

About Data

The data contains the information collected from the Mayo Clinic trial in primary biliary cirrhosis (PBC) of the liver conducted between 1974 and 1984. A description of the clinical background for the trial and the covariates recorded here is in Chapter 0, especially Section 0.2 of Fleming and Harrington, Counting Processes and Survival Analysis, Wiley, 1991. A more extended discussion can be found in Dickson, et al., Hepatology 10:1-7 (1989) and in Markus, et al., N Eng J of Med 320:1709-13 (1989).

A total of 424 PBC patients, referred to Mayo Clinic during that ten-year interval, met eligibility criteria for the randomized placebo-controlled trial of the drug D-penicillamine. The first 312 cases in the dataset participated in the randomized trial and contain largely complete data. The additional 112 cases did not participate in the clinical trial but consented to have basic measurements recorded and to be followed for survival. Six of those cases were lost to follow-up shortly after diagnosis, so the data here are on an additional 106 cases as well as the 312 randomized participants.

The dataset consists of following columns :

1. ID: unique identifier
2. N_Days: number of days between registration and the earlier of death, transplantation, or study analysis time in July 1986
3. Status: status of the patient C (censored), CL (censored due to liver tx), or D (death)
4. Drug: type of drug D-penicillamine or placebo
5. Age: age in [days]
6. Sex: M (male) or F (female)
7. Ascites: presence of ascites N (No) or Y (Yes)
8. Hepatomegaly: presence of hepatomegaly N (No) or Y (Yes)
9. Spiders: presence of spiders N (No) or Y (Yes)
10. Edema: presence of edema N (no edema and no diuretic therapy for edema), S (edema present without diuretics, or edema resolved by diuretics), or Y (edema despite diuretic therapy)
11. Bilirubin: serum bilirubin in [mg/dl]
12. Cholesterol: serum cholesterol in [mg/dl]
13. Albumin: albumin in [gm/dl]
14. Copper: urine copper in [ug/day]
15. Alk_Phos: alkaline phosphatase in [U/liter]
16. SGOT: SGOT in [U/ml]
17. Triglycerides: triglycerides in [mg/dl]
18. Platelets: platelets per cubic [ml/1000]
19. Prothrombin: prothrombin time in seconds [s]
20. Stage: histologic stage of disease (1, 2, 3, or 4)

Imports

In [1]:

```
# %load_ext google.colab.data_table
import numpy as np
import pandas as pd
import matplotlib.pyplot as plt
import seaborn as sns
```

In [2]:

```
##@title Default title text
df = pd.read_csv(r'C:\Users\91701\Desktop\liver prediction/cirrhosis.csv', index_col='ID')
df.head()
```

Out[2]:

	N_Days	Status	Drug	Age	Sex	Ascites	Hepatomegaly	Spiders	Edema	Bilirubi
ID										
1	400	D	D- penicillamine	21464	F	Y	Y	Y	Y	14.
2	4500	C	D- penicillamine	20617	F	N	Y	Y	N	1.
3	1012	D	D- penicillamine	25594	M	N	N	N	S	1.
4	1925	D	D- penicillamine	19994	F	N	Y	Y	S	1.
5	1504	CL	Placebo	13918	F	N	Y	Y	N	3.

In [3]:

df.info()

```

<class 'pandas.core.frame.DataFrame'>
Int64Index: 418 entries, 1 to 418
Data columns (total 19 columns):
 #   Column                Non-Null Count  Dtype
---  -
 0   N_Days                418 non-null    int64
 1   Status                418 non-null    object
 2   Drug                  312 non-null    object
 3   Age                   418 non-null    int64
 4   Sex                   418 non-null    object
 5   Ascites               312 non-null    object
 6   Hepatomegaly          312 non-null    object
 7   Spiders               312 non-null    object
 8   Edema                 418 non-null    object
 9   Bilirubin             418 non-null    float64
10   Cholesterol            284 non-null    float64
11   Albumin               418 non-null    float64
12   Copper                310 non-null    float64
13   Alk_Phos              312 non-null    float64
14   SGOT                  312 non-null    float64
15   Tryglicerides         282 non-null    float64
16   Platelets             407 non-null    float64
17   Prothrombin           416 non-null    float64
18   Stage                 412 non-null    float64
dtypes: float64(10), int64(2), object(7)
memory usage: 65.3+ KB

```

This data set has about 19 features. These features are related to the patient's details like age, sex, etc. and patient's blood tests like prothrombin, triglycerides, platelets levels, etc. All these factors help in understanding a patient's chances of liver cirrhosis.

we have some NA values in our data, lets look at some statistical summary of numerical columns in our dataset.

In [4]:

df.describe()

Out[4]:

	N_Days	Age	Bilirubin	Cholesterol	Albumin	Copper	Alk_Ph
count	418.000000	418.000000	418.000000	284.000000	418.000000	310.000000	312.0000
mean	1917.782297	18533.351675	3.220813	369.510563	3.497440	97.648387	1982.6557
std	1104.672992	3815.845055	4.407506	231.944545	0.424972	85.613920	2140.3888
min	41.000000	9598.000000	0.300000	120.000000	1.960000	4.000000	289.0000
25%	1092.750000	15644.500000	0.800000	249.500000	3.242500	41.250000	871.5000
50%	1730.000000	18628.000000	1.400000	309.500000	3.530000	73.000000	1259.0000
75%	2613.500000	21272.500000	3.400000	400.000000	3.770000	123.000000	1980.0000
max	4795.000000	28650.000000	28.000000	1775.000000	4.640000	588.000000	13862.4000

We have some missing values in our data, lets see how many and in which columns.

In [5]:

```
df.isna().sum()
```

Out[5]:

N_Days	0
Status	0
Drug	106
Age	0
Sex	0
Ascites	106
Hepatomegaly	106
Spiders	106
Edema	0
Bilirubin	0
Cholesterol	134
Albumin	0
Copper	108
Alk_Phos	106
SGOT	106
Tryglicerides	136
Platelets	11
Prothrombin	2
Stage	6

dtype: int64

Handling Missing Values

This is a problem, we could just get rid of all examples with NA values, but in this case our case of small dataset we cannot afford that.

We will impute the missing entries with some statistical calculations.

We have two different types of data

1. Numerical data (Age, Cholesterol, Platelets.. etc)
2. Categorical Data (Drug, Sex, Spiders..etc)

We will have to use different imputation for each type

1. For the numerical type we can use mean or median. In this case we will go with median to avoid skewing in the presence of outliers
2. For Categorical type we will impute the most frequent class.

In [6]:

```
# For Numerical Type
df.select_dtypes(include=['int64', 'float64']).isna().sum()
```

Out[6]:

```
N_Days      0
Age          0
Bilirubin    0
Cholesterol  134
Albumin      0
Copper       108
Alk_Phos     106
SGOT         106
Tryglicerides 136
Platelets    11
Prothrombin   2
Stage        6
dtype: int64
```

In [7]:

```
df.select_dtypes(include=['int64', 'float64']).isna().sum()
df_num_col = df.select_dtypes(include=['int64', 'float64']).columns
for c in df_num_col:
    df[c].fillna(df[c].median(), inplace=True)

df.select_dtypes(include=['int64', 'float64']).isna().sum()
```

Out[7]:

```
N_Days      0
Age          0
Bilirubin    0
Cholesterol  0
Albumin      0
Copper       0
Alk_Phos     0
SGOT         0
Tryglicerides 0
Platelets    0
Prothrombin  0
Stage        0
dtype: int64
```

In [8]:

```
# For Categorical type
df.select_dtypes(include=('object')).isna().sum()
```

Out[8]:

```
Status      0
Drug        106
Sex          0
Ascites     106
Hepatomegaly 106
Spiders     106
Edema        0
dtype: int64
```

In [9]:

```
df_cat_col = df.select_dtypes(include=('object')).columns
for c in df_cat_col:
    df[c].fillna(df[c].mode().values[0], inplace=True)

df.select_dtypes(include=('object')).isna().sum()
```

Out[9]:

```
Status      0
Drug         0
Sex          0
Ascites      0
Hepatomegaly 0
Spiders      0
Edema        0
dtype: int64
```

Exploratory Data Analysys

Lets dive into the data and visualize it, this often reveals interesting patterns.

First lets take a look at how many examples per calss do we have in our dataset.

In [10]:

```
df['Stage'].value_counts()
```

Out[10]:

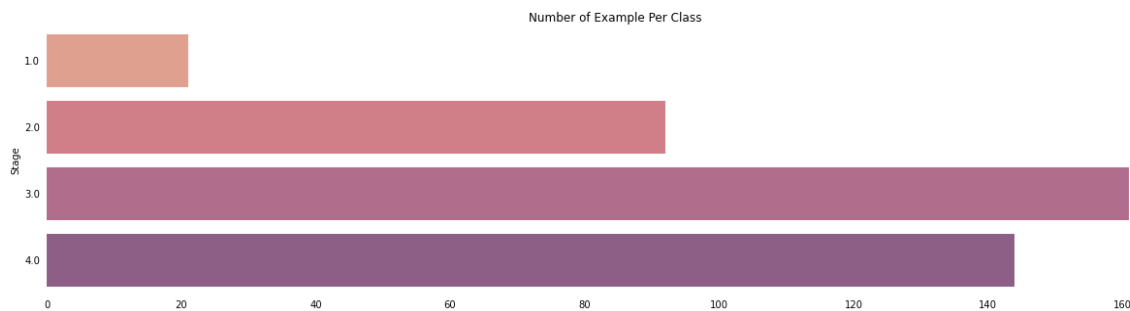
```
3.0    161
4.0    144
2.0     92
1.0     21
Name: Stage, dtype: int64
```

In [11]:

```
plt.figure(figsize=(21,5))
sns.countplot(y=df['Stage'], palette="flare", alpha=0.8, )
sns.despine(top=True, right=True, bottom=True, left=True)
plt.tick_params(axis='both', which='both', bottom=False, top=False, left=False)
plt.xlabel('')
plt.title('Number of Example Per Class')
```

Out[11]:

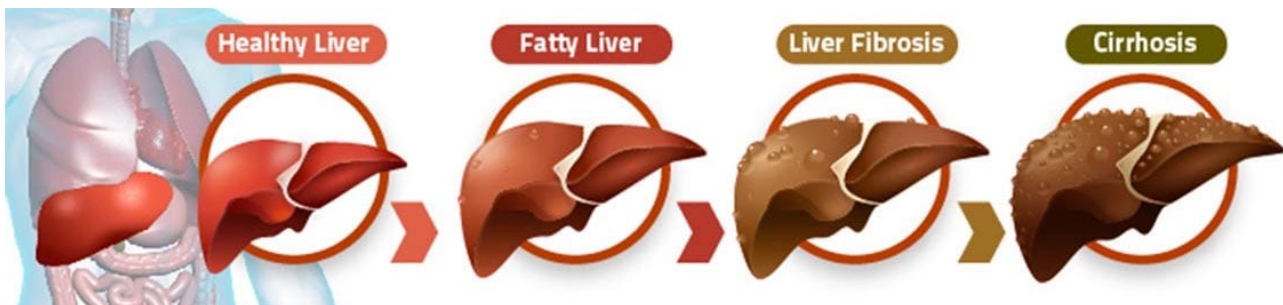
Text(0.5, 1.0, 'Number of Example Per Class')



As we can observe we have class imbalances in our dataset i.e some classes have more examples than other. This could make it difficult for our model to train and achieve desired score. No worries, we can fix that later.

Setting up Target and Features

For this demonstration we will keep things simple by predicting one of the two classes i.e (Cirrhosis and No Cirrhosis). I have another project that predicts the stage of the disease, converting this problem into a multiclass classification.



In [12]:

```
# Converting Target categories into integers 1 for Cirrhosis, 0 otherwise
df['Stage'] = np.where(df['Stage'] == 4, 1, 0)
```

Lets observe some Features with their relation with the disease

In [13]:

```
plt.figure(figsize=(21.2,10))

plt.subplot(2,3,1)
sns.countplot(x=df['Stage'], hue=df['Sex'], palette='Blues', alpha=0.9)
sns.despine(top=True, right=True, bottom=True, left=True)
plt.tick_params(axis='both', which='both', bottom=False, top=False, left=False)
plt.xlabel('')
plt.title('Disease Stage Across Gender')

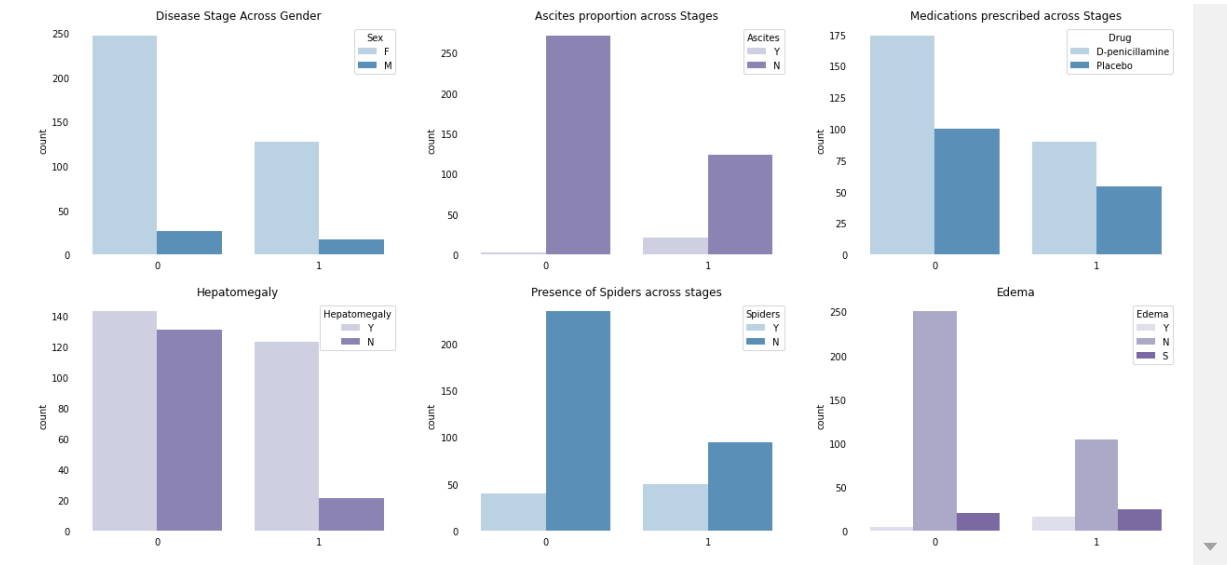
plt.subplot(2,3,2)
sns.countplot(x=df['Stage'], hue=df['Ascites'], palette='Purples', alpha=0.9)
sns.despine(top=True, right=True, bottom=True, left=True)
plt.tick_params(axis='both', which='both', bottom=False, top=False, left=False)
plt.xlabel('')
plt.title('Ascites proportion across Stages')

plt.subplot(2,3,3)
sns.countplot(x=df['Stage'], hue=df['Drug'], palette='Blues', alpha=0.9)
sns.despine(top=True, right=True, bottom=True, left=True)
plt.tick_params(axis='both', which='both', bottom=False, top=False, left=False)
plt.xlabel('')
plt.title('Medications prescribed across Stages');

plt.subplot(2,3,4)
sns.countplot(x=df['Stage'], hue=df['Hepatomegaly'], palette='Purples', alpha=0.9)
sns.despine(top=True, right=True, bottom=True, left=True)
plt.tick_params(axis='both', which='both', bottom=False, top=False, left=False)
plt.xlabel('')
plt.title('Hepatomegaly');

plt.subplot(2,3,5)
sns.countplot(x=df['Stage'], hue=df['Spiders'], palette='Blues', alpha=0.9)
sns.despine(top=True, right=True, bottom=True, left=True)
plt.tick_params(axis='both', which='both', bottom=False, top=False, left=False)
plt.xlabel('')
plt.title('Presence of Spiders across stages');

plt.subplot(2,3,6)
sns.countplot(x=df['Stage'], hue=df['Edema'], palette='Purples', alpha=0.9)
sns.despine(top=True, right=True, bottom=True, left=True)
plt.tick_params(axis='both', which='both', bottom=False, top=False, left=False)
plt.xlabel('')
plt.title('Edema');
```



There are some interesting insights if we observe closely. Take the case at Ascites, we observe that the risk of disease is higher with increase in Ascites. Also, presence of spiders has a positive relation with disease risk.

In [14]:

```
#@title Distribution Plots
plt.figure(figsize=(20,15))

plt.subplot(3,3,1)
sns.kdeplot(df['Cholesterol'], hue=df['Stage'], fill=True, palette='Purples')
sns.despine(top=True, right=True, bottom=True, left=True)
plt.tick_params(axis='both', which='both', bottom=False, top=False, left=False)
plt.xlabel('')
plt.title('Cholesterol Distribution in stages');

plt.subplot(3,3,2)
sns.kdeplot(df['Bilirubin'], hue=df['Stage'], fill=True, palette='Blues', common_norm=True)
sns.despine(top=True, right=True, bottom=True, left=True)
plt.tick_params(axis='both', which='both', bottom=False, top=False, left=False)
plt.xlabel('')
plt.title('Bilirubin');

plt.subplot(3,3,3)
sns.kdeplot(df['Tryglicerides'], hue=df['Stage'], fill=True, palette='Purples', common_norm=True)
sns.despine(top=True, right=True, bottom=True, left=True)
plt.tick_params(axis='both', which='both', bottom=False, top=False, left=False)
plt.xlabel('')
plt.title('Tryglicerides');

plt.subplot(3,3,4)
sns.kdeplot(df['Age'], hue=df['Stage'], fill=True, palette='Blues', common_norm=True)
sns.despine(top=True, right=True, bottom=True, left=True)
plt.tick_params(axis='both', which='both', bottom=False, top=False, left=False)
plt.xlabel('')
plt.title('Age Distribution in stages');

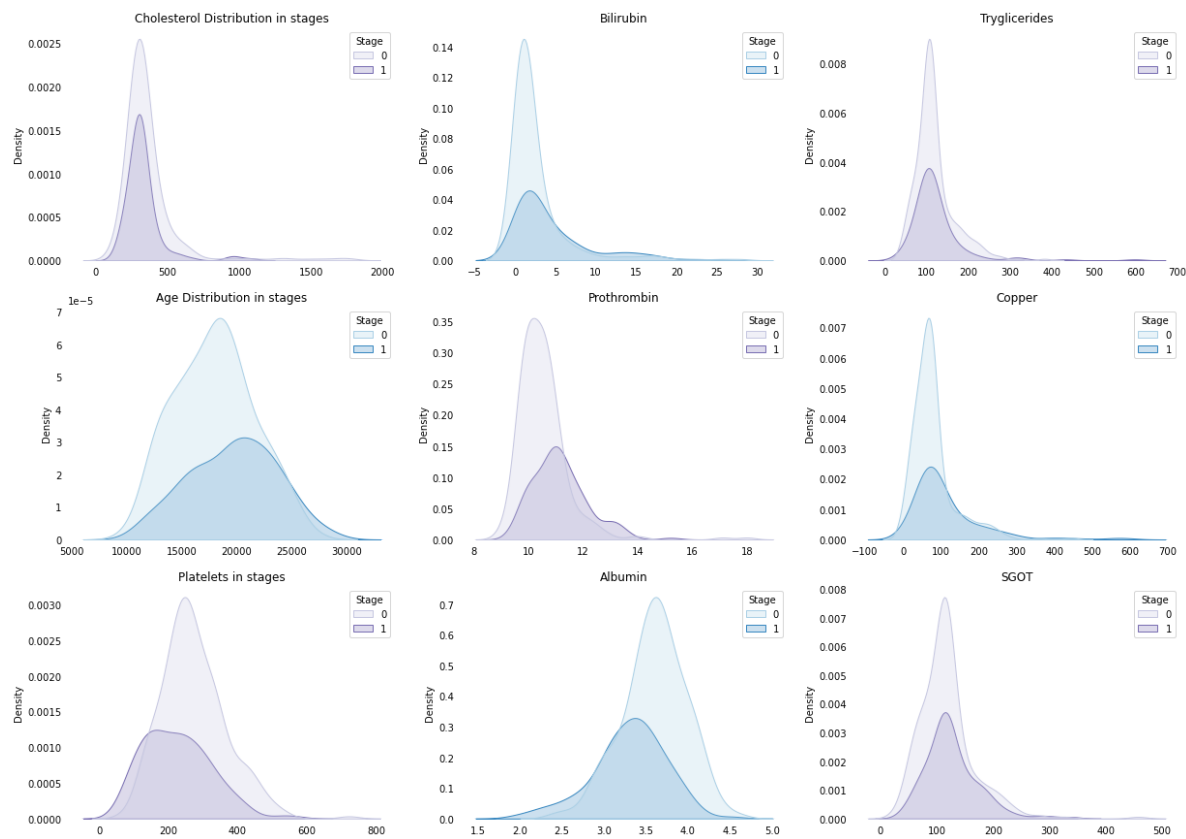
plt.subplot(3,3,5)
sns.kdeplot(df['Prothrombin'], hue=df['Stage'], fill=True, palette='Purples', common_norm=True)
sns.despine(top=True, right=True, bottom=True, left=True)
plt.tick_params(axis='both', which='both', bottom=False, top=False, left=False)
plt.xlabel('')
plt.title('Prothrombin');

plt.subplot(3,3,6)
sns.kdeplot(df['Copper'], hue=df['Stage'], fill=True, palette='Blues', common_norm=True)
sns.despine(top=True, right=True, bottom=True, left=True)
plt.tick_params(axis='both', which='both', bottom=False, top=False, left=False)
plt.xlabel('')
plt.title('Copper');

plt.subplot(3,3,7)
sns.kdeplot(df['Platelets'], hue=df['Stage'], fill=True, palette='Purples')
sns.despine(top=True, right=True, bottom=True, left=True)
plt.tick_params(axis='both', which='both', bottom=False, top=False, left=False)
plt.xlabel('')
plt.title('Platelets in stages');

plt.subplot(3,3,8)
sns.kdeplot(df['Albumin'], hue=df['Stage'], fill=True, palette='Blues', common_norm=True)
sns.despine(top=True, right=True, bottom=True, left=True)
plt.tick_params(axis='both', which='both', bottom=False, top=False, left=False)
plt.xlabel('')
plt.title('Albumin');
```

```
plt.subplot(3,3,9)
sns.kdeplot(df['SGOT'], hue=df['Stage'], fill=True, palette='Purples', common_norm=True)
sns.despine(top=True, right=True, bottom=True, left=True)
plt.tick_params(axis='both', which='both', bottom=False, top=False, left=False)
plt.xlabel('')
plt.title('SGOT');
```



Looking at the feature distribution we can observe that in features such as Age, Prothrombin, Copper the risk of the disease increase with increase in feature value, thus having a positive co-relation on with the disease probability. Lets fit a regression line to check.

In [15]:

```

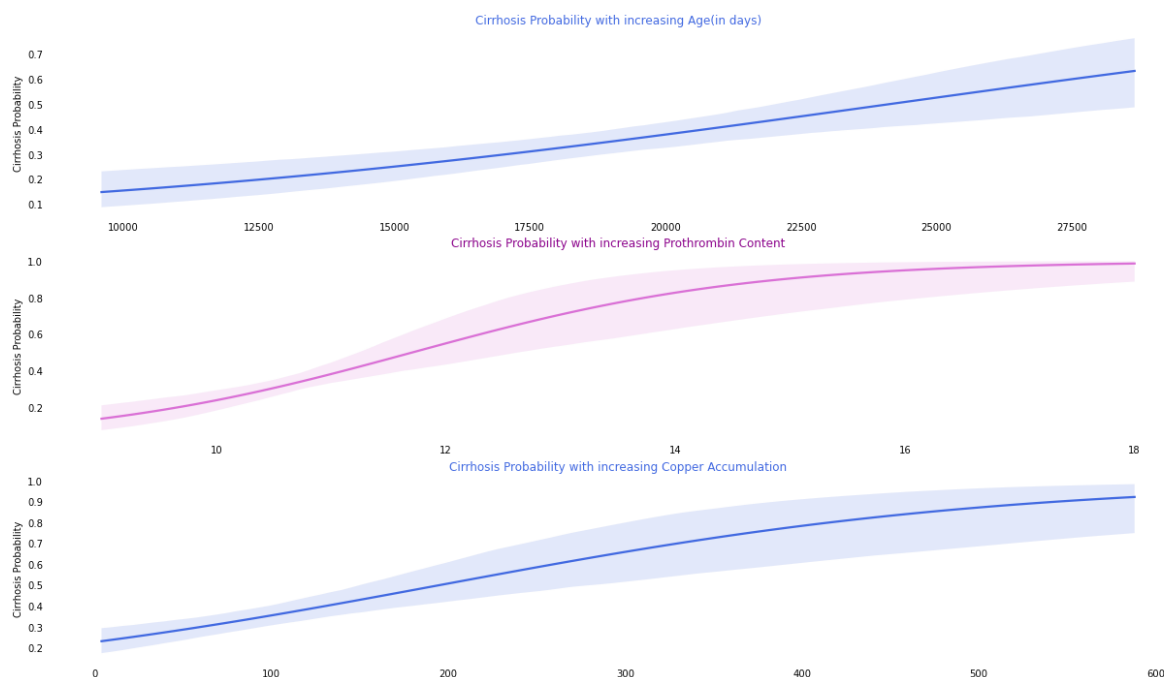
#@title Regression Plots of Positive Correlated Features.
plt.figure(figsize=(21,12))

plt.subplot(3,1,1)
sns.regplot(x=df['Age'], y=df['Stage'], scatter=False, logistic=True, color='royalblue')
sns.despine(fig=None, ax=None, top=True, right=True, left=True, bottom=True, offset=None, t
plt.tick_params(axis='both', which='both', bottom=False, top=False, left=False)
plt.xlabel('');
plt.ylabel('Cirrhosis Probability');
plt.setp(plt.title('Cirrhosis Probability with increasing Age(in days)'), color='royalblue'

plt.subplot(3,1,2)
sns.regplot(x=df['Prothrombin'], y=df['Stage'], scatter=False, logistic=True, color='orchid')
sns.despine(fig=None, ax=None, top=True, right=True, left=True, bottom=True, offset=None, t
plt.tick_params(axis='both', which='both', bottom=False, top=False, left=False)
plt.xlabel('');
plt.ylabel('Cirrhosis Probability');
plt.setp(plt.title('Cirrhosis Probability with increasing Prothrombin Content'), color='dar

plt.subplot(3,1,3)
sns.regplot(x=df['Copper'], y=df['Stage'], scatter=False, logistic=True, color='royalblue')
sns.despine(fig=None, ax=None, top=True, right=True, left=True, bottom=True, offset=None, t
plt.tick_params(axis='both', which='both', bottom=False, top=False, left=False)
plt.xlabel('');
plt.ylabel('Cirrhosis Probability');
plt.setp(plt.title('Cirrhosis Probability with increasing Copper Accumulation'), color='roy

```



Looks like the data checks with our intuition. These parameters indeed increase the risk of the disease.

We can also observe some features such as Platelets, Albumin, Cholesterol where the probability of disease decrease with increase in feature value. Lets tally that with some more regression plots.

In [16]:

```

#@title Regression Plots of negatively correlated Features.
plt.figure(figsize=(21,12))

plt.subplot(3,1,1)
sns.regplot(x=df['Platelets'], y=df['Stage'], scatter=False, logistic=True, color='orchid')
sns.despine(fig=None, ax=None, top=True, right=True, left=True, bottom=True, offset=None, t
plt.tick_params(axis='both', which='both', bottom=False, top=False, left=False)
plt.xlabel('');
plt.ylabel('Cirrhosis Probability');
plt.setp(plt.title('Cirrhosis Probability with Platelets'), color='darkmagenta');

plt.subplot(3,1,2)
sns.regplot(x=df['Albumin'], y=df['Stage'], scatter=False, logistic=True, color='royalblue')
sns.despine(fig=None, ax=None, top=True, right=True, left=True, bottom=True, offset=None, t
plt.tick_params(axis='both', which='both', bottom=False, top=False, left=False)
plt.xlabel('');
plt.ylabel('Cirrhosis Probability');
plt.setp(plt.title('Cirrhosis Probability with Albumin Content'), color='royalblue');

plt.subplot(3,1,3)
sns.regplot(x=df['Cholesterol'], y=df['Stage'], scatter=False, logistic=True, color='orchid')
sns.despine(fig=None, ax=None, top=True, right=True, left=True, bottom=True, offset=None, t
plt.tick_params(axis='both', which='both', bottom=False, top=False, left=False)
plt.xlabel('');
plt.ylabel('Cirrhosis Probability');
plt.setp(plt.title('Cirrhosis Probability Cholesterol'), color='darkmagenta') ;

```



Platelets, Albumin checks with our logic the findings about Cholesterol seems interesting! Looks like people with high Cholesterol have lower risk of Cirrhosis, this might not sound correct but our data certainly shows so.

This should help our model predict the target. We will be looking at what features contribute the most in later part of the project.

Preprocessing data

In [17]:

```
# replacing catagorical data with intigers.
df['Sex'] = df['Sex'].replace({'M':0, 'F':1})
df['Ascites'] = df['Ascites'].replace({'N':0, 'Y':1})
df['Drug'] = df['Drug'].replace({'D-penicillamine':0, 'Placebo':1})
df['Hepatomegaly'] = df['Hepatomegaly'].replace({'N':0, 'Y':1})
df['Spiders'] = df['Spiders'].replace({'N':0, 'Y':1})
df['Edema'] = df['Edema'].replace({'N':0, 'Y':1, 'S':-1})
df['Status'] = df['Status'].replace({'C':0, 'CL':1, 'D':-1})
```

```
# Male : 0 , F
# N : 0, Y : 1
# D-penicillam
# N : 0, Y : 1
# N : 0, Y : 1
# N : 0, Y : 1
# 'C':0, 'CL':
```

We will not be using 'Status' and 'N_days' as our features since this will cause data Leakage.

In [18]:

```
# Setting up Features and Target
X = df.drop(['Status', 'N_Days', 'Stage'], axis=1)
y = df.pop('Stage')
```

Earlier while examining the distribution of target in the data, we found that the data is unevenly distributed, that is there are more examples of a certain class than other.

###To tackel this imbalance we will use **Stratified k-Fold** Cross Validation.

Model Selection.

lets first try out a quick Logistic regression classifier and see how it performs.

In [20]:

```
from sklearn.metrics import classification_report
log_model_predict = log_model.predict(test)
log_model_predict_proba = log_model.predict_proba(test)

print(classification_report(y.iloc[test_index], log_model_predict))
```

	precision	recall	f1-score	support
0	0.70	0.78	0.74	27
1	0.45	0.36	0.40	14
accuracy			0.63	41
macro avg	0.58	0.57	0.57	41
weighted avg	0.62	0.63	0.62	41

In [21]:

```

from sklearn.metrics import roc_auc_score
from sklearn.metrics import roc_curve, auc

fpr, tpr, threshold = roc_curve(y.iloc[test_index], log_model_predict_proba[:,1])
roc_auc = auc(fpr, tpr)

print('AUC : ', roc_auc_score(y.iloc[test_index], log_model_predict_proba[:,1]))

```

AUC : 0.6507936507936508

In [73]:

```

def trainer(X_train, y_train, X_test, y_test):

    models= [[' SVM ',SVC()],
              [' Decision Tree ', DecisionTreeClassifier()],
              [' Random Forest ', RandomForestClassifier()],
              [' Logistic Regression ', LogisticRegression(max_iter=200)],
              [' AdaBoost ', AdaBoostClassifier()],
              [' KNN ', KNeighborsClassifier()]]

    scores = []

    for model_name, model in models:

        model = model
        model.fit(X_train, y_train)
        pred = model.predict(X_test)
        cm_model = confusion_matrix(y_test, pred)
        scores.append(accuracy_score(y_test, model.predict(X_test)))

    print(Back.YELLOW + Fore.BLACK + Style.BRIGHT + model_name)
    print(Back.RESET)
    print(cm_model)
    print('\n' + Fore.BLUE + 'Training Acc. : ' + Fore.GREEN + str(round(accuracy_score(y_train, model.predict(X_train)), 2)))
    print(Fore.BLUE + 'Validation Acc.: ' + Fore.GREEN + str(round(accuracy_score(y_test, model.predict(X_test)), 2)))
    print(Fore.CYAN + classification_report(y_test, pred))

    visualizer = ROCAUC(model)
    visualizer.fit(X_train, y_train)
    visualizer.score(X_test, y_test)
    visualizer.show()

    print('\n' + Fore.BLACK + Back.WHITE + '*****')

    return scores

```


In [63]:

```
import warnings
import itertools

import pandas as pd
import numpy as np

import matplotlib.pyplot as plt
import seaborn as sns

from sklearn.model_selection import train_test_split
from sklearn.preprocessing import StandardScaler, LabelEncoder
from sklearn.metrics import confusion_matrix, accuracy_score, classification_report, roc_auc_score
from sklearn.pipeline import Pipeline
from sklearn.tree import DecisionTreeClassifier
from sklearn.linear_model import LogisticRegression
from sklearn.ensemble import RandomForestClassifier, AdaBoostClassifier
from sklearn.neighbors import KNeighborsClassifier
from sklearn.svm import SVC

from xgboost import XGBClassifier
from catboost import CatBoostClassifier

from colorama import Fore, Back, Style
#from yellowbrick.classifier import ROCAUC
```

In [62]:

```
conda install -c districtdatalabs yellowbrick
```

Note: you may need to restart the kernel to use updated packages.

usage: conda-script.py [-h] [-V] command ...

conda-script.py: error: unrecognized arguments: yellowbrick

In [43]:

```
X_train, X_test, y_train, y_test = train_test_split(X, y, test_size = 0.2, random_state = 2)
```

In [109]:

```
def score_vis(score):

    names = ['Random Forest', 'Knn']

    plt.rcParams['figure.figsize']=8,10
    ax = sns.barplot(x=names, y=score, palette = "plasma", saturation =2.0)
    plt.xlabel('Model', fontsize = 20 )
    plt.ylabel('Accuracy(%)', fontsize = 20)
    plt.title('Model Comparison - Test set(Random Forest vs KNN)', fontsize = 20)
    plt.xticks(fontsize = 12, horizontalalignment = 'center', rotation = 8)
    plt.yticks(fontsize = 12)
    for i in ax.patches:
        width, height = i.get_width(), i.get_height()
        x, y = i.get_xy()
        ax.annotate(f'{round(height,2)}%', (x + width/2, y + height*1.02), ha='center', fon
    plt.show()
```

In [110]:

```
def trainer(X_train, y_train, X_test, y_test):

    models= [ [' KNN ', KNeighborsClassifier()], [' Random Forest ', RandomForestClassifier
    scores = []

    for model_name, model in models:

        model = model
        model.fit(X_train, y_train)
        pred = model.predict(X_test)
        cm_model = confusion_matrix(y_test, pred)
        scores.append(accuracy_score(y_test, model.predict(X_test)))
        print(Back.YELLOW + Fore.BLACK + Style.BRIGHT + model_name)
        print(Back.RESET)
        print(cm_model)
        print('\n' + Fore.BLUE + 'Training Acc. : ' + Fore.GREEN + str(round(accuracy_scor
        print(Fore.BLUE + 'Validation Acc.: ' + Fore.GREEN + str(round(accuracy_score(y_tes
        print(Fore.CYAN + classification_report(y_test, pred))
        print('\n' + Fore.BLACK + Back.WHITE + '*****

    return scores
```

In [111]:

```
scores = trainer(X_train, y_train, X_test, y_test)
```

KNN

```
[[45  6]  
 [26  7]]
```

Training Acc. : 78.14%

Validation Acc.: 61.9%

	precision	recall	f1-score	support
0	0.63	0.88	0.74	51
1	0.54	0.21	0.30	33
accuracy			0.62	84
macro avg	0.59	0.55	0.52	84
weighted avg	0.60	0.62	0.57	84

Random Forest

```
[[49  2]  
 [20 13]]
```

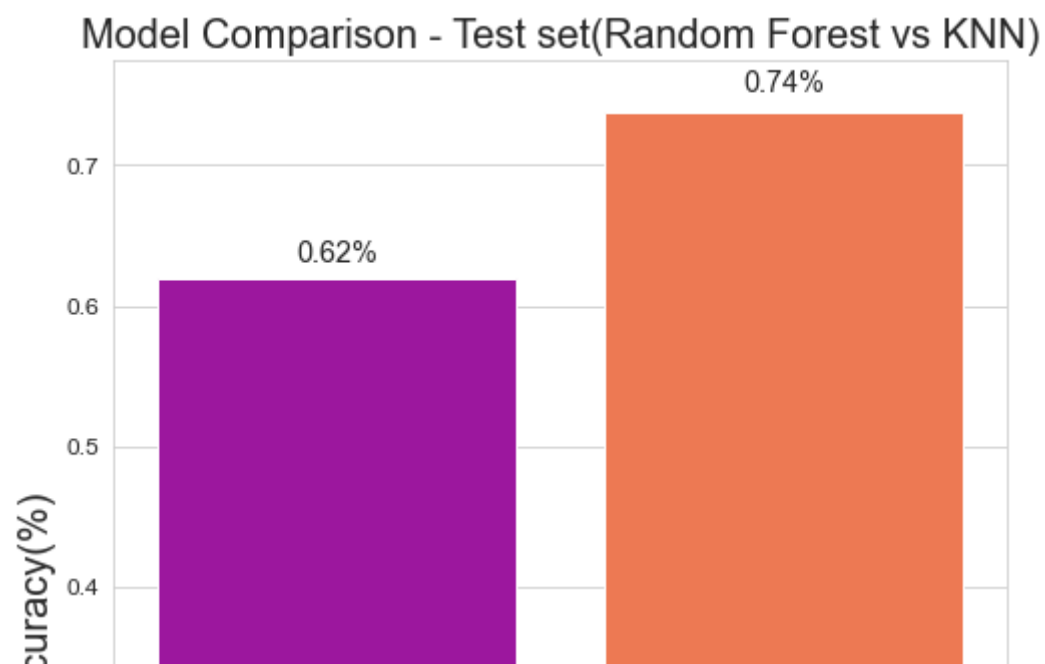
Training Acc. : 100.0%

Validation Acc.: 73.81%

	precision	recall	f1-score	support
0	0.71	0.96	0.82	51
1	0.87	0.39	0.54	33
accuracy			0.74	84
macro avg	0.79	0.68	0.68	84
weighted avg	0.77	0.74	0.71	84

In [112]:

```
score_vis(scores)
```



In []:

In []: