AML Lab Sessional 1 Jupyter Notebook

1. VS Code is a MUST

1. You should code in VS Code. Google Colab not allowed

2. Pre-requisites for lab in VS Code

 Install PDF export support https://saturncloud.io/blog/how-to-export-jupyternotebook-by-vscode-in-pdf-format/

3. At the end, export your notebook as html and ipynb

- 1. Open VS Code command palette Shift + Ctrl + P
- 2. Type "Export Jupyter Notebook" in the search bar and select "Export Jupyter Notebook to html"
- 3. Upload your html AND ipynb here: https://tinyurl.com/y9ptbej2 (In the prompt, put your name correctly else your submission will be rejected)

4. Lab Sessional Summary

- 1. A code template is given to you in lab sesional in this jupyter notebook
- 2. The template will follow a linear sequence of TODOs appropriately labelled with question marks
- 3. You will have to fill the TODO question marks to compile those notebook cells and proceed to next cells
- 4. You can think of the linear sequence of TODOs as a guided thought process.
- 5. Sessional is open book. Google, github, browse product documentation ChatGPT Do anything you want, except copying others. No discussing among yourselves (Sending questions to your seniors and seeking answers is prohibited). If you are caught carrying out these illegal activities, you will be reported for immediate action.
- 6. You CANNOT replace the TODO code template with some other code copied from stack overflow, ChatGPT etc. All your browsing and search should give you insights into finally how you can fit that into the framework I provide for the thought process of solving the problem. You will have to mandatorily fill the question marks and proceed with the lab problem.

```
In []: import numpy as np
   import pandas as pd
   import sklearn as sk

import matplotlib.pyplot as plt
   import seaborn as sns
%matplotlib inline

import warnings
warnings.filterwarnings("ignore")
```

```
In []: sk.__version__
Out[]: '1.3.0'
```

Problem Definition

You are provided with diabetes dataset. Your task is:

- 1. Part 1: To perform EDA and prediction without pipeline
- 2. Part 2: To move EDA and kNN prediction into a pipeline
- 3. Part 3: To combine the pipeline and gridsearch to get best K and weights hyperparameters
- 4. Part 4: Save the model as json file, load & use it to do prediction. This part is optional and will be used to accumulate bonus points as a buffer for the semester lab exam if there is any shortfall

Part 1 - Perform EDA and prediction without pipeline

```
In [ ]: # TODO: 1 Load the csv file
         df = pd.read_csv("d2.csv")
         df.head()
                                BloodPressure
                                              SkinThickness Insulin
                                                                   BMI DiabetesPedigreeFunction
Out[]:
            Pregnancies
                        Glucose
         0
                     2
                            138
                                                        35
                                                                   33.6
                                                                                          0.127
                                          62
                                                                0
                     0
                            84
                                          82
                                                        31
                                                               125
                                                                   38.2
                                                                                          0.233
                     0
         2
                                           0
                                                         0
                                                                0 44.2
                                                                                          0.630
                            145
         3
                                                                   42.3
                                                                                          0.365
                            135
                                          68
                                                              250
         4
                     1
                           139
                                          62
                                                        41
                                                              480 40.7
                                                                                          0.536
         # TODO: 2 Display information about dataframe
         df.info(verbose=True, show_counts=True)
         <class 'pandas.core.frame.DataFrame'>
         RangeIndex: 2000 entries, 0 to 1999
         Data columns (total 9 columns):
              Column
                                          Non-Null Count Dtype
         ---
             Pregnancies
          0
                                          2000 non-null
                                                           int64
                                          2000 non-null
          1
              Glucose
                                                           int64
              BloodPressure
                                          2000 non-null
                                                           int64
              SkinThickness
                                          2000 non-null
                                                           int64
          4
              Insulin
                                          2000 non-null
                                                           int64
          5
                                          2000 non-null
                                                           float64
          6
              DiabetesPedigreeFunction 2000 non-null
                                                           float64
          7
                                                           int64
                                          2000 non-null
              Age
```

2000 non-null

int64

Outcome

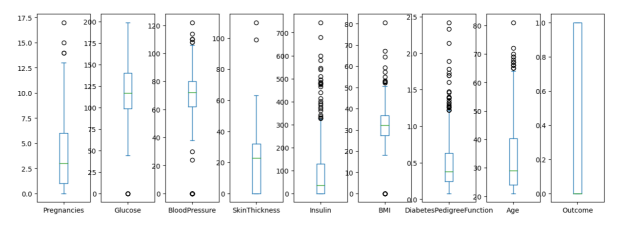
memory usage: 140.8 KB

dtypes: float64(2), int64(7)

```
# TODO: 3 Check if there are nulls and display their total for each feature
In [ ]:
        df.isna().sum()
        Pregnancies
                                    0
Out[]:
        Glucose
                                    0
        BloodPressure
                                    0
        SkinThickness
                                    0
        Insulin
                                    0
        BMT
                                    0
        DiabetesPedigreeFunction
                                    0
        Age
                                    0
        Outcome
                                    0
        dtype: int64
In [ ]: # TODO: 4 Check for duplicates
        dupsSeries = df.duplicated() # Should return a pandas Series with True False for el
        # Print the number of duplicates
        print(f"Number of duplicates = {dupsSeries.sum()}")
        Number of duplicates = 1256
In [ ]: # TODO: 5 Drop duplicates inplace
        df.drop_duplicates(inplace=True)
In [ ]: # TODO: 6 Display how many rows exist in dataframe after dropping duplicates
        df.info(verbose=True, show counts=True)
        <class 'pandas.core.frame.DataFrame'>
        Int64Index: 744 entries, 0 to 1568
        Data columns (total 9 columns):
         #
            Column
                                       Non-Null Count Dtype
            ____
                                       _____
         0
            Pregnancies
                                       744 non-null
                                                      int64
            Glucose
                                      744 non-null int64
         1
         2 BloodPressure
                                      744 non-null int64
         3 SkinThickness
                                      744 non-null int64
                                       744 non-null int64
         4
            Insulin
         5
             BMI
                                       744 non-null float64
             DiabetesPedigreeFunction 744 non-null float64
         6
                                       744 non-null int64
         7
             Age
             Outcome
                                       744 non-null
                                                      int64
        dtypes: float64(2), int64(7)
        memory usage: 58.1 KB
In [ ]: # TODO: 7 Identify candidate features for encoding by seeing how many distinct value
        df.nunique()
        Pregnancies
                                     17
Out[ ]:
        Glucose
                                    136
        BloodPressure
                                     47
        SkinThickness
                                     53
        Insulin
                                    182
        BMI
                                    247
        DiabetesPedigreeFunction
                                    505
                                     52
        Age
        Outcome
                                      2
        dtype: int64
In [ ]: # TODO: 8
        # Display all possible values that the column named "Pregnancies" has in this dataj
        df["Pregnancies"].unique()
        # Based on different possible values of Pregnancies, would you choose this feature
```

```
# State your reason as 1 sentence comment here
         # __it may not be necessary to encode it, as the values are already in a numerical
                               4,
                                    8, 3, 6, 5, 7, 10, 12, 9, 11, 13, 15, 17, 14],
         array([ 2, 0, 1,
Out[ ]:
                dtype=int64)
In [ ]: # TODO: 9 Find if this is an dataset is imbalanced wrt target variable classes
         zeroClassCount = df[df["Outcome"] == 0]["Outcome"].count()
         zeroClassCount/df["Outcome"].count()
         0.6599462365591398
Out[ ]:
In [ ]: # TODO: 10 Run this cell to display correlation matrix heat map
         # Would you consider dropping any feature based on their correlation?
         # Type your answers here: ____No_
         plt.figure(figsize=(12,6))
         sns.heatmap(df.corr(),cmap='plasma',fmt='.2g',annot=True,mask=np.triu(df.corr(),+1)
         plt.show()
                                                                                                  - 1.0
                  Pregnancies -
                                    1
                    Glucose
                                                                                                  - 0.8
                                            1
                 BloodPressure
                            0.14
                                    0.14
                                                                                                  - 0.6
                 SkinThickness
                            -0.063
                                   0.071
                                                   1
                     Insulin
                            -0.056
                                           0.092
                                                                                                  0.4
                            0.026
                       BMI
                                                                         1
         DiabetesPedigreeFunction
                            -0.025
                                           0.051
                                                  0.18
                                                                                                  0.2
                                                                 0.053
                                                         -0.064
                                                                        0.033
                                                                                1
                       Age
                                                                                                  0.0
                                           0.079
                    Outcome
                                                          0.11
                                                                                Age
                                    Glucose
                                            BloodPressure
                                                                 M
                                                                         DiabetesPedigreeFunction
In [ ]: # TODO: 11 Run this cell to see box plots.
         # Visually examine the box plot and answer this question
         # How do you identify the 1.5 IQR boundary?
         # Which two features have most outliers beyond 1.5 IQR?
         # (Increase the figure size if you want to see larger image)
         # Put your answer for both questions here
         # _IQR = Q3 (75th percentile) - Q1 (25th percentile)_
         # ____Insulin, DiabetesPedigreeFunction__
         df.plot(kind="box",subplots=True,figsize=(15,5),title="Data with Outliers");
```

Data with Outliers



```
In [ ]:
        # TODO: 12 Did you see any nulls in the dataset?
                          No
        # But look at how many features have zeros in them
        # Display zeroes in each column as percentage
        for col in df.columns:
            count = (df[col] == 0).sum()
            percentage = (count * 100)/df.shape[0]
            print(f'Count of zeros in Column {col} : {count}, percentage 0s: {percentage:.2
        Count of zeros in Column Pregnancies : 112, percentage 0s: 15.05%
        Count of zeros in Column Glucose : 5, percentage 0s: 0.67%
        Count of zeros in Column BloodPressure : 34, percentage 0s: 4.57%
        Count of zeros in Column SkinThickness: 215, percentage 0s: 28.90%
        Count of zeros in Column Insulin : 359, percentage 0s: 48.25%
        Count of zeros in Column BMI : 10, percentage 0s: 1.34%
        Count of zeros in Column DiabetesPedigreeFunction: 0, percentage 0s: 0.00%
        Count of zeros in Column Age : 0, percentage 0s: 0.00%
        Count of zeros in Column Outcome : 491, percentage 0s: 65.99%
```

- 1. After executing the above cell, you found many cells contain 0.
- 2. Some cells should never contain 0. For e.g. Glucose
- 3. Identify the cells that should never contain 0 and replace those 0 with Nan

```
In [ ]: # TODO: 13 Replace 0 with Nan for features that should never contain 0
         # Choose 5 columns you want to replace 0 with Nan
         df[["Glucose", "BloodPressure", "SkinThickness", "Insulin", "BMI"]] = \
             df[["Glucose", "BloodPressure", "SkinThickness", "Insulin", "BMI"]].replace(0,
In [ ]: # TODO: 14 Now do null check again to ensure the right featues have Nans. Otherwis
        df.isna().sum()
        Pregnancies
                                       0
Out[]:
        Glucose
                                       5
        BloodPressure
                                      34
        SkinThickness
                                     215
        Insulin
                                     359
        BMI
                                      10
        DiabetesPedigreeFunction
                                       0
        Age
                                       0
                                       0
        Outcome
        dtype: int64
In [ ]: y = df.pop("Outcome") #Setup target variable
         type(y)
```

```
Out[]: pandas.core.series.Series
```

```
In [ ]: # TODO: 15
        # Answer these questions with short one liner right in this cell as a comment
         # 1. What does the pop operation do?
         # pop operation will remove and returns the remaining elements in a list
         # 2. Is pop() idempotent or non-idempotent operation?
         # It is non-idempotent operation
         # 3. How do you check if an operation is idempotent/non-idempotent?
             An operation is idempotent if applying it multiple times has the same effect of
             An operation is non-idempotent if applying it multiple times can have a differ
In [ ]: X = df # Setup independent variables
In [ ]: from sklearn.model_selection import train_test_split
         # Fill this out
        X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.25, random_s1
In [ ]: # TODO: 16 Answer these questions
         # 1. Did you use stratify?
         # No, stratify was not used in train_test_split
         # 2. If yes, on which column and why? If not, why not?
         #
In [ ]: df.isna().sum()
        Pregnancies
                                       0
Out[]:
        Glucose
                                      5
        BloodPressure
                                      34
        SkinThickness
                                     215
        Insulin
                                     359
        BMI
                                      10
        {\tt DiabetesPedigreeFunction}
                                      0
                                       0
        dtype: int64
In [ ]: # TODO: 17 Display all records that have at least one Nan column value
         # If you cannot get this, you can leave this cell execution and proceed
         # Subsequent cells do not depend on this. If you cannot solve this, you can proceed
        X_train.loc[df.isna().any(axis=1), :]
```

Out[

]:		Pregnancies	Glucose	BloodPressure	SkinThickness	Insulin	ВМІ	DiabetesPedigreeFunction
	691	13	158.0	114.0	NaN	NaN	42.3	0.257
	294	0	161.0	50.0	NaN	NaN	21.9	0.254
	77	5	95.0	72.0	33.0	NaN	37.7	0.370
	352	3	61.0	82.0	28.0	NaN	34.4	0.243
	235	4	171.0	72.0	NaN	NaN	43.6	0.479
	•••							
	756	7	137.0	90.0	41.0	NaN	32.0	0.391
	9	2	89.0	90.0	30.0	NaN	33.5	0.292
	192	7	159.0	66.0	NaN	NaN	30.4	0.383
	592	3	132.0	80.0	NaN	NaN	34.4	0.402
	717	10	94.0	72.0	18.0	NaN	23.1	0.595

269 rows × 8 columns



Deciding on Imputation

- 1. Now you should decide which imputation to use whether SimpleImputer or IterativeImputer
- 2. Instead of following MCAR/MAR path to decide on SimpleImputer or IterativeImputer, you will take a simple alternative
- 3. Use IterativeImputer if a large number of rows have Nan for a column
- 4. Use SimpleImputer if the number of Nans is very small

```
print(f"X_train.shape={X_train.shape}")
         X_train.isna().sum()
        X_train.shape=(558, 8)
        Pregnancies
                                       0
Out[ ]:
        Glucose
                                       4
        BloodPressure
                                      23
        SkinThickness
                                     151
        Insulin
                                     267
        BMI
                                       7
        DiabetesPedigreeFunction
                                       0
        Age
                                       0
        dtype: int64
In [ ]: # TODO 18 Based on the results of previous cell which columns will you apply Simple
         # Select only those columns here
         X_train_mean_impute = X_train.loc[:,["Glucose", "BloodPressure", "BMI"]]
         X train mean impute
```

Out

[]:		Glucose	BloodPressure	ВМІ
	691	158.0	114.0	42.3
	599	109.0	38.0	23.1
	294	161.0	50.0	21.9
	77	95.0	72.0	37.7
	352	61.0	82.0	34.4
	•••			
	740	120.0	80.0	42.3
	192	159.0	66.0	30.4
	662	167.0	106.0	37.6
	592	132.0	80.0	34.4
	717	94.0	72.0	23.1

558 rows × 3 columns

```
In [ ]: # TODO: 19 Apply SimpleImputer to appropriate columns only
        from sklearn.impute import SimpleImputer
        mean_imputer = SimpleImputer(strategy="mean")
        #Fill code above to create SimpleImputer
        X_train_mean_imputed = mean_imputer.fit_transform(X_train_mean_impute)
        # Display first few
        X_train_mean_imputed[0:5,:]
        array([[158., 114., 42.3],
Out[ ]:
               [109., 38., 23.1],
               [161., 50., 21.9],
               [ 95. , 72. , 37.7],
               [ 61. , 82. , 34.4]])
In [ ]: # TODO: 20 Which columns will you apply IterativeImputer ?
        # Name those columns and provide 2 line short reason
        # ___SkinThickness, Insulin___
        X_train.loc[:,["SkinThickness", "Insulin"]]
```

Out[]:		SkinThickness	Insulin
	691	NaN	NaN
	599	18.0	120.0
	294	NaN	NaN
	77	33.0	NaN
	352	28.0	NaN
	•••		
	740	37.0	150.0
	192	NaN	NaN
	662	46.0	231.0
	592	NaN	NaN
	717	18.0	NaN

558 rows × 2 columns

Iterative Imputer uses all non null features of the dataset to impute. To apply Iteratative Imputer on the two columns identified above, you should have all other features combined into the dataframe or numpy nmatrix first.

You should do the following for that

- 1. Remove the columns of the dataframe that were subjected to mean imputation
- 2. Convert the remaining dataframe into numpy matrix
- 3. Combine the above numpy matrix with mean imputed columns earlier. The "combining" two matrices is achieved by concatenating those two numpy matrices by using an appropriate numpy stacking function
- 4. The stacked numpy matrix is then used for iterative imputation

```
In []: # TODO: 21 Remove the columns of X_train dataframe that were already subjected to n
        # Hold on to the rest as a Numpy matrix
        X_train_set_aside = X_train.drop(["Glucose", "BloodPressure", "BMI"], axis=1).to_n(
        X_train_set_aside
        array([[1.30e+01, nan,
                                        nan, 2.57e-01, 4.40e+01],
Out[ ]:
               [1.00e+00, 1.80e+01, 1.20e+02, 4.07e-01, 2.60e+01],
               [0.00e+00, nan, nan, 2.54e-01, 6.50e+01],
               [8.00e+00, 4.60e+01, 2.31e+02, 1.65e-01, 4.30e+01],
               [3.00e+00, nan, nan, 4.02e-01, 4.40e+01],
               [1.00e+01, 1.80e+01,
                                        nan, 5.95e-01, 5.60e+01]])
In [ ]: # TODO: 22 Concatenate the two numpy matrices by using an appropriate numpy stacking
        # Identify which are those two numpy matrices first and use them
        X train = np.hstack((X train mean imputed, X train set aside)) # this numpy matrix
        X train
```

```
Out[]: array([[1.58e+02, 1.14e+02, 4.23e+01, ...,
                                                         nan, 2.57e-01, 4.40e+01],
                [1.09e+02, 3.80e+01, 2.31e+01, ..., 1.20e+02, 4.07e-01, 2.60e+01],
               [1.61e+02, 5.00e+01, 2.19e+01, ..., nan, 2.54e-01, 6.50e+01],
                [1.67e+02, 1.06e+02, 3.76e+01, ..., 2.31e+02, 1.65e-01, 4.30e+01],
                [1.32e+02, 8.00e+01, 3.44e+01, ..., nan, 4.02e-01, 4.40e+01],
                [9.40e+01, 7.20e+01, 2.31e+01, ...,
                                                       nan, 5.95e-01, 5.60e+01]])
In [ ]: # TODO: 23 Perform Iterative Imputation on two columns with Nan data
         # At this point only two columns will have Nan - SkinThickness and Insulin
         # Those 2 will be imputed using all other columns
         # But to make life easy, this is equivalent to performing Iterative Imputation usiv
         from sklearn.experimental import enable iterative imputer # Enable the IterativeIn
         from sklearn.impute import IterativeImputer
         #Add IterativeImputer here and fit and transform appropriate columns
         iterative imputer = IterativeImputer()
         X_train_iter_imputed = iterative_imputer.fit_transform(X_train)
         X_train_iter_imputed # this variable should hold your imputer values
        array([[1.58000000e+02, 1.14000000e+02, 4.23000000e+01, ...,
Out[ ]:
                2.37599549e+02, 2.57000000e-01, 4.40000000e+01],
                [1.09000000e+02, 3.80000000e+01, 2.31000000e+01, ...,
                1.20000000e+02, 4.07000000e-01, 2.60000000e+01],
                [1.61000000e+02, 5.00000000e+01, 2.19000000e+01, ...,
                2.26669815e+02, 2.54000000e-01, 6.50000000e+01],
                [1.67000000e+02, 1.06000000e+02, 3.76000000e+01, ...,
                2.31000000e+02, 1.65000000e-01, 4.30000000e+01],
                [1.32000000e+02, 8.00000000e+01, 3.44000000e+01, ...,
                1.76808163e+02, 4.02000000e-01, 4.40000000e+01],
                [9.40000000e+01, 7.20000000e+01, 2.31000000e+01, ...,
                8.56658773e+01, 5.95000000e-01, 5.60000000e+01]])
In [ ]: # TODO: 24 Fill the question marks to do z transform
         from sklearn.preprocessing import StandardScaler
         scaler = StandardScaler()
         X_train_scaled = scaler.fit_transform(X_train_iter_imputed)
         X train scaled
        array([[ 1.18954035, 3.36441269, 1.31215916, ..., 0.85174053,
                 -0.63290132, 0.95454214],
                [-0.42357097, -2.78819233, -1.37708204, ..., -0.35994306,
                -0.17876717, -0.62389402],
                [1.28830226, -1.81672838, -1.54515962, ..., 0.73912633,
                -0.641984 , 2.79605101],
                [ 1.4858261 , 2.71677005, 0.65385532, ..., 0.78374227,
                -0.91143693, 0.86685124],
[ 0.33360373, 0.6119315 , 0.20564845, ..., 0.22537822,
                 -0.19390497, 0.95454214],
                [-0.91738055, -0.03571114, -1.37708204, ..., -0.71370372,
                  0.3904143 , 2.00683292]])
In [ ]: # TODO: 25 Train the KNN model in this cell
         from sklearn.neighbors import KNeighborsClassifier
         model = KNeighborsClassifier(n_neighbors=5)
         model.fit(X_train_scaled, y_train)
```

```
▼ KNeighborsClassifier
Out[ ]:
        KNeighborsClassifier()
In [ ]: # TODO: 26 Perform all necessary transformation on X_test in this cell to prepare
         \# look at all the transformation logic that was performed on X_train and repeat it
         # Put your code here. No template is provided for this section
         X_test_mean_impute = X_test.loc[:,["Glucose", "BloodPressure", "BMI"]]
         X_test_mean_imputed = mean_imputer.fit_transform(X_test_mean_impute)
         X_test_set_aside = X_test.drop(["Glucose", "BloodPressure", "BMI"], axis=1).to_num;
         X_test = np.hstack((X_test_mean_imputed, X_test_set_aside))
         X test iter imputed = iterative imputer.fit transform(X test)
         X_test_scaled = scaler.fit_transform(X_test_iter_imputed)
In [ ]: # TODO: 27
         # Write code here to predict using the model and calculate & display the accuracy of
         # You should get an accuracy of around 75%
         from sklearn.metrics import accuracy_score
```

Out[]: 0.7688172043010753

Do prediction using knn here

y_pred = model.predict(X_test_scaled)

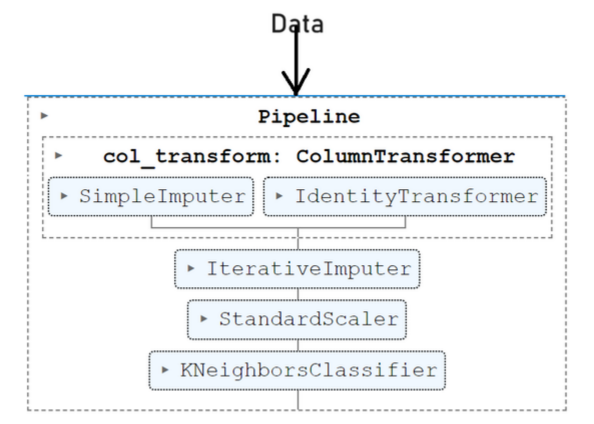
accuracy_score(y_pred=y_pred, y_true=y_test)

Part 2: Data processing and prediction using a pipeline

```
In []: # TODO: 28 Repeat activities that need to be done outside pipeline
    # Reading csv, dropping duplicates & other steps that you might have performed - bu
    df = pd.read_csv("d2.csv")
    df.drop_duplicates(inplace=True)
    #Add other code for dropping duplicates etc

In []: # TODO: 29 Split the data into X and y. Then do train test split here
    y = df.pop("Outcome")
    X = df
    X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.25, random_st
```

Design a pipeline that looks like following:



- 1. SimpleImputer performs mean imputation only on the relevant columns
- 2. IdentityTransformer (whose code is given below) is a custom sklearn transformer that performs no operation on the input data. Input data is passed through as is
- 3. The two are wrapped within a ColumnTransformer
- 4. This is followed by sequential execution of IterativeImputer, StandaScaler and the KNN based clasifier

```
In []: # This is a No-op transformer. Use this as is
from sklearn.base import BaseEstimator, TransformerMixin

class IdentityTransformer(BaseEstimator, TransformerMixin):
    def __init__(self):
        pass

def fit(self, X, y=None):
        return self

def transform(self, X, y=None):
        return X*1
```

Look at the columns of the dataframe and decide which ones need to go to SimpleImputer and which ones should go to IdentityTransformer

```
Out[]: ['Glucose', 'BloodPressure', 'SkinThickness', 'Insulin', 'BMI']
In [ ]: mean_impute_col_mask = df.columns.isin(mean_impute_cols)
         mean_impute_col_mask
        array([False, True, True, True, True, False, False])
Out[ ]:
In [ ]: # TODO: 31 Use the mean_impute_col_mask to select remaining columns from df.columns
         other_cols = df.columns[~mean_impute_col_mask].to_list()
         other cols
Out[]: ['Pregnancies', 'DiabetesPedigreeFunction', 'Age']
In [ ]: # TODO 32 Build the pipeline using the diagram shown earlier and the details provide
         from sklearn.pipeline import Pipeline
         from sklearn.compose import ColumnTransformer
         from sklearn.preprocessing import StandardScaler
         from sklearn.impute import SimpleImputer, IterativeImputer
         from sklearn.neighbors import KNeighborsClassifier
         from sklearn.base import BaseEstimator, TransformerMixin
         # Create transformers
         mean_imputer = SimpleImputer(strategy='mean', add_indicator=False)
         identity_transformer = IdentityTransformer()
         iterative_imputer = IterativeImputer()
         scaler = StandardScaler()
         # Create the ColumnTransformer
         preprocessor = ColumnTransformer(
            transformers=[
                 ('mean_imputer', mean_imputer, mean_impute_cols),
                 ('identity_transformer', identity_transformer, other_cols),
            ],
         )
         # Create the final pipeline
         pipeline = Pipeline([
             ('col_transform', preprocessor),
             ('iterative_imputer', iterative_imputer),
             ('scaler', scaler),
             ('model', KNeighborsClassifier(n_neighbors=5))
         1)
In [ ]: # TODO: 33 Run this cell to verify if indeed your pipeline looks like what was expe
         from sklearn import set config
         set config(display='diagram')
         pipeline # Put your pipeline variable name here to display pipeline as diagram
```

```
Pipeline
Out[ ]:
            col_transform: ColumnTransformer
          ▶ mean_imputer → identity_transformer
           ▶ SimpleImputer
                            ▶ IdentityTransformer
                     ▶ IterativeImputer
                      ▶ StandardScaler
                   KNeighborsClassifier
In [ ]: # TODO: 34 Train pipeline with data
        pipeline.fit(X_train, y_train)
                          Pipeline
Out[]:
          col_transform: ColumnTransformer
          ▶ mean_imputer → identity_transformer
           ▶ SimpleImputer
                            ▶ IdentityTransformer
                     ▶ IterativeImputer
                      StandardScaler
                   ▶ KNeighborsClassifier
In [ ]: # TODO: 35 Do predictions with pipeline and calculate the accuracy score
        # Predict using the trained pipeline
        y_pred = pipeline.predict(X_test)
        # Calculate the accuracy score
        accuracy = accuracy_score(y_true=y_test, y_pred=y_pred)
        accuracy
        0.7903225806451613
```

Part 3: Combine pipeline & gridsearch for hyperparam tuning

```
In [ ]: # TODO: 36. Use GridSearchCV with KFold = 3 for CV and train a model using the pipe
        from sklearn.model_selection import GridSearchCV, KFold
        # Create KFold here
        kf = KFold(n_splits=3, shuffle=True, random_state=0)
        # Define hyperparameters
        grid_params = {'model__n_neighbors': [2, 3, 4, 5, 6],
                        'model__weights': ['uniform', 'distance'],
                        'model__metric': ['euclidean', 'manhattan']}
```

Out[]:

```
#Create Grid SearchCV object here using KFold object and gridparams
        grid = GridSearchCV(estimator=pipeline, param_grid=grid_params, cv=kf, scoring='acc
        #Do hyperparameter tuning here with GridSearchCV object
        fitted_model = grid.fit(X_train, y_train)
In [ ]: # TODO: 37 Display the results for best score, best estimator and best params select
        # fitted_model is the result of hyperpaarameter tuning in the previous cell
        print(fitted_model.best_score_)
        print(fitted_model.best_estimator_)
        print(fitted_model.best_params_)
        0.7275985663082437
        Pipeline(steps=[('col_transform',
                          ColumnTransformer(transformers=[('mean_imputer',
                                                           SimpleImputer(),
                                                           ['Glucose', 'BloodPressure',
                                                             'SkinThickness', 'Insulin',
                                                            'BMI']),
                                                          ('identity_transformer',
                                                           IdentityTransformer(),
                                                           ['Pregnancies',
                                                             'DiabetesPedigreeFunction',
                                                            'Age'])])),
                         ('iterative_imputer', IterativeImputer()),
                         ('scaler', StandardScaler()),
                         ('model',
                          KNeighborsClassifier(metric='manhattan', n_neighbors=6,
                                               weights='distance'))])
        {'model__metric': 'manhattan', 'model__n_neighbors': 6, 'model__weights': 'distanc
        e'}
In [ ]: # TODO: 38 Do predictions with gridsearch and calculate the accuracy score
        y_pred_grid = fitted_model.predict(X_test)
        # Calculate the accuracy score
        accuracy_grid = accuracy_score(y_true=y_test, y_pred=y_pred_grid)
        accuracy grid
        0.7526881720430108
```

Confusion matrix calculation

Out[]:

The next 3 cells are meant for providing hints to calculating the confusion matrix

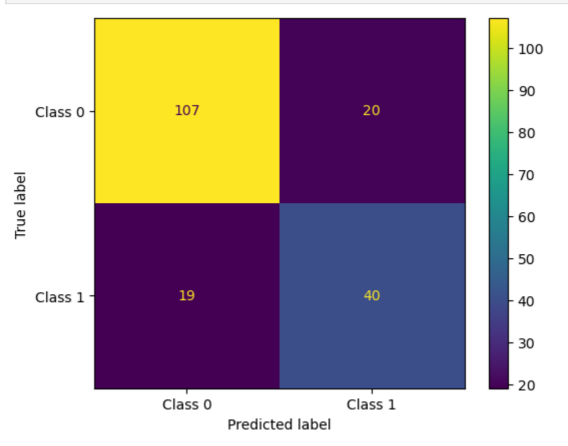
```
In [ ]: # TODO: 39 Put your pipeline name in place of question mark
         pipeline.named steps
        {'col_transform': ColumnTransformer(transformers=[('mean_imputer', SimpleImputer
Out[ ]:
         (),
                                           ['Glucose', 'BloodPressure', 'SkinThickness',
                                            'Insulin', 'BMI']),
                                          ('identity_transformer', IdentityTransformer(),
                                           ['Pregnancies', 'DiabetesPedigreeFunction',
                                             'Age'])]),
          'iterative_imputer': IterativeImputer(),
          'scaler': StandardScaler(),
          'model': KNeighborsClassifier()}
In [ ]: # TODO: 40 Put your pipeline name in place of first question mark
         # and appropriate value for second question mark to display the model
         pipeline.named_steps['model']
```

Out[]: v KNeighborsClassifier
KNeighborsClassifier()

In []: # TODO: 41 Put your pipeline name in place of first question mark
and appropriate value for second question mark to display the model
vars(pipeline.named_steps['model'])

```
Out[]: {'n_neighbors': 5,
          'radius': None,
         'algorithm': 'auto',
          'leaf_size': 30,
          'metric': 'minkowski',
          'metric params': None,
          'p': 2,
          'n_jobs': None,
          'weights': 'uniform',
          'n_features_in_': 8,
          'outputs_2d_': False,
          'classes_': array([0, 1], dtype=int64),
          '_y': array([1, 0, 0, 0, 0, 1, 0, 1, 0, 0, 1, 0, 0, 1, 0, 1, 0, 1, 0, 1, 0, 1,
                1, 1, 0, 0, 1, 1, 0, 1, 0, 0, 1, 1, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0,
                0, 1, 0, 0, 1, 1, 0, 0, 1, 0, 0, 1, 0, 1, 0, 0, 1, 1, 1, 0, 1, 0,
                1, 1, 0, 0, 1, 0, 0, 0, 1, 1, 0, 0, 0, 0, 1, 0, 0, 0, 0, 0, 1, 1,
                0, 1, 1, 0, 0, 0, 0, 1, 1, 0, 0, 0, 0, 0, 0, 1, 1, 0, 1, 0, 1,
                0, 0, 1, 0, 0, 0, 0, 1, 0, 0, 0, 1, 1, 0, 0, 0, 0, 0, 0, 0, 0,
                0, 1, 0, 0, 1, 0, 0, 0, 0, 1, 0, 0, 0, 1, 0, 0, 0, 0, 0, 0,
                1, 0, 0, 0, 1, 0, 1, 0, 1, 1, 0, 1, 1, 0, 0, 0, 0, 0, 1, 0, 1, 1,
                0, 1, 0, 0, 0, 1, 0, 1, 0, 1, 1, 1, 1, 0, 1, 0, 0, 1, 0, 0, 0,
                1, 1, 0, 1, 0, 0, 0, 1, 0, 0, 1, 0, 1, 1, 0, 0, 0, 0, 0, 0, 0, 0,
                0, 0, 0, 0, 0, 0, 1, 0, 0, 0, 1, 1, 0, 0, 0, 0, 1, 0, 0, 0, 0,
                0, 0, 0, 1, 0, 0, 0, 1, 1, 1, 0, 1, 0, 0, 1, 0, 0, 0, 0, 1, 0, 0,
                0, 0, 1, 0, 0, 0, 1, 0, 0, 1, 0, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0,
                0, 0, 0, 0, 0, 1, 0, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 0, 1, 1, 1, 0,
                1, 0, 0, 1, 1, 0, 0, 1, 1, 0, 1, 0, 0, 1, 0, 1, 1, 0, 1, 0, 0, 0,
                0, 1, 1, 0, 0, 0, 0, 0, 0, 1, 1, 1, 0, 0, 0, 1, 1, 0, 0, 1, 0,
                1, 1, 0, 0, 1, 0, 1, 1, 1, 1, 0, 0, 0, 1, 0, 0, 0, 0, 0, 0, 1, 0,
                0, 0, 1, 0, 0, 1, 0, 0, 1, 0, 1, 1, 1, 1, 0, 1, 0, 0, 0, 1, 0, 0,
                1, 0, 1, 0, 1, 0, 1, 0, 0, 0, 1, 0, 1, 1, 0, 0, 0, 1, 0, 0, 0,
                0, 1, 1, 1, 0, 0, 0, 1, 0, 0, 0, 1, 0, 0, 1, 1, 0, 0, 0, 0, 0, 0, 0,
                1, 1, 0, 0, 0, 0, 0, 1, 1, 0, 0, 0, 0, 0, 0, 1, 0, 0, 1, 1, 1, 0,
                0, 0, 0, 0, 1, 1, 0, 0, 1, 0, 1, 0, 1, 0, 0, 0, 1, 1, 1, 0, 1, 0,
                0, 0, 0, 1, 1, 1, 0, 0, 0, 0, 0, 1, 0, 1, 0, 1, 0, 0, 0, 1, 0,
                0, 0, 0, 1, 0, 1, 1, 0, 0, 0, 1, 0, 0, 1, 0, 1, 0, 0, 0, 0,
                1, 1, 1, 0, 0, 1, 1, 0, 0, 0, 0, 1, 0, 0, 1, 0, 0, 0, 1, 0, 0,
                0, 0, 1, 1, 1, 1, 0], dtype=int64),
          'effective_metric_params_': {},
          'effective_metric_': 'euclidean',
          '_fit_method': 'kd_tree',
          _fit_X': array([[ 1.15399459, 2.34780853, -1.34604126, ..., 2.68061558,
                 -0.63290132, 0.95454214],
                [-0.37397249, -1.65789891, -0.21865313, ..., -0.80741433,
                 -0.17876717, -0.62389402],
                [1.2475436, -1.02541879, -1.34604126, ..., -1.09808349,
                 -0.641984 , 2.79605101],
                [1.43464161, 1.92615512, 1.53506174, ..., 1.22726978,
                 -0.91143693, 0.86685124],
                 [0.34323655, 0.55578152, -1.34604126, ..., -0.22607601,
                 -0.19390497, 0.95454214],
                 [-0.84171751, 0.1341281, -0.21865313, ..., 1.8086081,
                  0.3904143 , 2.00683292]]),
          'n samples fit ': 558,
          '_tree': <sklearn.neighbors._kd_tree.KDTree at 0x22820aa9730>}
        # TODO: 42 Write code to display a confusion matrix by filling the question marks
         from sklearn.metrics import confusion matrix, ConfusionMatrixDisplay
         cm = confusion_matrix(y_test, y_pred, labels=[0, 1])
         disp = ConfusionMatrixDisplay(confusion_matrix=cm, display_labels=['Class 0', 'Clas'
```

```
disp.plot()
plt.show()
```



```
In []: # TODO: 43 Interpret the confusion matrix in your own words

# True Positives (TP): 40
# This means that the model correctly predicted 40 instances as Class 1, and they w

# True Negatives (TN): 107
# The model correctly predicted 107 instances as Class 0, and they were indeed Clas

# False Positives (FP): 19
# The model incorrectly predicted 19 instances as Class 1 when they were actually (

# False Negatives (FN): 20
# The model incorrectly predicted 20 instances as Class 0 when they were actually (
```

Part 4: Save the model as json file, load & use it to do prediction

This is optional and will be used to accumulate bonus points as a buffer for the semester lab exam if there is any shortfall

```
In []: import json
import numpy as np
from sklearn.neighbors import KNeighborsClassifier

# Create and fit a KNeighborsClassifier
model = KNeighborsClassifier(n_neighbors=5)
model.fit(X_train, y_train)

# Serialize the model and its parameters to a JSON file
```

```
model_info = {
    "model_params": model.get_params(),
    "X_train": X_train.to_numpy().tolist(), # Convert DataFrame to NumPy array and
    "y_train": y_train.to_list() # Convert Series to a List
}

with open("AML_Sessional_1_Students.json", "w") as json_file:
    json.dump(model_info, json_file)

# Load model information from the JSON file
with open("AML_Sessional_1_Students.json", "r") as json_file:
    loaded_model_info = json.load(json_file)

# Create a new KNeighborsClassifier with the Loaded parameters
loaded_model = KNeighborsClassifier(**loaded_model_info["model_params"])

# Fit the Loaded model using the Loaded training data
loaded_model.fit(np.array(loaded_model_info["X_train"]), np.array(loaded_model_info
# Perform predictions using the Loaded and fitted model
y_pred = loaded_model.predict(X_test)
```