection Prevalence : 0.5618
##
         Balanced Accuracy: 0.8527
##
          'Positive' Class : 0
##
##
```

```
print(RMSE(validation$Survived,gbm_pred))
```

```
## [1] 0.4105354
```