# GPH 2338 Project

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## Data Upload

```
library(haven)
library(psych)
library(caret)
library(tidyverse)
library(ggplot2)
library(psych)
library(pander)
library(corrplot)
library(pander)
library(readr)
library(r02pro)
library(plyr)
library(tree)
library(gbm)
library(caret)
library(leaps)
library(readr)
```

### Data Preparation

```
df<-readr::read_tsv("brca_metabric_clinical_data.tsv")</pre>
```

```
## Rows: 2509 Columns: 39
## -- Column specification ------
## Delimiter: "\t"
## chr (27): Study ID, Patient ID, Sample ID, Type of Breast Surgery, Cancer Ty...
## dbl (12): Age at Diagnosis, Cohort, Neoplasm Histologic Grade, Lymph nodes e...
##
## i Use 'spec()' to retrieve the full column specification for this data.
## i Specify the column types or set 'show_col_types = FALSE' to quiet this message.
```

#### colnames(df)

```
## [1] "Study ID" "Patient ID"
## [3] "Sample ID" "Age at Diagnosis"
## [5] "Type of Breast Surgery" "Cancer Type"
## [7] "Cancer Type Detailed" "Cellularity"
## [9] "Chemotherapy" "Pam50 + Claudin-low subtype"
```

```
## [11] "Cohort"
                                          "ER status measured by IHC"
## [13] "ER Status"
                                         "Neoplasm Histologic Grade"
                                         "HER2 Status"
## [15] "HER2 status measured by SNP6"
## [17] "Tumor Other Histologic Subtype"
                                         "Hormone Therapy"
## [19] "Inferred Menopausal State"
                                         "Integrative Cluster"
## [21] "Primary Tumor Laterality"
                                         "Lymph nodes examined positive"
## [23] "Mutation Count"
                                          "Nottingham prognostic index"
                                          "Overall Survival (Months)"
## [25] "Oncotree Code"
## [27] "Overall Survival Status"
                                         "PR Status"
                                         "Relapse Free Status (Months)"
## [29] "Radio Therapy"
## [31] "Relapse Free Status"
                                         "Number of Samples Per Patient"
## [33] "Sample Type"
## [35] "3-Gene classifier subtype"
                                         "TMB (nonsynonymous)"
## [37] "Tumor Size"
                                          "Tumor Stage"
## [39] "Patient's Vital Status"
```

## colSums(is.na(df))

	g. 1 TD	D TD
##	Study ID	Patient ID
##	0	0
##	Sample ID	Age at Diagnosis
##	0	11
##	Type of Breast Surgery	Cancer Type
##	554	0
##	Cancer Type Detailed	Cellularity
##	0	592
##	Chemotherapy	Pam50 + Claudin-low subtype
##	529	529
##	Cohort	ER status measured by IHC
##	11	83
##	ER Status	Neoplasm Histologic Grade
##	40	121
##	HER2 status measured by SNP6	HER2 Status
##	529	529
##	Tumor Other Histologic Subtype	Hormone Therapy
##	135	529
##	Inferred Menopausal State	Integrative Cluster
##	529	529
##	Primary Tumor Laterality	Lymph nodes examined positive
##	639	266
##	Mutation Count	Nottingham prognostic index
##	151	222
##	Oncotree Code	Overall Survival (Months)
##	0	528
##	Overall Survival Status	PR Status
##	528	529
##	Radio Therapy	Relapse Free Status (Months)
##	529	121
##	Relapse Free Status	Number of Samples Per Patient
##	21	0
##	Sample Type	Sex
##	0	0
##	3-Gene classifier subtype	TMB (nonsynonymous)
##	745	0

```
##
                       Tumor Size
                                                      Tumor Stage
##
                               149
                                                               721
##
           Patient's Vital Status
##
                               529
print(df)
  # A tibble: 2,509 x 39
##
      'Study ID'
                    Patie~1 Sampl~2 Age a~3 Type ~4 Cance~5 Cance~6 Cellu~7 Chemo~8
##
                             <chr>
                                                                              <chr>
      <chr>
                    <chr>
                                       <dbl> <chr>
                                                     <chr>
                                                              <chr>>
    1 brca_metabric MB-0000 MB-0000
##
                                        75.6 MASTEC~ Breast~ Sreast~ <NA>
                                                                              NO
    2 brca_metabric MB-0002 MB-0002
                                        43.2 BREAST~ Breast~ Breast~ High
                                                                              NO
##
    3 brca_metabric MB-0005 MB-0005
                                        48.9 MASTEC~ Breast~ Breast~ High
                                                                              YES
##
    4 brca_metabric MB-0006 MB-0006
                                        47.7 MASTEC~ Breast~ Breast~ Modera~ YES
    5 brca_metabric MB-0008 MB-0008
                                        77.0 MASTEC~ Breast~ Breast~ High
                                                                              YES
    6 brca_metabric MB-0010 MB-0010
                                        78.8 MASTEC~ Breast~ Breast~ Modera~ NO
    7 brca_metabric MB-0014 MB-0014
                                        56.4 BREAST~ Breast~ Breast~ Modera~ YES
    8 brca_metabric MB-0020 MB-0020
                                        70
                                             MASTEC~ Breast~ Breast~ High
    9 brca_metabric MB-0022 MB-0022
                                        89.1 BREAST~ Breast~ Breast~ Modera~ NO
## 10 brca_metabric MB-0025 MB-0025
                                                     Breast~ Breast~ <NA>
                                                                              <NA>
                                        76.2 <NA>
     ... with 2,499 more rows, 30 more variables:
       'Pam50 + Claudin-low subtype' <chr>, Cohort <dbl>,
##
       'ER status measured by IHC' <chr>, 'ER Status' <chr>,
       'Neoplasm Histologic Grade' <dbl>, 'HER2 status measured by SNP6' <chr>,
## #
## #
       'HER2 Status' <chr>, 'Tumor Other Histologic Subtype' <chr>,
       'Hormone Therapy' <chr>, 'Inferred Menopausal State' <chr>,
## #
## #
       'Integrative Cluster' <chr>, 'Primary Tumor Laterality' <chr>, ...
df1 <- df %>% na.omit()
colSums(is.na(df1))
                         Study ID
                                                       Patient ID
                         Sample ID
                                                 Age at Diagnosis
```

```
##
##
##
##
           Type of Breast Surgery
##
                                                         Cancer Type
##
                                  0
                                                                    0
##
              Cancer Type Detailed
                                                         Cellularity
##
                                  0
##
                      Chemotherapy
                                        Pam50 + Claudin-low subtype
##
                                  0
                                                                    0
##
                             Cohort
                                          ER status measured by IHC
##
##
                          ER Status
                                          Neoplasm Histologic Grade
##
##
     HER2 status measured by SNP6
                                                         HER2 Status
##
                                                                    0
   Tumor Other Histologic Subtype
                                                     Hormone Therapy
##
                                  0
                                                                    0
##
        Inferred Menopausal State
                                                Integrative Cluster
##
                                  0
##
                                     Lymph nodes examined positive
         Primary Tumor Laterality
##
                                  0
```

```
##
                    Mutation Count
                                      Nottingham prognostic index
##
##
                     Oncotree Code
                                        Overall Survival (Months)
##
##
          Overall Survival Status
                                                         PR Status
##
                                                                  0
##
                     Radio Therapy
                                     Relapse Free Status (Months)
##
##
              Relapse Free Status
                                    Number of Samples Per Patient
##
                                 0
                                                                  0
##
                       Sample Type
                                                                Sex
##
                                                                  0
                                               TMB (nonsynonymous)
##
        3-Gene classifier subtype
##
##
                        Tumor Size
                                                       Tumor Stage
##
                                                                  0
##
           Patient's Vital Status
##
df2<-df1[,-c(1, 2, 3, 6)] # remove StudyID, Patient ID, Sample ID, Cancer Type
print(df2)
## # A tibble: 1,092 x 35
      Age at Diagn~1 Type ~2 Cance~3 Cellu~4 Chemo~5 Pam50~6 Cohort ER st~7 ER St~8
##
##
                                               <chr>>
                                                       <chr>>
                                                                 <dbl> <chr>
               <dbl> <chr>
                              <chr>
                                      <chr>>
##
   1
                43.2 BREAST~ Breast~ High
                                               NO
                                                       LumA
                                                                     1 Positve Positi~
##
                77.0 MASTEC~ Breast~ High
                                               YES
                                                       LumB
                                                                     1 Positve Positi~
##
                78.8 MASTEC~ Breast~ Modera~ NO
                                                       LumB
                                                                     1 Positve Positi~
##
   4
                86.4 BREAST~ Breast~ Modera~ NO
                                                       LumB
                                                                     1 Positve Positi~
                84.2 MASTEC~ Breast~ High
                                                       Her2
                                                                     1 Negati~ Positi~
##
    6
                85.5 MASTEC~ Breast~ Modera~ NO
                                                       LumA
                                                                     1 Positve Positi~
##
    7
                45.4 BREAST~ Breast~ High
                                               YES
                                                       LumB
                                                                     1 Positve Positi~
##
    8
                61.5 BREAST~ Breast~ High
                                               NO
                                                       LumB
                                                                     1 Positve Positi~
##
                68.7 MASTEC~ Breast~ Low
                                               YES
                                                       Basal
                                                                     1 Negati~ Negati~
                46.9 MASTEC~ Breast~ Modera~ NO
## 10
                                                       Normal
                                                                     1 Positve Positi~
     ... with 1,082 more rows, 26 more variables:
       'Neoplasm Histologic Grade' <dbl>, 'HER2 status measured by SNP6' <chr>,
## #
       'HER2 Status' <chr>, 'Tumor Other Histologic Subtype' <chr>,
## #
## #
       'Hormone Therapy' <chr>, 'Inferred Menopausal State' <chr>,
## #
       'Integrative Cluster' <chr>, 'Primary Tumor Laterality' <chr>,
       'Lymph nodes examined positive' <dbl>, 'Mutation Count' <dbl>,
## #
       'Nottingham prognostic index' <dbl>, 'Oncotree Code' <chr>, ...
## #
dim(df2)
## [1] 1092
              35
is_categorical <- sapply(df2, is.character)</pre>
is categorical
##
                  Age at Diagnosis
                                            Type of Breast Surgery
##
                             FALSE
                                                              TRUE
```

```
##
             Cancer Type Detailed
                                                        Cellularity
                               TRUE
                                                                TRUE
##
                                       Pam50 + Claudin-low subtype
##
                      Chemotherapy
                              TRUE
##
##
                            Cohort
                                         ER status measured by IHC
##
                              FALSE
                                                                TRUE
                         ER Status
                                         Neoplasm Histologic Grade
##
##
                               TRUE
                                                               FALSE
##
     HER2 status measured by SNP6
                                                        HER2 Status
##
                               TRUE
                                                                TRUE
##
   Tumor Other Histologic Subtype
                                                    Hormone Therapy
##
                               TRUE
                                                                TRUE
                                                Integrative Cluster
##
        Inferred Menopausal State
##
                              TRUE
                                                                TRUE
##
         Primary Tumor Laterality
                                     Lymph nodes examined positive
##
                               TRUE
                                                               FALSE
                    Mutation Count
##
                                       Nottingham prognostic index
##
                              FALSE
                                                               FALSE
##
                     Oncotree Code
                                         Overall Survival (Months)
##
                              TRUE
                                                               FALSE
          Overall Survival Status
                                                          PR Status
##
##
                                                                TRUE
                     Radio Therapy
##
                                      Relapse Free Status (Months)
##
                               TRUE
##
              Relapse Free Status
                                     Number of Samples Per Patient
##
                              TRUE
                                                               FALSE
                                                                 Sex
##
                       Sample Type
                                                                TRUE
##
                               TRUE
##
        3-Gene classifier subtype
                                                TMB (nonsynonymous)
                               TRUE
##
                                                               FALSE
##
                        Tumor Size
                                                        Tumor Stage
##
                              FALSE
                                                               FALSE
##
           Patient's Vital Status
##
                               TRUE
```

Create Dummy Variable for Categorical Variable

```
df2$Type_of_Breast_Surgery = ifelse(df2$`Type of Breast Surgery` =="BREAST CONSERVING",1,0)
df2$new_Chemotherapy = ifelse(df2$Chemotherapy =="NO",0,1)
for (i in 1:nrow(df2)) {
    if (df2[i,]$Cellularity == "High") {
        df2$new_Cellularity[i] <- 3
    }
    else if (df2[i,]$Cellularity == "Moderate") {
        df2$new_Cellularity[i] <- 2
    }
    else {
        df2$new_Cellularity[i] <- 1
    }
}
df2$Pam50_Claudin_low_subtype_Luma = ifelse(df2$`Pam50 + Claudin-low subtype` == "LumA",1,0)
df2$Pam50_Claudin_low_subtype_LumB = ifelse(df2$`Pam50 + Claudin-low subtype` == "LumB",1,0)
df2$Pam50_Claudin_low_subtype_Her2 = ifelse(df2$`Pam50 + Claudin-low subtype` == "Her2",1,0)</pre>
```

```
df2$Pam50_Claudin_low_subtype_Basal = ifelse(df2$`Pam50 + Claudin-low subtype` == "Basal",1,0)
df2$Pam50_Claudin_low_subtype_Normal = ifelse(df2$`Pam50 + Claudin-low subtype` == "Normal",1,0)
df2$Pam50_Claudin_low_subtype_claudin = ifelse(df2$`Pam50 + Claudin-low subtype` == "claudin-low",1,0)
df2$Pam50_Claudin_low_subtype_NC = ifelse(df2$`Pam50 + Claudin-low_subtype` == "NC",1,0)
df2$ER_status_measured_by_IHC = ifelse(df2$`ER status measured by IHC` =="Positve",1,0)
df2$ER Status Positive = ifelse(df2$`ER Status` =="Positive",1,0)
for (i in 1:nrow(df2)) {
  if (df2[i,]$`HER2 status measured by SNP6` == "NEUTRAL") {
   df2$HER2_status_measured_by_SNP6[i] <- 4</pre>
  else if (df2[i,]$`HER2 status measured by SNP6` == "GAIN") {
   df2$HER2_status_measured_by_SNP6[i] <- 3</pre>
  else if (df2[i,]$`HER2 status measured by SNP6` == "LOSS"){
   df2$HER2_status_measured_by_SNP6[i] <- 2</pre>
  else if (df2[i,]$`HER2 status measured by SNP6` == "UNDEF"){
   df2$HER2_status_measured_by_SNP6[i] <- 1</pre>
}
df2$HER2_Status_Positive = ifelse(df2$`HER2 Status` =="Positive",1,0)
df2$Tumor_Other_Histologic_Subtype_Ductal = ifelse(df2$`Tumor Other Histologic Subtype` == "Ductal/NST"
df2$Tumor_Other_Histologic_Subtype_Tubular = ifelse(df2$`Tumor Other Histologic Subtype` == "Tubular/ c
df2$Tumor_Other_Histologic_Subtype_Medullary = ifelse(df2$`Tumor Other Histologic Subtype` == "Medullar
#df2$Oncotree_Code_IDC = ifelse(df2$`Oncotree Code` == "IDC",1,0)
#df2$Oncotree_Code_MLDC = ifelse(df2$`Oncotree Code` == "MLDC",1,0)
#df2$Oncotree_Code_ILC = ifelse(df2$`Oncotree Code` == "ILC",1,0)
#df2$Oncotree_Code_IMMC = ifelse(df2$`Oncotree Code` == "IMMC",1,0)
#df2$Oncotree_Code_BREAST = ifelse(df2$`Oncotree Code` == "BREAST",1,0)
df2$Hormone_Therapy = ifelse(df2$`Hormone Therapy` =="NO",0,1)
df2$Inferred_Menopausal_State = ifelse(df2$`Inferred_Menopausal_State` =="Pre",0,1)
df2$Primary_Tumor_Laterality = ifelse(df2$`Primary Tumor Laterality` =="Right",0,1)
df2$0verall_Survival_Status = ifelse(df2$\cdot \text{Overall Survival Status} =="1:DECEASED",1,0)
df2$PR_Status = ifelse(df2$`PR Status` =="Positive",1,0)
df2$Radio_Therapy = ifelse(df2$`Radio Therapy` =="NO",0,1)
df2$Relapse_Free_Status = ifelse(df2$`Relapse Free Status` =="0:Not Recurred",0,1)
df2$Gene_classifier_subtype_ERH = ifelse(df2$^3-Gene classifier subtype^ == "ER+/HER2- High Prolif",1,0
df2$Gene_classifier_subtype_ERL = ifelse(df2$^3-Gene_classifier_subtype^ == "ER+/HER2- Low Prolif",1,0)
df2$Gene_classifier_subtype_ERM = ifelse(df2$^3-Gene classifier subtype^ == "ER-/HER2-",1,0)
df2$Gene_classifier_subtype_ERP = ifelse(df2$^3-Gene_classifier_subtype^ == "HER2+",1,0)
df2$Patients_Vital_Status_Living = ifelse(df2$^Patient's Vital_Status^ == "Living",1,0)
df2$Patients_Vital_Status_Died = ifelse(df2$`Patient's Vital Status` == "Died of Disease",1,0)
df2$Patients_Vital_Status_Do = ifelse(df2$`Patient's Vital Status` == "Died of Other Causes",1,0)
for (i in 1:nrow(df2)) {
```

```
if (df2[i,]$`Integrative Cluster` == "4ER+") {
    df2$Integrative_Cluster[i] <- 4.5</pre>
  else if (df2[i,]$`Integrative Cluster` == "9") {
    df2$Integrative_Cluster[i] <- 9</pre>
  else if (df2[i,]$`Integrative Cluster` == "7"){
    df2$Integrative_Cluster[i] <- 7</pre>
  else if (df2[i,]$`Integrative Cluster` == "3"){
    df2$Integrative_Cluster[i] <- 3</pre>
  }
  else if (df2[i,]$`Integrative Cluster` == "10"){
    df2$Integrative_Cluster[i] <- 10</pre>
  else if (df2[i,]$`Integrative Cluster` == "8"){
    df2$Integrative_Cluster[i] <- 8</pre>
  else if (df2[i,]$`Integrative Cluster` == "6"){
    df2$Integrative_Cluster[i] <- 6</pre>
  else if (df2[i,]$`Integrative Cluster` == "1"){
    df2$Integrative_Cluster[i] <- 1</pre>
  else if (df2[i,]$`Integrative Cluster` == "2"){
    df2$Integrative_Cluster[i] <- 2</pre>
  else if (df2[i,]$`Integrative Cluster` == "5"){
    df2$Integrative_Cluster[i] <- 5</pre>
  }
  else if (df2[i,]$`Integrative Cluster` == "4ER-"){
    df2$Integrative_Cluster[i] <- 3.5</pre>
  }
}
names(df2) [names(df2) == "Cancer Type Detailed"] <- "Cancer_Type_Detailed"</pre>
df2<- df2[df2$Cancer_Type_Detailed == "Breast Invasive Ductal Carcinoma",]
df2
## # A tibble: 859 x 67
##
      Age at Diagn~1 Type ~2 Cance~3 Cellu~4 Chemo~5 Pam50~6 Cohort ER st~7 ER St~8
##
                <dbl> <chr>
                               <chr>
                                       <chr>>
                                                <chr>
                                                         <chr>
                                                                   <dbl> <chr>
                                                                                  <chr>>
## 1
                 43.2 BREAST~ Breast~ High
                                                NO
                                                         LumA
                                                                       1 Positve Positi~
```

```
## 2
                78.8 MASTEC~ Breast~ Modera~ NO
                                                     LumB
                                                                  1 Positve Positi~
                86.4 BREAST~ Breast~ Modera~ NO
## 3
                                                     LumB
                                                                  1 Positve Positi~
## 4
               85.5 MASTEC~ Breast~ Modera~ NO
                                                                  1 Positve Positi~
                                                     LumA
## 5
                45.4 BREAST~ Breast~ High
                                             YES
                                                     LumB
                                                                  1 Positve Positi~
## 6
                61.5 BREAST~ Breast~ High
                                             NO
                                                     LumB
                                                                  1 Positve Positi~
##
   7
                68.7 MASTEC~ Breast~ Low
                                             YES
                                                     Basal
                                                                  1 Negati~ Negati~
## 8
               49.9 MASTEC~ Breast~ Modera~ YES
                                                     LumA
                                                                  1 Positve Positi~
                                                     LumA
                                                                  1 Positve Positi~
## 9
               54.2 MASTEC~ Breast~ High
                                             NO
               48.6 MASTEC~ Breast~ Low
## 10
                                             NO
                                                     LumA
                                                                  1 Positve Positi~
```

```
## # ... with 849 more rows, 58 more variables: 'Neoplasm Histologic Grade' <dbl>,
       'HER2 status measured by SNP6' <chr>, 'HER2 Status' <chr>,
       'Tumor Other Histologic Subtype' <chr>, 'Hormone Therapy' <chr>,
       'Inferred Menopausal State' <chr>, 'Integrative Cluster' <chr>,
## #
       'Primary Tumor Laterality' <chr>, 'Lymph nodes examined positive' <dbl>,
## #
## #
       'Mutation Count' <dbl>, 'Nottingham prognostic index' <dbl>,
       'Oncotree Code' <chr>, 'Overall Survival (Months)' <dbl>, ...
## #
Create Dummy Variable for Numeric Variable
names(df2)[names(df2) == "Age at Diagnosis"] <- "Age"</pre>
names(df2)[names(df2) == "Neoplasm Histologic Grade"] <- "Neo_Grade"</pre>
names(df2)[names(df2) == "Lymph nodes examined positive"] <- "Lymph"</pre>
names(df2)[names(df2) == "Mutation Count"] <- "Mutation"</pre>
names(df2)[names(df2) == "Nottingham prognostic index"] <- "Nottingham"</pre>
names(df2)[names(df2) == "Overall Survival (Months)"] <- "Overall_Month"</pre>
names(df2) [names(df2) == "Relapse Free Status (Months)"] <- "Relapse Month"
names(df2)[names(df2) == "TMB (nonsynonymous)"] <- "TMB"</pre>
names(df2)[names(df2) == "Tumor Size"] <- "Tumor_Size"</pre>
names(df2)[names(df2) == "Tumor Stage"] <- "Tumor_Stage"</pre>
df3 \leftarrow df2[,-c(2,3,4:6,8,9,11:17,21,23:25,27:32,35)]
## # A tibble: 859 x 42
        Age Cohort Neo_Grade Lymph Mutation Notti~1 Overa~2 Relap~3 Tumor~4 Tumor~5
##
                        <dbl> <dbl>
                                       <dbl>
                                                <dbl>
                                                        <dbl>
                                                                <dbl>
                                                                         <dbl>
                                                                                 <dbl>
      <dbl> <dbl>
##
   1 43.2
                            3
                                  0
                                           2
                                                 4.02
                                                        84.6
                                                                83.5
                                                                            10
                                                                                     1
                 1
## 2 78.8
                                  0
                                                4.06
                                                        7.8
                                                                 2.89
                                                                                     4
                            3
                                           4
                                                                            31
## 3 86.4
                 1
                            3
                                  1
                                           4
                                                5.03
                                                        36.6
                                                                36.1
                                                                            16
                                                                                     2
## 4 85.5
                 1
                            2
                                  0
                                           1
                                                3.04 132.
                                                               123.
                                                                            22
                                                                                     4
                                                4.05 141.
## 5 45.4
                            3
                                  0
                                                                            23
                                                                                     2
                 1
                                           5
                                                               139.
                                                                                     2
## 6 61.5
                 1
                            2
                                  1
                                           3
                                                4.03 157.
                                                               155.
                                                                            16
## 7 68.7
                            3
                                  0
                                                4.08
                                                         8.07
                                                                            39
                                                                                     2
                                           1
                                                                 7.83
                 1
   8 49.9
                                                                                     3
##
                 1
                            1
                                  5
                                           4
                                                 4.14
                                                        85.3
                                                                84.2
                                                                            70
## 9 54.2
                            1
                                  0
                                           4
                                                 2.05 127.
                                                               125.
                                                                            27
                                                                                     2
                 1
## 10 48.6
                 1
                                  0
                                           3
                                                 3.06
                                                        13.4
                                                                13.2
                                                                            30
## # ... with 849 more rows, 32 more variables: Type_of_Breast_Surgery <dbl>,
       new_Chemotherapy <dbl>, new_Cellularity <dbl>,
## #
       Pam50_Claudin_low_subtype_Luma <dbl>, Pam50_Claudin_low_subtype_LumB <dbl>,
## #
       Pam50_Claudin_low_subtype_Her2 <dbl>,
## #
       Pam50_Claudin_low_subtype_Basal <dbl>,
## #
       Pam50_Claudin_low_subtype_Normal <dbl>,
## #
```

Table 1: Descriptive Statistics of The Data (continued below)

Pam50\_Claudin\_low\_subtype\_claudin <dbl>, ...

pander(summary(df3),caption='Descriptive Statistics of The Data')

## #

Age	Cohort	Neo_Grade	Lymph
Min. :26.36	Min. :1.000	Min. :1.000	Min.: 0.000

Age	Cohort	$Neo\_Grade$	Lymph
1st Qu.:49.99	1st Qu.:1.000	1st Qu.:2.000	1st Qu.: 0.000
Median $:60.62$	Median $:2.000$	Median $:3.000$	Median: 0.000
Mean $:60.05$	Mean $:2.191$	Mean $:2.517$	Mean: $1.916$
3rd Qu.:69.75	3rd Qu.:3.000	3rd Qu.:3.000	3rd Qu.: 2.000
Max. :96.29	Max. $:5.000$	Max. $:3.000$	Max. :41.000

Table 2: Table continues below

Mutation	Nottingham	$Overall\_Month$	$Relapse\_Month$
Min.: 1.000	Min. :2.018	Min.: 0.10	Min.: 0.10
1st Qu.: 3.000	1st Qu.:3.080	1st Qu.: 58.05	1st Qu.: 40.10
Median: 5.000	Median $:4.050$	Median: 115.30	Median: 98.42
Mean: 5.423	Mean $:4.216$	Mean : $124.05$	Mean $:109.35$
3rd Qu.: 7.000	3rd Qu.:5.050	3rd Qu.:186.32	3rd Qu.:172.12
Max. :46.000	Max. :6.360	Max. :337.03	Max. :296.91

Table 3: Table continues below

Tumor_Size	${\bf Tumor\_Stage}$	${\bf Type\_of\_Breast\_Surgery}$	new_Chemotherapy
Min.: 1.0	Min. :1.000	Min. :0.0000	Min. :0.0000
1st Qu.: 17.0	1st Qu.:1.000	1st Qu.:0.0000	1st Qu.:0.0000
Median: 22.0	Median : 2.000	Median: 0.0000	Median: 0.0000
Mean: 25.7	Mean $:1.767$	Mean $:0.4319$	Mean $:0.2538$
3rd Qu.: 30.0	3rd Qu.:2.000	3rd Qu.:1.0000	3rd Qu.:1.0000
Max. $:180.0$	Max. :4.000	Max. $:1.0000$	Max. $:1.0000$

Table 4: Table continues below

new_Cellularity	$Pam50\_Claudin\_low\_subtype\_Luma$
Min. :1.000	Min. :0.0000
1st Qu.:2.000	1st Qu.:0.0000
Median : 3.000	Median: 0.0000
Mean $:2.421$	Mean $:0.3667$
3rd Qu.:3.000	3rd Qu.:1.0000
Max. :3.000	Max. :1.0000

Table 5: Table continues below

$Pam50\_Claudin\_low\_subtype\_LumB$	$Pam50\_Claudin\_low\_subtype\_Her2$
Min. :0.0000	Min. :0.0000
1st Qu.:0.0000	1st Qu.:0.0000
Median: 0.0000	Median: 0.0000
Mean $:0.2608$	Mean $: 0.1036$
3rd Qu.:1.0000	3rd Qu.:0.0000
Max. :1.0000	Max. :1.0000

Table 6: Table continues below

Pam50_Claudin_low_subtype_Basal	$Pam 50 \_Claudin \_low \_subtype \_Normal$
Min. :0.0000	Min. :0.00000
1st Qu.:0.0000	1st Qu.:0.00000
Median: 0.0000	Median $:0.00000$
Mean $:0.1141$	Mean $:0.05122$
3rd Qu.:0.0000	3rd Qu.:0.00000
Max. $:1.0000$	Max. :1.00000

Table 7: Table continues below

Pam50_Claudin_low_subtype_claudin	$Pam50\_Claudin\_low\_subtype\_NC$
Min. :0.0000	Min. :0.000000
1st Qu.:0.0000	1st Qu.:0.000000
Median: 0.0000	Median $:0.000000$
Mean $:0.1013$	Mean $:0.002328$
3rd Qu.:0.0000	3rd Qu.:0.000000
Max. :1.0000	Max. :1.000000

Table 8: Table continues below

ER_status_measured_by_IHC	ER_Status_Positive	HER2_status_measured_by_SNP6
Min. :0.0000	Min. :0.0000	Min. :1.000
1st Qu.:0.0000	1st Qu.:0.0000	1st Qu.:3.000
Median: 1.0000	Median $:1.0000$	Median: 4.000
Mean $:0.7404$	Mean $:0.7404$	Mean $: 3.631$
3rd Qu.:1.0000	3rd Qu.:1.0000	3rd Qu.:4.000
Max. $:1.0000$	Max. $:1.0000$	Max. $:4.000$

Table 9: Table continues below

HER2_Status_Positive	$Tumor\_Other\_Histologic\_Subtype\_Ductal$
Min. :0.0000	Min. :0.0000
1st Qu.:0.0000	1st Qu.:1.0000
Median: 0.0000	Median: 1.0000
Mean $: 0.1444$	Mean $: 0.9674$
3rd Qu.:0.0000	3rd Qu.:1.0000
Max. :1.0000	Max. :1.0000

Table 10: Table continues below

Tumor_Other_Histologic_Subtype_Tubular
Min. :0.00000
1st Qu.:0.00000
Median : 0.00000
Mean $: 0.01746$

Tumor_Other_Histologic_Subtype_Tubular
3rd Qu.:0.00000 Max. :1.00000
Max. :1.00000

Table 11: Table continues below

$Tumor\_Other\_Histologic\_Subtype\_Medullary$	$Hormone\_Therapy$
Min. :0.00000	Min. :0.0000
1st Qu.:0.00000	1st Qu.:0.0000
Median $:0.00000$	Median : 1.0000
Mean $:0.01513$	Mean $:0.6042$
3rd Qu.:0.00000	3rd Qu.:1.0000
Max. :1.00000	Max. :1.0000

Table 12: Table continues below

$Inferred\_Menopausal\_State$	${\bf Primary\_Tumor\_Laterality}$	$Overall\_Survival\_Status$
Min. :0.0000	Min. :0.000	Min. :0.0000
1st Qu.:0.5000	1st Qu.:0.000	1st Qu.:0.0000
Median: 1.0000	Median: 1.000	Median: 1.0000
Mean $: 0.7497$	Mean $:0.525$	Mean $:0.5565$
3rd Qu.:1.0000	3rd Qu.:1.000	3rd Qu.:1.0000
Max. $:1.0000$	Max. $:1.000$	Max. $:1.0000$

Table 13: Table continues below

PR_Status	Radio_Therapy	$Relapse\_Free\_Status$
Min. :0.0000	Min. :0.0000	Min. :0.0000
1st Qu.:0.0000	1st Qu.:0.0000	1st Qu.:0.0000
Median: 0.0000	Median $:1.0000$	Median: 0.0000
Mean $:0.4994$	Mean $:0.6799$	Mean $:0.4214$
3rd Qu.:1.0000	3rd Qu.:1.0000	3rd Qu.:1.0000
Max. :1.0000	Max. :1.0000	Max. :1.0000

Table 14: Table continues below

${\tt Gene\_classifier\_subtype\_ERH}$	Gene_classifier_subtype_ERL
Min. :0.000	Min. :0.0000
1st Qu.:0.000	1st Qu.:0.0000
Median: 0.000	Median: 0.0000
Mean: 0.362	Mean $:0.3132$
3rd Qu.:1.000	3rd Qu.:1.0000
Max. :1.000	Max. :1.0000

Table 15: Table continues below

${\tt Gene\_classifier\_subtype\_ERM}$	${\tt Gene\_classifier\_subtype\_ERP}$
Min. :0.0000	Min. :0.0000
1st Qu.:0.0000	1st Qu.:0.0000
Median: 0.0000	Median: 0.0000
Mean $:0.1921$	Mean $:0.1327$
3rd Qu.:0.0000	3rd Qu.:0.0000
Max. $:1.0000$	Max. :1.0000

Table 16: Table continues below

Patients_Vital_Status_Living	$Patients\_Vital\_Status\_Died$
Min. :0.0000	Min. :0.0000
1st Qu.:0.0000	1st Qu.:0.0000
Median: 0.0000	Median: 0.0000
Mean $:0.4435$	Mean $:0.3481$
3rd Qu.:1.0000	3rd Qu.:1.0000
Max. :1.0000	Max. :1.0000

Patients_Vital_Status_Do	Integrative_Cluster
Min. :0.0000	Min.: 1.000
1st Qu.:0.0000	1st Qu.: 3.500
Median: 0.0000	Median: 5.000
Mean $:0.2084$	Mean: 5.861
3rd Qu.:0.0000	3rd Qu.: 8.000
Max. $:1.0000$	Max. $:10.000$

pander(head(df3),caption='Head of data selection')

Table 18: Head of data selection (continued below)

Age	Cohort	Neo_Grade	Lymph	Mutation	Nottingham	Overall_Month
43.19	1	3	0	2	4.02	84.63
78.77	1	3	0	4	4.062	7.8
86.41	1	3	1	4	5.032	36.57
85.49	1	2	0	1	3.044	132
45.43	1	3	0	5	4.046	140.9
61.49	1	2	1	3	4.032	157.4

Table 19: Table continues below

$Relapse\_Month$	${\bf Tumor\_Size}$	${\bf Tumor\_Stage}$	Type_of_Breast_Surgery
83.52	10	1	1
2.89	31	4	0
36.09	16	2	1

Relapse_Month	${\bf Tumor\_Size}$	${\bf Tumor\_Stage}$	Type_of_Breast_Surgery
123.3	22	4	0
139	23	2	1
155.4	16	2	1

Table 20: Table continues below

${\bf new\_Chemotherapy}$	${\bf new\_Cellularity}$	$Pam 50 \_Claudin \_low \_subtype \_Luma$
0	3	1
0	2	0
0	2	0
0	2	1
1	3	0
0	3	0

Table 21: Table continues below

Pam50_Claudin_low_subtype_LumB	Pam50_Claudin_low_subtype_Her2
0	0
1	0
1	0
0	0
1	0
1	0

Table 22: Table continues below

$Pam 50 \_Claudin \_low \_subtype \_Basal$	$Pam 50 \_Claudin \_low \_subtype \_Normal$
0	0
0	0
0	0
0	0
0	0
0	0

Table 23: Table continues below

$Pam 50\_Claudin\_low\_subtype\_claudin$	$Pam50\_Claudin\_low\_subtype\_NC$
0	0
0	0
0	0
0	0
0	0
0	0

Table 24: Table continues below

ER_status_measured_by_IHC	ER_Status_Positive	HER2_status_measured_by_SNP6
1	1	4
1	1	4
1	1	3
1	1	4
1	1	4
1	1	4

Table 25: Table continues below

HER2_Status_Positive	${\bf Tumor\_Other\_Histologic\_Subtype\_Ductal}$
0	1
0	1
0	1
0	1
0	1
0	1

Table 26: Table continues below

${\bf Tumor\_Other\_Histologic\_Subtype\_Tubular}$	
0	
0	
0	
0	
U	
U	

Table 27: Table continues below

${\bf Tumor\_Other\_Histologic\_Subtype\_Medullary}$	${\bf Hormone\_The rapy}$
0	1
0	1
0	1
0	1
0	1
0	1

Table 28: Table continues below

_			
	$Inferred\_Menopausal\_State$	${\bf Primary\_Tumor\_Laterality}$	$Overall\_Survival\_Status$
	0	0	0
	1	1	1
	1	0	1
	1	1	1

Inferred_Menopausal_State	Primary_Tumor_Laterality	Overall_Survival_Status
0	0	0
1	1	0

Table 29: Table continues below

PR_Status	Radio_Therapy	$Relapse\_Free\_Status$	${\tt Gene\_classifier\_subtype\_ERH}$
1	1	0	1
1	1	1	1
0	1	0	1
1	1	1	0
1	1	0	1
1	1	0	1

Table 30: Table continues below

${\it Gene\_classifier\_subtype\_ERL}$	${\tt Gene\_classifier\_subtype\_ERM}$
0	0
0	0
0	0
1	0
0	0
0	0

Table 31: Table continues below

${\tt Gene\_classifier\_subtype\_ERP}$	Patients_Vital_Status_Living
0	1
0	0
0	0
0	0
0	1
0	1

Patients_Vital_Status_Died	Patients_Vital_Status_Do	Integrative_Cluster
0	0	4.5
1	0	7
0	1	9
1	0	3
0	0	10
0	0	7

```
df3 %>%
  keep(is.numeric) %>%
  gather() %>%
  ggplot(aes(value)) +
  facet_wrap(~ key, scales = "free") +
  geom_histogram()
```

## 'stat\_bin()' using 'bins = 30'. Pick better value with 'binwidth'.

```
Age
                         Cohort
                                       measure
                                                      Status Po
                                                                     sifier_sub
                                                                                    ssifier_sub
                                                                                                   sifier_sub
          406080100
                        12345
                                     0.000256071500
                                                   0.0002550071500
                                                                   0.000256071500
                                                                                  0.000255071500
                                                                                                 0.000255071500
                                       _Status_P
         sifier_sub
                                                      none_The
                                                                     Menopaus
                                                                                    grative_Clu
                                                                                                     Lymph
                        s_measure
                                     0.000256071500
       0.000256071500
                        1 2 3 4
                                                    0.000256071500
                                                                   0.000256071500
                                                                                     2.55.07.50.0
                                                                                                   010203040
         Mutation
                        leo Grade
                                       w Cellulai
                                                      Chemothe
                                                                     Nottinghan
                                                                                    rerall Mor
                                                                                                    Survival
                                                                                    01000000
        010203040
                       1.0.52.02.53.0
                                      1.0.52.02.53.0
                                                    0.00255071500
                                                                     23456
                                                                                                  0.00255071500
 count
         din_low_si
                        in_low_su
                                       din_low_s
                                                      din_low_s
                                                                     le_wol_nit
                                                                                    udin_low_:
                                                                                                   in_low_su
       0.000255071500
                     0.000255071500 0.000255071500 0.000255071500
                                                                   0.000255071500
                                                                                  0.00255071500
                                                                                                 0.000255071500
                                       Vital_Stat
                                                      R_Status
                                                                                    dio_Thera
         Vital Star
                        _Vital_Sta
                                                                      Tumor_L
                                                                                                    se_Free_
       0.000256071500
                      0.000256071500
                                     0.0002550071500
                                                    0.000256071500
                                                                   0.000256071500
                                                                                  0.000256071500
                                                                                                  0.0002550071500
         lapse_Mo
                        listologic_
                                       tologic_Si
                                                      istologic_5
                                                                     umor_Siz
                                                                                    ımor_Stag
                                                                                                   f_Breast_$
         0 10 20 300
                     0.000256071500 0.000256071500
                                                    0.0002550071500
                                                                     0 501 0050
                                                                                    1 2 3 4
                                                                                                  0.00255071500
                                                       value
tr_ind <- 1:(nrow(df3) * 0.7)</pre>
df3_tr <- df3[tr_ind, ]</pre>
nrow(df3_tr)
## [1] 601
df3_te <- df3[-tr_ind, ]</pre>
nrow(df3_te)
## [1] 258
variable selection
set.seed(0)
fit_BIC <- regsubsets(Overall_Survival_Status ~ ., data = df3_tr)</pre>
## Warning in leaps.setup(x, y, wt = wt, nbest = nbest, nvmax = nvmax, force.in =
## force.in, : 4 linear dependencies found
## Reordering variables and trying again:
```

## summary\_BIC <- summary(fit\_BIC)</pre>

## ## Warning in log(vr): NaNs produced

#### summary\_BIC

```
## Subset selection object
## Call: regsubsets.formula(Overall_Survival_Status ~ ., data = df3_tr)
## 41 Variables (and intercept)
##
                                             Forced in Forced out
## Age
                                                 FALSE
                                                             FALSE
## Cohort
                                                 FALSE
                                                             FALSE
## Neo_Grade
                                                 FALSE
                                                             FALSE
## Lymph
                                                 FALSE
                                                             FALSE
## Mutation
                                                 FALSE
                                                             FALSE
## Nottingham
                                                 FALSE
                                                             FALSE
## Overall_Month
                                                 FALSE
                                                             FALSE
## Relapse_Month
                                                 FALSE
                                                             FALSE
## Tumor_Size
                                                 FALSE
                                                             FALSE
## Tumor_Stage
                                                 FALSE
                                                            FALSE
## Type_of_Breast_Surgery
                                                 FALSE
                                                             FALSE
## new Chemotherapy
                                                 FALSE
                                                             FALSE
## new Cellularity
                                                 FALSE
                                                             FALSE
## Pam50 Claudin low subtype Luma
                                                 FALSE
                                                            FALSE
## Pam50_Claudin_low_subtype_LumB
                                                 FALSE
                                                            FALSE
## Pam50_Claudin_low_subtype_Her2
                                                 FALSE
                                                             FALSE
## Pam50_Claudin_low_subtype_Basal
                                                             FALSE
                                                 FALSE
## Pam50_Claudin_low_subtype_Normal
                                                 FALSE
                                                             FALSE
## Pam50_Claudin_low_subtype_claudin
                                                 FALSE
                                                             FALSE
## ER_status_measured_by_IHC
                                                 FALSE
                                                             FALSE
## ER_Status_Positive
                                                 FALSE
                                                             FALSE
## HER2_status_measured_by_SNP6
                                                 FALSE
                                                             FALSE
## HER2_Status_Positive
                                                 FALSE
                                                             FALSE
## Tumor_Other_Histologic_Subtype_Ductal
                                                 FALSE
                                                             FALSE
## Tumor_Other_Histologic_Subtype_Tubular
                                                 FALSE
                                                             FALSE
## Hormone_Therapy
                                                 FALSE
                                                             FALSE
## Inferred_Menopausal_State
                                                             FALSE
                                                 FALSE
## Primary_Tumor_Laterality
                                                 FALSE
                                                             FALSE
## PR_Status
                                                 FALSE
                                                             FALSE
## Radio_Therapy
                                                 FALSE
                                                             FALSE
## Relapse Free Status
                                                 FALSE
                                                             FALSE
## Gene_classifier_subtype_ERH
                                                 FALSE
                                                             FALSE
## Gene classifier subtype ERL
                                                 FALSE
                                                            FALSE
## Gene_classifier_subtype_ERM
                                                 FALSE
                                                             FALSE
## Patients_Vital_Status_Living
                                                 FALSE
                                                             FALSE
## Patients_Vital_Status_Died
                                                 FALSE
                                                             FALSE
## Integrative_Cluster
                                                             FALSE
                                                 FALSE
## Pam50_Claudin_low_subtype_NC
                                                 FALSE
                                                             FALSE
## Tumor_Other_Histologic_Subtype_Medullary
                                                 FALSE
                                                             FALSE
## Gene_classifier_subtype_ERP
                                                 FALSE
                                                             FALSE
## Patients_Vital_Status_Do
                                                 FALSE
                                                             FALSE
## 1 subsets of each size up to 9
```

```
## Selection Algorithm: exhaustive
##
           Age Cohort Neo_Grade Lymph Mutation Nottingham Overall_Month
     (1)""""
                               11 11
                                    11 11
                                             11 11
     (1)""""
## 2
                     11 11
                               11 11
                                    11 11
                                                       11 11
## 3
     (1)"*"""
                     11 11
                               11 11
## 4
     (1)"*"""
     (1)"*""*"
                     "*"
                               11 11
                                    11 11
                     "*"
                               "*"
     (1) "*" "*"
## 6
## 7
     (1) "*" "*"
                     "*"
                               "*"
     (1)"*""*"
                     "*"
                               "*"
                                    "*"
## 8
## 9 (1) "*" "*"
                     "*"
                               "*"
                                    "*"
                                             "*"
                                                       "*"
##
           Relapse_Month Tumor_Size Tumor_Stage Type_of_Breast_Surgery
     (1)""
                        11 11
                                  11 11
## 1
## 2 (1)""
## 3 (1)""
                        11 11
                                  11 11
     (1)""
## 4
## 5
     (1)""
                        11 11
                                  11 11
     (1)""
## 6
     (1)""
                        11 11
                                  11 11
## 7
                                  11 11
## 8 (1)""
## 9 (1)""
                        "*"
                                  11 11
                                              11 11
           new_Chemotherapy new_Cellularity Pam50_Claudin_low_subtype_Luma
## 1 (1)""
                           11 11
     (1)""
                           11 11
                                          .. ..
## 2
## 3 (1)""
     (1)""
                           11 11
     (1)""
## 5
     (1)""
                           11 11
## 6
     (1)""
## 7
## 8 (1) " "
                           11 11
                                          11 11
     (1)""
## 9
##
           Pam50_Claudin_low_subtype_LumB Pam50_Claudin_low_subtype_Her2
     (1)""
## 1
     (1)""
                                        .. ..
## 2
     (1)""
## 3
     (1)""
## 4
## 5 (1)""
## 6 (1) " "
     (1)""
## 7
## 8 (1)""
                                        .. ..
## 9 (1)""
##
           Pam50_Claudin_low_subtype_Basal Pam50_Claudin_low_subtype_Normal
     (1)""
## 1
## 2 (1)""
## 3 (1)""
     (1)""
## 4
     (1)""
## 5
     (1)""
## 6
    (1)"*"
## 7
     (1)""
## 8
     (1)""
## 9
##
           Pam50_Claudin_low_subtype_claudin Pam50_Claudin_low_subtype_NC
## 1 (1)""
## 2 (1)""
                                           11 11
```

```
## 3 (1)""
                                     11 11
## 4 (1)""
## 5 (1)""
## 6 (1) " "
    (1)""
## 7
## 8 (1) " "
## 9 (1)""
##
         ER_status_measured_by_IHC ER_Status_Positive
## 1 (1) " "
## 2 (1)""
## 3 (1)""
    (1)""
## 4
   (1)""
## 5
## 6 (1) " "
## 7 (1)""
## 8 (1)""
## 9 (1)""
                               11 11
         HER2_status_measured_by_SNP6 HER2_Status_Positive
## 1 (1)""
    (1)""
## 2
## 3 (1)""
                                 11 11
## 4 (1)""
## 5 (1)""
## 6
    (1)""
## 7 (1)""
## 8 (1)""
                                 11 11
## 9 (1)""
##
          Tumor_Other_Histologic_Subtype_Ductal
## 1 (1)""
## 2 (1)""
    (1)""
## 3
## 4 (1)""
## 5 (1)""
## 6 (1) " "
    (1)""
## 7
## 8 (1)""
## 9 (1)""
##
          Tumor_Other_Histologic_Subtype_Tubular
## 1 (1)""
## 2 (1)""
## 3 (1) " "
    (1)""
## 4
## 5
    (1)""
## 6 (1)""
## 7 (1)""
## 8 (1)""
    (1)""
## 9
##
          Tumor_Other_Histologic_Subtype_Medullary Hormone_Therapy
## 1 (1)""
    (1)""
## 2
## 3 (1)""
## 4 (1)""
## 5 (1)""
## 6 (1) " "
                                           11 11
```

```
11 11
## 7 (1)""
## 8 (1)""
                                                11 11
## 9 (1)""
                                                11 11
##
           Inferred_Menopausal_State Primary_Tumor_Laterality PR_Status
                                  11 11
## 1 (1) " "
## 2 (1)""
                                                         11 11
## 3 (1) " "
                                  11 11
                                                         11 11
    (1)""
## 4
## 5
     (1)""
## 6 (1)""
## 7 (1)""
                                  11 11
## 8 (1)""
                                  11 11
                                                         11 11
    (1)""
## 9
           Radio_Therapy Relapse_Free_Status Gene_classifier_subtype_ERH
##
## 1 (1)""
                       11 11
     (1)""
                       "*"
## 2
                                         .. ..
## 3 (1)""
                       "*"
## 4 (1)""
                       "*"
## 5 (1)""
    (1)""
## 6
    (1)""
## 7
## 8 (1)""
## 9 (1)""
           {\tt Gene\_classifier\_subtype\_ERL\ Gene\_classifier\_subtype\_ERM}
## 1 (1)""
                                    11 11
## 2 (1)""
## 3 (1)""
## 4
    (1)""
## 5 (1)""
## 6 (1) " "
    (1)""
## 7
                                    11 11
## 8 (1) " "
## 9 (1)""
##
           Gene_classifier_subtype_ERP Patients_Vital_Status_Living
## 1 (1)""
                                    "*"
## 2 (1)""
                                    "*"
## 3 (1)""
                                    "*"
## 4 (1) " "
                                    "*"
     (1)""
## 5
## 6 (1) " "
## 7 (1)""
## 8 (1) " "
## 9
     (1)""
                                    "*"
##
           Patients_Vital_Status_Died Patients_Vital_Status_Do
## 1 (1)""
                                   11 11
    (1)""
## 2
                                   11 11
    (1)""
## 3
## 4 (1)""
## 5 (1)"*"
    (1)"*"
## 6
    (1)"*"
## 7
                                   "*"
## 8 (1) "*"
## 9 (1)""
                                   11 11
##
           Integrative_Cluster
```

```
## 1 (1)""
## 2 (1)""
     (1)""
## 4
     (1)""
     (1)""
## 5
## 6 (1) " "
## 7 (1)""
## 8 (1) " "
## 9 (1) " "
min_BIC <- which.min(summary_BIC$bic)</pre>
min_BIC
## [1] 8
coef_BIC = coef(fit_BIC,min_BIC)
## Warning in log(vr): NaNs produced
coef_BIC
##
                                 (Intercept)
                                -0.677629271
##
##
                                          Age
                                 0.010622924
##
##
                                      Cohort
                                 0.145974676
##
##
                                   Neo_Grade
##
                                 0.028670848
##
                                       Lymph
##
                                 0.018462621
##
                                    Mutation
##
                                 0.004245253
##
                                  Nottingham
##
                                 0.037713480
## Tumor_Other_Histologic_Subtype_Medullary
##
                                 0.166695765
                Gene_classifier_subtype_ERP
##
##
                                 0.103582951
#Forward Stepwise Selection with Adjusted R squared
\#fit\_FORWARD \leftarrow regsubsets(Overall\_Survival\_Status \sim ., data = df3\_tr, method = "forward", numax = 10)
#as.factor(df3_tr$Cancer_Type_Detailed)
#summary_FORWARD <- summary(fit_FORWARD)</pre>
#max_FORWARD <- which.max(summary_FORWARD$adjr2)</pre>
#max_FORWARD
\#coef\_FORWARD = coef(fit\_FORWARD, max\_FORWARD)
#coef_FORWARD
```

Backward Stepwise Selection with Cp

```
fit_BACKWARD <- regsubsets(Overall_Survival_Status ~ ., data = df3_tr, method = "backward", nvmax = nco</pre>
## Warning in leaps.setup(x, y, wt = wt, nbest = nbest, nvmax = nvmax, force.in =
## force.in, : 4 linear dependencies found
## Reordering variables and trying again:
## Warning in rval$lopt[] <- rval$vorder[rval$lopt]: number of items to replace is
## not a multiple of replacement length
summary_BACKWARD <- summary(fit_BACKWARD)</pre>
mix_BACKWARD <- which.min(summary_BACKWARD$cp)</pre>
mix_BACKWARD
## [1] 19
coef_BACKWARD = coef(fit_BACKWARD, mix_BACKWARD)
coef_BACKWARD
##
                                 (Intercept)
                                 0.101529777
##
##
                                          Age
##
                                 0.008725082
##
                                       Cohort
##
                                 0.205473176
##
                                        Lymph
##
                                 0.014394673
##
                               Overall_Month
##
                                -0.001669157
##
                               Relapse_Month
##
                                -0.001376028
##
                                 Tumor_Stage
##
                                -0.037640439
##
             Pam50_Claudin_low_subtype_Luma
##
                                 -0.118693753
##
             Pam50_Claudin_low_subtype_LumB
##
                                 -0.069504986
##
            Pam50_Claudin_low_subtype_Basal
##
                                 -0.065317856
##
          Pam50_Claudin_low_subtype_claudin
##
                                 -0.117549258
##
                        HER2_Status_Positive
##
                                 0.022130151
##
     Tumor_Other_Histologic_Subtype_Tubular
                                 0.033264067
##
##
                             Hormone_Therapy
##
                                 -0.057071936
##
                                    PR_Status
##
                                 0.021133168
##
                Gene classifier subtype ERH
```

0.090267981

##

```
##
                                       Gene_classifier_subtype_ERL
##
                                                                             -0.003118641
##
                                       Gene classifier subtype ERM
                                                                               0.006531345
##
##
                                    Pam50_Claudin_low_subtype_NC
                                                                               0.412900231
##
## Tumor_Other_Histologic_Subtype_Medullary
                                                                               0.146141437
##
set.seed(0)
predict BIC = glm(Overall Survival Status~Age + Cohort + Neo Grade + Lymph + Mutation + Nottingham + T
pred_BIC = round(predict(predict_BIC,df3_te,type = "response"))
error_BIC = mean((df3_te$Overall_Survival_Status - pred_BIC)^2)
error_BIC
## [1] 0.3333333
\#predict\_FORWARD = glm(Overall\_Survival\_Status \sim Cohort + Neo\_Grade + Overall\_Month + Relapse\_Month + neo\_Grade + Overall\_Month + Relapse\_Month + neo\_Grade + Overall\_Month + neo\_Grade + Overall\_Month + neo\_Grade + Overall\_Month + neo\_Grade + ne
#pred FORWARD = round(predict(predict FORWARD, df3 te, type = "response"))
#pred FORWARD
#error_FORWARD = mean((df3_te$Overall_Survival_Status - pred_FORWARD)^2)
#error_FORWARD
predict_BACKWARD = glm(Overall_Survival_Status ~ Age+ Cohort + Lymph + Overall_Month + Relapse_Month
pred_BACKWARD = round(predict(predict_BACKWARD,df3_tr,type = "response"))
error_BACKWARD = mean((df3_te$Overall_Survival_Status - pred_BACKWARD)^2)
error_BACKWARD
## [1] 0.5291181
which.min(data.frame(error_BIC, error_BACKWARD)) # All the errors listed
## error BIC
##
                           1
formula = Overall_Survival_Status~Age + Cohort + Neo_Grade + Lymph + Mutation + Nottingham + Tumor_Oth
Model 1
set.seed(0)
library(randomForest)
rf_model <- randomForest(formula, data = df3_tr,ntree = 10,importance = T)
predictions <- predict(rf_model, df3_te)</pre>
error_rf <- mean((df3_te$Overall_Survival_Status - predictions)^2)</pre>
error_rf
## [1] 0.217086
```

Model 2

```
set.seed(0)
knn_test_error <- vector()</pre>
new <- data.frame(k=numeric(), knn_test_error)</pre>
for(i in seq(from = 1, to = 100, by = 5)) {
  knmodel <- knn3(formula, df3_tr, k = i)</pre>
  knn_test <- predict(knmodel, newdata = df3_te)</pre>
  knn_test_error <- mean((knn_test - df3_te$0verall_Survival_Status)^2)</pre>
  new[i,] <- c(i, knn_test_error)</pre>
frame = data.frame(knn_test_error,1:24)
error_knn <- data.frame(knn_test_error,1:24)[which.min(frame$knn_test_error),]</pre>
error_knn
   knn_test_error X1.24
##
          0.2786025
## 1
Model 3 Ridge
x_tr<-as.matrix(df3_tr[,c(1:23, 25:ncol(df3_tr))])</pre>
y_tr<-as.matrix(df3_tr[,24])</pre>
x_te<-as.matrix(df3_te[,c(1:23, 25:ncol(df3_te))])
y te<-as.matrix(df3 te[,24])</pre>
library(glmnet)
## Warning: package 'glmnet' was built under R version 4.2.2
## Warning: package 'Matrix' was built under R version 4.2.2
set.seed(0)
ridge<-glmnet(x=x_tr,y=y_tr,alpha=0)</pre>
#plot(ridge, xvar='lambda')
ridge_cv<-cv.glmnet(x=x_tr,y=y_tr,type.measure='mse',nfold=10,alpha=0)
best_ridge<-coef(ridge_cv, s = ridge_cv$lambda.min)</pre>
result_ridge<-predict(ridge_cv,newx=x_te,interval='prediction')</pre>
error_ridge <- (err_ridge<-mean((y_te-result_ridge)^2))</pre>
error_ridge
## [1] 0.04608354
Model 4 Lasso
set.seed(0)
lasso<-glmnet(x=x_tr,y=y_tr,alpha=1)</pre>
#plot(lasso,xvar='lambda')
```

```
lasso_cv<-cv.glmnet(x=x_tr,y=y_tr,type.measure='mse',nfold=10,alpha=1,keep=T)
#lasso_cv$lambda.min

result_la<-predict(lasso_cv,newx=x_te,interval='prediction')
error_lasso <- (err_la<-mean((y_te-result_la)^2))
error_lasso

## [1] 0.04679185

Model Selection

which.min(data.frame(error_rf, error_knn, error_ridge, error_lasso)) # All the errors listed

## error_ridge
## error_ridge
## 4</pre>
```

Model Present cross validation