SurvPredPipe

SurvPredPipe: a R-Package for the Pipeline to Predict Survival Probability of Cancer Patients

Introduction:

The **SurvPredPipe** package is about the pipeline for the prediction Survival Probability of Cancer Patients. It performs various steps: Data Processing, Split data into training and test subset, Data Normalization, Select Significant features based on Univariate survival, Generate LASSO PI Score, Develop Prediction model for survival probability based on different features and draw survival curve based on predicted survival probability values and barplots for predicted mean and median survival time of patients.

Key steps:

- **1. Data Processing:** Removing samples where OS time information is missing and convert OS time in days into months
- 2. Train-Test Split: For Feature Selection and Model Development
- 3. Data Normalization: Log-scale transformation followed by Quantile Normalization
- 4. Univariate Feature Selection
- 5. Quantification of PI index per sample based on LASSO (glmnet method)
- 6. MTLR model Development based on:
 - a. Clinical features
 - b PI Score
 - c. PI index + Clin feature
 - d Univariate
 - e. Univariate + Clin features
- 7. Survival Curve for Patients
- 8. BarPlots for Mean/Median Survival time
- 9. Nomogram

Final Results:

- 1. Table representing IBS score and C-Index for training model
- 2. Survival Probability curve plot per patient for test data
- 3. Mean and median Survival time value per patient
- 4. Table for Survival Probability
- 5. Nomogram to predict death risk, 1-year, 3-year and 5-year survival probability of patients based on user- selected features

Follow the Steps to Install package

```
"" #Step1: First Install remote package
install.packages("remotes")

#load remortes package
library("remotes")

#Step2: install SurvPredPipe package
remotes::install_github("hks5august/SurvPredPipe", local = TRUE)

# load package
library("SurvPredPipe") ""
```

Workflow with an Example Data

#Load package

library(SurvPredPipe)

#Load other required packages

```
library(caret)
library(preprocessCore)
library(ggfortify)
library(survival)
library(survminer)
library(dplyr)
library(ggplot2)
library(ggfortify)
library (MASS)
library (MTLR)
library(dplyr)
library(SurvMetrics)
library(pec)
library(glmnet)
library(reshape2)
library(rms)
library(Matrix)
library(Hmisc)
library(survivalROC)
library(ROCR)
```

#set seed

set.seed(7)

#set path of the working directory where input data available
"" setwd() ""

Input Data

Input data: Example_TCGA_LGG_FPKM_data.txt

Example Input data: "Example_TCGA_LGG_FPKM_data.txt" is a tab separated file. It contains Samples (184 LGG Cancer Samples) in the rows and Features in the columns. Gene Expression is available in terms of FPKM values in the data.

- **Features information**: In the data there are 11 clinical + demographic, 4 types survival with time and event information and 19,978 protein coding genes.
- Clinical and demographic features: Clinical demographic features that are present in this example data include Age, subtype, gender, race, ajcc_pathologic_tumor_stage, histological_type, histological_grade, treatment_outcome_first_course, radiation treatment adjuvant, sample type, type.
- Types of Survival: 4 types of Survival include OS (overall survival), PFS (progression-free survival), DSS (disease-specific survival), DFS (Disease-free survival). In the data, column names OS, PFS, DSS and DFS represent event information, while OS.time, PFS.time, DSS.time and DFS.time indicate survival time in days.

Let's Check input Data:

```
> #Check dimensions of input data
> dim(data)
     184 19997
[1]
> #check top 3 rows and first 25 columns of the input data
> head(data[1:25],3)
                 Age subtype gender race ajcc pathologic tumor stage
histological type histological grade treatment outcome first course
TCGA-TM-A7CA-01 44.94
                         PN
                              Male WHITE
                                                                 NA
Astrocytoma
                          G2
                                Complete Remission/Response
TCGA-DU-A6S3-01 60.34
                          PN
                              Male WHITE
                                                                 NA
Oligodendroglioma
                                G2
                                                   Stable Disease
TCGA-CS-5390-01 47.80
                          PN Female WHITE
                                                                 NA
Oligodendroglioma
                                                             <NA>
                                G2
               radiation treatment adjuvant sample type type OS OS.time DSS
DSS.time DFI DFI.time PFI PFI.time A1BG
                                          A1CF
                                                   A2M A2ML1 A3GALT2 A4GALT
TCGA-TM-A7CA-01
                                        NO
                                               Primary LGG 0
                                                                 1058
                   0 1058 0.1166 0.0073 65.8414 0.6225 0.2736 0.4515
1058 0
            1058
TCGA-DU-A6S3-01
                                        NO
                                               Primary LGG 0
                       656 0.0782 0.0070 42.7621 0.6587 0.1310 1.7230
656 NA
             NA 0
                                       YES
TCGA-CS-5390-01
                                               Primary LGG 0
               0 NA 0.0877 0.0000 90.7590 0.7980 0.0653 1.0989
NA O
            NA
```

Step 1- Data Processing: This function converts OS time (in days) into months and then removes samples where OS/OS.time information is missing.

Here, we need to provide input data in tsv or txt format. Further, we needs to define col_num (column number at which clinical/demographic and survival information ends,e.g. 20, surv_time (name of column which contain survival time (in days) information, e.g. OS.time) and output file name, e.g. "New data.txt"

#Data Processing:

```
SurvPredPipe::data_process_f(data="Example_TCGA_LGG_FPKM_data.txt",col_num=20,
surv_time="OS.time" , output="New_data.txt")
```

After data processing, data_process_f function will give us a new output file "New_data.txt", which contains 176 samples. Thus, data_process_f function removes 8 samples where OS/OS time information is missing. Besides, here is a new 21st column in the data with column name "OS_month" where OS time is available in months.

Let's Check output of data process f function:

```
> #check output data
> output <- read.table("New data.txt", sep="\t", header=TRUE, row.names=1,
check.names=FALSE)
> dim(output)
    176 19998
[1]
> head(output[1:25],3)
                 Age subtype gender race ajcc pathologic tumor stage
histological type histological grade treatment outcome first course
TCGA-TM-A7CA-01 44.94
                          PN
                               Male WHITE
                                                                   NA
Astrocytoma
                           G2
                                 Complete Remission/Response
                               Male WHITE
TCGA-DU-A6S3-01 60.34
                          PN
                                                                   NA
                                 G2
Oligodendroglioma
                                                    Stable Disease
TCGA-DU-8158-01 57.85
                        <NA> Female WHITE
                                                                   NA
Astrocytoma
                                                        <NA>
               radiation treatment adjuvant sample type OS OS.time DSS
DSS.time DFI DFI.time PFI PFI.time OS month
                                             A1BG
                                                             A2M A2ML1
                                                    A1CF
A3GALT2
TCGA-TM-A7CA-01
                                         NO
                                                Primary LGG 0
                                                                   1058
                         1058
                                    35 0.1166 0.0073 65.8414 0.6225
1058 0
            1058
TCGA-DU-A6S3-01
                                                Primary LGG 0
                                                                    656
                         656
                                   22 0.0782 0.0070 42.7621 0.6587
                                                                    0.1310
656 NA
TCGA-DU-8158-01
                                                Primary LGG 1
                                                                    155
155 NA
                         155
                                    5 0.2789 0.0052 91.1555 0.8302 0.0613
```

Step 2 - Split Data into Training and Test Subset: Before proceeding further, we need to split our data into training and test subset for the purpose of feature selection and model development. Here, we need output from the previous step as an input (which was "New_data.txt"). Next we need to define the fraction (e.g. 0.9) by which we want to split data into training and test. Thus, fraction=0.9 will split data into 90% training and 10% as test set. Besides, we also need to provide training and set output names (e.g. train FPKM.txt,test FPKM.txt)

Split Data into Training and Test subset

```
SurvPredPipe::tr_test_f(data="New_data.txt",fraction=0.9,
train_data="train_FPKM.txt", test_data="test_FPKM.txt")
```

After the train-test split, we got two new outputs: "train_FPKM.txt", "test_FPKM.txt", where, train_FPKM.txt contains 158 samples and test_FPKM.txt contains 18 samples. Thus, tr_test_f function splits data into a 90:10 ratio.

Let's Check output of tr test f function:

```
> #load train_data output
> train_data <- read.table("train_FPKM.txt", sep="\t", header=TRUE,
row.names=1, check.names=FALSE)
> #check dimension of train_data output
> dim(train_data)
[1]    158    19998
> #load test_data output
> test_data <- read.table("test_FPKM.txt", sep="\t", header=TRUE, row.names=1,
check.names=FALSE)
#check dimension of train_data output
> dim(test_data)
[1]    18    19998
```

Step 3 - Data Normalization: Next to select features and develop ML models, data must be normalized. Since, expression is available in terms of FPKM values. Thus, `train_test_normalization_f` function will first convert FPKM value into log scale [log2(FPKM+1) followed by quantile normalization using the "preprocessCore" package. Here, training data will be used as a target matrix for quantile normalization. Here, we need to provide training and test datasets (that we obtained from the previous step of Train/Test Split). Further, we need to provide column number where clinical information ends (e.g. 21) in the input

datasets. Besides, we also need to provide output files names (train_clin_data (which contains only Clinical information of training data), test_clin_data (which contains only Clinical information of training data), train_Normalized_data_clin_data (which contains Clinical information and normalized values of genes of training samples), test_Normalized_data_clin_data (which contains Clinical information and normalized values of genes of test samples).

Data Normalization

```
``SurvPredPipe::train_test_normalization_f(train_data="train_FPKM.txt",test_dat a="test_FPKM.txt", col_num=21, train_clin_data="Train_Clin.txt", test_clin_data="Test_Clin.txt", train_Normalized_data_clin_data="Train_Norm_data.txt", test_Normalized_data_clin_data="Test_Norm_data.txt")``
```

After, running the function, we obtained 4 outputs:

- 1. Train Clin.txt Contains only Clinical features,
- 2. Test Clin.txt- contains only Clinical features of Test samples;
- 3. Train Norm data.txt- Clinical features with normalized values of genes for training samples;
- 4. Test Norm data.txt Clinical features with normalized values of genes for test samples.

```
Let's Check output of train test normalization f function:
> # load normalized training data with clinical information
> train Normalized data clin data <- read.table("Train Norm data.txt",
sep="\t", header=TRUE, row.names=1, check.names=FALSE)
> # Check dimensions of normalized training data
> dim(train Normalized data clin data)
     158 19998
> # View top 2 rows of training data with first 25 columns
> head(train Normalized data clin data[1:25],2)
                  Age subtype gender race ajcc pathologic tumor stage
histological type histological grade treatment outcome first course
TCGA-CS-5396-01 53.11
                           PN Female WHITE
                                                                     NA
Oligodendroglioma
                                  G3
                                                                <NA>
TCGA-DU-A76L-01 54.27
                           ME
                                Male WHITE
                                                                     NA
Oligodendroglioma
                                  G3
                                                Progressive Disease
                radiation treatment adjuvant sample type type OS OS.time DSS
DSS.time DFI DFI.time PFI PFI.time OS month A1BG A1CF
                                                           A2M A2ML1 A3GALT2
TCGA-CS-5396-01
                                         YES
                                                 Primary LGG 0
                                                                     1631
                                                 0 71.912 0.286
                                                                    0.114
1631 NA
              NA
                    0
                          1631
                                     54 0.294
TCGA-DU-A76L-01
                                        <NA>
                                                 Primary LGG 1
                                                                      814
814 NA
                          410
                                    27 0.144
                                                0 233.872 0.108
```

> #load normalized test data with clinical information

```
> test Normalized data clin data<- read.table("Test Norm data.txt", sep="\t",
header=TRUE, row.names=1, check.names=FALSE)
> # Check dimensions of normalized test data
> dim(test Normalized data clin data)
      18 19998
> # View top 2 rows of test data with first 25 columns
> head(test Normalized data clin data[1:25],2)
                 Age subtype gender race ajcc pathologic tumor stage
histological type histological grade treatment outcome first course
TCGA-DH-A669-02 70.66
                          PN
                              Male WHITE
                                                                  NA
Oligodendroglioma
                                 G3
                                                   Stable Disease
TCGA-WY-A859-01 34.58
                        <NA> Female WHITE
                                                                  NA
Astrocytoma
                           G2
                                             Stable Disease
               radiation treatment adjuvant sample type type OS OS.time DSS
DSS.time DFI DFI.time PFI PFI.time OS month A1BG A1CF
                                                         A2M A2ML1 A3GALT2
                                             Recurrent LGG 1
TCGA-DH-A669-02
                                       <NA>
919 NA
             NA
                         260
                                   30 0.301 0.009 60.121 0.401
                                                                0.000
TCGA-WY-A859-01
                                       <NA>
                                               Primary LGG 0
                         1213 40 0.081 0.000 31.696 1.748
                                                                 0.095
1213 NA
             NA
                   0
```

Step 4a - Prognostic Index (PI) Score Calculation:

Next to create a survival model, we will create a Prognostic Index (PI) Score. PI score is calculated based on the expression of the features selected by the LASSO regression model and their beta coefficients. For instance, 5 features (G1, G2, G3, G4, and G5 and their coefficient values are B1, B2, B3, B4, and B5, respectively) selected by the LASSO method. Then PI score will be computed as following:

$$PI score = G1*B1 + G2*B2 + G3 * B3 + G4*B4 + G5*B5$$

Here, we need to provide Normalized training (Train_Norm_data.txt) and test data (Test_Norm_data.txt)as input data that we have obtained from the previous function "train_test_normalization_f". Further, we need to provide col_num n column number at which clinical features ends (e.g. 21), nfolds (number of folds e.g. 5) for the LASSO regression method to select features. We implemented LASSO using the "glmnet" package. Further, we need to provide surv_time (name of column containing survival time in months, e.g. OS_month) and surv_event (name of column containing survival event information, e.g. OS) information in the data. Besides, we also need to provide names and training and test output file names to store data containing LASSO genes and PI values.

Feature Selection using LASSO Regression and Prognostic Index (PI) Score Calculation

```
SurvPredPipe::Lasso_PI_scores_f(train_data="Train_Norm_data.txt",test_data="Test
t_Norm_data.txt", nfolds=5, col_num=21, surv_time="OS_month", surv_event="OS"
, train PI data="Train PI data.txt", test PI data="Test PI data.txt")
```

Thus, the Lasso PI scores f gave us following outputs:

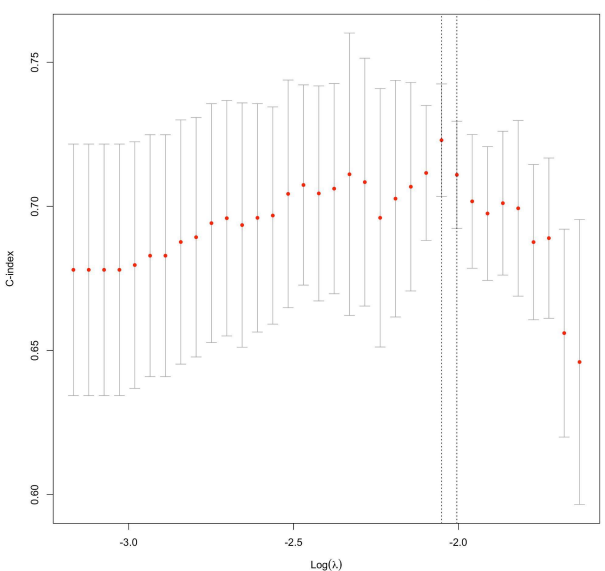
1. **Train_Lasso_key_variables.txt**: List of features selected by LASSO and their beta coefficient values

Let's Check output of Lasso PI scores f function:

```
> # Load LASSO selected variables file
> Lasso key variables <- read.table("Train Lasso key variables.txt", sep="\t",
header=TRUE, row.names=1, check.names=FALSE)
> # Check dimension
> dim(Lasso key variables)
[1] 12 1
> # view top of file
> head(Lasso key variables)
         coeff
         0.037
ALG6
ARHGAP11A 0.096
DESI1 -0.028
         0.101
GALNT7
GJD3 0.284
GPC1 0.005
```

2. Train Cox Lasso Regression lamda plot.jpeg: Lasso Regression Lambda plot.





- 3. **Train_PI_data.txt**: It contains expression of genes selected by LASSO and PI score in the last column for training samples.
- 4. **Test_PI_data.tx**t: It contains expression of genes selected by LASSO and PI score in the last column for test samples.

```
> # Load train data containing LASSO selected genes and PI Value
> train_PI_data <- read.table("Train_PI_data.txt", sep="\t", header=TRUE,
row.names=1, check.names=FALSE)</pre>
```

> #check dimensions
> dim(train_PI_data)
[1] 158 15

```
> #View top 2 rows
> head(train PI data,2)
               OS OS month ALG6 ARHGAP11A DESI1 GALNT7 GJD3
                                                                GPC1
H2BC5 HOXD12 RNF185 TANGO2
                            UNG ZNF648
                                    1.241 24.866 5.760 0.074 20.434
TCGA-CS-5396-01 0
                        54 4.705
4.004 0.019 19.033 1.660 11.262 0.115 0.287790
TCGA-DU-A76L-01 1
                        27 3.392
                                    3.124 9.048 2.927 2.521 61.482
5.335 0.087 9.067 0.897 14.827 0.055 1.617081
> # Load test data containing LASSO selected genes and PI Value
> test PI data<- read.table("Test PI data.txt", sep="\t", header=TRUE,
row.names=1, check.names=FALSE)
> #check dimensions
> dim(test PI data)
[1] 18 15
> #View top 2 rows
> head(test PI data,2)
               OS OS month ALG6 ARHGAP11A DESI1 GALNT7 GJD3
                                                                GPC1
H2BC5 HOXD12 RNF185 TANGO2
                             UNG ZNF648
                                              PΙ
TCGA-DH-A669-02 1
                                   1.288 18.533 2.323 0.295 54.726
                        30 2.919
8.651 0.136 11.046 1.780 20.197 0.089 0.486198
TCGA-WY-A859-01 0
                        40 1.433
                                   0.279 23.926 1.066 0.117 39.250
4.240 0.006 13.179 1.829 9.860 0.003 -0.275303
```

Step 4b - Univariate Survival Significant Feature Selection:

Besides PI score, with the "Univariate_sig_features_f" function of SurvPredPipe package, we can select significant (p-value <0.05) features based on univariate survival analysis. These features are selected based on their capability to stratify high-risk and low-risk survival groups using the cut off value of their median expression.

Here, we need to provide Normalized training (Train_Norm_data.txt) and test data (Test_Norm_data.txt)as input data that we have obtained from the previous function "train_test_normalization_f". Further, we need to provide a "col_num" (e.g 21) column number at which clinical features end. Further, we need to provide surv_time (name of column containing survival time in months, e.g. OS_month) and surv_event (name of column containing survival event information, e.g. OS) information in the data. Besides, we also

need to provide names and training and test output file names to store data containing expression of selected genes.

Feature selection using Univariate Survival Analysis

```
`SurvPredPipe::Univariate_sig_features_f(train_data="Train_Norm_data.txt", test_data="Test_Norm_data.txt", col_num=21, surv_time="OS_month", surv_event="OS", output_univariate_train="Train_Uni_sig_data.txt", output_univariate_test="Test_Uni_sig_data.txt")`
```

Thus, the Univariate sig features f function gave us following outputs:

1. **Univariate_Survival_Significant_genes_List.txt**: a table of univariate significant genes along with their corresponding coefficient values, HR value, P-values, C-Index values.

Let's Check output of Lasso_PI_scores_f function:

```
> #Load list of significant genes selected by Univariate Survival analysis
> Univariate Survival Significant genes List<-
read.table("Univariate Survival Significant genes List.txt", sep="\t",
header=TRUE, row.names=1, check.names=FALSE)
> # Check dimensions
> dim(Univariate Survival Significant genes List)
[1] 2391
> #View top 5 rows of Univariate Significant genes results
> head(Univariate Survival Significant genes List,5)
             Beta HR P-value GP1 GP2 Hr-Inv-lst Concordance
Std Error
A2ML1 -0.9255326 0.3963203 0.016055099 20 138 2.5232117 0.5578119
0.03819547
AADACL2 1.3245266 3.7604047 0.001723308 148 10 0.2659288
                                                           0.5620198
0.03521353
AARS1 -0.7793207 0.4587175 0.016107789 74 84 2.1799910
                                                           0.6337358
0.04249702
ABCA12 1.3690240 3.9315117 0.004805718 150 8 0.2543551 0.5539700
0.03202036
      0.9601084 2.6119796 0.038305760 136 22 0.3828514 0.5554336
ABCA6
0.03656689
```

- 2. **Train_Uni_sig_data.txt:** It contains expression of significant genes selected by univariate survival analysis for training samples.
- > #Load training data with univariate significant genes

```
> train Uni sig data <- read.table("Train Uni sig data.txt", sep="\t",
header=TRUE, row.names=1, check.names=FALSE)
> # Check dimensions of data
> dim(train Uni sig data )
[1] 158 2391
> # View top rows of training data
> head(train Uni sig data[1:20],2)
              A2ML1 AADACL2 AARS1 ABCA12 ABCA6 ABCC11 ABCC3 ABCC9 ABHD10
      ABI1 ABI3BP ABLIM1 AC004997.1 AC005832.4 AC006030.1 AC011455.2
ABHD11
AC012309.1
TCGA-CS-5396-01 0.286 0 33.197 0.214 0.403 0.157 1.152 0.630 12.924
6.461 22.471 0.541 14.608 0.000 0
                                               0.006
0.015
TCGA-DU-A76L-01 0.108 0 26.215 0.298 0.231 0.059 5.263 0.371 9.812
7.890 14.744 0.877 9.228 0.005
                                    0
                                                 0.010
0.007
              AC091057.5 AC093323.1
TCGA-CS-5396-01 0 5.222
TCGA-DU-A76L-01
                     0
                             8.101
```

3. **Test_Uni_sig_data.txt:** It contains expression of significant genes selected by univariate survival analysis for test samples.

```
> #Load test data with univariate significant genes
> test Uni sig data<- read.table("Test Uni sig data.txt", sep="\t",
header=TRUE, row.names=1, check.names=FALSE)
> # Check dimensions of data
> dim(test Uni sig data)
[1] 18 2391
> # View top rows of test data
> head(test Uni sig data[1:20],2)
            A2ML1 AADACL2 AARS1 ABCA12 ABCA6 ABCC11 ABCC3 ABCC9 ABHD10
       ABI1 ABI3BP ABLIM1 AC004997.1 AC005832.4 AC006030.1 AC011455.2
ABHD11
TCGA-DH-A669-02 0.401 0 37.720 0.078 0.011 0.073 0.338 0.194 14.795
17.986 20.666 0.272 13.782 0.000 0
                                           0
6.817 31.915 0.341 17.683 0.002 0
                                              Ω
            AC091057.5 AC093323.1
                   0
TCGA-DH-A669-02
                         4.955
TCGA-WY-A859-01
                0
                        10.609
```

Step 5 - Prediction model development for survival probability of patients

After selecting significant or key features using LASSO or Univariate survival analysis, next we want to develop an ML prediction model to predict survival probability of patients.

MTLR_pred_model_f function of SurvPredPipe give us multiple options to develop models including Only Clinical features (Model_type=1), PI score (Model_type=2), PI Score + Clinical features (Model_type=3), Significant Univariate features (Model_type=4), Significant Univariate features Clinical features (Model_type=5) using MTLR package. Further, here, we were interested in developing a model based on PI score. Thus, we need to provide following inputs:

(1) Training data with only clinical features, (2) Test data with only clinical features, (3) Model type (e.g. 2, since we want to develop model based on PI score), (4) Training data with PI score, (5) Test data with PI score, (6) Clin_Feature_List (e.g. Key_PI_list.txt), a list of features which will be used to build model. Furthermore, we also need to provide surv_time (name of column containing survival time in months, e.g. OS_month) and surv_event (name of column containing survival event information, e.g. OS) information in the clinical data

Survival Model Development and Prediction of Survival, survival probability for Individual Patients using Selected Features employing MTLR

Survival Model Development and Prediction of Survival, survival probability for Individual Patients using Selected Features employing MTLR

```
# Model for only Clinical features
SurvPredPipe::MTLR pred model f(train clin data = "Train Clin.txt",
test clin data = "Test Clin.txt", Model type = 1, train features data =
"Train Clin.txt", test features data = "Test Clin.txt",
Clin Feature List="Key Clin feature list.txt", surv time="OS month",
surv event="OS")
# Model for PI
SurvPredPipe::MTLR pred model f(train clin data = "Train Clin.txt",
test clin data = "Test Clin.txt", Model type = 2, train features data =
"Train PI data.txt", test features data = "Test PI data.txt",
Clin Feature List="Key PI list.txt", surv time="OS month", surv event="OS")
# Model for Clinical features + PI
SurvPredPipe::MTLR pred model f(train clin data = "Train Clin.txt",
test clin data = "Test Clin.txt", Model type = 3, train features data =
"Train PI data.txt", test features data = "Test PI data.txt",
Clin Feature List="Key Clin features with PI list.txt", surv time="OS month",
surv event="OS")
```

```
# Model for univariate features
SurvPredPipe::MTLR_pred_model_f(train_clin_data = "Train_Clin.txt",
test_clin_data = "Test_Clin.txt", Model_type = 4, train_features_data =
"Train_Uni_sig_data.txt", test_features_data = "Test_Uni_sig_data.txt",
Clin_Feature_List="Key_univariate_features_list.txt", surv_time="OS_month",
surv_event="OS")

# Model for Univariate + Clinical features
SurvPredPipe::MTLR_pred_model_f(train_clin_data = "Train_Clin.txt",
test_clin_data = "Test_Clin.txt", Model_type = 5, train_features_data =
"Train_Uni_sig_data.txt", test_features_data = "Test_Uni_sig_data.txt",
Clin_Feature_List="Key_univariate_features_with_Clin_list.txt",
surv_time="OS_month", surv_event="OS")
```

After, implementing MTLR_pred_model_f function, we got following outputs:

- 1. Model with PI.RData: Model on training data
- 2. survCurves_data.txt: Table containing predicted survival probability of each patient at different time points. This data can be further used to plot the survival curve of patients.

Let's check output of MTLR_pred_model_f function:

- > # Load Survival curve data
 > survCurves_data<- read.table("survCurves_data.txt", sep="\t", header=TRUE, check.names=FALSE)</pre>
- > #Check dimension of Survival curve data
 > dim(survCurves_data)
 [1] 15 19
- > #View top 5 rows of Survival curve data
- > head(survCurves data,5)

time point TCGA-DH-A669-02 TCGA-WY-A859-01 TCGA-TQ-A7RN-01 TCGA-FG-A6IZ-01 TCGA-HW-8319-01 TCGA-TM-A7CF-02 TCGA-DU-7010-01 TCGA-HT-7686-01 TCGA-WY-A858-01 1 0.000000 1.0000000 1.0000000 1.0000000 1.0000000 1.0000000 1.000000 1.0000000 1.0000000 1.0000000 0.9909155 3.466667 0.9972704 0.9773009 0.9955466 0.9961896 0.9905094 0.9960141 0.9939210 0.9959694 3 8.000000 0.9719876 0.9916419 0.9295280 0.9863267 0.9877695 0.9883111 0.9707273 0.9813015 0.9876316 4 14.000000 0.9514804 0.9852768 0.8798802 0.9760716 0.9795008 0.9493315 0.9785641 0.9674200 0.9783257 5 18.000000 0.9398923 0.9814651 0.8533639 0.9700671 0.9731460 0.9743045 0.9372695 0.9594131 0.9728513 TCGA-TQ-A8XE-01 TCGA-DU-6408-01 TCGA-HT-7874-01 TCGA-DU-8162-01

TCGA-KT-A7W1-01 TCGA-VV-A829-01 TCGA-VM-A8CA-01 TCGA-HT-A61C-01 TCGA-S9-A6WN-01

```
1.0000000 1.0000000 1.0000000 1.0000000
1.0000000
            1.0000000
                         1.0000000
                                      1.0000000
                                                   1.0000000
      0.9921240 0.9962126 0.9966324
                                            0.9942035
0.9683777 0.9979307 0.9971536
                                      0.9765579
                                                   0.9931295
                  0.9883821
                               0.9896764
      0.9757360
                                            0.9821754
0.9014997
            0.9936732
                         0.9912822
                                      0.9271999
                                                   0.9788516
      0.9578822
                   0.9796236
                                0.9818651
                                             0.9689217
0.8333833
         0.9888145 0.9846516 0.8759960
                                                   0.9632163
                  0.9744564
                                0.9772317
                                             0.9612589
      0.9477176
                                                  0.9542526
0.7979822
            0.9858691
                         0.9806885
                                      0.8487143
```

3. mean_median_survival_time_data.txt: Table containing predicted mean and median survival time of each patient in the test data. This data can be further used for bar plots.

```
> #load mean median survival time data
> mean median survival time data<-
read.table("mean median survival time data.txt", sep="\t", header=TRUE,
check.names=FALSE)
> # Check dimension of mean median survival time data
> dim(mean median survival time data)
[1] 18 3
> #View top 5 rows of mean median survival time data
> head(mean median survival time data,5)
              IDs
                      Mean
                           Median
1 TCGA-DH-A669-02 67.84700 60.52883
2 TCGA-WY-A859-01 90.91569 85.75383
3 TCGA-TQ-A7RN-01 46.94777 36.31438
4 TCGA-FG-A6IZ-01 81.98710 81.61567
5 TCGA-HW-8319-01 84.05714 82.71186
```

4. Error_mat_for_Model.txt: Table containing performance parameters obtained on test data based on prediction model. It contains IBS score (Integrated Brier Score) =0.192, C-Index =0.81.

Step 6 - Survival curves/plots for individual patient

Next to visualize survival of patients, we will plot survival curve plots using the surv_curve_plots_f function based on the data "survCurves_data.txt" that we obtained from the previous step after running the MTLR_pred_model_f function. Further, the surv_curve_plots_f function also allows highlighting a specific patient on the curve. Thus the function needs only two inputs: 1) Surv_curve_data, (2) Sample ID of a specific patient (e.g. TCGA-TQ-A8XE-01) that needs to be highlighted.

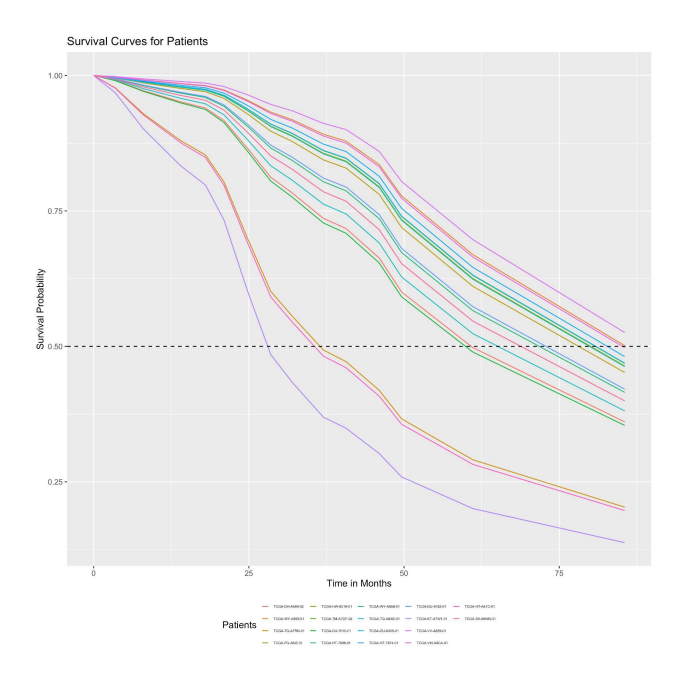
#Create Survival curves/plots for individual patients

```
SurvPredPipe::surv_curve_plots_f(Surv_curve_data="survCurves_data.txt",
selected sample="TCGA-TQ-A8XE-01")
```

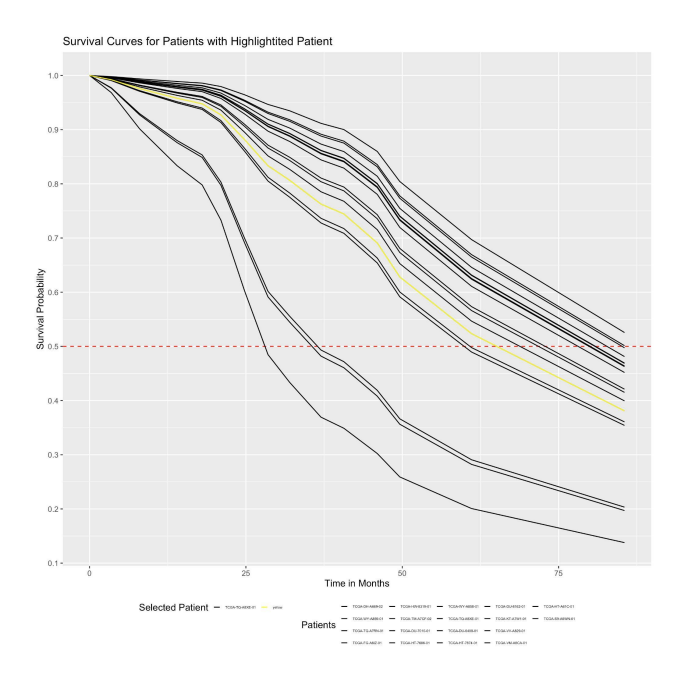
Let's check output of surv_curve_plots_f function:

Here, we obtained two output plots:

1. Survival curves for all patients in the test data with different colors



2. Survival curves for all patients (in black) and highlighted patient (yellow) in the test data



Step 7 - Bar Plot for predicted mean and median survival time of individual patients

Next to visualize predicted survival time of patients, we will plot barplot for mean/median using "mean_median_surv_barplot_f" function based on the data that we obtained from step 5 after running the MTLR_pred_model_f function. Further, the mean_median_surv_barplot_f function also allows highlighting a specific patient on the curve. Thus the function needs only two inputs: 1) surv_mean_med_data, (2) Sample ID of a specific patient (e.g. TCGA-TQ-A8XE-01) that needs to be highlighted.

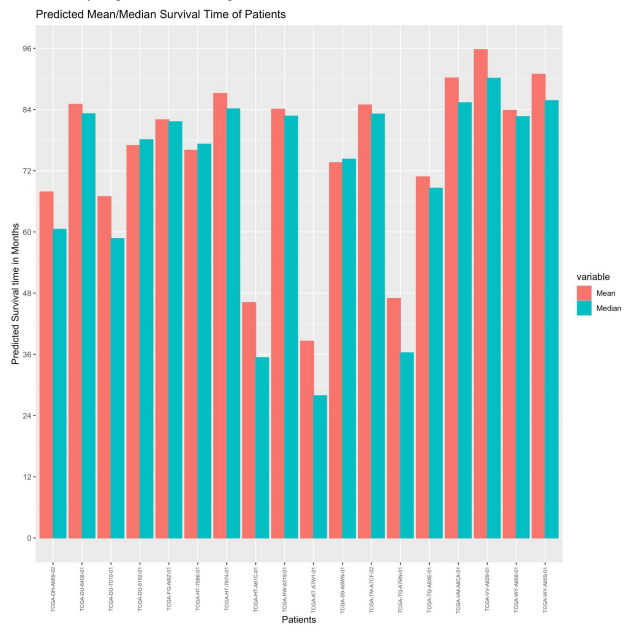
#Create Bar Plot for predicted mean and median survival time

SurvPredPipe::mean_median_surv_barplot_f(surv_mean_med_data="mean_median_surviv
al_time_data.txt", selected_sample="TCGA-TQ-A8XE-01")

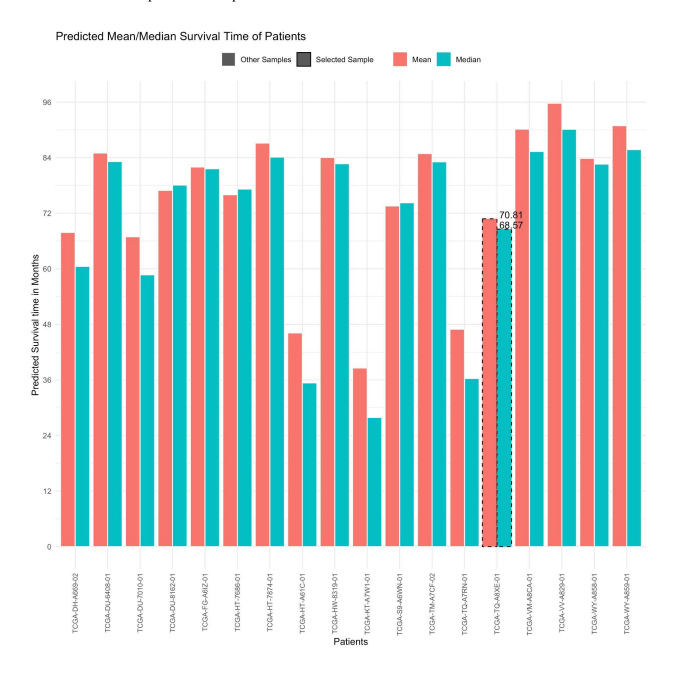
Let's check output of mean median surv barplot f function:

Here, we obtained two output plots:

1. Barplot for all patients in the test data, where the red color bar represents mean survival and cyan/green color bar represents median survival time.



2. Barplot for all patients with a highlighted patient (dashed black outline) in the test data. It shows this patient has a predicted mean and median survival is 81.58 and 75.50 months.



Step 8- Nomogram based on Key features

Next, the Nomogram_generate_f function of SurvPredPipe also provides an option to generate a nomogram plot based on user defined clinical and other features in the data. For instance, we will generate a nomogram based on 6 features (Age, gender, race, histological_type, sample_type, PI). Here, we will provide data containing all the features (Samples in rows and features in columns) (e.g. Train_Data_Nomogram_input.txt) and a list of features (feature_list_for_Nomogram.txt) based on which we want to generate a nomogram. Further, we

also need to provide surv_time (name of column containing survival time in months, e.g. OS_month) and surv_event (name of column containing survival event information, e.g. OS) information in the data.

#Create Nomogram based on Key features

```
SurvPredPipe::Nomogram_generate_f(data="Train_Data_Nomogram_input.txt",
Feature_List="feature_list_for_Nomogram.txt", surv_time="OS_month",
surv_event="OS")
```

Let's check output of Nomogram_generate_f function:

Here, we will get a Nomogram based on features that we provide. This nomogram can predict Risk (Event risk, eg, Death), 1-year, 3-year, 5-year and 10 years survival of patients.

