Final Project

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```
!pip3 install ucimlrepo
```

Requirement already satisfied: ucimlrepo in /Users/NoahRipstein/miniconda3/envs/3d_39/lib/pythouse

```
import pandas as pd
import numpy as np
from sklearn.preprocessing import StandardScaler
import matplotlib.pyplot as plt
import matplotlib as mpl
import matplotlib.cm as cm
from sklearn.impute import SimpleImputer
import seaborn as sns
from patsy import dmatrices, dmatrix
from sklearn.preprocessing import StandardScaler
from scipy.stats import zscore
from sklearn.model_selection import train_test_split
from sklearn.cluster import KMeans
from sklearn import metrics
from sklearn.linear_model import LogisticRegression, LinearRegression
from sklearn.decomposition import PCA, TruncatedSVD
from sklearn.ensemble import RandomForestClassifier
from sklearn.metrics import confusion_matrix, classification_report, silhouette_samples, silhouette_sample
from sklearn.metrics.cluster import rand_score
import statsmodels.api as sm
```

```
from mlxtend.feature_selection import ExhaustiveFeatureSelector as EFS

from mlxtend.feature_selection import SequentialFeatureSelector as SFS

from mlxtend.plotting import plot_sequential_feature_selection as plot_sfs

from sklearn.neighbors import KNeighborsClassifier
```

```
from ucimlrepo import fetch_ucirepo

chronic_kidney_disease = fetch_ucirepo(id=336)

df = pd.concat([chronic_kidney_disease.data.features, chronic_kidney_disease.data.targets], ax

df.head()
```

	age	bp	sg	al	su	rbc	pc	pcc	ba	bgr	 pcv	wbcc	rbo
0	48.0	80.0	1.020	1.0	0.0	NaN	normal	notpresent	notpresent	121.0	 44.0	7800.0	5.2
1	7.0	50.0	1.020	4.0	0.0	NaN	normal	notpresent	notpresent	NaN	 38.0	6000.0	Na
2	62.0	80.0	1.010	2.0	3.0	normal	normal	notpresent	notpresent	423.0	 31.0	7500.0	Na
3	48.0	70.0	1.005	4.0	0.0	normal	abnormal	present	notpresent	117.0	 32.0	6700.0	3.9
4	51.0	80.0	1.010	2.0	0.0	normal	normal	notpresent	notpresent	106.0	 35.0	7300.0	4.6

1. Classification Problem Identification: Define and describe a classification problem based on the dataset.

Our goal is to predict whether individuals have Chronic Kidney Disease (CKD) based on various medical predictor variables. This classification problem involves distinguishing between two categories: individuals diagnosed with CKD and those without. We will use a clinical dataset to build a predictive model that can accurately identify these two states based on patient medical data.

2. Variable Transformation: Implement any transformations chosen or justify the absence of such modifications.

df.describe()

	age	bp	sg	al	su	bgr	bu	sc
count	391.000000	388.000000	353.000000	354.000000	351.000000	356.000000	381.000000	383.000000
mean	51.483376	76.469072	1.017408	1.016949	0.450142	148.036517	57.425722	3.072454
std	17.169714	13.683637	0.005717	1.352679	1.099191	79.281714	50.503006	5.741126
min	2.000000	50.000000	1.005000	0.000000	0.000000	22.000000	1.500000	0.400000
25%	42.000000	70.000000	1.010000	0.000000	0.000000	99.000000	27.000000	0.900000
50%	55.000000	80.000000	1.020000	0.000000	0.000000	121.000000	42.000000	1.300000
75%	64.500000	80.000000	1.020000	2.000000	0.000000	163.000000	66.000000	2.800000
max	90.000000	180.000000	1.025000	5.000000	5.000000	490.000000	391.000000	76.000000

df.dtypes

age	float64
bp	float64
sg	float64
al	float64
su	float64
rbc	object
рс	object
pcc	object
ba	object
bgr	float64
bu	float64
sc	float64
sod	float64
pot	float64
hemo	float64
pcv	float64
wbcc	float64

```
float64
rbcc
          object
htn
dm
          object
cad
          object
          object
appet
          object
ре
          object
ane
          object
class
dtype: object
float64_columns = df.select_dtypes(
    include=['float64']
    ).columns
float64_columns
scaler = StandardScaler()
df[float64_columns] = scaler.fit_transform(df[float64_columns])
cat_columns = df.select_dtypes(
    include=['object']
    ).columns
for col in cat_columns:
    print(df[col].value_counts(normalize=True))
rbc
            0.810484
normal
abnormal
            0.189516
Name: proportion, dtype: float64
рс
            0.773134
normal
abnormal
            0.226866
Name: proportion, dtype: float64
```

рсс

notpresent 0.893939

present 0.106061

Name: proportion, dtype: float64

ba

notpresent 0.944444

present 0.055556

Name: proportion, dtype: float64

htn

no 0.630653

yes 0.369347

Name: proportion, dtype: float64

dm

no 0.653266

yes 0.344221

\tno 0.002513

Name: proportion, dtype: float64

cad

no 0.914573

yes 0.085427

Name: proportion, dtype: float64

appet

good 0.794486

poor 0.205514

Name: proportion, dtype: float64

ре

no 0.809524

yes 0.190476

Name: proportion, dtype: float64

ane

no 0.849624

yes 0.150376

Name: proportion, dtype: float64

class

ckd 0.620 notckd 0.375 ckd\t 0.005

Name: proportion, dtype: float64

```
for col in cat_columns:
    df[col] = df[col].astype('category').cat.codes

df.head(5)
```

	age	bp	sg	al	su	rbc	pc	pcc	ba	bgr	 pcv
0	-0.203139	0.258373	0.454071	-0.012548	-0.410106	-1	1	0	0	-0.341498	 0.569881 -
1	-2.594124	-1.936857	0.454071	2.208413	-0.410106	-1	1	0	0	NaN	 -0.098536 -
2	0.613295	0.258373	-1.297699	0.727772	2.323069	1	1	0	0	3.473064	 -0.878356 -
3	-0.203139	-0.473370	-2.173584	2.208413	-0.410106	1	0	1	0	-0.392022	 -0.766953 -
4	-0.028189	0.258373	-1.297699	0.727772	-0.410106	1	1	0	0	-0.530963	 -0.432744 -

Here, we performed two data transformation steps: 1. Transformation 1: Standardizing Numerical Features.

This step Z-transformed the numerical features to make them come from a distribution closer to a standard normal distribution with mean 0 and variance 1. This can improve performance of some classification algorithms, including logistic regression.

2. Transformation 2: Encoding Categorical Features.

This step converted columns containing categorical features into categorical variables within pandas. This is needed so that when we use our pandas dataframe as input to our classification models later, the libraries recognize the variables as categorical, rather than continuous.

3. Dataset Overview: Provide a detailed description of the dataset, covering variables, summaries, observation counts, data types, and distributions (at least three statements).

df.info()

RangeIndex: 400 entries, 0 to 399 Data columns (total 25 columns): Column Non-Null Count Dtype _____ 391 non-null float64 0 age 388 non-null float64 1 bp 2 353 non-null float64 sg 3 354 non-null al float64 351 non-null 4 float64 su 5 rbc 400 non-null int8 400 non-null 6 рс int8 400 non-null 7 рсс int8 8 ba 400 non-null int8 356 non-null float64 bgr 10 bu 381 non-null float64 383 non-null float64 11 sc 12 sod 313 non-null float64 pot 312 non-null float64 13 348 non-null 14 hemo float64 15 pcv 329 non-null float64 294 non-null 16 wbcc float64 17 rbcc 269 non-null float64 18 htn 400 non-null int8 19 dm400 non-null int8 400 non-null 20 cad int8 appet 400 non-null 21 int8 22 ре 400 non-null int8 ane 400 non-null 23 int8 24 class 400 non-null int8

<class 'pandas.core.frame.DataFrame'>

dtypes: float64(14), int8(11)

memory usage: 48.2 KB

df.describe()

	age	bp	sg	al	su	rbc	pc	p
count	3.910000e+02	3.880000e+02	3.530000e+02	354.000000	351.000000	400.00000	400.000000	40
mean	9.994847e-17	-2.380684e-16	2.415443e-15	0.000000	0.000000	0.12250	0.485000	0.
std	1.001281e+00	1.001291e+00	1.001419e+00	1.001415	1.001428	0.93256	0.759089	0.
min	-2.885708e+00	-1.936857e+00	-2.173584e+00	-0.752868	-0.410106	-1.00000	-1.000000	-1
25%	-5.530393e-01	-4.733701e-01	-1.297699e+00	-0.752868	-0.410106	-1.00000	0.000000	0.
50%	2.050779e-01	2.583733e-01	4.540705e-01	-0.752868	-0.410106	1.00000	1.000000	0.
75%	7.590867e-01	2.583733e- 01	4.540705e- 01	0.727772	-0.410106	1.00000	1.000000	0.
max	2.246163e+00	7.575807e + 00	1.329955e+00	2.948733	4.145186	1.00000	1.000000	1.

df["class"].value_counts()

```
class
```

0 248

2 150

1 2

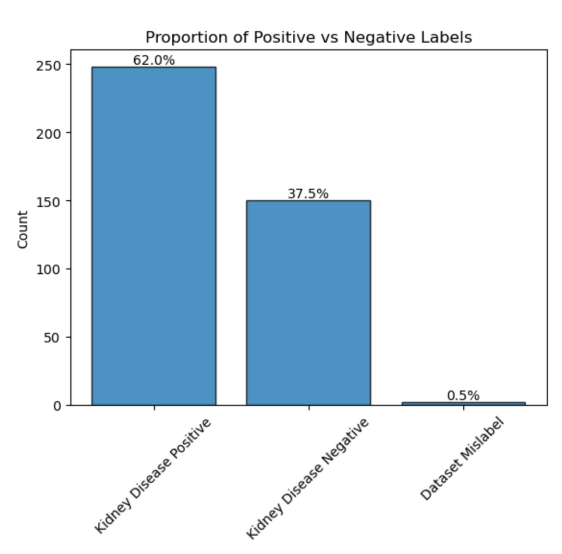
Name: count, dtype: int64

```
fig, ax = plt.subplots(1, 1)
bar_data = df["class"].value_counts()
ax.bar(range(len(bar_data)), bar_data, edgecolor="black", alpha=0.8)

ax.set_xticks([0, 1, 2])
ax.set_xticklabels(["Kidney Disease Positive", "Kidney Disease Negative", "Dataset Mislabel"],

for i, count in enumerate(bar_data):
    percentage = count / bar_data.sum() * 100
```

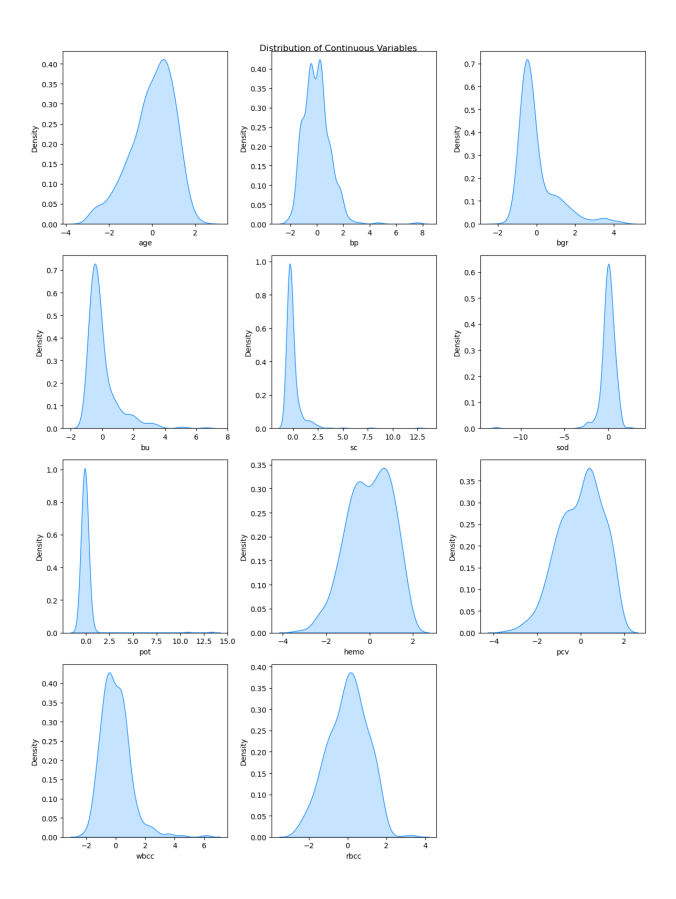
```
ax.text(i, count, f"{percentage:.1f}%", ha="center", va="bottom")
ax.set_ylabel("Count")
ax.set_title("Proportion of Positive vs Negative Labels")
plt.show()
```



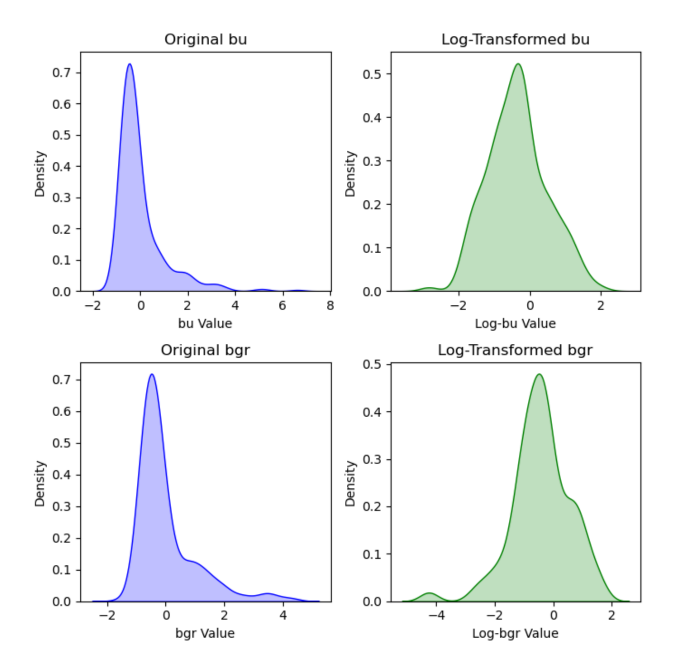
Visualizing distribution of continuous variables with Kernel Density Estimation

```
num_vars = ['age', 'bp', 'bgr', 'bu', 'sc', 'sod', 'pot', 'hemo', 'pcv', 'wbcc', 'rbcc']
num_features = len(num_vars)
num_rows = 4  # Number of rows in the subplot grid
```

```
num_cols = 3  # Number of columns in the subplot grid
fig, axes = plt.subplots(num_rows, num_cols, figsize=(4 * num_cols, 4 * num_rows))
for i, cont_feature in enumerate(df[num_vars]):
   row = i // num_cols # Calculate the row index for the subplot
    col = i % num_cols # Calculate the column index for the subplot
   ax_kde = axes[row, col]
   # Plot KDE for the feature
    sns.kdeplot(df[cont_feature], ax=ax_kde, fill=True, color="dodgerblue")
# Remove empty subplots
for i in range(num_features, num_rows * num_cols):
   fig.delaxes(axes.flatten()[i])
plt.suptitle("Distribution of Continuous Variables")
plt.tight_layout()
plt.show()
```



```
# Create a new DataFrame with selected variables and their transformations
data_log_vis = pd.DataFrame({
    'bu': df['bu'],
    'log_bu': np.log(df['bu'] + 1), # Log transform with handling zero values
    'bgr': df['bgr'],
    'log_bgr': np.log(df['bgr'] + 1)
})
# Variables to plot
variables = ['bu', 'bgr']
# Create a figure with 2 rows and 2 columns
fig, axes = plt.subplots(2, 2, figsize=(7, 7))
axes = axes.flatten() # Flatten to simplify indexing
for i, var in enumerate(variables):
    # Original Data Plot
    sns.kdeplot(data_log_vis[var], ax=axes[2*i], fill=True, color="blue")
    axes[2*i].set_title(f"Original {var}")
    axes[2*i].set_xlabel(f"{var} Value")
    axes[2*i].set_ylabel("Density")
    # Log-Transformed Data Plot
    sns.kdeplot(data_log_vis[f'log_{var}'], ax=axes[2*i+1], fill=True, color="green")
    axes[2*i+1].set_title(f"Log-Transformed {var}")
    axes[2*i+1].set_xlabel(f"Log-{var} Value")
    axes[2*i+1].set_ylabel("Density")
plt.tight_layout()
plt.show()
```



Observations: 1. The dataset has an imbalance in the number of kidney disease positive vs negative examples. Our visual exploratory data analysis also revealed that there are two mislabeled variables in the dataset's target column. The column in the dataset should include only "positive" or "negative" Kidney disease status, but there were a few examples with a third label. We discuss this more in the outliers section. 2. Many of the variables look roughly noramlly distributed, except that the blood glucode random and blood urea features are long-tailed. This has implications for feature engineering: we expect that log-transforming these features will make them closer to a normal distribution; this is likely to improve performance on classifiers such as logistic regression. We

visualized these variables log-transformed to confirm that they look closer to a normal distribution after the transformation 3. Most variables are continuous, although the specific gravity, albumin and sugar levels are categorical.

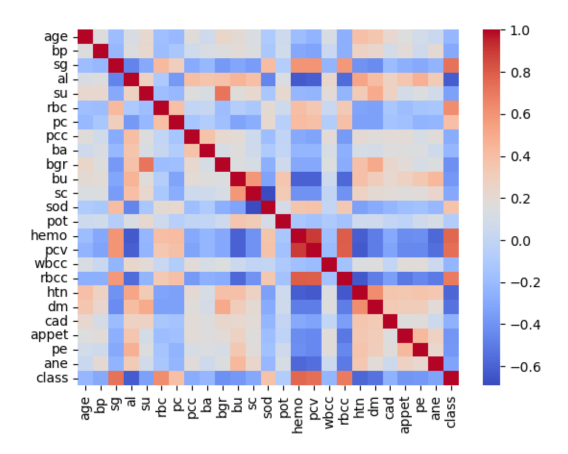
In this stage of exploratory data analysis, we are unsure whether we will end up using the log-transformed variables. This will depend on our results we obtain later.

4. Association Between Variables: Analyze variable relationships and their implications for feature selection or extraction (at least three statements)

```
correlation = df.corr()
sns.heatmap(correlation, cmap='coolwarm')
correlation
```

	age	bp	sg	al	su	rbc	pc	pcc	ba
age	1.000000	0.159480	-0.191096	0.122091	0.220866	-0.181683	-0.209743	0.169865	0.065425
bp	0.159480	1.000000	-0.218836	0.160689	0.222576	-0.194643	-0.129873	0.074018	0.126518
sg	-0.191096	-0.218836	1.000000	-0.469760	-0.296234	0.421101	0.299093	-0.290210	-0.220317
al	0.122091	0.160689	-0.469760	1.000000	0.269305	-0.110803	-0.375461	0.403257	0.366845
su	0.220866	0.222576	-0.296234	0.269305	1.000000	-0.187230	-0.221037	0.156997	0.115534
rbc	-0.181683	-0.194643	0.421101	-0.110803	-0.187230	1.000000	0.393821	0.002845	0.019199
pc	-0.209743	-0.129873	0.299093	-0.375461	-0.221037	0.393821	1.000000	-0.136040	-0.088435
pcc	0.169865	0.074018	-0.290210	0.403257	0.156997	0.002845	-0.136040	1.000000	0.376102
ba	0.065425	0.126518	-0.220317	0.366845	0.115534	0.019199	-0.088435	0.376102	1.000000
bgr	0.244992	0.160193	-0.374710	0.379464	0.717827	-0.193079	-0.175899	0.215386	0.109492
bu	0.196985	0.188517	-0.314295	0.453528	0.168583	-0.071404	-0.323372	0.192276	0.167696
sc	0.132531	0.146222	-0.361473	0.399198	0.223244	-0.122191	-0.279445	0.060680	0.063784
sod	-0.100046	-0.116422	0.412190	-0.459896	-0.131776	0.197653	0.218343	-0.183387	-0.100474
pot	0.058377	0.075151	-0.072787	0.129038	0.219450	0.061364	-0.058745	-0.003962	0.001224
hemo	-0.192928	-0.306540	0.602582	-0.634632	-0.224775	0.402049	0.418814	-0.295985	-0.233115
pcv	-0.242119	-0.326319	0.603560	-0.611891	-0.239189	0.350038	0.391230	-0.326328	-0.230173
wbcc	0.118339	0.029753	-0.236215	0.231989	0.184893	0.029804	-0.079035	0.184171	0.115111
									Į.

	age	bp	sg	al	su	rbc	pc	pcc	ba
rbcc	-0.268896	-0.261936	0.579476	-0.566437	-0.237448	0.339400	0.390282	-0.371968	-0.266713
htn	0.389724	0.277324	-0.410243	0.525234	0.321166	-0.321229	-0.344689	0.206843	0.111083
dm	0.354065	0.235513	-0.436692	0.406456	0.500133	-0.345661	-0.345482	0.173907	0.099610
cad	0.221807	0.098398	-0.195717	0.272713	0.276542	-0.129224	-0.154193	0.184861	0.157115
appet	0.148648	0.184732	-0.268856	0.359009	0.089770	-0.190258	-0.172015	0.193949	0.155157
pe	0.085726	0.062676	-0.298504	0.477127	0.144712	-0.143371	-0.244199	0.113742	0.141271
ane	0.041271	0.204279	-0.243082	0.322958	0.077908	-0.135308	-0.233601	0.178299	0.064608
class	-0.222361	-0.297019	0.729117	-0.625585	-0.345589	0.630148	0.397401	-0.283455	-0.222438



Observations: 1. Correlations between the following: White bloodcell count-packed cell volume (Hemo and pcv features), red bloodcell count-hemoglobin (hemo and rbcc features), Packed cell volume-red blood cell count (pcv and rbcc features) have the three highest positive correlations. 2. Correlations between the following: Serum creatinine and sodium (sc and sod features), Hemoglobin and hypertension (hemo and htm features), Packed cell volume and hypertension (pcv and htm

features), Hemoglobin and anemia (hemo and ane features), packed cell volume and anemia (pcv and ane features) have the highest negative correlations. 3. Highly correlated features can lead to overfitting or redundant information. We can get rid of redundant features which leads to simpler models. 4. The packed cell volume feature is highly correlated with many variables. This means that on it's own, it adds very little information. For that reason, we opted to remove that feature. We did some initial testing to verify that it did not degrade performance of our classifiers, and found that it had no impact on performance. Including it, then, would simply add what amounts to random noise, making the model less robust for future predictions.

```
df = df.drop(["pcv"], axis=1)
```

5. Missing Value Analysis and Handling: Implement your strategy for identifying and addressing missing values in the dataset, or provide reasons for not addressing them.

```
# Missing Value Analysis
missing_values = df.isnull().sum()
print(missing_values)
```

9 age 12 bp 47 sg 46 al S11 49 0 rbc рс 0 0 рсс 0 ba 44 bgr 19 bu sc 17 87 sod 88 pot

```
52
hemo
          106
wbcc
rbcc
          131
htn
            0
dm
            0
            0
cad
appet
            0
ре
            0
ane
class
            0
dtype: int64
```

```
# Mean imputer for numerical values and most frequent imputer for categorical values
num_vars = ['age', 'bp', 'bgr', 'bu', 'sc', 'sod', 'pot', 'hemo', 'pcv', 'wbcc', 'rbcc']
num_vars = ['age', 'bp', 'bgr', 'bu', 'sc', 'sod', 'pot', 'hemo', 'wbcc', 'rbcc']
cat_vars = ['sg', 'al', 'su']
imputer_num = SimpleImputer(strategy='mean')
imputer_cat = SimpleImputer(strategy='most_frequent')

df[num_vars] = imputer_num.fit_transform(df[num_vars])
df[cat_vars] = imputer_cat.fit_transform(df[cat_vars])
```

- For numerical features (age, blood pressure, blood glucose random, blood urea, serum creatinine, sodium, potassium, hemoglobin, packed cell volume, white blood cell count, red blood cell count), we'll use mean imputation.
- For categorical features (specific gravity, albumin, sugar), we'll use mode imputation.
- Binary features (red blood cells, pus cell, pus cell clumps, bacteria, hypertension, diabetes
 mellitus, coronary artery disease, appetite, pedal edema, anemia) already have no missing
 values.
- 6. Outlier Analysis: Implement your approach for identifying and managing outliers, or provide reasons for not addressing them.

```
display(df["dm"].value_counts())
display(df["class"].value_counts())
dm
 1
      260
 2
      137
-1
        2
        1
Name: count, dtype: int64
class
0
     248
2
     150
       2
1
Name: count, dtype: int64
\mbox{\tt\#}\ \mbox{\tt I} noticed dm has 1s and 2s, so I converted them to 0s and 1s
# Class has Os and 2s, so I converted them to Os and 1s
df['dm'] = df['dm'].replace({2:1, 1:0, -1:0})
# df['dm'] = df['dm'].replace({'2':1, '1':0,})
df['class'] = df['class'].replace({2:1})
display(df["dm"].value_counts())
display(df["class"].value_counts())
dm
0
     263
1
     137
Name: count, dtype: int64
class
```

```
0 248
```

1 152

Name: count, dtype: int64

We found two errors in the dataset: features which are reported as binary in the data card which have more than two values in the dataset.

We found that for both the target variable and diabetes mellitus, which were each supposed to be binary variables, there were a few examples of additional categories. These additional categories were merged into the most frequent category.

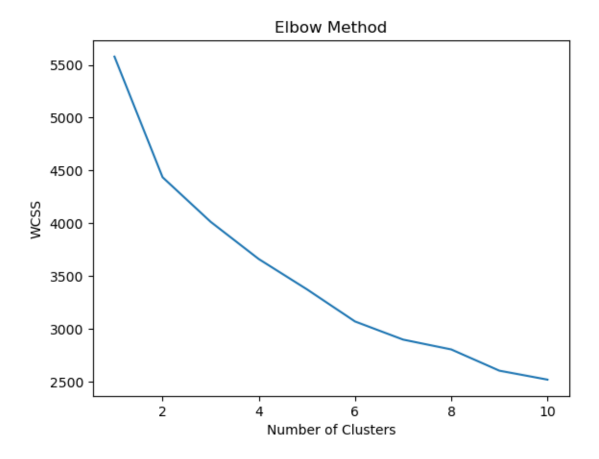
We ran our classification algorithms without any in-depth analysis regarding removal of outliers. Before doing so, we decided that if our classification performance is strong (accuracy above 90%), then we would likely not include any outlier analysis. We did find strong performance, and below is our justification for not removing outliers.

Since both our Random Forest and Logistic Regression classifiers are achieving extremley high accuracy, sensitivity, and specificity without any loss of performance due to the presence of outliers, we believe it's best to keep the outliers in the dataset. Removing them might cause me to lose valuable information and reduce the variability of the data, potentially affecting my ability to generalize to unseen data. Additionally, as Random Forest and Logistic Regression models are quite robust to outliers, we don't think their presence would significantly impact my performance. Also, we are cautious about making decisions regarding outlier removal to avoid introducing bias into the dataset and ensuring that any data manipulation doesn't compromise the integrity of my analysis results. Overall, if removing outliers doesn't lead to noticeable improvements in my performance, we would prefer to stick with the original dataset and keep the outliers intact.

7. Sub-group Analysis: Explore potential sub-groups within the data, employing appropriate data science methods to find the sub-groups of patients and visualize the sub-groups. The sub-group analysis must not include the labels (for CKD patients and healthy controls).

```
# Split data into features and target variable
X = df.drop('class', axis=1)
y = df['class']
```

```
# Determine the optimal number of clusters using the elbow method
wcss = []
for i in range(1, 11):
    kmeans = KMeans(n_clusters=i, init='k-means++', max_iter=300, n_init=10, random_state=0)
   kmeans.fit(X)
    wcss.append(kmeans.inertia_)
# Plot the elbow method graph to find the optimal number of clusters
plt.plot(range(1, 11), wcss)
plt.title('Elbow Method')
plt.xlabel('Number of Clusters')
plt.ylabel('WCSS')
plt.show()
# Based on the elbow method, choose the number of clusters
num_clusters = 2  # Adjust as needed
kmeans = KMeans(n_clusters=num_clusters, n_init=20, random_state=0)
kmeans.fit(X)
silhouette_avg = silhouette_score(X, kmeans.labels_)
print("Average Silhouette Score:", silhouette_avg)
```



Average Silhouette Score: 0.21162760671350317

```
# Determine optimal number of clusters using silhouette scores
range_n_clusters = [2, 3, 4]

optimal_k = (0,0)

for n_clusters in range_n_clusters:
    km = KMeans(n_clusters = n_clusters, n_init = 20, random_state=0)
    cluster_labels_km = km.fit_predict(X)

# average silhouette score

silhouette_avg_km = silhouette_score(X, cluster_labels_km)

# compute the silhouette scores for each sample

sample_silhouette_values = silhouette_samples(X, cluster_labels_km)

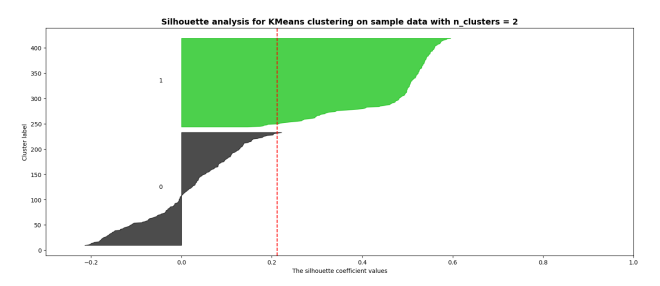
fig, ax1 = plt.subplots(1, 1)

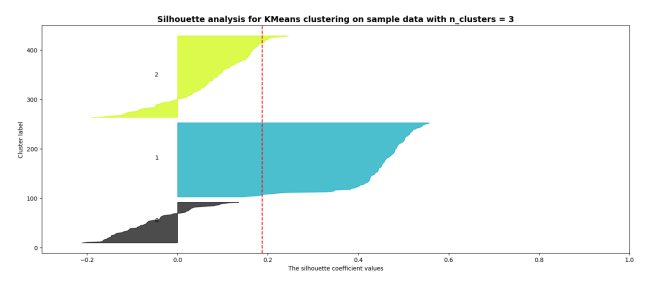
fig.set_size_inches(18, 7)

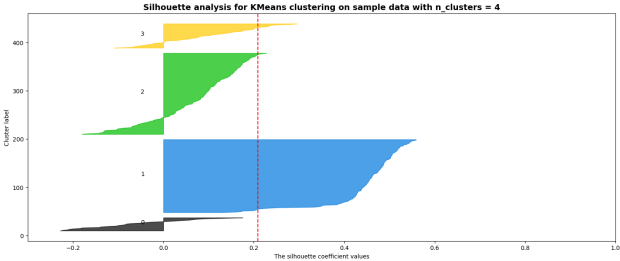
ax1.set_xlim([-0.3, 1])# change this based on the silhouette range
```

```
y_lower = 10
for i in range(n_clusters):
    # Aggregate the silhouette scores for samples belonging to
   # cluster i, and sort them
    ith_cluster_silhouette_values = sample_silhouette_values[cluster_labels_km == i]
    ith_cluster_silhouette_values.sort()
    size_cluster_i = ith_cluster_silhouette_values.shape[0]
   y_upper = y_lower + size_cluster_i
    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,
    )
    # label the silhouette plots with their cluster numbers at the middle
    ax1.text(-0.05, y_lower + 0.5 * size_cluster_i, str(i))
    # Compute the new y_lower for next cluster silhouette scores
   y_lower = y_upper + 10
ax1.set_title("The silhouette plot for various cluster")
ax1.set_xlabel("The silhouette coefficient values")
ax1.set_ylabel("Cluster label")
```

```
# vertical line for average silhouette score of all the values
ax1.axvline(x=silhouette_avg_km, color="red", linestyle="--")
plt.title(
    "Silhouette analysis for KMeans clustering on sample data with n_clusters = %d"
    % n_clusters,
    fontsize=14,
    fontweight="bold",
)
    optimal_k = (n_clusters, silhouette_avg_km) if optimal_k[1] < silhouette_avg_km else optimal
plt.show()
print("Optimal number of clusters:", optimal_k[0])</pre>
```







Optimal number of clusters: 2

```
# Apply k-means clustering with optimal k
kmeans_optimal = KMeans(n_clusters=optimal_k[0], n_init=20, random_state=0)
kmeans_optimal.fit(X)
cluster_counts = pd.Series(kmeans_optimal.labels_).value_counts().sort_index()
print("Number of observations within each cluster:")
print(cluster_counts)

# Perform PCA
pca = PCA()
```

```
df2_plot = pd.DataFrame(pca.fit_transform(X))
print("Variances: ",df2_plot.iloc[:,:].std(axis=0, ddof=0).to_numpy())

df2_plot.iloc[:,:].var(axis=0, ddof=0).plot(kind='bar', rot=0)
plt.ylabel('Variances')
```

Number of observations within each cluster:

0 224

1 176

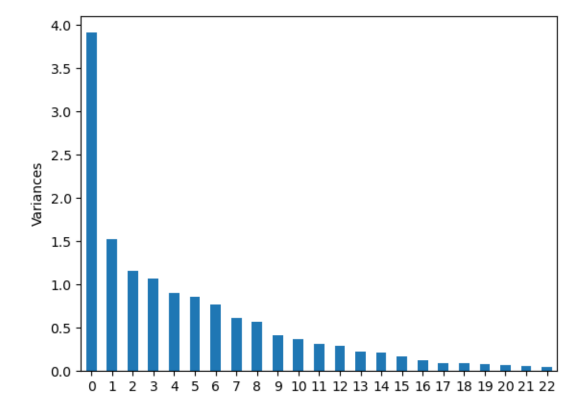
Name: count, dtype: int64

Variances: [1.97824553 1.23616004 1.07685887 1.03490043 0.9498406 0.92415524

0.88020906 0.78520753 0.75648553 0.64444749 0.60909821 0.56257017

0.54068757 0.47359118 0.45735511 0.40608239 0.35604038 0.30341904

Text(0, 0.5, 'Variances')



	PC1	PC2	PC3	PC4	PC5	PC6	PC7	PC8
Standard Deviation	1.978246	1.236160	1.076859	1.034900	0.949841	0.924155	0.880209	0.785208
Proportion of Variance	0.280628	0.109577	0.083155	0.076801	0.064695	0.061244	0.055558	0.044212
Cumulative Proportion	0.280628	0.390206	0.473361	0.550162	0.614857	0.676101	0.731658	0.775870

```
fig , (ax1,ax2) = plt.subplots(1,2, figsize=(15,5))

# Left plot
ax1.plot(pca.explained_variance_ratio_, '-o')
ax1.set_ylabel('Proportion of Variance Explained')
ax1.set_ylim(ymin=-0.01)

# Right plot
ax2.plot(np.cumsum(pca.explained_variance_ratio_), '-ro')
ax2.set_ylabel('Cumulative Proportion of Variance Explained')
ax2.set_ylim(ymax=1.05)

for ax in fig.axes:
    ax.set_xlabel('Principal Component')
    ax.set_xlim(0,15)
```

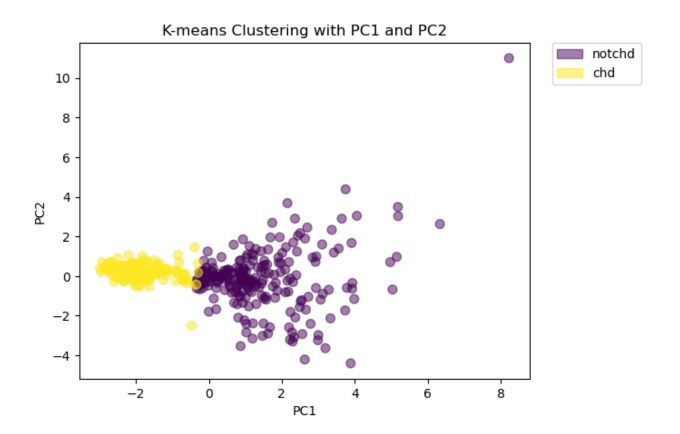
```
0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.25 - 0.
```

```
# Visualization of k-means cluster assignments using first two principal components
cmap = plt.cm.viridis

plt.scatter(df2_plot.iloc[:, 0], df2_plot.iloc[:, 1], c=kmeans_optimal.labels_, cmap=cmap, alpi
plt.xlabel('PC1')
plt.ylabel('PC2')
plt.title('K-means Clustering with PC1 and PC2')
handles = []
labels = pd.factorize(y.unique())
norm = mpl.colors.Normalize(vmin=0.0, vmax=1.0)

for i, v in zip(labels[0], labels[1]):
    handles.append(mpl.patches.Patch(color=cmap(norm(i)), label='chd' if v else 'notchd', alphe
plt.legend(handles=handles, bbox_to_anchor=(1.05, 1), loc=2, borderaxespad=0.)

plt.show()
```



loadings = pd.DataFrame(pca.components_.T, index=['PC'+str(i) for i in range(1,24)], columns=X
loadings

	age	bp	sg	al	su	rbc	pc	pcc	ba
PC1	0.187481	-0.142848	-0.332067	0.342348	-0.028598	-0.715559	-0.219458	-0.331568	0.109907
PC2	0.191835	-0.057524	-0.263973	0.177803	-0.525457	0.477178	-0.572327	-0.009737	0.045376
PC3	-0.305444	0.194334	-0.061244	0.324916	-0.001024	-0.046148	-0.122113	0.226060	-0.616412
PC4	0.301262	-0.142218	0.419900	-0.296534	-0.048340	-0.035415	-0.086280	-0.301424	-0.160720
PC5	0.213803	-0.459014	0.169075	0.371432	0.167873	0.190477	0.047790	0.135095	0.025224
PC6	-0.191137	0.055962	0.597954	0.125475	-0.031671	-0.155125	-0.416759	-0.115325	-0.207036
PC7	-0.172196	-0.094542	0.256790	-0.023357	-0.006085	0.093823	-0.200085	-0.306737	0.306594
PC8	0.059672	-0.035402	0.046848	-0.049790	0.000174	-0.045076	-0.035190	-0.046063	-0.030545
PC9	0.034915	-0.014265	0.036999	-0.035364	-0.017840	-0.005851	-0.031697	-0.030322	-0.017143
PC10	0.243771	-0.441432	0.049703	0.239993	0.296319	0.158132	0.091798	0.026811	-0.281345
PC11	0.331909	0.350027	0.212712	0.168158	-0.175769	-0.133963	0.065876	0.102201	-0.165886

	age	bp	sg	al	su	rbc	pc	pcc	ba
PC12	0.276794	0.468200	0.062802	0.256852	0.323272	0.086364	-0.127712	0.118008	0.228832
PC13	-0.224488	-0.290847	0.044785	0.015266	-0.518689	-0.168231	0.238112	0.132806	-0.097497
PC14	0.096974	0.099964	0.312560	0.418159	-0.381446	-0.015415	0.408705	0.071475	0.324271
PC15	-0.379467	-0.085367	0.057243	0.263912	0.169933	-0.005894	-0.104212	-0.031622	0.206046
PC16	0.091764	-0.195640	0.128557	-0.230602	0.019990	-0.307274	-0.339152	0.751861	0.243807
PC17	-0.300522	-0.069126	0.005050	0.188518	0.120493	0.063612	-0.043900	0.018577	0.110443
PC18	0.170662	-0.025807	-0.073268	-0.008285	-0.014891	-0.063309	-0.017943	-0.045158	-0.134117
PC19	0.154400	-0.092539	-0.076945	0.045352	0.043479	-0.031860	0.027395	0.025995	-0.089955
PC20	0.054064	-0.007259	-0.011827	0.016842	0.027190	-0.017680	-0.007664	-0.035311	-0.049205
PC21	0.089668	0.000769	-0.017763	-0.070699	-0.038262	-0.026814	-0.029931	0.006906	-0.082575
PC22	0.087658	0.013662	0.037247	-0.074564	-0.028892	-0.042924	0.027939	0.012466	-0.080151
PC23	0.083520	0.055551	0.004774	-0.037824	-0.067158	0.018269	0.015566	0.026223	-0.108663

```
# Identify variable with most significant influence on all PCs
sub_groups = set()
for i in range(0,23):
    most_influential_variable = loadings.iloc[i,:].idxmax()
    print("Variable with most significant influence " + '(PC'+str(i+1)+'):', most_influential_sub_groups.add(most_influential_variable)

print("\nSub Groups: ",sub_groups)
```

Variable with most significant influence (PC1): al
Variable with most significant influence (PC2): rbc
Variable with most significant influence (PC3): bu
Variable with most significant influence (PC4): bgr
Variable with most significant influence (PC5): al
Variable with most significant influence (PC6): sg
Variable with most significant influence (PC7): ba
Variable with most significant influence (PC8): htn

```
Variable with most significant influence (PC9): ane
Variable with most significant influence (PC10): sc

Variable with most significant influence (PC11): bp

Variable with most significant influence (PC12): hemo

Variable with most significant influence (PC13): hemo

Variable with most significant influence (PC13): bu

Variable with most significant influence (PC14): bu

Variable with most significant influence (PC15): pot

Variable with most significant influence (PC16): pcc

Variable with most significant influence (PC17): bgr

Variable with most significant influence (PC18): wbcc

Variable with most significant influence (PC20): pe

Variable with most significant influence (PC20): pe

Variable with most significant influence (PC21): rbcc

Variable with most significant influence (PC22): wbcc

Variable with most significant influence (PC22): pe
```

Sub Groups: {'rbc', 'bgr', 'bu', 'al', 'bp', 'pcc', 'hemo', 'pe', 'ane', 'sc', 'rbcc', 'sg',

- Variable Influence on Principal Components: After performing PCA, each principal component represents a linear combination of the original variables. The loadings of the original variables on each principal component indicate their influence on that component. The code iterates through each principal component and identifies the variable with the highest loading for that component. This variable is considered to have the most significant influence on that principal component.
- Identification of Subgroups: By identifying the variable with the highest loading for each principal component, we essentially identify the key features that contribute the most to the variance captured by each component. These identified variables serve as the subgroups. Each subgroup represents a set of variables that are most influential in defining the variance along a particular principal component axis.
- Interpretation of Subgroups: The identified subgroups provide insights into the underlying structure of the data. They represent the dimensions along which the data vary the most. For example, if 'al' (albumin) is identified as the most influential variable for a principal component, it suggests that variations in albumin levels contribute significantly to the variance

captured by that component. These subgroups help in understanding the key factors driving the patterns observed in the data and aid in interpretation. For example, variables like 'al' (albumin) and 'rbc' (red blood cell count) are highlighted as influential across multiple principal components, suggesting their importance in distinguishing different clusters.

```
# Compare true labels with k-means cluster assignments
adjusted_rand_index = round(adjusted_rand_score(y, kmeans_optimal.labels_), 2)
rand = rand_score(kmeans_optimal.labels_, y).round(2)
print("Rand Index:", rand)
print("Adjusted Rand Index:", adjusted_rand_index)
```

Rand Index: 0.87

Adjusted Rand Index: 0.74

Adjusted Rand Index/Rand Index: The comparison of true labels with KMeans cluster assignments using the Adjusted Rand Index and Rand Index provides a measure of clustering quality. A higher value indicates better agreement between the true labels and the clusters identified by KMeans.

8. Data Splitting: Segregate 30% of the data for testing, using a random seed of 1. Use the remaining 70% for training and model selection.

```
np.random.seed(1)

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

9. Classifier Choices: Identify the two classifiers you have chosen and justify your selections.

```
# Classifier Choices
rf = RandomForestClassifier()
lr = LogisticRegression()

rf.fit(X_train, y_train)
lr.fit(X_train, y_train)

# Model Evaluation, Presenting Additional Info
```

```
rf_pred = rf.predict(X_test)
rf_y_prob = rf.predict_proba(X_test)
lr_pred = lr.predict(X_test)
lr_y_prob = lr.predict_proba(X_test)
probT_rf = pd.DataFrame(
    data = {'prob0': rf_y_prob[:,1], 'y_test': y_test}
    )
probT_lr = pd.DataFrame(
    data = {'prob0': lr_y_prob[:,1], 'y_test': y_test}
    )
probT_rf['y_test_pred'] = probT_rf.prob0.map(lambda x: 1 if x>0.5 else 0)
probT_lr['y_test_pred'] = probT_lr.prob0.map(lambda x: 1 if x>0.5 else 0)
```

We chose K-Nearest Neighbours (KNN) because it is a relatively simple classification algorithm, which we wanted to use as a baseline to compare to other, more sophisticated methods.

We chose logistic regression because it is relatively simple and easy to interpret, but more sophisticated and powerful than KNN. It was important that we chose an interpretable model so that we could understand what factors cause the model to make predictions, and ensure that they make sense.

Beyond what we learned in class we also chose to include a random forest classifier because it is known for not overfitting and being able to handle multi dimensional data. It also works well when there is both numerical and catagorical data which we have in this case. Random forests are similar to the decision trees we learned about in class, but they use bootstrapping and random feature selection to make predictions more robust. A random forest is a collection of decision trees which are generated using bootstrapping. Each decision tree is trained on a random subset of the data and a random subset of features. During prediction, the results from all trees are majority-voted for classification to determine the predicted class.

10. Performance Metrics: Outline the two metrics for comparing the performance of the classifiers.

```
# Accuracy, Sensitivity, Specificity

def evaluate(y_test, y_test_pred):
    cm = confusion_matrix(y_test,y_test_pred)
    print('Confusion Matrix : \n', cm)

    total = sum(sum(cm))
    accuracy = (cm[0,0]+cm[1,1])/total
    print ('Accuracy : ', accuracy)
    sensitivity = cm[0,0]/(cm[0,0]+cm[0,1])
    print('Sensitivity : ', sensitivity )
    specificity = cm[1,1]/(cm[1,0]+cm[1,1])
    print('Specificity : ', specificity)
    print(classification_report(y_test, y_test_pred, zero_division=0.0))
```

Accuracy, sensitivity, and specificity are essential metrics derived from the confusion matrix used to evaluate the performance of classification models.

Accuracy measures the overall correctness of the model's predictions by calculating the ratio of correctly predicted observations to the total observations in the dataset.

Sensitivity, also known as recall or true positive rate, assesses the model's ability to correctly identify positive cases, calculated as the proportion of true positives to the sum of true positives and false negatives.

Specificity evaluates the model's capacity to correctly identify negative cases, represented as the ratio of true negatives to the sum of true negatives and false positives.

These metrics provide valuable insights into different aspects of the classifiers's behavior, helping to compare classifiers.

11. Feature Selection/Extraction: Implement methods to enhance the performance of at least one classifier in (9). The answer for this question can be included in (12).

```
# Feature selection using Backward selection (called Recursive Feature Elimination (RFE) in sk
from sklearn.feature_selection import RFE

rfe_selector = RFE(lr, n_features_to_select=10, step=1)

rfe_selector.fit(X_train, y_train)

selected_features_rfe = X_train.columns[rfe_selector.support_]
```

```
print("Are all selected in sub groups?", all([True for i in selected_features_rfe if i in sub_selected features Using Backward Selection
```

['bp', 'sg', 'al', 'su', 'rbc', 'bgr', 'sc', 'sod', 'hemo', 'pe']

print("Selected Features Using Backward Selection")

print(selected_features_rfe.to_list())

Are all selected in sub groups? True

We opted to use backward selection to select a subset of the most important features to retrain our models on.

Backward selection starts with all candidate features fits a logistic regression model to the full dataset. It then iteratively removes the least significant feature at each step. The process continues until a predefined stopping condition is met, in this case, that there are 10 features remaining (James et al. (2023), page 235).

12. Classifier Comparison: Utilize the selected metrics to compare the classifiers based on the test set. Discuss your findings (at least two statements).

Using all the features

```
# Classifier Choices and Instantiation

rf = RandomForestClassifier()

lr = LogisticRegression()

# Fit models to training data
```

```
rf.fit(X_train, y_train)
lr.fit(X_train, y_train)
# Model Evaluation
rf_pred = rf.predict(X_test)
rf_y_prob = rf.predict_proba(X_test)
lr_pred = lr.predict(X_test)
lr_y_prob = lr.predict_proba(X_test)
probT_rf = pd.DataFrame(
   data = {'prob0': rf_y_prob[:,1], 'y_test': y_test}
    )
probT_lr = pd.DataFrame(
    data = {'prob0': lr_y_prob[:,1], 'y_test': y_test}
probT_rf['y_test_pred'] = probT_rf.prob0.map(lambda x: 1 if x>0.5 else 0)
probT_lr['y_test_pred'] = probT_lr.prob0.map(lambda x: 1 if x>0.5 else 0)
print('Random Forest Classifier:\n')
evaluate(probT_rf.y_test, rf_pred)
print('Logistic Regression Classifier:\n')
evaluate(probT_lr.y_test, lr_pred)
Random Forest Classifier:
Confusion Matrix :
 [[70 0]
 [ 0 50]]
Accuracy: 1.0
Sensitivity: 1.0
Specificity: 1.0
             precision recall f1-score support
```

0	1.00	1.00	1.00	70
1	1.00	1.00	1.00	50
accuracy			1.00	120
macro avg	1.00	1.00	1.00	120
weighted avg	1.00	1.00	1.00	120

Logistic Regression Classifier:

Confusion Matrix :

[[70 0] [0 50]]

Accuracy: 1.0

Sensitivity: 1.0 Specificity: 1.0

support	f1-score	recall	precision	
70	1.00	1.00	1.00	0
50	1.00	1.00	1.00	1
120	1.00			accuracy
120	1.00	1.00	1.00	macro avg
120	1.00	1.00	1.00	weighted avg

Before performing KNN classification, we will first select the optimal value of K. This will be done using the technique learned in class: we will try a number of values, and select the one which yields best performance

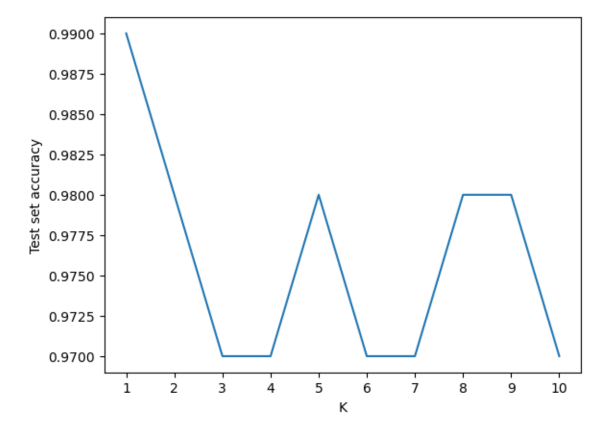
```
k_range = range(1, 11)
scores = []

for k in k_range:
```

```
knn = KNeighborsClassifier(n_neighbors=k)
knn.fit(X_train, y_train)
y_pred = knn.predict(X_test)
scores.append(round(metrics.accuracy_score(y_test, y_pred),2))
print(f"Best value of k: {k_range[np.argmax(np.array(scores))]}")
```

Best value of k: 1

```
plt.plot(k_range, scores)
plt.xlabel('K')
plt.ylabel('Test set accuracy')
plt.xticks(k_range)
plt.show()
```



K=1 has the best performance, so we select it

```
# KNN:
# KNN Classifier
knn = KNeighborsClassifier(n_neighbors=1)
# Fit model to training data
knn.fit(X_train, y_train)
# Model Evaluation
knn_pred = knn.predict(X_test)
knn_y_prob = knn.predict_proba(X_test)
probT_knn = pd.DataFrame(
   data={'prob0': knn_y_prob[:, 1], 'y_test': y_test}
)
probT_knn['y_test_pred'] = probT_knn.prob0.map(lambda x: 1 if x > 0.5 else 0)
print('KNN Classifier:\n')
evaluate(probT_knn.y_test, knn_pred)
KNN Classifier:
Confusion Matrix :
 [[69 1]
 [ 0 50]]
Accuracy: 0.991666666666667
Sensitivity: 0.9857142857142858
Specificity: 1.0
              precision recall f1-score
                                              support
           0
                   1.00
                             0.99
                                                   70
                                       0.99
           1
                   0.98
                             1.00
                                       0.99
                                                   50
```

accuracy			0.99	120
macro avg	0.99	0.99	0.99	120
weighted avg	0.99	0.99	0.99	120

Now using the selected features

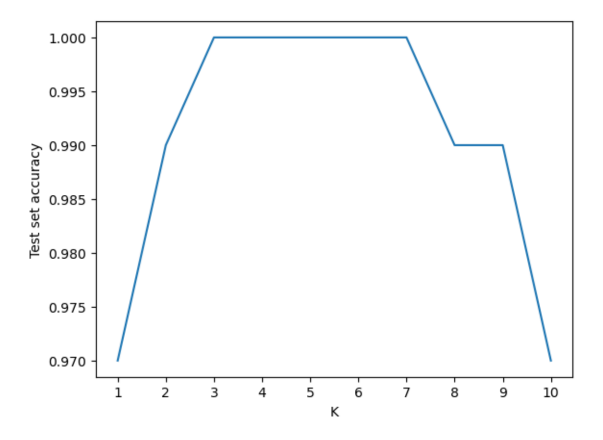
```
k_range = range(1, 11)
scores = []

for k in k_range:
    knn = KNeighborsClassifier(n_neighbors=k)
    knn.fit(X_train[selected_features_rfe], y_train)
    y_pred = knn.predict(X_test[selected_features_rfe])
    scores.append(round(metrics.accuracy_score(y_test, y_pred),2))

print(f"Best value of k: {k_range[np.argmax(np.array(scores))]}")

plt.plot(k_range, scores)
plt.xlabel('K')
plt.ylabel('Test set accuracy')
plt.xticks(k_range)
plt.show()
```

Best value of k: 3



This time, K=3 has the best performance, so we select it

```
# KNN:
# KNN2 Classifier
knn2 = KNeighborsClassifier(n_neighbors=3)

# Fit model to training data
knn2.fit(X_train[selected_features_rfe], y_train)

# Model Evaluation
knn2_pred = knn2.predict(X_test[selected_features_rfe])
knn2_y_prob = knn2.predict_proba(X_test[selected_features_rfe])

probT_knn2 = pd.DataFrame(
    data={'prob0': knn2_y_prob[:, 1], 'y_test': y_test})
```

```
probT_knn2['y_test_pred'] = probT_knn2.prob0.map(lambda x: 1 if x > 0.5 else 0)
print('KNN2 Classifier:\n')
evaluate(probT_knn2.y_test, knn2_pred)
KNN2 Classifier:
Confusion Matrix:
 [[70 0]
 [ 0 50]]
Accuracy: 1.0
Sensitivity: 1.0
Specificity: 1.0
             precision recall f1-score
                                             support
           0
                  1.00
                            1.00
                                      1.00
                                                  70
                  1.00
                            1.00
                                      1.00
                                                  50
           1
                                      1.00
                                                 120
   accuracy
  macro avg
                  1.00
                            1.00
                                      1.00
                                                 120
weighted avg
                  1.00
                            1.00
                                      1.00
                                                 120
# Classifier Choices and Instantiation
rf2 = RandomForestClassifier()
lr2 = LogisticRegression()
# Fit models to training data
rf2.fit(X_train[selected_features_rfe], y_train)
lr2.fit(X_train[selected_features_rfe], y_train)
```

Model Evaluation

```
rf_pred = rf2.predict(X_test[selected_features_rfe])
rf_y_prob = rf2.predict_proba(X_test[selected_features_rfe])
lr_pred = lr2.predict(X_test[selected_features_rfe])
lr_y_prob = lr2.predict_proba(X_test[selected_features_rfe])
probT_rf = pd.DataFrame(
    data={'prob0': rf_y_prob[:, 1], 'y_test': y_test}
probT_lr = pd.DataFrame(
    data={'prob0': lr_y_prob[:, 1], 'y_test': y_test}
probT_rf['y_test_pred'] = probT_rf.prob0.map(lambda x: 1 if x > 0.5 else 0)
probT_lr['y_test_pred'] = probT_lr.prob0.map(lambda x: 1 if x > 0.5 else 0)
print('Random Forest Classifier:\n')
evaluate(probT_rf.y_test, rf_pred)
print('Logistic Regression Classifier:\n')
evaluate(probT_lr.y_test, lr_pred)
Random Forest Classifier:
```

```
Confusion Matrix:

[[70 0]

[ 0 50]]

Accuracy: 1.0

Sensitivity: 1.0

Specificity: 1.0

precision recall f1-score support
```

1.00

1.00

1.00

1.00

0

1

1.00

1.00

70

50

accuracy			1.00	120
macro avg	1.00	1.00	1.00	120
weighted avg	1.00	1.00	1.00	120

Logistic Regression Classifier:

Confusion Matrix :

[[70 0] [0 50]]

Accuracy: 1.0

Sensitivity: 1.0

Specificity: 1.0

	precision	recall	f1-score	support
0	1.00	1.00	1.00	70
1	1.00	1.00	1.00	50
accuracy			1.00	120
macro avg	1.00	1.00	1.00	120
weighted avg	1.00	1.00	1.00	120

We were happy to see very strong performance with all of the methods we tested. We achieved perfect classification on the test set using logistic regression and random forest classification when we used all of the features in the dataset. The KNN classifier had one misclassification, leading to the following metrics:

Accuracy: 0.99 Sensitivity: 0.99 Specificity: 1.00

We found that in this case, the optimal value of K is 1. When training on the full dataset, increasing the value beyond 1 decreases accuracy. There are some fluctuations in accuracy as K increases, but the trend remains downwards. When K equals 1, the target datapoint is classified based on its nearest neighbor. While K=1 may offer optimal performance, it increases the classifier's sensitivity to errors caused by outliers in the training set. This occurs because an outlier might be in close

proximity to the target datapoint, despite the majority of nearby datapoints belonging to a different class.

When using only the subset of features obtained from backward selection, we found that the logistic regression and random forest classifiers maintained their optimal performance. This is a sign that the backward selection process worked well, as no valuable information was lost. Our KNN classifier also achived perfect performance on the test set once we trained it only on the features selected using backward selection. Additionally, the optimal value of K changed under these features. We found an optimal value of k=3. This is promising in terms of generalization performance because a higher k value than 1 reduces the model's sensitivity to noise and outliers in the training data, enhancing its ability to generalize to unseen data.

13. Interpretable Classifier Insight: After re-training the interpretable classifier with all available data, analyze and interpret the significance of predictor variables in the context of the data and the challenge (at least two statements).

```
# retrain Logistic regression on whole dataset:
lr = LogisticRegression()
lr.fit(X[selected_features_rfe], y)

# Interpret Logistic Regression coefficients
coefficients = pd.DataFrame(lr.coef_, columns=X_train[selected_features_rfe].columns)
coefficients_transposed = coefficients.transpose()
coefficients_transposed.columns = ['Coefficient']
coefficients_transposed = coefficients_transposed.assign(Abs_Coefficient=abs(coefficients_transposed = coefficients_transposed.sort_values(by='Abs_Coefficient', ascending=coefficients_transposed = coefficients_transposed.drop('Abs_Coefficient', axis=1)
print("Interpretable Classifier Insight:")
print(coefficients_transposed.head(10))
```

```
Interpretable Classifier Insight:
```

Coefficient

hemo 2.010680

```
-1.849457
al
          1.796728
rbc
         -1.080013
ре
          1.055752
sg
          0.897158
sod
         -0.852279
sc
bgr
         -0.674604
         -0.587153
bp
         -0.490523
su
```

For logistic regression, the value and the sign of the predictor variables are important in interpreting the results. In order to best interpret the results, we will first provide a brief overview of how to interpret them, and then point to a specific example and highlight what conclusions we may draw from our interpretation of it.

A positive coefficient for a given predictor variable implies a positive relationship between that variable and the outcome: when predictor variable increases so does the outcome. Similarly, negative implies a negative relationship. The particular value of a variable's coefficient indicates how strongly the predictor affects the outcome. A higher magnitude indicates that this variable implies a higher probability of either a positive or negative outcome (depending on the sign). Additionally, in logistic regression, we can use p-values to show whether each predictor variable is statistically significant. The p-value in logistic regression comes from a hypothesis test, where the null hypothesis is that the true slope of the variable's coefficient is 0. This null hypothesis holds that changing the variable has no impact on the probability of a patient having kidney disease. The lower the p-value, the more likely the variable is to not be zero and be useful in predicting the outcome.

In this case, for example, the albumin predictor (al feature) has a coefficient -1.849457. This means that if patient albumin levels increases by one unit then the log-odds of the patient having CKD decreases (since sign is negative) by -1.849457.

From an interpretability perspective this highlights that our logistic regression model predicts that as a patient's levels of albumin decrease, the risk of the kidney disease becomes higher. Low Albumin is an established marker for kidney damage within the biomedial literature (Cheng et al.

(2023)). We take it as a promising sign that our model identified a pattern which is well-established through other means.

Similar interpretations can be drawn with respect to the hemoglobin (hemo) feature, which has a positive coefficient of 2.010680. For a patient, each increase in 1 unit of hemoglobin results in our logistic regression model predicting that the log-odds of the patient having CKD increases (since sign is positive) by 2.010680. Other research has shown that those with higher hemoglobin are at higher risk of CKD (Oh et al. (2012)).

Similar conclusions can be drawn for other features based on their sign and magnitude.

14. Sub-group Improvement Strategy: If sub-groups were identified, propose and implement a method to improve one classifier performance further. Compare the performance of the new classifier with the results in (12).

```
from sklearn.ensemble import RandomForestClassifier
from sklearn.model_selection import cross_val_score
from sklearn.metrics import accuracy_score

X_subgroup1 = X_train[selected_features_rfe].iloc[:5]
y_subgroup2 = y_train.iloc[:5]

X_subgroup2 = X_train[selected_features_rfe].iloc[5:]
y_subgroup2 = y_train.iloc[5:]

# Train a separate Random Forest Classifier for each subgroup
rf_subgroup1 = RandomForestClassifier()
rf_subgroup1.fit(X_subgroup1, y_subgroup1)

rf_subgroup2 = RandomForestClassifier()
rf_subgroup2.fit(X_subgroup2, y_subgroup2)

# Combine predictions from both subgroups
def combined_predictions(X):
```

```
predictions_subgroup1 = rf_subgroup1.predict(X)
    predictions_subgroup2 = rf_subgroup2.predict(X)
    combined_predictions = []
    for pred1, pred2 in zip(predictions_subgroup1, predictions_subgroup2):
        # Combine predictions using a voting scheme, for example
        combined_predictions.append(max(pred1, pred2))
    return combined_predictions

# Evaluate the performance of the combined model
    combined_predictions_train = combined_predictions(X_train[selected_features_rfe])
    combined_predictions_test = combined_predictions(X_test[selected_features_rfe])

print("Combined Model Performance:")

print("Train Accuracy:", accuracy_score(y_train, combined_predictions_train))

print("Test Accuracy:", accuracy_score(y_test, combined_predictions_test))
```

Combined Model Performance:

Train Accuracy: 0.9821428571428571 Test Accuracy: 0.9916666666666667

- Creating Subgroups: The dataset X_train is split into two subgroups (X_subgroup1 and X_subgroup2) along with their corresponding target variables (y_subgroup1 and y_subgroup2). In this example, the first five instances are considered as subgroup 1 (X_subgroup1, y_subgroup1), and the remaining instances are considered as subgroup 2 (X_subgroup2, y_subgroup2).
- Training Separate Classifiers: Two random forest classifiers (rf_subgroup1 and rf_subgroup2) are trained separately on each subgroup. This allows each classifier to learn the patterns specific to its subgroup.
- Combining Predictions: A custom function combined_predictions is defined to combine the predictions from both subgroup classifiers. In this example, a simple maximum function is used where the final prediction is the max of the predictions from rf_subgroup1 and rf_subgroup2.

• Evaluating Combined Model Performance: The combined model's performance is evaluated on both the training and test datasets. The accuracy of the combined model on both datasets is printed to assess its performance. The output shows the accuracy of the combined model on the training set (98.21%) and the test set (99.17%). This approach demonstrates how training separate models for subgroups and combining their predictions can potentially improve overall model performance, especially in scenarios where different subgroups exhibit distinct patterns or characteristics.

Contributions

Jenna: - Created/set up repository and jupyter notebook - started working on questions 1-4 - started working on 11, - made general edits - q11-13 with Noah

Viransh: - References added, - done questions 5-10 - done question 14

Noah: - Finished question 3, - added visualizations and discussion of normality/log-transformation, - added writing and detail for questions 1, 2 and 4 - q11-13 with Jenna - Proofread and edited all writing in final stages

Github Link

Github link (https://github.com/JennaOrvitz/Stats3DA3FinalProject/tree/main)

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