Import Libraries

```
import numpy as np # linear algebra
import pandas as pd # data processing, CSV file I/O (e.g. pd.read_csv)
import random
import copy
from sklearn.model_selection import LeaveOneOut, GridSearchCV
from sklearn.neighbors import KNeighborsClassifier
from sklearn.metrics import matthews_corrcoef, confusion_matrix, accuracy_score
from scipy.stats import fisher_exact
from sklearn.preprocessing import StandardScaler
```

Read In Data

```
gene df = pd.read excel('/kaggle/input/pomeroy-et-al-brain-data/Dataset C MD outcome2.gct (6).xlsx')
```

Create a list of tuples formated ('patientIDs', 'patientStatus') where patientIDs is Brain_MD_X and patientStatus is 0 (dead) or 1 (alive)

```
# Brin_MD_X
patientIDs = gene_df.columns[2:].to_list()
# dead is 0, alive is 1
patientStatus = [0 if i <= 21 else 1 for i in range(1, 61)]
# Create the list of tuples
patients = list(zip(patientIDs, patientStatus))
#print(patients)</pre>
```

Create Training and Testing Datasets.

```
temp trainingSet = patients[:30]
    # set number of dead patients in set variable to 0
    dead, alive = 0, 0
    # Count how many dead patients were selected for the set
    for patient in temp_trainingSet:
        if patient[1] == 0:
            dead+=1
        else:
            alive+=1
    # To stop infinite loops
    if attemptNumber == 10:
        break
    # If about half of the 39 alive patients are in this data set, populate the training and testing datas
    if 17 <= alive <= 22:
        print(alive, 'patients who responded and ', dead, 'patients who did not respond in testing dataset
        patients_trainingSet = patients[:30]
        patients_testingSet = patients[30:]
        goodData = True
    else:
        # If attempt fails
        print("Failed attempt", attemptNumber)
# print training and datasets
#print('Training Dataset:\n', patients_trainingSet,'\n\n')
#print('Testing Dataset:\n', patients_testingSet)
     18 patients who responded and 12 patients who did not respond in testing dataset
```

Ensure data sets are good

- 1. Ensure the correct number of patients are in the datasets
- 2. Ensure there is no overlap between data sets (no duplicate patients)

```
if len(patients_trainingSet) == 30:
    if len(patients_testingSet) == 30:
        print('The correct number of patients per dataset')
    else:
        print('panic', len(patients_trainingSet), len(patients_testingSet))

for train_patient in patients_trainingSet:
    for test_patient in patients_testingSet:
        if train_patient[0] == test_patient[0]:
            print('Panic, ', test_patiet[0], ' is duplicated')
            sys.exit()

print("No duplicate patients across training and testing datasets")

The correct number of patients per dataset
    No duplicate patients across training and testing datasets
```

K-Nearest Neighbor analysis

Craete a input

- 1. outcome -> a list of the outcomes for all the patients in the training data set
- 2. expressionLevel -> a list of lists. For every probe, each patient's expression level for that probe is consolidated into a list.

Function to find the best K-Nearest Neighbor classifier and parameters

```
def knn leave one out(X, y):
    loo = LeaveOneOut()
    # Define parameter grid for grid search
    param_grid = {'n_neighbors': range(1, 20)} # This can be adjusted up to the number of patients in the
    # Initialize KNN classifier
    knn = KNeighborsClassifier()
    # Perform grid search with leave-one-out cross-validation
    grid_search = GridSearchCV(knn, param_grid, scoring='accuracy', cv=loo)
    grid_search.fit(X, y)
    # Get the best parameters
    best params = grid search.best params
    # Train the best KNN classifier on the entire dataset
    best knn = KNeighborsClassifier(n neighbors=best params['n neighbors'])
    best knn.fit(X, y)
    return best_knn, best_params
# Perform leave-one-out testing
best_knn, best_params = knn_leave_one_out(train_expressionLevel, trainOutcomes)
print("Best Parameters:", best_params)
print(best knn)
     Best Parameters: {'n neighbors': 1}
     KNeighborsClassifier(n_neighbors=1)
```

Apply the classifier to the testing set and evaluate the accuracy

```
# Create a copy of the entire dataset
test = gene_df.copy()

# Get a list of patients that will be removed to create the training dataset
patTest = [lisTest[0] for lisTest in patients_trainingSet]

# for each patient that is in the testing set, drop it out of the training set
for patient in patTest:
    test = test.drop(columns=[patient])

# A list of the outcomes (dead/alive) for all the patients in the training data set
testOutcomes = [lis[1] for lis in patients_trainingSet]

# A list of lists. For every probe, each patient's expression level for that probe is consolidated into a
test_expressionLevel = test.drop(test.iloc[:, 0:2], axis=1)

scaler.fit(test_expressionLevel)

#train_expressionLevel = train_expressionLevel.transpose()
test_expressionLevel = scaler.transform(test_expressionLevel).transpose()
```

```
# Use the trained KNN classifier to make predictions on the test data
testPredictions = best_knn.predict(test_expressionLevel)
# Calculate the confusion matrix
confMatrix = confusion_matrix(testOutcomes, testPredictions)
# Fisher's exact test
oddsRatio, pValue = fisher_exact(confMatrix)
print("Fisher's exact test p-value:", pValue)
#Assuming you have predictions and true labels
mccPhi = matthews_corrcoef(testOutcomes, testPredictions)
print("Matthews' correlation coefficient Phi:", mccPhi)
     Fisher's exact test p-value: 0.03415061699919271
     Matthews' correlation coefficient Phi: 0.43082021842766455
predictions 0 = best knn.predict(test expressionLevel)
# Evaluate the performance using accuracy and other metrics
accuracy = accuracy_score(testOutcomes, predictions_0)
print("Accuracy on new data:", accuracy)
```