# STATS 3DA3

#### Homework Assignment 6

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 $Github\ URL:\ https://github.com/Xxxxcd/STATS3DA3.git$ 

```
from ucimlrepo import fetch_ucirepo
import matplotlib.pyplot as plt
import matplotlib.cm as cm
import pandas as pd
import numpy as np
import seaborn as sns
from sklearn.cluster import KMeans
from sklearn.datasets import make_circles
from sklearn.impute import SimpleImputer
from sklearn.metrics import mean_squared_error, confusion_matrix, classification_report
from sklearn.metrics import accuracy_score, classification_report
from sklearn.metrics import silhouette_samples, silhouette_score
from sklearn.model_selection import train_test_split, cross_val_score
from sklearn.model_selection import train_test_split
from sklearn.preprocessing import StandardScaler
from sklearn.preprocessing import scale
from sklearn.svm import SVC
from sklearn.tree import DecisionTreeClassifier, DecisionTreeRegressor, plot_tree
from sklearn.tree import plot_tree
```

```
# fetch dataset
chronic_kidney_disease = fetch_ucirepo(id=336)
# data (as pandas dataframes)
X = chronic_kidney_disease.data.features
y = chronic_kidney_disease.data.targets
for i in range(len(y)):
    if y.loc[i, "class"] == "ckd\t":
        y.loc[i, "class"] = "ckd"
X["dm"].isna().sum()
for i in range(len(y)):
    if X.loc[i, "dm"] == "\tno":
        X.loc[i, "dm"] = "no"
distinct = set(y["class"])
print(distinct)
```

```
{'ckd', 'notckd'}
```

The dataset has some issues converting it into Pandas dataframe as it contains TAB/ spaces in the data which I correct them in by remove "" which represents tab.

#### 1.

This dataset contains medical information for each record, including whether an individual has chronic kidney disease ("ckd" or "notckd"). We would like to utilize this dataset to predict the presence of chronic kidney disease, In generial, we want to use model for classification, essentially classifying groups of individuals who may potentially have the condition of chronic kidney disease.

```
X = pd.DataFrame(X)
y = pd.DataFrame(y)
X.head(5)
```

_													
	age	bp	sg	al	su	rbc	pc	pcc	ba	bgr	 hemo	pcv	wbc
0	48.0	80.0	1.020	1.0	0.0	NaN	normal	notpresent	notpresent	121.0	 15.4	44.0	7800
1	7.0	50.0	1.020	4.0	0.0	NaN	normal	notpresent	notpresent	NaN	 11.3	38.0	6000
2	62.0	80.0	1.010	2.0	3.0	normal	normal	notpresent	notpresent	423.0	 9.6	31.0	7500
3	48.0	70.0	1.005	4.0	0.0	normal	abnormal	present	notpresent	117.0	 11.2	32.0	6700
4	51.0	80.0	1.010	2.0	0.0	normal	normal	notpresent	notpresent	106.0	 11.6	35.0	7300

### 2.

We decide to use Decision Tree and Support Vector Machine (SVM) to classify the data. For Decision Tree, it is unnecessary to transform the variables, because they are not sensitive to feature scaling. Doing so will not change the result. However, normalize the data could be useful for SVM, we will standardlize numerical columns after we have an overview of the dataset and use them when we fit SVM model for the data in later codes.

```
print(X.dtypes)
print(y.dtypes)
```

age float64
bp float64
sg float64

al float64 float64 su object rbc object рс object рсс ba object float64 bgr float64 bu sc float64 float64 sod float64 pot hemo float64 float64 pcv float64 wbcc float64 rbcc object htn object dmobject cad object appet object ре object ane dtype: object object class dtype: object

### X.isna().sum()

age 9
bp 12
sg 47
al 46
su 49

```
65
рс
           4
рсс
           4
ba
bgr
          44
          19
bu
          17
sc
          87
sod
          88
pot
hemo
          52
         71
pcv
         106
wbcc
rbcc
         131
           2
htn
dm
           2
cad
appet
           1
ре
ane
dtype: int64
cat = ['rbc', 'pc', 'pcc', 'ba', 'htn', 'dm', 'cad', 'appet', 'pe', 'ane', 'al', 'su']
#al and su are categorical variables, though their type is float64.
for col in cat:
    X[col] = pd.Categorical(X[col]).codes
#After the previous processing, categorical variables are now switched into numbers for model ;
for col in cat:
```

rbc

152

X[col] = X[col].replace(-1, np.nan)

### X.isna().sum()

age	9
bp	12
sg	47
al	46
su	49
rbc	152
pc	65
pcc	4
ba	4
bgr	44
bu	19
sc	17
sod	87
pot	88
hemo	52
pcv	71
wbcc	106
rbcc	131
htn	2
dm	2
cad	2
appet	1
pe	1
ane	1
dtype:	int64

### 3. Dataset Overview

```
class
0 ckd
1 ckd
2 ckd
3 ckd
4 ckd
... ...
395 notckd
```

```
[400 rows x 1 columns]
```

396

397

398

399

notckd

notckd

notckd

notckd

We manually check the dictionary from the data source webstite. For variable "al" and "su", although they have a type of "float", they are actually discreate numbers, so we treat them as categorical datas.

```
categorcal_cols = ['rbc', 'pc', 'pcc', 'ba', 'htn', 'dm', 'cad', 'appet', 'pe', 'ane', 'al', 'su']
numerical_cols = ["age", "bp", "sg", "bgr", "bu", "sc", "sod", "pot", "hemo", "pcv", "wbcc", "rbcc"]
print(X.shape)
print(y.shape)
```

```
(400, 24)
```

(400, 1)

### X[numerical\_cols].describe().T

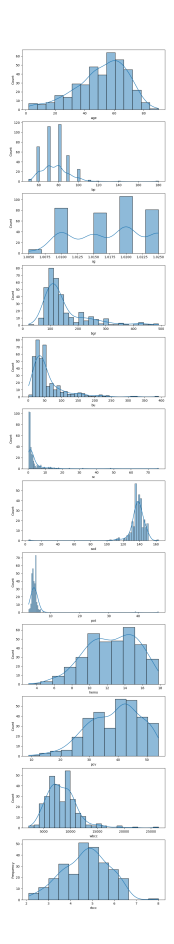
	count	mean	std	min	25%	50%	75%	max
age	391.0	51.483376	17.169714	2.000	42.00	55.00	64.50	90.000
bp	388.0	76.469072	13.683637	50.000	70.00	80.00	80.00	180.000
sg	353.0	1.017408	0.005717	1.005	1.01	1.02	1.02	1.025
bgr	356.0	148.036517	79.281714	22.000	99.00	121.00	163.00	490.000
bu	381.0	57.425722	50.503006	1.500	27.00	42.00	66.00	391.000
sc	383.0	3.072454	5.741126	0.400	0.90	1.30	2.80	76.000
$\operatorname{sod}$	313.0	137.528754	10.408752	4.500	135.00	138.00	142.00	163.000
pot	312.0	4.627244	3.193904	2.500	3.80	4.40	4.90	47.000
hemo	348.0	12.526437	2.912587	3.100	10.30	12.65	15.00	17.800
pcv	329.0	38.884498	8.990105	9.000	32.00	40.00	45.00	54.000
wbcc	294.0	8406.122449	2944.474190	2200.000	6500.00	8000.00	9800.00	26400.000
rbcc	269.0	4.707435	1.025323	2.100	3.90	4.80	5.40	8.000

### X[categorcal\_cols]

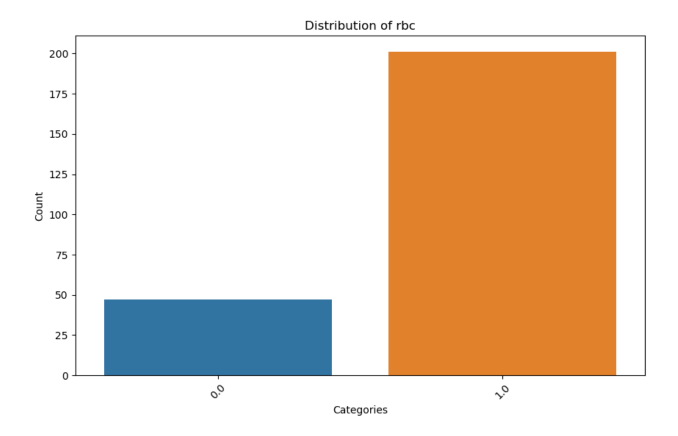
	rbc	pc	pcc	ba	htn	dm	cad	appet	pe	ane	al	su
0	NaN	1.0	0.0	0.0	1.0	1.0	0.0	0.0	0.0	0.0	1.0	0.0
1	NaN	1.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	4.0	0.0
2	1.0	1.0	0.0	0.0	0.0	1.0	0.0	1.0	0.0	1.0	2.0	3.0
3	1.0	0.0	1.0	0.0	1.0	0.0	0.0	1.0	1.0	1.0	4.0	0.0
4	1.0	1.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	2.0	0.0
395	1.0	1.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
396	1.0	1.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
397	1.0	1.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
398	1.0	1.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
399	1.0	1.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0

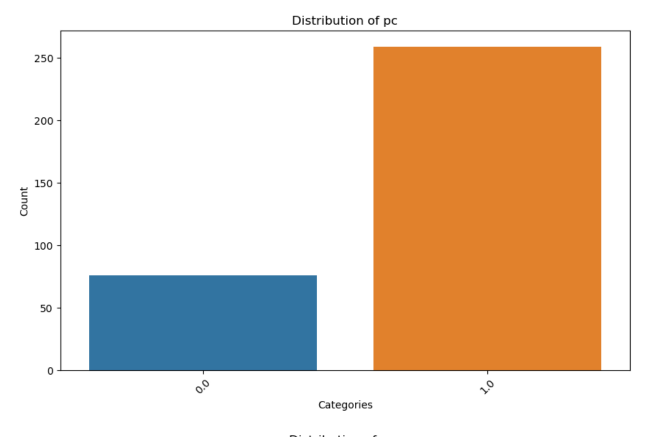
### numeric\_X = X.select\_dtypes(include=['float'])

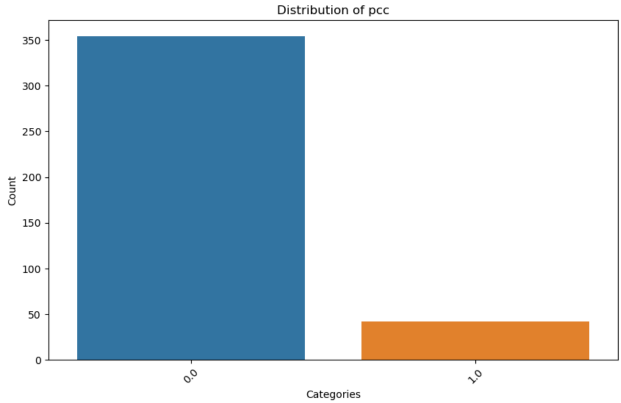
```
def plot_dist(data, features):
    fig, axes = plt.subplots(nrows=len(features), ncols=1, figsize=(8, 4 * len(features)))
    if len(features) == 1:
        axes = [axes]
    for ax, feature in zip(axes, features):
        sns.histplot(data[feature], kde=True, ax=ax)
    ax.set_xlabel(feature)
    ax.set_ylabel('Frequency')
plt.show()
plot_dist(X,numerical_cols)
```

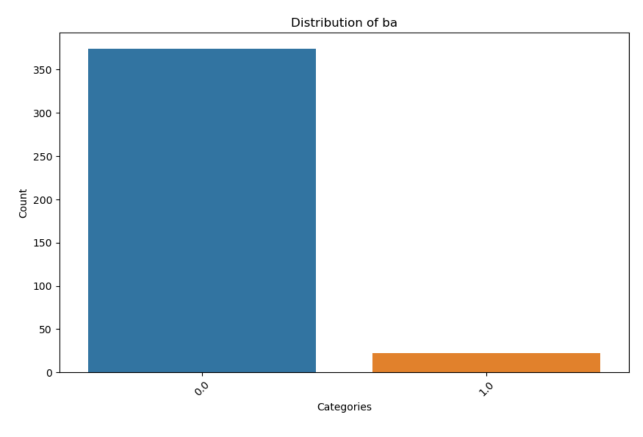


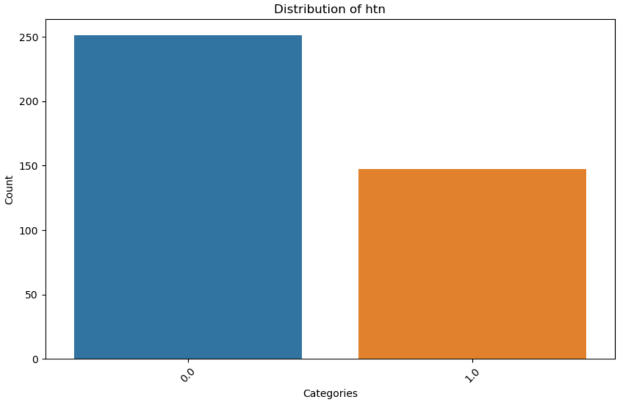
```
for i in range(len(categorcal_cols)):
    plt.figure(figsize=(10, 6))
    sns.countplot(x=categorcal_cols[i], data=X)
    plt.title('Distribution of '+ categorcal_cols[i])
    plt.xlabel('Categories')
    plt.ylabel('Count')
    plt.xticks(rotation=45)  # Rotate x-axis labels for better readability if needed
    plt.show()
```

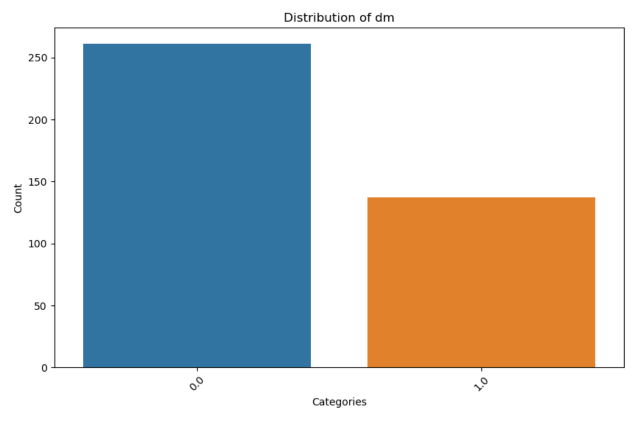


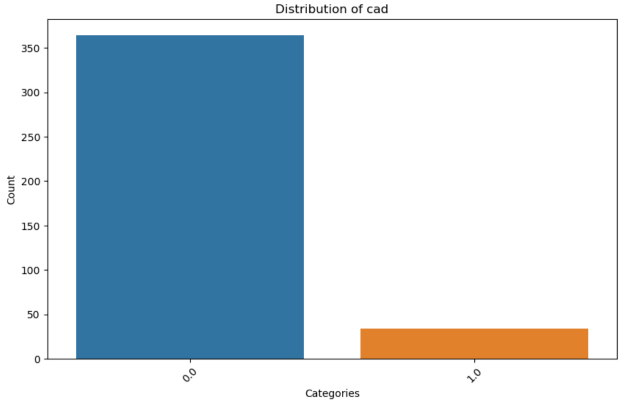


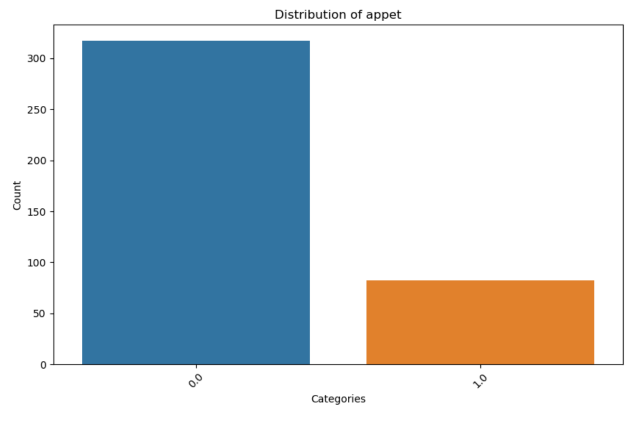


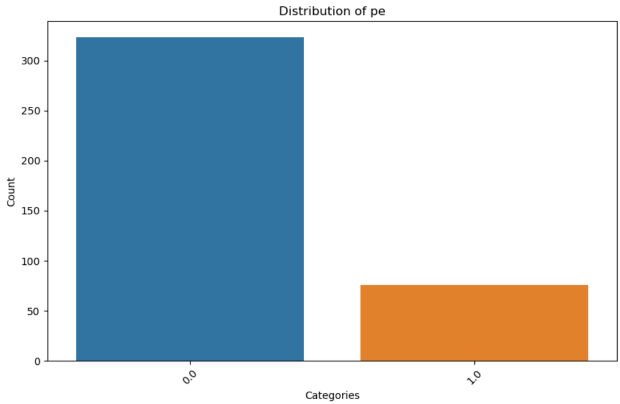


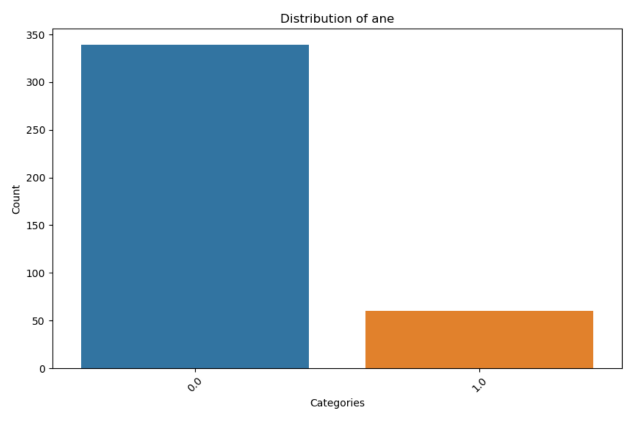


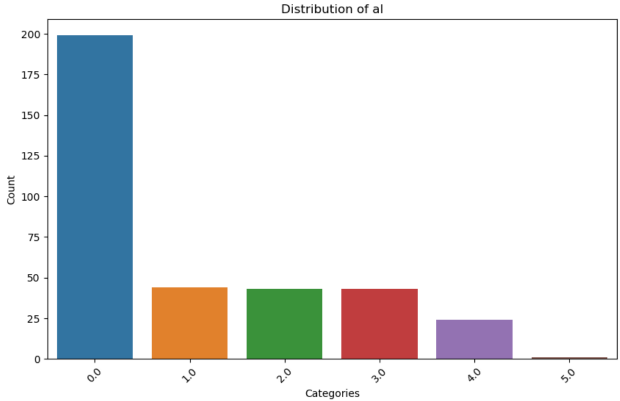


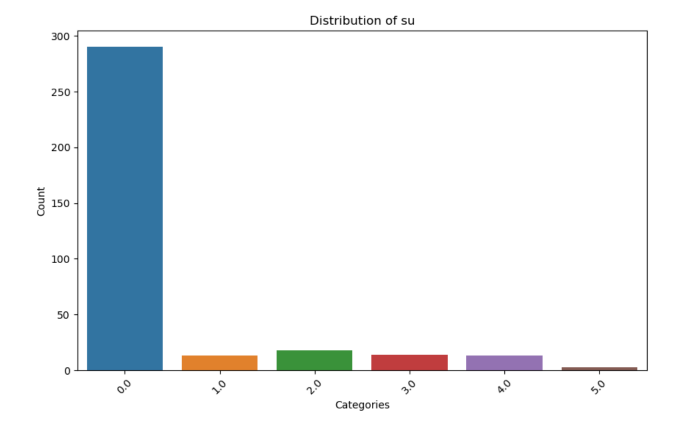












We standardlize numerical columns here, and we will not use them untill we train our SVM model.

```
from sklearn.preprocessing import StandardScaler
scaler = StandardScaler()
StandardCol=X
StandardCol[numerical_cols] = scaler.fit_transform(StandardCol[numerical_cols])
StandardCol[numerical_cols].describe().T
```

	count	mean	std	min	25%	50%	75%	max
age	391.0	9.994847e-17	1.001281	-2.885708	-0.553039	0.205078	0.759087	2.246163
bp	388.0	-2.380684e-16	1.001291	-1.936857	-0.473370	0.258373	0.258373	7.575807
sg	353.0	2.415443e-15	1.001419	-2.173584	-1.297699	0.454071	0.454071	1.329955
bgr	356.0	-1.796316e-16	1.001407	-1.591967	-0.619380	-0.341498	0.189004	4.319341
bu	381.0	-3.729883e-17	1.001315	-1.108830	-0.603246	-0.305843	0.170001	6.613723
sc	383.0	0.000000e+00	1.001308	-0.466102	-0.378897	-0.309133	-0.047519	12.719271

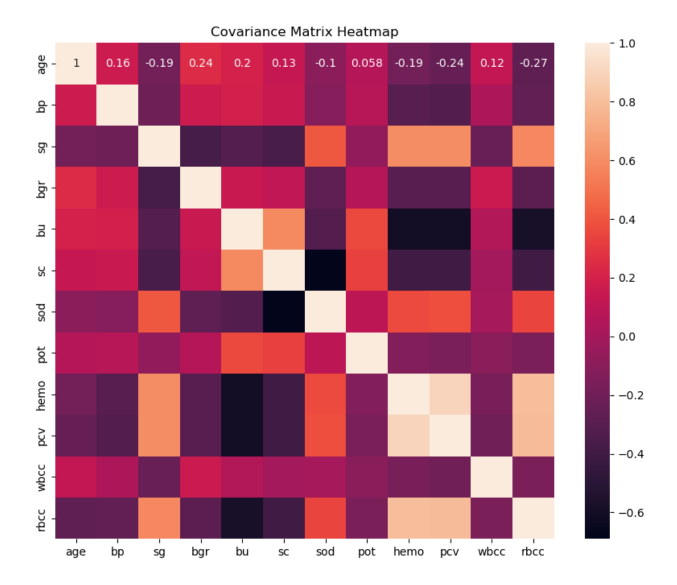
	count	mean	std	min	25%	50%	75%	max
$\operatorname{sod}$	313.0	2.270105e-17	1.001601	-12.800936	-0.243334	0.045347	0.430254	2.451017
pot	312.0	-7.970832e-17	1.001606	-0.667102	-0.259423	-0.071263	0.085536	13.288071
hemo	348.0	4.083579e-17	1.001440	-3.241109	-0.765520	0.042485	0.850490	1.813219
pcv	329.0	1.295823e-16	1.001523	-3.329218	-0.766953	0.124270	0.681284	1.683910
wbcc	294.0	1.450087e-16	1.001705	-2.111312	-0.648460	-0.138162	0.474195	6.121486
rbcc	269.0	8.452553e-16	1.001864	-2.547777	-0.788961	0.090447	0.676719	3.217231

There are a total of 400 records. The age of these records are mostly distributed among 40 to 80. For variable "sc", "bu", "bgr" and "wbcc", they are left-skewed. For variable "pcv", and "hemo", they are right-skewed.

### 4

The heatmap below shows the corrlations of the combinations of the variables.

```
# Plot heatmap of covariance matrix
plt.figure(figsize=(10, 8))
sns.heatmap(X[numerical_cols].corr(),annot=True)
plt.title('Covariance Matrix Heatmap')
plt.show()
```



Relation between "sod" and "sc" has the most negative corrleation;

Relation between "hemo" and "rbcc" has the most positive correlation.;

Relation between "wbcc" and "bp" has the smallest correlation, which means they are close to a independent relation.

### 5

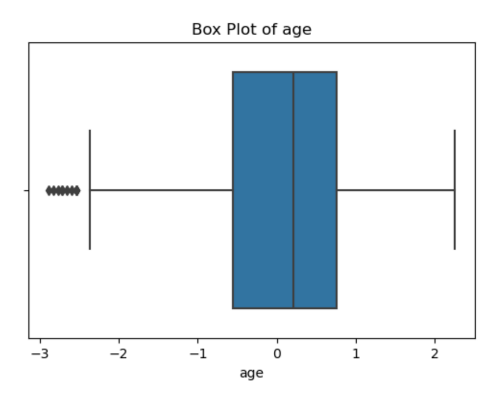
For quantitative variables, we replace the missing values by the mean of each column. For categorical variables, we replace the missing values by the most frequently appeared value in each column.

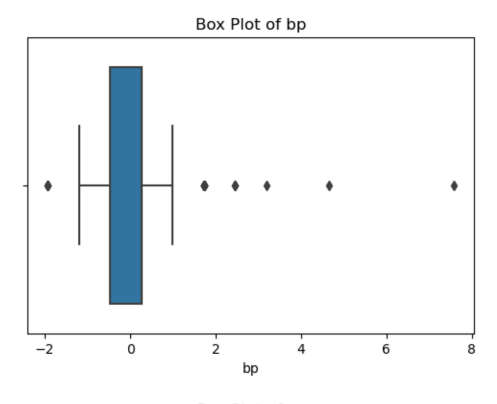
However, we will not process the missing values now since Decision Tree can accept missing values. We will impute missing values before we use SVM.

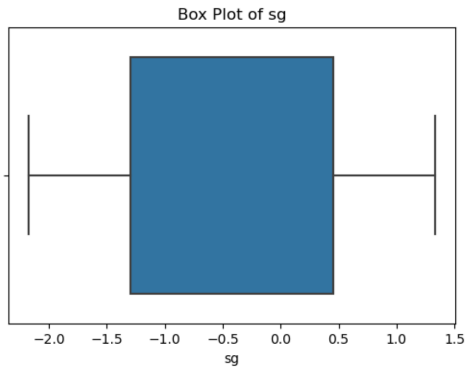
### 6 Outlier Analysis

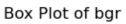
From the histogrms below we obtain that some outliers do exist. However, we decide not to remove them because a certain value in a certain column may be influenced by other variables, we cannot find a clear method or rule to remove the outliers though we got a heatmap. Also, some of the medical data may be conclusive in determing the disease, so we think keep these outlier here is not necessary.

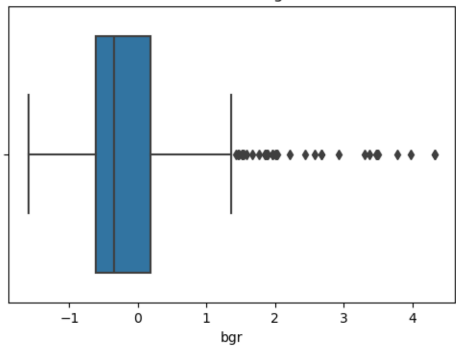
```
for col in numerical_cols:
   plt.figure(figsize=(6, 4))
   sns.boxplot(x=X[col])
   plt.title(f'Box Plot of {col}')
   plt.xlabel(col)
   plt.show()
```



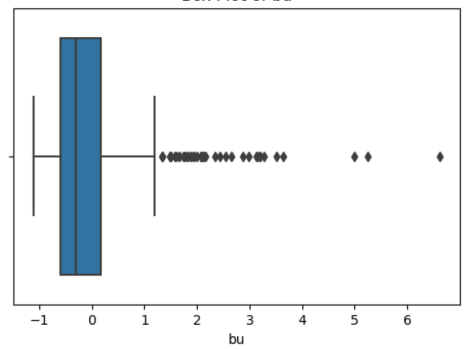




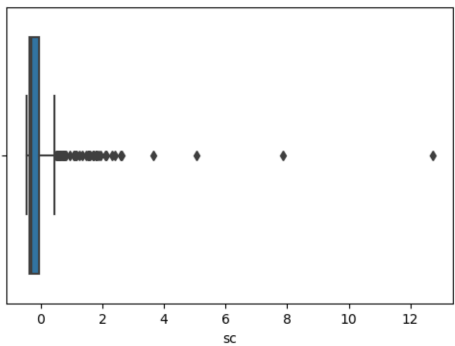




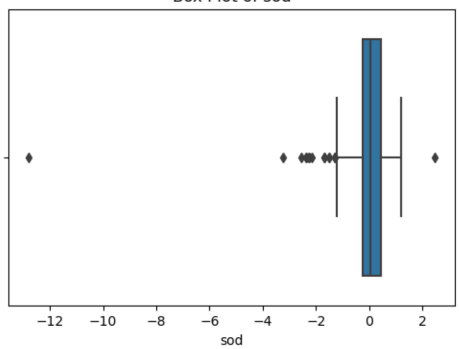
# Box Plot of bu

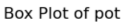


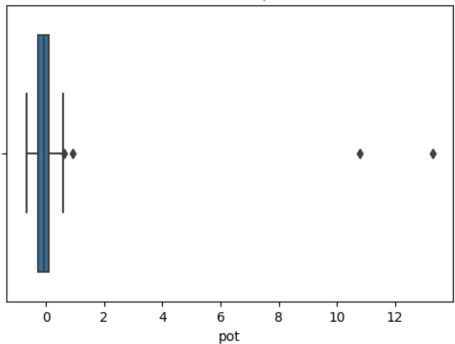




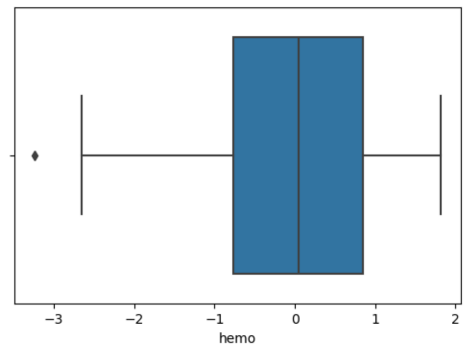
# Box Plot of sod

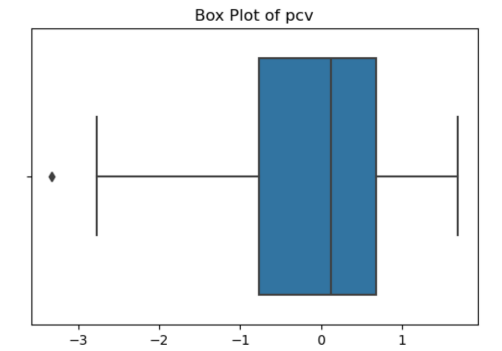


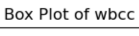




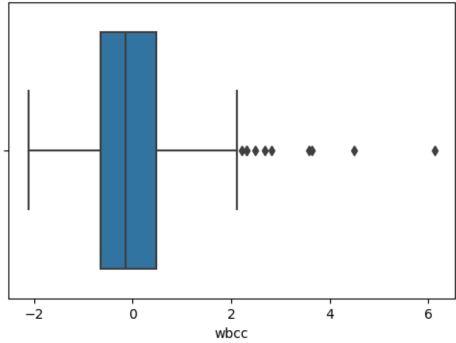
# Box Plot of hemo

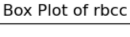


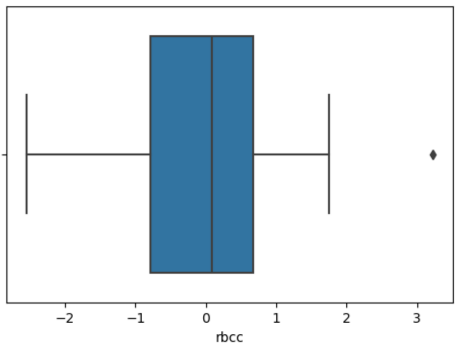




pcv







# 7. Subgrouping

```
combined_df = pd.concat([X, y], axis=1)
# Drop rows with missing values
combined_df = combined_df.dropna()
new_y=combined_df["class"]
new_X = combined_df.drop(columns=["class"])
```

```
print(new_y.shape)
print(new_X.shape)
```

```
(158,)
(158, 24)
```

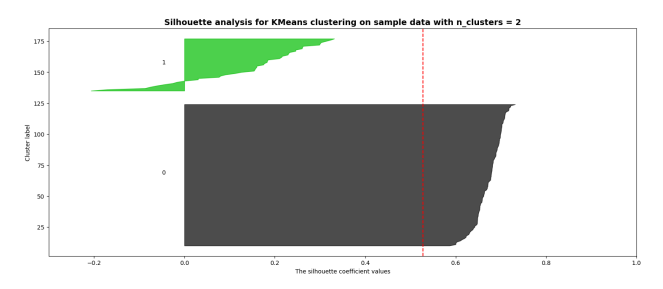
```
range_n_clusters = [2, 3, 4, 5, 6]
for n_clusters in range_n_clusters:
   km = KMeans(n_clusters=n_clusters, n_init=20, random_state=0)
   labels = km.fit_predict(new_X)
   silhouette_avg = silhouette_score(new_X, labels)
   sample_silhouette_values = silhouette_samples(new_X, labels)
   fig, ax1 = plt.subplots(1, 1)
   fig.set_size_inches(18, 7)
   ax1.set_xlim([-0.3, 1])
   y_lower = 10
   for i in range(n_clusters):
        ith_cluster_silhouette_values = sample_silhouette_values[labels == i]
        ith_cluster_silhouette_values.sort()
        size_cluster_i = ith_cluster_silhouette_values.shape[0]
       y_upper = y_lower + size_cluster_i
       # Use the colormap for coloring
        color = cm.nipy_spectral(float(i) / n_clusters)
        ax1.fill_betweenx(
           y=np.arange(y_lower, y_upper),
           x1=0,
           x2=ith_cluster_silhouette_values,
           facecolor=color,
            edgecolor=color,
           alpha=0.7,
        )
```

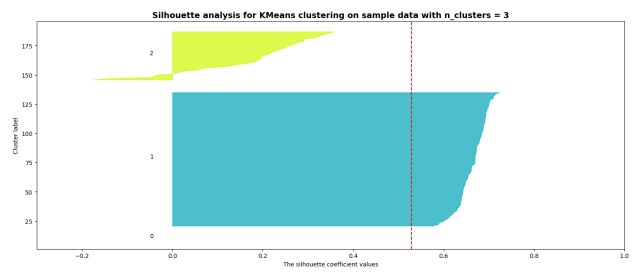
```
ax1.text(-0.05, y_lower + 0.5 * size_cluster_i, str(i))

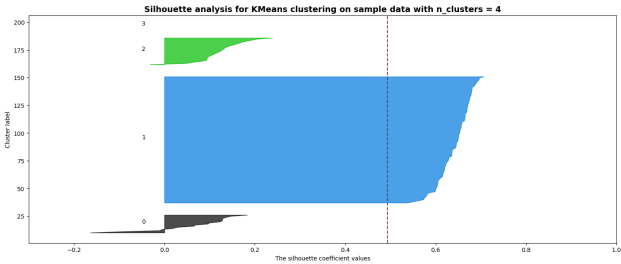
y_lower = y_upper + 10

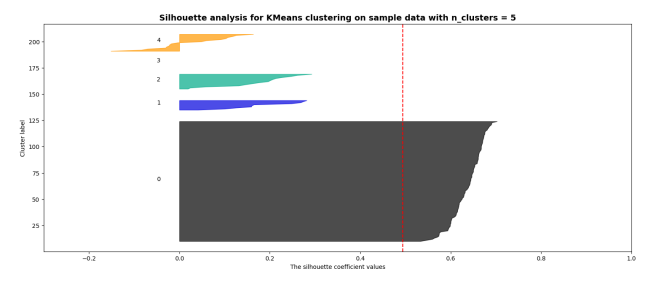
ax1.set_title("The silhouette plot for various clusters")
ax1.set_xlabel("The silhouette coefficient values")
ax1.set_ylabel("Cluster label")

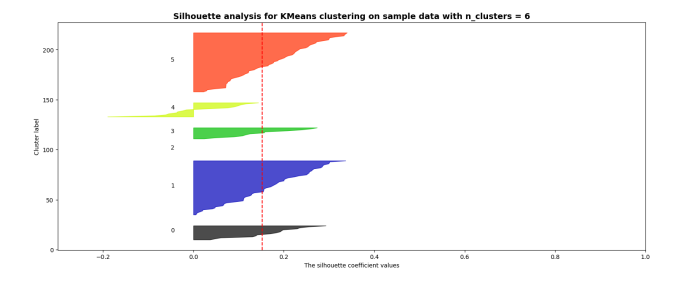
ax1.axvline(x=silhouette_avg, color="red", linestyle="--")
plt.title(
    "Silhouette analysis for KMeans clustering on sample data with n_clusters = %d"
    % n_clusters,
    fontsize=14,
    fontweight="bold",
)
```



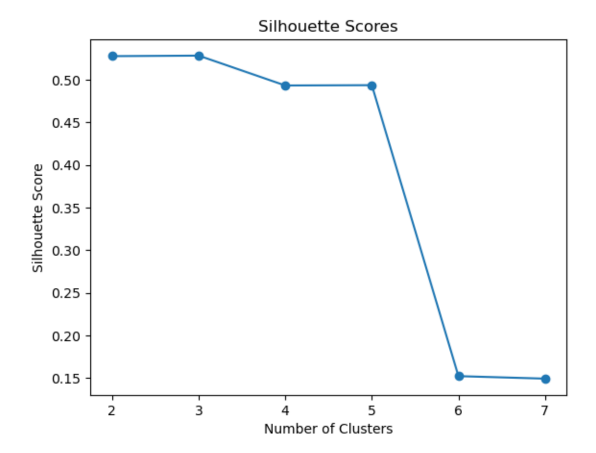








```
k_values = range(2, 8) # You can adjust this range as needed
# Initialize lists to store silhouette scores
silhouette_scores = []
# Iterate through different values of k
for k in k_values:
# Fit KMeans clustering to the data
    kmeans = KMeans(n_clusters=k, n_init = 20, random_state=0)
    kmeans.fit(new_X)
# Compute the silhouette score
    silhouette_avg = silhouette_score(new_X, kmeans.labels_)
    silhouette_scores.append(silhouette_avg)
# Plot silhouette scores against k
plt.plot(k_values, silhouette_scores, marker='o')
plt.xlabel('Number of Clusters')
plt.ylabel('Silhouette Score')
plt.title('Silhouette Scores')
plt.show()
```



We observe that when k=2, it has the best silhouette score. This result also aligns with the common sense the dataset.

```
combined_df = pd.concat([X, y], axis=1)

# Drop rows with missing values
combined_df = combined_df.dropna()

combined_df["class"]
```

3 ckd
 9 ckd
 11 ckd
 14 ckd

```
20 ckd
...
395 notckd
396 notckd
397 notckd
398 notckd
399 notckd
Name: class, Length: 158, dtype: object
```

### 8. Data Splitting

```
X_train, X_test, y_train, y_test = train_test_split(
    X, y, test_size=0.3, random_state=1, stratify=y)
```

### 9.

Like we mentioned before, we will be using Decision Tree and SVM for classification. We choose these two methods after we explore the whole dataset.

```
cs_dt = DecisionTreeClassifier(
   max_depth =30,
   random_state=1
)
```

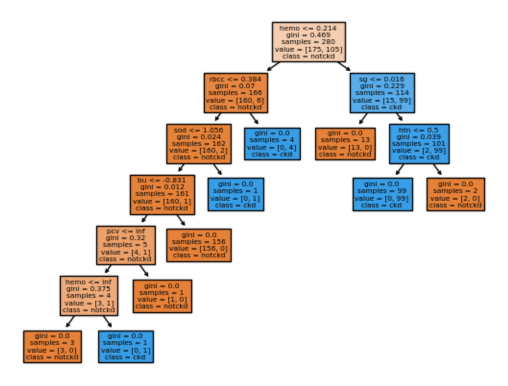
```
cs_dt.fit(X_train, y_train)

plot_tree(
    cs_dt,
    max_depth= 30,
    feature_names = X_train.columns.tolist(),
    class_names=['notckd', 'ckd'],
```

```
filled=True
)

pred_DT = cs_dt.predict(X_test)
pred_DT[:5]
```

array(['ckd', 'ckd', 'notckd', 'notckd'], dtype=object)



### 10. Performance Metrics

We use the confucsion matrix as well as classification\_report to measure our accuracy.

```
cm_DT = pd.DataFrame(confusion_matrix(y_test, pred_DT), index=['No', 'Yes'], columns=['No', 'Yes']
cm_DT.index.name = 'True'
cm_DT.columns.name = 'Predicted'
```

```
print(cm_DT)
print(cs_dt.score(X_test, y_test))
print(classification_report(y_test, pred_DT))
```

```
Predicted No Yes
True
No 73 2
Yes 3 42
0.9583333333333333334
```

	precision	recall	f1-score	support
ckd	0.96	0.97	0.97	75
notckd	0.95	0.93	0.94	45
accuracy			0.96	120
macro avg	0.96	0.95	0.96	120
weighted avg	0.96	0.96	0.96	120

### 11.

Here we use two methods to improve the performance on decision tree. First method we prune the tree. Second Method, we fit our data with criterion='entropy', we did some research, and we agree on using entropy instead of "gini" is more suitable for this dataset, and its performance also proves that.

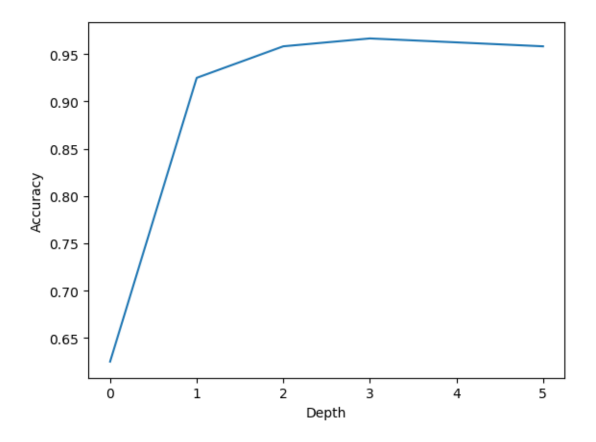
```
path = cs_dt.cost_complexity_pruning_path(
    X_train,
    y_train
)
ccp_alphas, impurities = path.ccp_alphas, path.impurities
```

```
clfs = [] # save fitted trees with different alphas
for ccp_alpha in ccp_alphas:
    clf = DecisionTreeClassifier(
        random_state=0,
        ccp_alpha=ccp_alpha
        )
    clf.fit(X_train, y_train)
    clfs.append(clf)
```

```
depth = [clf.tree_.max_depth for clf in clfs]
depth
```

[5, 5, 3, 2, 2, 1, 0]

```
test_score = [clf.score(X_test, y_test) for clf in clfs]
plt.plot(depth, test_score)
plt.xlabel('Depth')
plt.ylabel('Accuracy')
plt.show()
```



```
cs_dt_best = DecisionTreeClassifier(
    max_depth = 3,
    random_state=1
    )
cs_dt_best.fit(X_train, y_train)
```

DecisionTreeClassifier(max\_depth=3, random\_state=1)

```
X_train, X_test, y_train, y_test = train_test_split(
    X, y, test_size=0.3, random_state=1, stratify=y)
plot_tree(
    cs_dt_best,
    max_depth= 3,
    feature_names = X_train.columns.tolist(),
    filled=True,
    class_names=['notckd', 'ckd']
```

```
pred_BDT = cs_dt.predict(X_test)

print(classification_report(y_test, pred_BDT))

cm_BDT = pd.DataFrame(confusion_matrix(y_test, pred_BDT), index=['No', 'Yes'], columns=['No',

cm_BDT.index.name = 'True'

cm_BDT.columns.name = 'Predicted'

print(cm_BDT)

cs_dt.score(X_test, y_test)

print("Accuracy:", accuracy_score(y_test, pred_BDT))
```

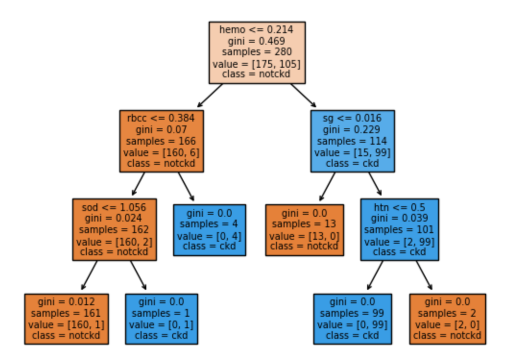
	precision	recall	f1-score	support
ckd	0.96	0.97	0.97	75
notckd	0.95	0.93	0.94	45
accuracy			0.96	120
macro avg	0.96	0.95	0.96	120
weighted avg	0.96	0.96	0.96	120

Predicted No Yes

True

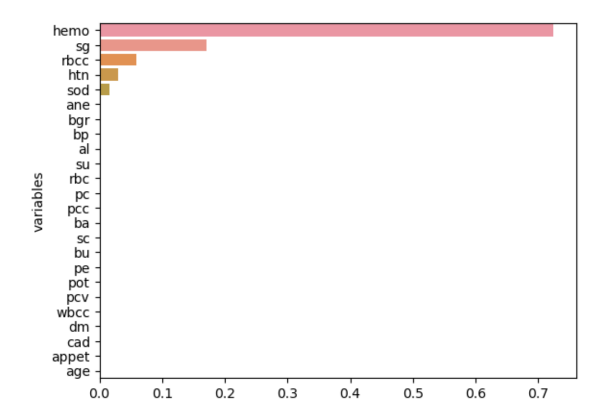
No 73 2 Yes 3 42

Accuracy: 0.95833333333333334



```
fea_imp = cs_dt_best.feature_importances_
sorted_indices = fea_imp.argsort()[::-1]
sorted_feature_names = X_train.columns[sorted_indices]
sorted_importances = fea_imp[sorted_indices]
sns.barplot(x = sorted_importances, y = sorted_feature_names)
plt.ylabel("variables")

plt.show()
```



From above plot, we can observe variable "hemo" has a significant influce on the model, next is "sg" and "rbcc". "htn" and "sod" also have some influence. Rest of the variables do not have a crucial impact on the model. After pruning the tree, we can see the acuracy did not increase. That is mostly because the previous one has a really large depth, but even with a max depth of 3, the model can achieve the same accuracy. Simplifing the model can also be seen as an improvement.

```
X_train, X_test, y_train, y_test = train_test_split(
    X, y, test_size=0.3, random_state=1, stratify=y)

clf = DecisionTreeClassifier(max_depth =30,
    random_state=1, criterion='entropy')

clf.fit(X_train, y_train)

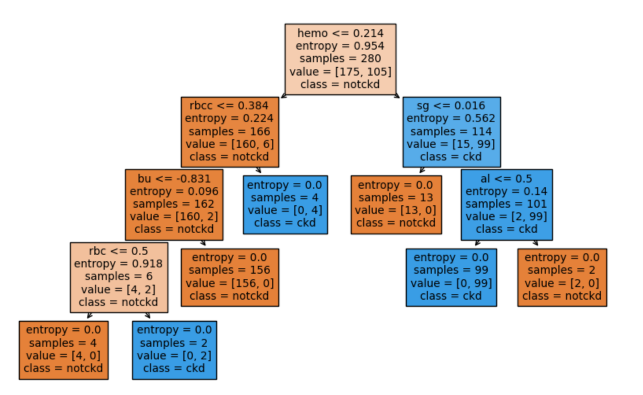
pred_clf = clf.predict(X_test)
```

```
print("Accuracy:", accuracy_score(y_test, pred_clf))
print(classification_report(y_test, pred_clf))

plt.figure(figsize=(10,6))
plot_tree(clf, filled=True,feature_names = X_train.columns.tolist(),class_names=['notckd', 'cke']
plt.show()
```

#### Accuracy: 0.966666666666667

	precision	recall	f1-score	support
ckd	0.96	0.99	0.97	75
notckd	0.98	0.93	0.95	45
accuracy			0.97	120
macro avg	0.97	0.96	0.96	120
weighted avg	0.97	0.97	0.97	120



The next method we used for improving the model is to change the crierion. The improvement of accuracy scores indicates a better result, which implies we have a better model.

```
cm_clf = pd.DataFrame(confusion_matrix(y_test, pred_clf), index=['No', 'Yes'], columns=['No',
cm_clf.index.name = 'True'
cm_clf.columns.name = 'Predicted'
cm_clf
```

Predicted	No	Yes
True		
No	74	1
Yes	3	42

Like we mentioned above, we will use mean for missing numerical datas, and most frequent more categorical data.

```
float_col = X[numerical_cols].columns.tolist()
num_imputer = SimpleImputer(missing_values=np.nan,strategy='mean')
X.loc[:, float_col] = num_imputer.fit_transform(X.loc[:, float_col])

# For categorical features
obj_col = X[categorcal_cols].columns.tolist()
cat_imputer = SimpleImputer(strategy='most_frequent',missing_values=np.nan)
X.loc[:, obj_col] = cat_imputer.fit_transform(X.loc[:, obj_col])
```

## X.isna().sum()

```
age 0
bp 0
sg 0
al 0
su 0
```

```
0
rbc
         0
рс
рсс
         0
ba
         0
bgr
         0
         0
bu
         0
sc
sod
         0
pot
hemo
         0
pcv
wbcc
rbcc
         0
htn
         0
dm
         0
cad
         0
appet
         0
ре
         0
ane
dtype: int64
X_train, X_test, y_train, y_test = train_test_split(
   X, y, test_size=0.3, random_state=1, stratify=y)
# Initialize and train SVM model
svm = SVC(kernel='linear')
y_train = np.ravel(y_train)
svm.fit(X_train, y_train)
```

SVC(kernel='linear')

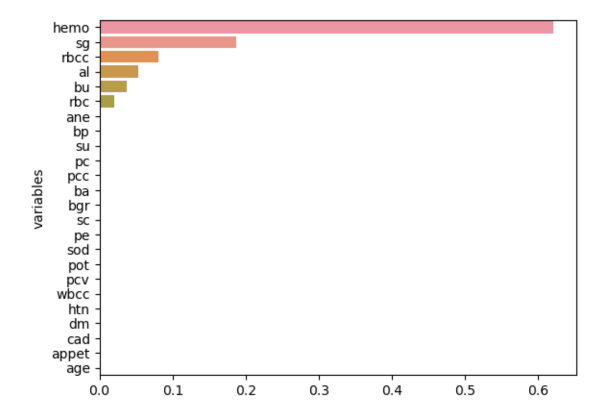
```
cm = confusion_matrix(y_test, pred_clf)
print('Confusion Matrix from Decicion Tree: \n', cm)
accuracyDT = accuracy_score(y_test, pred_clf)
print("Accuracy from SVM:", accuracyDT)
Confusion Matrix from Decicion Tree:
 [[74 1]
[ 3 42]]
Accuracy from SVM: 0.966666666666667
y_pred_SVM = svm.predict(X_test)
accuracy = accuracy_score(y_test, y_pred_SVM)
cm_SVM = confusion_matrix(y_test, y_pred_SVM)
print('Confusion Matrix from Decicion Tree: \n', cm_SVM)
print("Accuracy from SVM:", accuracy)
Confusion Matrix from Decicion Tree:
 [[75 0]
 [ 2 43]]
Accuracy from SVM: 0.9833333333333333
```

By comparing the confusion matrix and accuarcy score, we can conclude that linear SVC is a more suitable model for this dataset.

#### 13

```
fea_imp = clf.feature_importances_
sorted_indices = fea_imp.argsort()[::-1]
```

```
sorted_feature_names = X_train.columns[sorted_indices]
sorted_importances = fea_imp[sorted_indices]
sns.barplot(x = sorted_importances, y = sorted_feature_names)
plt.ylabel("variables")
plt.show()
```



### 14

```
from sklearn.preprocessing import PolynomialFeatures
from sklearn.metrics import accuracy_score

X_train, X_test, y_train, y_test = train_test_split(
    new_X, new_y, test_size=0.3, random_state=1, stratify=new_y)
```

New Model Accuracy: 0.979166666666666

The improved model does not bring us a better result. It is may because polynomial features introduce interaction terms between features, and sometimes there may be cases these interactions may not be relevant or may even introduce noise, leading to overfitting. It also increase the dimension of the model where a overly complicated model may not be as accurate as a simplier one.

#### 15 Distribution

```
Tianmu Li (400371751): Question 2, 3, 4, 5, 6, 11
Chengdai Xu (400397089): Question 5, 7, 8, 9, 10, 11,14
Jiajun Zhang (400359213): Question 1, 3, 7, 11, 12, 13
```

We have done a lot of work together by sharing the jupyter file itself instead of using github, but all of the questions are done under continuously disscussion and communication in the group. The contribution of each team members are equal and fair.