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Drug Classification using Supervised Machine Learning

Overview

Data has been collected from a set of patients that all suffer from the same illness. Throughout their treatment, each patient responded to one of five different medications. The purpose of this exercise, is to develop a model that could help predict the appropriate drug for a future patient with the same illness based on certain characteristics about their health.

The model will be developed using three different supervised machine learning models: a decision tree, logistic regression, and a random forest.

This dataset came from Kaggle: https://www.kaggle.com/datasets/prathamtripathi/drugclassification See the citations section for detailed citation of this dataset.

The license of this dataset is a public domain with over 261 different code contributions and the size of the dataset is 2 kB.

The data will be imported using pandas.

Loading and Exploring Raw Dataset

```
In [ ]: import pandas as pd
        import numpy as np
        data = pd.read_csv('drugdata200.csv')
```

Now that the dataset has been imported, it's time to understand what is in this dataset.

```
data.info
In [ ]:
Out[]: <bound method DataFrame.info of
                                                      BP Cholesterol Na to K
                                                                              Drug
                                         Age Sex
                                 HIGH 25.355 DrugY
             23
                      HIGH
             47
                       LOW
                                 HIGH 13.093 drugC
        2
             47 M
                                 HIGH 10.114 drugC
                       LOW
       3
             28 F NORMAL
                                 HIGH 7.798 drugX
             61 F
                       LOW
                                 HIGH 18.043 DrugY
                                  . . .
            . . .
                       . . .
                                           . . .
             56 F
                                 HIGH 11.567 drugC
       195
                       LOW
        196
             16 M
                       LOW
                                 HIGH 12.006 drugC
       197
             52 M NORMAL
                                 HIGH 9.894 drugX
       198
             23 M NORMAL
                               NORMAL 14.020 drugX
                               NORMAL 11.349 drugX
        199
             40 F
                       LOW
        [200 rows x \in columns]>
```

```
data.shape
Out[]: (200, 6)
        data.dtypes
Out[]: Age
                          int64
                         object
        Sex
        BP
                        object
        Cholesterol
                        object
                        float64
        Na_to_K
                         object
        Drug
        dtype: object
```

Looking at the columns, this dataset uses a variety of factors including age, sex, blood pressure (BP), Cholesterol, and Na/k ratio (NA_to_K). Na is sodium and K is potassium. In recent years, doctors have been using the Na/K as a ratio to measure a person's diet quality and overall health according to the European Journal of Medical Research:

https://eurjmedres.biomedcentral.com/articles/10.1186/s40001-020-00476-5#citeas.

The drug column is the label and ultimately the target.

Data cleaning

First check to see if there are any missing values:

```
In [ ]: if data.isnull().values.any():
    print("Dataframe contains null values")
else:
    print("Dataframe has no null values")
```

Dataframe has no null values

Looking at the Na/K column, the number of decimal places will be reduce in order to make the data easier to look at.

```
In [ ]: #Rounds the decimal places
data["Na_to_K"] = data["Na_to_K"].round(2)
data.info
```

```
Out[]: <bound method DataFrame.info of
                                              Age Sex
                                                            BP Cholesterol Na_to_K
                                                                                       Drug
               23
                    F
                         HIGH
                                     HIGH
                                              25.36
                                                     DrugY
        1
              47
                          LOW
                                     HIGH
                                              13.09
                                                     drugC
                    Μ
        2
              47
                    Μ
                          LOW
                                     HIGH
                                              10.11
                                                     drugC
        3
              28
                    F
                       NORMAL
                                     HIGH
                                               7.80
                                                     drugX
        4
              61
                    F
                          LOW
                                     HIGH
                                              18.04
                                                     DrugY
         . .
              . . .
                   . .
                          . . .
                                      . . .
                                                . . .
                                                       . . .
        195
              56
                   F
                          LOW
                                     HIGH
                                              11.57
                                                     drugC
        196
                          LOW
                                              12.01
                                                    drugC
              16
                  Μ
                                     HIGH
                  M NORMAL
        197
              52
                                     HIGH
                                              9.89
                                                     drugX
        198
              23
                  М
                       NORMAL
                                   NORMAL
                                              14.02
                                                     drugX
        199
              40
                  F
                          LOW
                                   NORMAL
                                              11.35 drugX
        [200 rows x 6 columns]>
```

Look at how each drug is labeled:

```
In [ ]: unique_drugs = data["Drug"].unique()
    for drug in unique_drugs:
        print(drug)

DrugY
    drugC
    drugX
    drugA
    drugB
```

To reduce redundancy, each drug variable will be renamed.

```
In [ ]:
        #Create a mapping dictionary
        mapping = {
             'DrugY': 'Y',
             'drugC': 'C',
             'drugX': 'X',
             'drugA': 'A',
             'drugB': 'B'
        #Now use the dictionary to replace
        data["Drug"] = data["Drug"].replace(mapping)
        print(data)
            Age Sex
                          BP Cholesterol
                                          Na_to_K Drug
       0
             23
                       HIGH
                                    HIGH
                                             25.36
                                                      Υ
       1
                        LOW
                                    HIGH
                                            13.09
                                                      C
             47
                  Μ
       2
             47
                         LOW
                                    HIGH
                                             10.11
                                                      C
                  Μ
       3
             28
                  F NORMAL
                                    HIGH
                                            7.80
                                                      Χ
       4
             61
                        LOW
                                    HIGH
                                            18.04
                                                      Υ
                         . . .
                                    . . .
                                              . . .
            . . .
                 . .
                                                   . . .
       . .
                         LOW
                                            11.57
       195
             56
                 F
                                    HIGH
                                                      C
       196
                  Μ
                         LOW
                                    HIGH
                                             12.01
                                                      C
             16
       197
             52
                  M NORMAL
                                    HIGH
                                             9.89
                                                      Χ
       198
             23
                  Μ
                     NORMAL
                                  NORMAL
                                            14.02
                                                      Χ
       199
             40
                         LOW
                                  NORMAL
                                            11.35
                                                      Χ
```

[200 rows x 6 columns]

Let's also take a look at the statistics of the dataset

```
In [ ]: #Statistical Analysis
       statistics = data.describe()
       print(statistics)
                   Age
                          Na_to_K
      count 200.000000 200.000000
             44.315000 16.084900
      mean
      std
             16.544315 7.224417
      min
            15.000000 6.270000
            31.000000 10.447500
      50%
            45.000000 13.940000
             58.000000
      75%
                        19.382500
             74.000000
                        38.250000
      max
```

The numeric values all seem to be reasonable and no outliers are detected. It's important to check if there are any unique values for the other remaining categorical values to ensure a clean dataset.

```
In [ ]: # Unique Blood Pressure
        unique_BP = data["BP"].unique()
        for i in unique_BP:
            print(i)
        # Unique Cholesterol
        unique_cholesterol = data["Cholesterol"].unique()
        for i in unique_cholesterol :
            print(i)
        # Unique Sex
        unique_sex = data["Sex"].unique()
        for i in unique_sex :
            print(i)
       HIGH
       LOW
       NORMAL
       HIGH
       NORMAL
```

All of the categorical values have the expected values. There are not any typos or misplaced data labels.

The cleaning that was done for this dataset was done in order to make working with decimals in the "Na_to_K" easier and reduce the redundancy of the "Drug" column.

Exploratory Data Analysis (EDA)

The data looks good enought work with and explore at a higher lever. The purpose of doing this is to detect any outliers that might mess with the data or be a mistake.

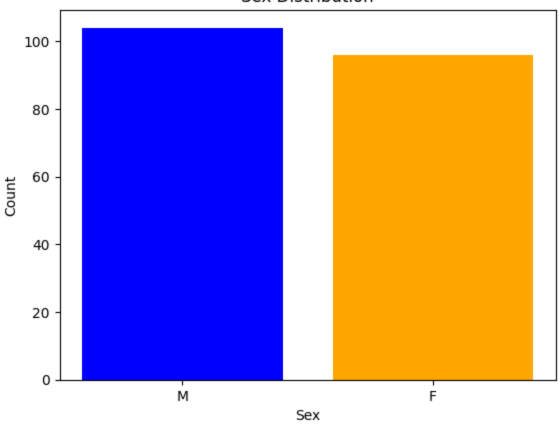
Additionally, deeper analysis could result in uncovering relationships between variables or variables and the label.

```
import matplotlib.pyplot as plt
# Age distribution
plt.hist(data["Age"], bins = 10, edgecolor = 'black')
plt.title("Age Distribution")
plt.xlabel("Age")
plt.ylabel("Frequency")
plt.show()
```

Age Distribution 25 20 10 5 Age Age

```
In []: # Sex distribution
    sex_counts = data["Sex"].value_counts()
    colors = ['blue', 'orange']
    plt.bar(sex_counts.index, sex_counts.values, color = colors)
    plt.title("Sex Distribution")
    plt.xlabel("Sex")
    plt.ylabel("Count")
    plt.show()
```

Sex Distribution



```
import seaborn as sns
# Blood pressure distribution

bp_counts = data["BP"].value_counts()

colors = sns.color_palette("colorblind")

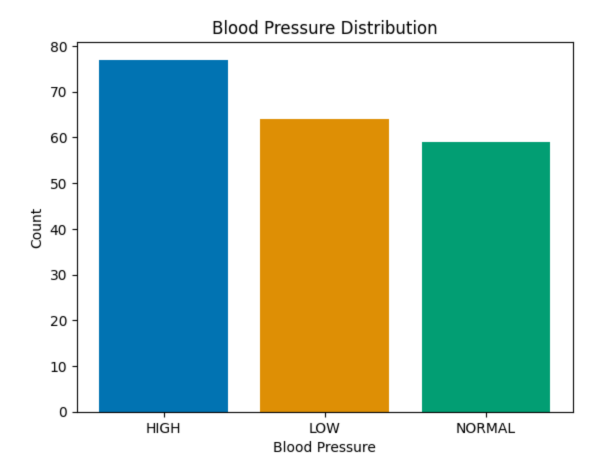
plt.bar(bp_counts.index, bp_counts.values, color = colors)

plt.title("Blood Pressure Distribution")

plt.xlabel("Blood Pressure")

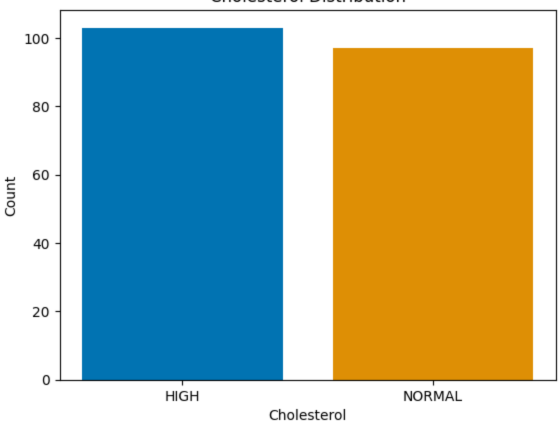
plt.ylabel("Count")

plt.show()
```

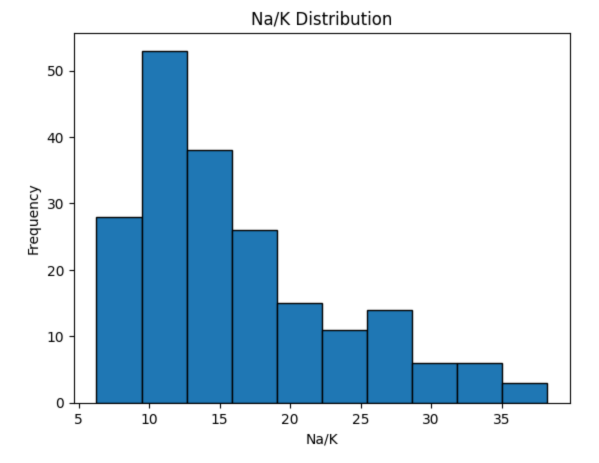


```
In []: # Cholesterol distribution
    Cholesterol_counts = data["Cholesterol"].value_counts()
    colors = sns.color_palette("colorblind")
    plt.bar(Cholesterol_counts.index, Cholesterol_counts.values, color = colors)
    plt.title("Cholesterol Distribution")
    plt.xlabel("Cholesterol")
    plt.ylabel("Count")
    plt.show()
```

Cholesterol Distribution



```
In [ ]: # Na/K distribution
plt.hist(data["Na_to_K"], bins = 10, edgecolor = 'black')
plt.title("Na/K Distribution")
plt.xlabel("Na/K")
plt.ylabel("Frequency")
plt.show()
```



Age, cholesterol, blood pressure, and sex all seem to have a good distribution but Na/K appears a little lopsided. Let's look at how this may affect things by comparing the N/K ratio across age, cholesterol, blood pressure, and sex.

```
In []: #Create plot comparing Na/K vs age by sex
    # Create the scatter plot
    plt.scatter(data[data['Sex'] == 'M']['Age'], data[data['Sex'] == 'M']['Na_to_K'], c
    plt.scatter(data[data['Sex'] == 'F']['Age'], data[data['Sex'] == 'F']['Na_to_K'], c

    plt.title("Scatter Plot: Na_to_K vs. Age by Sex")
    plt.ylabel("Age")
    plt.ylabel("Na_to_K")

plt.legend()

#Create plot comparing Na/K vs age by Cholesterol
    # Create the scatter plot
    plt.scatter(data[data['Cholesterol'] == 'HIGH']['Age'], data[data['Cholesterol'] == plt.scatter(data[data['Cholesterol'] == 'NORMAL']['Age'], data[data['Cholesterol']    plt.title("Scatter Plot: Na_to_K vs. Age by Cholesterol")
    plt.xlabel("Age")
```

```
plt.ylabel("Na_to_K")
plt.legend()

plt.show()

#Create plot comparing Na/K vs age by Blood Pressure
# Create the scatter plot

plt.scatter(data[data['BP'] == 'HIGH']['Age'], data[data['BP'] == 'HIGH']['Na_to_K']

plt.scatter(data[data['BP'] == 'NORMAL']['Age'], data[data['BP'] == 'NORMAL']['Na_t]

plt.scatter(data[data['BP'] == 'LOW']['Age'], data[data['BP'] == 'LOW']['Na_to_K'],

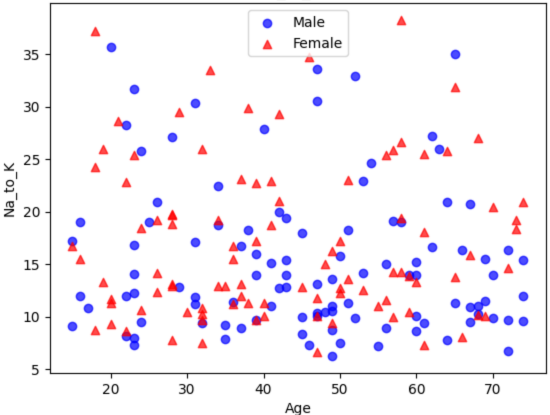
plt.title("Scatter Plot: Na_to_K vs. Age by Blood Pressure")

plt.xlabel("Age")
plt.ylabel("Na_to_K")

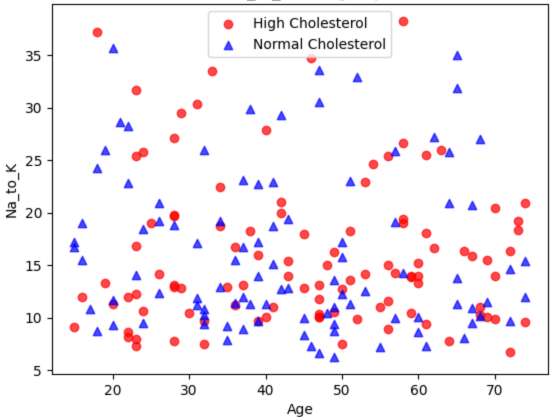
plt.legend()

plt.show()
```

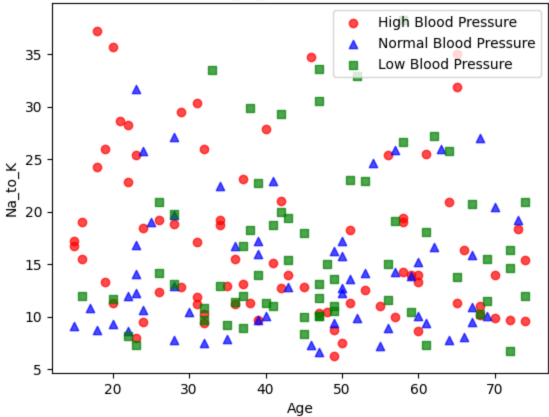
Scatter Plot: Na_to_K vs. Age by Sex







Scatter Plot: Na_to_K vs. Age by Blood Pressure



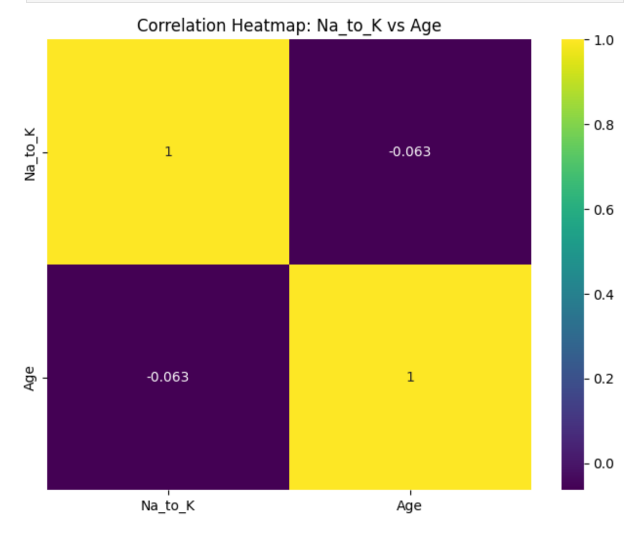
Based on the scatter plots above, the biggest takeaway is that there doesn't seem to be any relation among the variables. Further analysis will be needed to confirm if this is the case.

Since sex, cholesterol, and blood pressure are all categorical data some scatter plots were made since these values aren't useful for making a correlation heatmap or correlation matrix. This will create further confirmation about the relationship between variables.

```
In []: # Select the "Na_to_K" and "Age" columns
    columns = ["Na_to_K", "Age"]
    subset_data = data[columns]
    # Compute the correlation matrix
    correlation_matrix = subset_data.corr()

# Create the heatmap
    plt.figure(figsize=(8, 6))
    sns.heatmap(correlation_matrix, annot=True, cmap="viridis")

plt.title("Correlation Heatmap: Na_to_K vs Age")
    plt.show()
```



As expected based on the scatterplots, there is a very weak negative correlation between the two variables.

Overall it does not look like the variables will influence one another very much.

Exploring the variables and their relation to the drug type

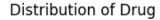
Continuing the exploration, it is important to take a look at how each drug type may relate to each variable.

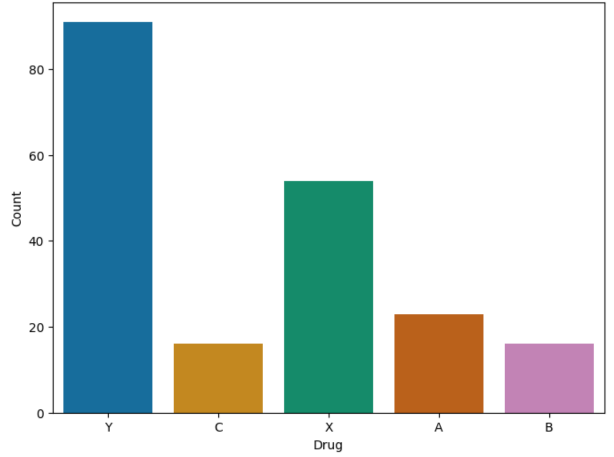
First, let's take a look at the distribution of the drug type.

```
In []: # Count the occurrences of each unique value in the "Drug" column
drug_counts = data["Drug"].value_counts()

# Create the bar plot
plt.figure(figsize=(8, 6))
sns.countplot(x="Drug", data=data, palette="colorblind")

plt.title("Distribution of Drug")
plt.xlabel("Drug")
plt.ylabel("Count")
plt.show()
```





There is a noticeable amount of Drug Y compared to the other drugs in this dataset. This will become important before any classification begins, but for now more analysis is required.

Violin plots will be used to examine the distribution of drugs across age and Na/K ratio becuase they are numeric.

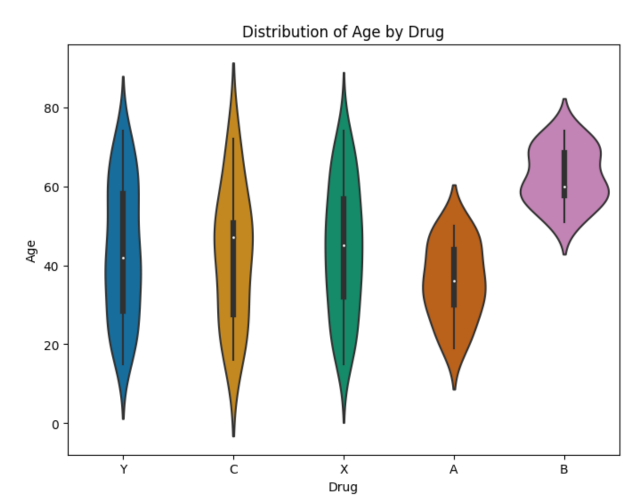
```
In []: # Violin Plots
# Drug and age
plt.figure(figsize=(8, 6))
sns.violinplot(x="Drug", y="Age", data=data, palette="colorblind")

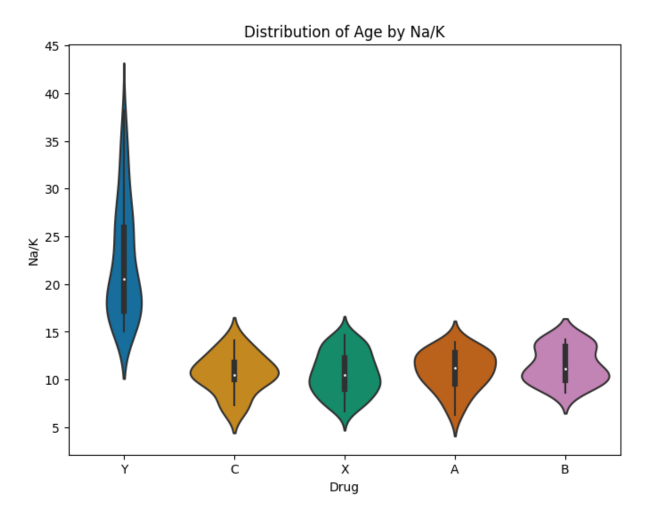
plt.title("Distribution of Age by Drug")
plt.xlabel("Drug")
plt.ylabel("Age")

plt.show()

# Drug and Na/K
plt.figure(figsize=(8, 6))
sns.violinplot(x="Drug", y="Na_to_K", data=data, palette="colorblind")

plt.title("Distribution of Age by Na/K")
plt.xlabel("Drug")
plt.ylabel("Na/K")
```





Now bar plots can be used comparing the drugs vs Blood Pressure, Cholesterol, and sex.

```
In []: #Create plot for Blood Pressure
    plt.figure(figsize=(8, 6))
    sns.countplot(x="BP", hue="Drug", data=data, palette="colorblind")

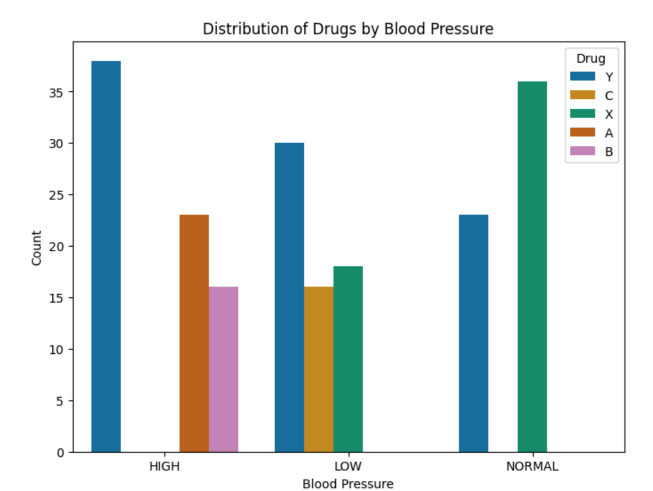
# Set plot title and axis labels
    plt.title("Distribution of Drugs by Blood Pressure")
    plt.xlabel("Blood Pressure")
    plt.ylabel("Count")

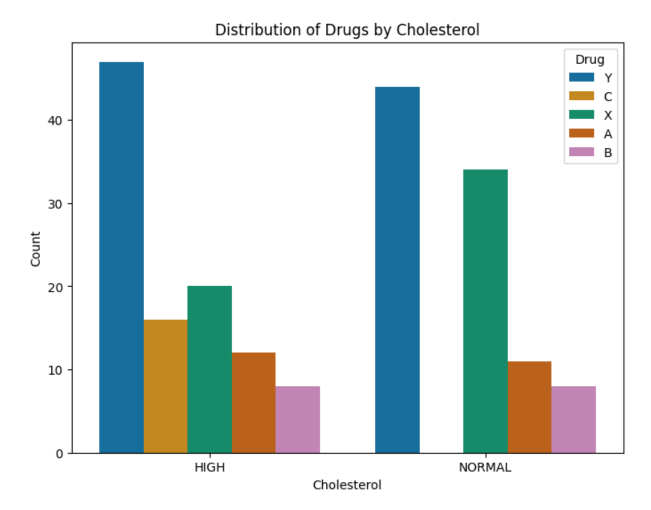
# Add a Legend
    plt.legend(title="Drug", loc="upper right")

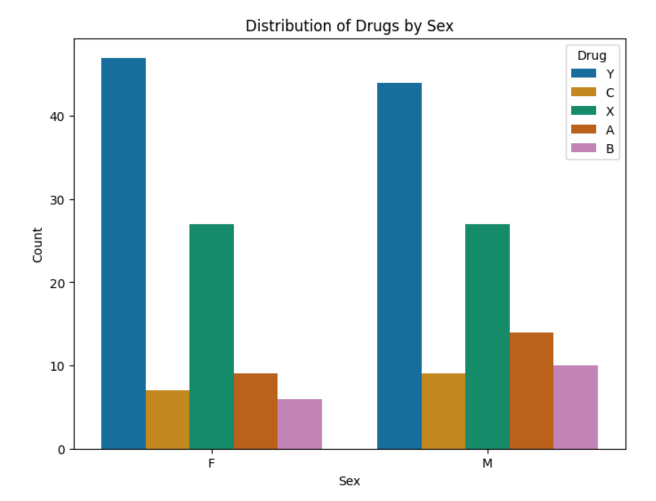
# Display the plot
    plt.show()

#Create plot for Cholesterol
    plt.figure(figsize=(8, 6))
    sns.countplot(x="Cholesterol", hue="Drug", data=data, palette="colorblind")
```

```
# Set plot title and axis labels
plt.title("Distribution of Drugs by Cholesterol")
plt.xlabel("Cholesterol")
plt.ylabel("Count")
# Add a Legend
plt.legend(title="Drug", loc="upper right")
# Display the plot
plt.show()
#Create plot for Sex
plt.figure(figsize=(8, 6))
sns.countplot(x="Sex", hue="Drug", data=data, palette="colorblind")
# Set plot title and axis labels
plt.title("Distribution of Drugs by Sex")
plt.xlabel("Sex")
plt.ylabel("Count")
# Add a Legend
plt.legend(title="Drug", loc="upper right")
# Display the plot
plt.show()
```







Here are the takeaway for comparing the drugs vs variables:

- -Age: Drugs Y, X, and C appear to be prescribed across all ages. However, drug A does not appear to be for young or old people while drug B appears to be exclusively for older people.
- -Na/K: Drug Y appears to be prescribed to people with higher Na/K ratios. Really anything with a ratio greater than 15. The other drugs, appear to all be for low ratios.
- -Blood Pressure: Drug Y appears across all 3 levels of BP, drug C is only found with the low BP population, drugs A and B are only found with the high BP population, and drug X is found with the low and normal BP populations.
- -Cholesterol: Drug C is only found in the high cholesterol population while the other 4 drugs are evenly dispersed across high and low cholestorol populations.
- -Sex: All drugs are evenly distriubted across both sexes.

Models

Dataset Preparation for Models

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> After going through the EDA and looking at the results of the correlation matrix as well as all the other plots, there is some potential for colinearity. This is indidcated in the plots done in the previous section as well as the takeaways. While collinearity may not be inherently bad, it could result in overfitting. As a result, feature engineering will be done later to address this concern.

First step is to split the dataset into training and testing data. The data will be split into 80% training and 20% testing. The sklearn library will be important for this.

```
In [ ]: # Import important libraries
        from sklearn.metrics import confusion_matrix
        from sklearn.metrics import classification_report
        from sklearn.model selection import train test split
        # Remove the features and set the target variable
        # X will remove the features, y will be the target variable
        X = data.drop("Drug", axis = 1)
        y = data["Drug"]
        # Split the dataset
        X_train, X_test, y_train, y_test = train_test_split(X, y, test_size = 0.2, random_s
        # To understand the training and testing a little better it's best to see the shape
        print(X_train.shape, y_train.shape)
        print(X_test.shape, y_test.shape)
       (160, 5) (160,)
```

(40, 5) (40,)

The shapes appear to be what is expected.

Now, some feature engineering needs to be done on the training and testing datasets to improve the overall performance of the machine learning models. This is because much of the data is categorical values.

```
In [ ]: # Import from the sklearn library
        from sklearn.preprocessing import LabelEncoder
        # Apply label encoding to the categorical columns
        label_encoder = LabelEncoder()
        categorical_columns = ["BP", "Cholesterol", "Sex"]
        for column in categorical columns:
            X[column] = label_encoder.fit_transform(X[column])
        # Split the dataset again
        X_train, X_test, y_train, y_test = train_test_split(X, y, test_size = 0.2, random_s
        X_train.head()
```

Out[]:		Age	Sex	BP	Cholesterol	Na_to_K
	79	32	0	1	1	10.84
	197	52	1	2	0	9.89
	38	39	0	2	1	9.71
	24	33	0	1	0	33.49
	122	34	1	2	0	22.46

```
In [ ]: X_test.head()
```

Out[]:		Age	Sex	ВР	Cholesterol	Na_to_K
	95	36	1	1	1	11.42
	15	16	0	0	1	15.52
	30	18	0	2	1	8.75
	158	59	0	1	0	10.44
	128	47	1	1	1	33.54

The last thing needed to be done for the models, is to prevent overfitting.

As stated previously, "Y" drug is far more prevelant than the other drugs. Since there's just one drug that is higher than the others, undersampling will be the technique of choice.

```
In []: # Import the necessary libraries
!pip install imbalanced-learn
from imblearn.under_sampling import RandomUnderSampler

# Use an instance of the under-sampler
undersampler = RandomUnderSampler(random_state = 42)
X_train, y_train = undersampler.fit_resample(X_train, y_train)
```

Requirement already satisfied: imbalanced-learn in /usr/local/python/3.10.8/lib/pyth on3.10/site-packages (0.10.1)

Requirement already satisfied: numpy>=1.17.3 in /home/codespace/.local/lib/python3.1 0/site-packages (from imbalanced-learn) (1.24.3)

Requirement already satisfied: scipy>=1.3.2 in /home/codespace/.local/lib/python3.1 0/site-packages (from imbalanced-learn) (1.10.1)

Requirement already satisfied: scikit-learn>=1.0.2 in /home/codespace/.local/lib/pyt hon3.10/site-packages (from imbalanced-learn) (1.2.2)

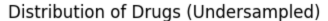
Requirement already satisfied: joblib>=1.1.1 in /home/codespace/.local/lib/python3.1 0/site-packages (from imbalanced-learn) (1.2.0)

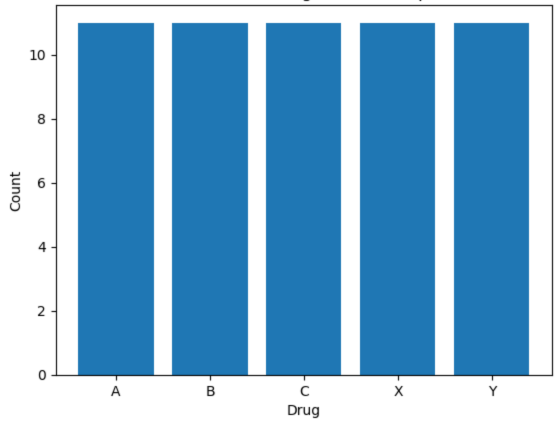
Requirement already satisfied: threadpoolctl>=2.0.0 in /home/codespace/.local/lib/py thon3.10/site-packages (from imbalanced-learn) (3.1.0)

```
In [ ]: import matplotlib.pyplot as plt
import seaborn as sns
# Calculate the count of each drug category in the undersampled data
```

```
drug_counts = y_train.value_counts()

# Create a bar plot to visualize the drug distribution
plt.bar(drug_counts.index, drug_counts.values)
plt.xlabel('Drug')
plt.ylabel('Count')
plt.title('Distribution of Drugs (Undersampled)')
plt.show()
```





Model Selection

Three models will be used on this dataset.

Decision tree classifier: this model was chosen because there does not seem to be much of any linear relationship in the data and decision tree classifiers handle randomization very well.

Logistic Regression: this model was chosen because it is reliant on linear relationships in order to do a good job of predicting. This will be a good way to compare and contrast against the other chosen models. Also, while there does not appear to be a strong linear relationship in the data, there is some. Potentially that means this model could be good at predicting things the other models aren't.

Random forest: this model was chosen for the same reason the decision tree classifier was except that random forests are a more complex version of a decision tree classifier. This

should mean that the random forest will perform as well or better than the decision tree.

The first model used for this dataset will be a decision tree classifier.

```
In [ ]: from sklearn.tree import DecisionTreeClassifier
        from sklearn.metrics import accuracy score
        # Make the DTC. Include parameters in order to introduce some randomness. Otherwi
        clf = DecisionTreeClassifier(max_depth=3, min_samples_split=6, min_samples_leaf=8,
        # Training data
        clf.fit(X_train, y_train)
        # Predicitons on testing data
        y_pred_dt = clf.predict(X_test)
        # Create a classification report
        dt_classification_report = classification_report(y_test, y_pred_dt)
        # Show the accuracy
        accuracy_dt = accuracy_score(y_test, y_pred_dt)
        print(dt_classification_report)
        print("Accuracy:", accuracy_dt)
        # Get feature importances
        feature_importances = clf.feature_importances_
        # Get feature names
        feature_names = X.columns
        # Sort feature importances in descending order
        sorted_indices = feature_importances.argsort()[::-1]
        sorted_importances = feature_importances[sorted_indices]
        sorted_feature_names = feature_names[sorted_indices]
```

	precision	recall	f1-score	support
Α	1.00	1.00	1.00	6
В	1.00	1.00	1.00	3
С	0.45	1.00	0.62	5
X	1.00	0.45	0.62	11
Υ	1.00	1.00	1.00	15
accuracy			0.85	40
macro avg	0.89	0.89	0.85	40
weighted avg	0.93	0.85	0.85	40

Accuracy: 0.85

The next model will be made using logistical regression

```
In [ ]: from sklearn.linear_model import LogisticRegression
   logisticial_regression = LogisticRegression(solver = 'sag', max_iter = 200)
```

```
# Train the model
logisticial_regression.fit(X_train, y_train)

# Predict the testing data
y_pred_lr = logisticial_regression.predict(X_test)

# Create a classification report
lr_classification_report = classification_report(y_test, y_pred_lr)

# Summarize the accuracy
accuracy_lr = accuracy_score(y_test, y_pred_lr)
print(lr_classification_report)
print("Accuracy:", accuracy_lr)
```

	precision	recall	f1-score	support
А	0.67	0.33	0.44	6
В	0.38	1.00	0.55	3
С	1.00	0.60	0.75	5
Х	1.00	0.55	0.71	11
Υ	0.70	0.93	0.80	15
accuracy			0.70	40
macro avg	0.75	0.68	0.65	40
weighted avg	0.79	0.70	0.70	40

Accuracy: 0.7

```
/home/codespace/.local/lib/python3.10/site-packages/sklearn/linear_model/_sag.py:35
0: ConvergenceWarning: The max_iter was reached which means the coef_ did not conver ge
   warnings.warn(
```

The next model to be implemented will be Random Forest

```
In [ ]: from sklearn.ensemble import RandomForestClassifier

#Create instance of Random Forest. Make sure to include parameters to introduce ra
    rfc = RandomForestClassifier(n_estimators = 10, max_depth = 12, min_samples_split =

# Fit model to the training data
    rfc.fit(X_train, y_train)

# Make predictions
y_pred_rf = rfc.predict(X_test)

# Create a classification report
rf_classification_report = classification_report(y_test, y_pred_rf)

# Summarize the accuracy
accuracy_rf = accuracy_score(y_test, y_pred_rf)
print(rf_classification_report)
print("Accuracy:", accuracy_rf)
```

	precision	recall	f1-score	support
А	0.86	1.00	0.92	6
В	1.00	1.00	1.00	3
C	0.62	1.00	0.77	5
X	1.00	0.73	0.84	11
Υ	1.00	0.93	0.97	15
accuracy			0.90	40
macro avg	0.90	0.93	0.90	40
weighted avg	0.93	0.90	0.90	40

Accuracy: 0.9

Results and Summary

Here are the basic accuracy results of each model:

```
In [ ]: pip install tabulate
```

Requirement already satisfied: tabulate in /usr/local/python/3.10.8/lib/python3.10/s ite-packages (0.9.0)

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

Model	Accuracy
Decision Tree	0.85
Logistic Regression	0.7
Random Forest	0.9 ++

Here are some additional metrics to measure the performance of each model:

Decision Tree Classification Report

<pre>In []: print(dt_cl</pre>	assification _.	_report)		
	precision	recall	f1-score	support
А	1.00	1.00	1.00	6
В	1.00	1.00	1.00	3
С	0.45	1.00	0.62	5
Х	1.00	0.45	0.62	11
Υ	1.00	1.00	1.00	15
accuracy			0.85	40
macro avg	0.89	0.89	0.85	40
weighted avg	0.93	0.85	0.85	40

Logistic Regression Classification Report

<pre>[n []: print(lr_cla</pre>	assification_	_report)		
	precision	recall	f1-score	support
А	0.67	0.33	0.44	6
В	0.38	1.00	0.55	3
С	1.00	0.60	0.75	5
X	1.00	0.55	0.71	11
Υ	0.70	0.93	0.80	15
accuracy			0.70	40
macro avg	0.75	0.68	0.65	40
weighted avg	0.79	0.70	0.70	40

Random Forest Classification Report

<pre>In []: print(rf_cl</pre>	assification _.	_report)		
	precision	recall	f1-score	support
А	0.86	1.00	0.92	6
В	1.00	1.00	1.00	3
C	0.62	1.00	0.77	5
X	1.00	0.73	0.84	11
Υ	1.00	0.93	0.97	15
accuracy			0.90	40
macro avg	0.90	0.93	0.90	40
weighted avg	0.93	0.90	0.90	40

Decision Tree

The overall accuracy of the decision tree is 85%, which is commendable. The model is also precise when looking at drugs A, B, X, and Y but performs poorly with drug C. The f1-score and recall perform well with drugs A, B, and Y but fail to perform well with drugs C and X.

Logistic Regression

The logistic regression model had an overall accuracy score of 70% which is the lowest of the three models. Generally speaking this model did not perform well in most metrics regardless of the drug type. Interestingly though, it was most precise with drug C unlike the other two models.

Random Forest

The random forest performed the best of any model with an accuracy of 90%. In fact, this model performed consistently well across all metrics and drugs with the exception of the precision score of drug C.

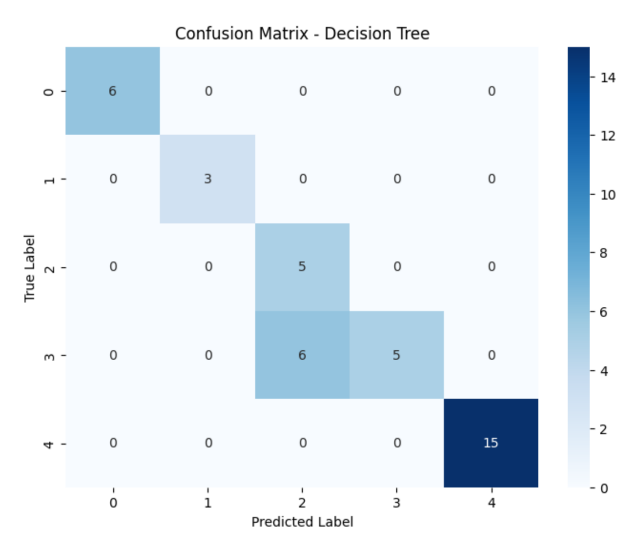
Confusion Matrix

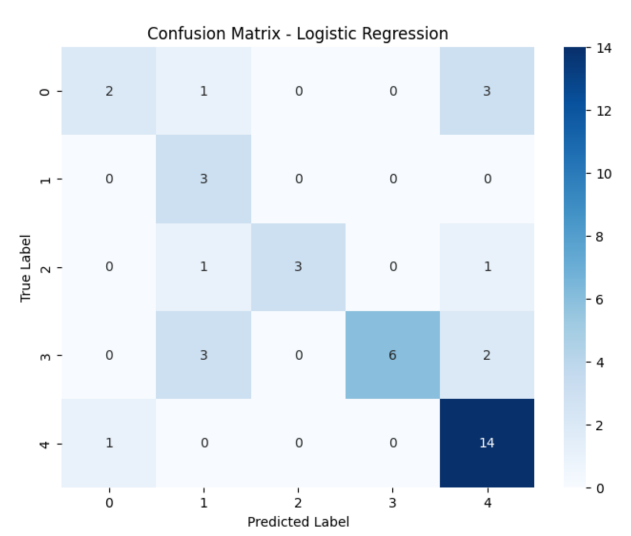
```
In []: from sklearn.metrics import confusion_matrix

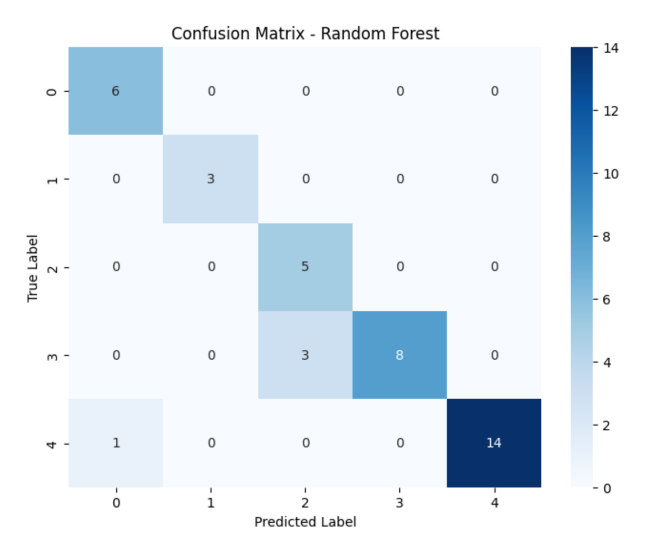
# Create a list of models and each prediciton they had
models = ["Decision Tree", "Logistic Regression", "Random Forest"]
predictions = [y_pred_dt, y_pred_lr, y_pred_rf]

# Now, go through each model using the zipping method
for model, pred in zip(models, predictions):
    cm = confusion_matrix(y_test, pred)

# Heatmap
    plt.figure(figsize = (8,6))
    sns.heatmap(cm, annot = True, fmt = 'd', cmap = "Blues")
    plt.title(f"Confusion Matrix - {model}")
    plt.xlabel("Predicted Label")
    plt.ylabel("True Label")
    plt.show()
```







The confusion matricies further illustrate that the decision tree and random forest models perform well at predicting each drug label. The confusion matrix for the logistic regression also reinforces the prediction limits of the model. Interestingly, the logistic regression model did do a good job predicting drug C unlike the other two models.

Model Improvement

This section is dedicated to trying to improve model performance

```
In [ ]: # Model improvement attempt for DTC
    # Original model (max_depth=3, min_samples_split=5, min_samples_leaf=2, max_feature

# Make the DTC. Include parameters in order to introduce some randomness. Otherwi
    # CLF introduces two new parameters with no performance improvement
    clf = DecisionTreeClassifier(max_depth=3, min_samples_split=6, min_samples_leaf=8,
        clf2 = DecisionTreeClassifier(max_depth=2, min_samples_split=7, min_samples_leaf=4,
        # Training data
    clf.fit(X_train, y_train)
    clf2.fit(X_train, y_train)
    # Predicitons on testing data
    y_pred_dt = clf.predict(X_test)
    y_pred_dt_2 = clf2.predict(X_test)
```

```
# Create a classification report
 dt_classification_report = classification_report(y_test, y_pred_dt)
 dt_classification_report_2 = classification_report(y_test, y_pred_dt_2)
 # Show the accuracy
 accuracy_dt = accuracy_score(y_test, y_pred_dt)
 print(dt_classification_report)
 print("Accuracy:", accuracy_dt)
 accuracy_dt_2 = accuracy_score(y_test, y_pred_dt_2)
 print(dt_classification_report_2)
 print("Accuracy:", accuracy_dt_2)
              precision
                           recall f1-score
                                              support
           Α
                   1.00
                             1.00
                                       1.00
                                                     6
           В
                   1.00
                             1.00
                                       1.00
                                                     3
           C
                                                    5
                   0.45
                             1.00
                                       0.62
           Χ
                   1.00
                             0.45
                                       0.62
                                                   11
           Υ
                   1.00
                             1.00
                                       1.00
                                                   15
                                       0.85
                                                   40
   accuracy
  macro avg
                   0.89
                             0.89
                                       0.85
                                                   40
weighted avg
                   0.93
                             0.85
                                       0.85
                                                   40
Accuracy: 0.85
              precision
                           recall f1-score
                                              support
                   0.67
                             1.00
           Α
                                       0.80
                                                     6
                   0.00
                             0.00
                                       0.00
                                                     3
           В
           C
                   0.31
                                                    5
                             1.00
                                       0.48
           Χ
                   0.00
                             0.00
                                       0.00
                                                   11
                   1.00
                             1.00
                                       1.00
                                                   15
                                       0.65
                                                   40
    accuracy
  macro avg
                   0.40
                             0.60
                                       0.46
                                                   40
                                       0.55
weighted avg
                   0.51
                             0.65
                                                   40
Accuracy: 0.65
/home/codespace/.local/lib/python3.10/site-packages/sklearn/metrics/_classification.
py:1344: UndefinedMetricWarning: Precision and F-score are ill-defined and being set
to 0.0 in labels with no predicted samples. Use `zero_division` parameter to control
this behavior.
 _warn_prf(average, modifier, msg_start, len(result))
/home/codespace/.local/lib/python3.10/site-packages/sklearn/metrics/_classification.
py:1344: UndefinedMetricWarning: Precision and F-score are ill-defined and being set
to 0.0 in labels with no predicted samples. Use `zero_division` parameter to control
this behavior.
 _warn_prf(average, modifier, msg_start, len(result))
/home/codespace/.local/lib/python3.10/site-packages/sklearn/metrics/_classification.
py:1344: UndefinedMetricWarning: Precision and F-score are ill-defined and being set
```

Two new parameters were introduced into the model which ultimately did not improve performance. In fact, by tuning some of these parameters it degraded the overall

_warn_prf(average, modifier, msg_start, len(result))

to 0.0 in labels with no predicted samples. Use `zero_division` parameter to control

this behavior.

performance.

```
In [ ]: #ModeL
        logisticial_regression = LogisticRegression(solver = 'sag', max_iter = 200)
        logisticial_regression2 = LogisticRegression(solver = 'liblinear', max_iter = 200,
        # Train the model
        logisticial_regression.fit(X_train, y_train)
        logisticial_regression2.fit(X_train, y_train)
        # Predict the testing data
        y_pred_lr = logisticial_regression.predict(X_test)
        y_pred_lr2 = logisticial_regression.predict(X_test)
        # Create a classification report
        lr_classification_report = classification_report(y_test, y_pred_lr)
        lr_classification_report2 = classification_report(y_test, y_pred_lr2)
        # Summarize the accuracy
        accuracy_lr = accuracy_score(y_test, y_pred_lr)
        print(lr_classification_report)
        print("Accuracy:", accuracy_lr)
        accuracy_lr2 = accuracy_score(y_test, y_pred_lr2)
        print(lr_classification_report2)
        print("Accuracy:", accuracy_lr2)
                     precision
                                  recall f1-score
                                                     support
                  Α
                          0.67
                                    0.33
                                              0.44
                                                           6
                  В
                          0.38
                                    1.00
                                              0.55
                                                           3
                  C
                          1.00
                                    0.60
                                              0.75
                                                           5
                          1.00
                                    0.55
                  Χ
                                              0.71
                                                          11
                          0.70
                                    0.93
                                              0.80
                                                          15
                                              0.70
                                                          40
           accuracy
                          0.75
                                    0.68
                                              0.65
                                                          40
         macro avg
      weighted avg
                          0.79
                                    0.70
                                              0.70
                                                          40
      Accuracy: 0.7
                     precision
                                  recall f1-score
                                                     support
                          0.67
                                    0.33
                                              0.44
                  Α
                                                           6
                  В
                          0.38
                                    1.00
                                              0.55
                                                           3
                  C
                          1.00
                                    0.60
                                              0.75
                                                           5
                  Χ
                          1.00
                                    0.55
                                              0.71
                                                          11
                  Υ
                          0.70
                                    0.93
                                              0.80
                                                          15
                                              0.70
                                                          40
          accuracy
         macro avg
                          0.75
                                    0.68
                                              0.65
                                                          40
      weighted avg
                          0.79
                                    0.70
                                              0.70
                                                          40
      Accuracy: 0.7
```

```
/home/codespace/.local/lib/python3.10/site-packages/sklearn/linear_model/_sag.py:35
0: ConvergenceWarning: The max_iter was reached which means the coef_ did not conver
ge
    warnings.warn(
```

These changes to the parameters also did not improve performance of the model.

```
In [ ]: #Create instance of Random Forest. Make sure to include parameters to introduce ra
        rfc = RandomForestClassifier(n_estimators = 10, max_depth = 12, min_samples_split =
        rfc2 = RandomForestClassifier(n_estimators = 1000, max_depth = 1, min_samples_split
        # Fit model to the training data
        rfc.fit(X_train, y_train)
        rfc2.fit(X_train, y_train)
        # Make predictions
        y_pred_rf = rfc.predict(X_test)
        y_pred_rf2 = rfc.predict(X_test)
        # Create a classification report
        rf_classification_report = classification_report(y_test, y_pred_rf)
        rf_classification_report2 = classification_report(y_test, y_pred_rf2)
        # Summarize the accuracy
        accuracy_rf = accuracy_score(y_test, y_pred_rf)
        print(rf_classification_report)
        print("Accuracy:", accuracy_rf)
        accuracy_rf2 = accuracy_score(y_test, y_pred_rf2)
        print(rf_classification_report)
        print("Accuracy:", accuracy_rf2)
```

	precision	recall	f1-score	support
А	0.86	1.00	0.92	6
В	1.00	1.00	1.00	3
С	0.62	1.00	0.77	5
X	1.00	0.73	0.84	11
Υ	1.00	0.93	0.97	15
accuracy			0.90	40
macro avg	0.90	0.93	0.90	40
weighted avg	0.93	0.90	0.90	40
Accuracy: 0.9				
Accuracy: 0.9	precision	recall	f1-score	support
Accuracy: 0.9		recall	f1-score	support 6
•	precision			
А	precision 0.86	1.00	0.92	6
А В	precision 0.86 1.00	1.00 1.00	0.92 1.00	6
A B C	0.86 1.00 0.62	1.00 1.00 1.00	0.92 1.00 0.77	6 3 5
A B C X	0.86 1.00 0.62 1.00	1.00 1.00 1.00 0.73	0.92 1.00 0.77 0.84	6 3 5 11
A B C X	0.86 1.00 0.62 1.00	1.00 1.00 1.00 0.73	0.92 1.00 0.77 0.84	6 3 5 11
A B C X Y	0.86 1.00 0.62 1.00	1.00 1.00 1.00 0.73	0.92 1.00 0.77 0.84 0.97	6 3 5 11 15

Accuracy: 0.9

Similarly, changing the parameters did not improve the performance of this model.

Discussion and Conclusions

The biggest conclusion in comparing these models is that the random forest performed the best by all metrics and should be the recommneded model if trying to use this in practice. The decision tree also performed well but despite attempts at improving the model there does not seem to be room for improvement. The logistic regression model performed the worst out of all three models and could not be improved.

These machine learning models were all relatively easy to set up and work with. There were not too many issues when it came to adjusting the models in order to perform well.

One of the most noteable things about these models was despite turning the parameters, there was not any improvement in any model. This is probably due to the fact that there was feature engineering done prior to setting the testing and training data.

Based on the EDA done earlier, the most likely conclusion for the logistic regression model performing poorly is that there was not a strong sense of linear relationships in the dataset. Logistic regression models are heavily reliant on having linear relationships in the data in order to help predict. However, it should be noted that this model did the best at identifying drug C.

On the other hand, decision trees and random forests both handle a lack of linear relationships well. Random forests in particular are good at capturing more complex relationships in order to make accurate predictions.

The best way to improve these models overall may be to make the dataset bigger. Larger datasets always bode well for making models more robust and have a better representation of how well they may perform. With larger datasets it is possible more linear relationships will be seen which would help the logistic regression model. Both the decision trees and random forests are already capable of handling complex data so this would just make the models more robust.

Citations

Thanks to Pratham Tripathi who supplied this dataset on Kaggle: Pratham Tripathi. (June 2020). Drug Classification, version 1 https://www.kaggle.com/datasets/prathamtripathi/drug-classification

For information about Na/K relationships in the human body:

Mirmiran, P., Gaeini, Z., Bahadoran, Z., Ghasemi, A., Norouzirad, R., Tohidi, M., & Azizi, F. (2021, January 6). Urinary sodium-to-potassium ratio: A simple and useful indicator of diet quality in population-based studies - European Journal of Medical Research. BioMed Central. https://eurjmedres.biomedcentral.com/articles/10.1186/s40001-020-00476-5#citeas