

Proj_04_Breast_Cancer_SVM_Full_4

October 17, 2024

1 EE915: Week-4 - Project-4 - SVM - Breast Cancer Prediction

Name: Venkateswar Reddy Melachervu **Roll No.:** 23156022 **Email:** vmela23@iitk.ac.in **Dataset:** Breast Cancer Data Set from sklearn **Professor:** Prof. Aditya K. Jagganatham **TAs:** Meesam and Priyanka

This project implements a Support Vector Machine (SVM) classifier to predict the class of breast cancer from the provided dataset.

The dataset is `load_breast_cancer()` from sklearn. The breast cancer dataset consists of 569 samples, each representing a patient with a set of features.

The dataset is used for training a SVM classifier and evaluate its performance using various metrics such as accuracy, precision, recall, F1-score.

Additionally, various other classifiers, such as Logistic Regression, Naive Bayes, and K-Nearest Neighbors are also implemented and evaluated for comparison.

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```
[1]: # Importing necessary libraries for breast cancer SVM classifier
from sklearn import datasets
from sklearn.model_selection import train_test_split
from sklearn.preprocessing import StandardScaler
from sklearn.metrics import accuracy_score
from sklearn.svm import SVC
import pandas as pd # for dataframe manipulation
from pandas.plotting import parallel_coordinates # for parallel coordinates
    ↪plot of breast cancer data set
import matplotlib.pyplot as plt # for plotting graphs
from sklearn.manifold import TSNE # for t-SNE plot
from sklearn.ensemble import RandomForestClassifier
from sklearn.metrics import auc, confusion_matrix, precision_score,
    ↪recall_score, roc_curve
from sklearn.model_selection import cross_val_score
from sklearn.metrics import f1_score
from sklearn.linear_model import LogisticRegression
from sklearn.naive_bayes import GaussianNB
from sklearn.neighbors import KNeighborsClassifier
from sklearn.tree import DecisionTreeClassifier
```

```

import seaborn as sns
from collections import Counter
import numpy as np
from sklearn.metrics import accuracy_score, f1_score, roc_curve, auc,
precision_recall_curve, confusion_matrix, ConfusionMatrixDisplay
from matplotlib.colors import ListedColormap

```

```

[2]: # Define roll number, name, email
roll_number = "23156022"
name = "Venkateswar Reddy Melachervu"
email = "vmela23@iitk.ac.in"

# Load the breast cancer dataset from sklearn
bcancer = datasets.load_breast_cancer()
X = bcancer.data
Y = bcancer.target

# Scale the data using standard scaler
scaler = StandardScaler()
standard_scaled_X = scaler.fit_transform(X)

# Display metadata about the breast cancer dataset
df = pd.DataFrame(data=X, columns=bcancer.feature_names)
df['target'] = Y

# Set display option to show all columns
pd.set_option('display.max_columns', None)
pd.set_option('display.width', 1000)
print('Breast Cancer Data Set Details:')
print(f'\tComprises of {df.shape[0]} data sample vectors')
print(f'\tEach vector is of {df.shape[1]-1} features/dimensions and a target\u
scalar response.')

# Count the data samples for each target value
target_counts = df['target'].value_counts()

# Print the counts separately
count_0 = target_counts[0]
count_1 = target_counts[1]

print(f"\tNumber of samples with target value 1 - \"Benign/Non-cancerous\u
\tTumors\": {count_1}")
print(f"\tNumber of samples with target value 0 - \"Malignant/Cancerous\u
\tTumors\": {count_0}")
print(f"\tTotal number of data samples - \"Breast Cancer\" data set: {count_0}")

# Capitalize the first letter of each word in the feature names

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df.columns = [col.title() for col in df.columns]

# Filter out the 'Target' column after capitalizing
feature_columns = [col for col in df.columns if col != 'Target']

# Convert to list
column_names = feature_columns
print(f"\tThe feature names are:")
print(f'\t{column_names}')

```

Breast Cancer Data Set Details:

Comprises of 569 data sample vectors
 Each vector is of 30 features/dimensions and a target scalar response.
 Number of samples with target value 1 - "Benign/Non-cancerous Tumors":
 357
 Number of samples with target value 0 - "Malignant/Cancerous Tumors":
 212
 Total number of data samples - "Breast Cancer" data set: 212
 The feature names are:
 ['Mean Radius', 'Mean Texture', 'Mean Perimeter', 'Mean Area', 'Mean Smoothness', 'Mean Compactness', 'Mean Concavity', 'Mean Concave Points', 'Mean Symmetry', 'Mean Fractal Dimension', 'Radius Error', 'Texture Error', 'Perimeter Error', 'Area Error', 'Smoothness Error', 'Compactness Error', 'Concavity Error', 'Concave Points Error', 'Symmetry Error', 'Fractal Dimension Error', 'Worst Radius', 'Worst Texture', 'Worst Perimeter', 'Worst Area', 'Worst Smoothness', 'Worst Compactness', 'Worst Concavity', 'Worst Concave Points', 'Worst Symmetry', 'Worst Fractal Dimension']

[3]: # Display the first row of the data set

```

print('First row of standard scaled data sample in the Breast Cancer data set is:')
first_sample = df.iloc[0]
for idx, (feature, value) in enumerate(first_sample.items(), start=1):
    print(f'{idx}. {feature}: {value:.6f}')

```

First row of standard scaled data sample in the Breast Cancer data set is:

1. Mean Radius	17.990000
2. Mean Texture	10.380000
3. Mean Perimeter	122.800000
4. Mean Area	1001.000000
5. Mean Smoothness	0.118400
6. Mean Compactness	0.277600
7. Mean Concavity	0.300100
8. Mean Concave Points	0.147100
9. Mean Symmetry	0.241900
10. Mean Fractal Dimension	0.078710
11. Radius Error	1.095000
12. Texture Error	0.905300

13. Perimeter Error	8.589000
14. Area Error	153.400000
15. Smoothness Error	0.006399
16. Compactness Error	0.049040
17. Concavity Error	0.053730
18. Concave Points Error	0.015870
19. Symmetry Error	0.030030
20. Fractal Dimension Error	0.006193
21. Worst Radius	25.380000
22. Worst Texture	17.330000
23. Worst Perimeter	184.600000
24. Worst Area	2019.000000
25. Worst Smoothness	0.162200
26. Worst Compactness	0.665600
27. Worst Concavity	0.711900
28. Worst Concave Points	0.265400
29. Worst Symmetry	0.460100
30. Worst Fractal Dimension	0.118900
31. Target	0.000000

```
[4]: # Lets plot t-SNE plot of the breast cancer data set

# Perform t-SNE on pristine data
tsne_pristine = TSNE(n_components=2, random_state=42)
X_tsne_pristine = tsne_pristine.fit_transform(X)

# Perform t-SNE on standard scaled data
tsne_scaled = TSNE(n_components=2, random_state=42)
X_tsne_scaled = tsne_scaled.fit_transform(standard_scaled_X)

# Print the derived X and Y values
print("t-SNE on Pristine Data (first 5 points):")
print(X_tsne_pristine[:5])
print("\nt-SNE on Standard Scaled Data (first 5 points):")
print(X_tsne_scaled[:5])

# Define colors and line width for the plot
colors = ['red', 'green']
lw = 2

# Plot t-SNE results
fig, axes = plt.subplots(1, 2, figsize=(16, 6))

# t-SNE on pristine data
axes[0].set_title('t-SNE of Pristine Breast Cancer Dataset')
for color, i, target_name in zip(colors, [0, 1], bcancer.target_names):
```

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    axes[0].scatter(X_tsne_pristine[Y == i, 0], X_tsne_pristine[Y == i, 1], color=color, alpha=.8, lw=lw, label=target_name.capitalize())
    axes[0].legend(loc='best', shadow=False, scatterpoints=1)

# t-SNE on standard scaled data
axes[1].set_title('t-SNE of Standard Scaled Breast Cancer Dataset')
for color, i, target_name in zip(colors, [0, 1], bcancer.target_names):
    axes[1].scatter(X_tsne_scaled[Y == i, 0], X_tsne_scaled[Y == i, 1], color=color, alpha=.8, lw=lw, label=target_name.capitalize())
    axes[1].legend(loc='best', shadow=False, scatterpoints=1)

# Add watermark with roll number
for ax in axes:
    ax.text(0.5, 0.5, roll_number, fontsize=50, color='gray', alpha=0.3, ha='center', va='center', rotation=45, transform=ax.transAxes)
plt.show()

```

t-SNE on Pristine Data (first 5 points):

```

[[ 42.337463 -12.640097 ]
 [ 42.50024   -9.92803  ]
 [ 37.894382 -10.039126 ]
 [-17.654438   1.4511237]
 [ 36.764866  -7.9306464]]

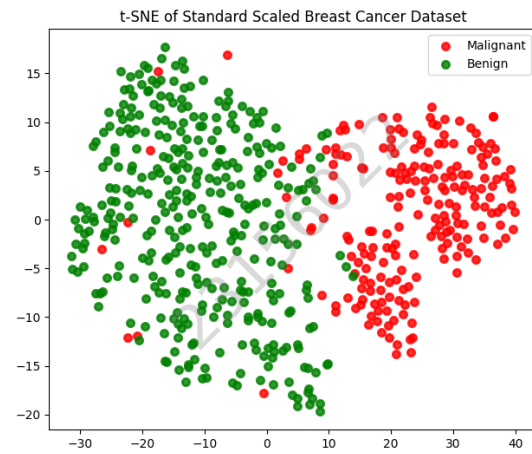
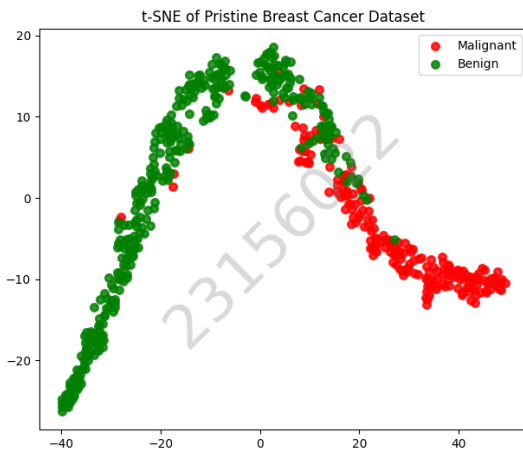
```

t-SNE on Standard Scaled Data (first 5 points):

```

[[ 37.784595    1.2305161]
 [ 23.71192    7.3292117]
 [ 31.739286    3.551559 ]
 [ 20.865301   -13.742583 ]
 [ 26.342352   -1.7316735]]

```



```
[5]: # Parallel coordinates plot of the breast cancer data set for pristine and standard scaled data

# Create DataFrames from the data
df_pristine = pd.DataFrame(data=X, columns=bcancer.feature_names)
df_pristine['target'] = Y

df_scaled = pd.DataFrame(data=standard_scaled_X, columns=bcancer.feature_names)
df_scaled['target'] = Y

# Convert target to string for color coding
df_pristine['target'] = df_pristine['target'].astype(str)
df_scaled['target'] = df_scaled['target'].astype(str)

# Define colors for each class explicitly
color_dict = {'0': 'red', '1': 'green'}
# Create a colormap from the color dictionary
cmap = plt.cm.colors.ListedColormap([color_dict['0'], color_dict['1']])

# Identify features with mean value above 100 - Exclude the 'target' column
mean_values = df_pristine.drop('target', axis=1).mean()
features_above_100 = mean_values[mean_values > 100].index

# Create subplots for parallel coordinates
fig, axes = plt.subplots(1, 2, figsize=(16, 8))

# Parallel coordinates plot for pristine data
parallel_coordinates(df_pristine, 'target', colormap=cmap, ax=axes[0])
axes[0].set_title('Parallel Coordinates Plot of Pristine Data')
axes[0].set_xlabel('Features')
axes[0].set_ylabel('Feature values')

# Highlight the dominant features
for feature in features_above_100:
    axes[0].axvline(df_pristine.columns.get_loc(feature), color='blue', linestyle='--', linewidth=1)

# Parallel coordinates plot for standard scaled data
parallel_coordinates(df_scaled, 'target', colormap=cmap, ax=axes[1])
axes[1].set_title('Parallel Coordinates Plot of Standard Scaled Data')
axes[1].set_xlabel('Features')
axes[1].set_ylabel('Feature values')

# Highlight the dominant features
for feature in features_above_100:
    axes[1].axvline(df_scaled.columns.get_loc(feature), color='blue', linestyle='--', linewidth=1)
```

```

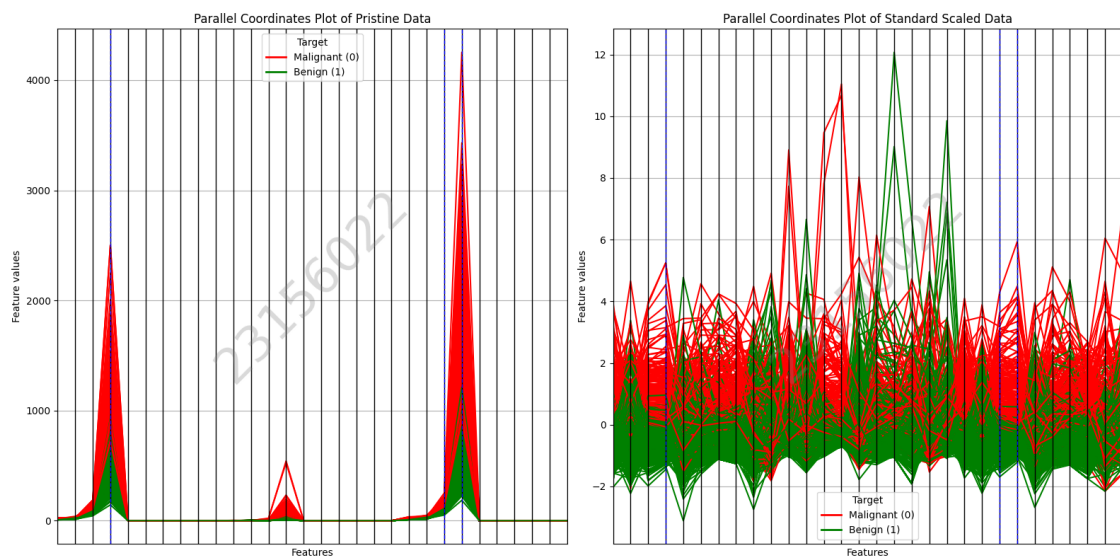
# Add custom legend
handles = [plt.Line2D([0], [0], color=color_dict['0'], lw=2), plt.Line2D([0], [0], color=color_dict['1'], lw=2)]
axes[0].legend(handles, ['Malignant (0)', 'Benign (1)'], title='Target')
axes[1].legend(handles, ['Malignant (0)', 'Benign (1)'], title='Target')

# Remove x-axis labels
axes[0].set_xticks([])
axes[1].set_xticks([])

# Add watermark with roll number
for ax in axes:
    ax.text(0.5, 0.5, roll_number, fontsize=50, color='gray', alpha=0.3,
            ha='center', va='center', rotation=45, transform=ax.transAxes)

plt.tight_layout()
plt.show()

```



[6]: # Plot curves for all classifiers - accuracy, confusion matrix, ROC curve, PR curve, feature importance, F1 score

```

# Define the formulae as strings
precision_formula = "Precision = True Positives / (True Positives + False Positives)"
recall_formula = "Recall = True Positives / (True Positives + False Negatives)"
f1_score_formula = "F1 Score = 2 * (Precision * Recall) / (Precision + Recall)"

```

```

# Print the formulae
print(precision_formula)
print(recall_formula)
print(f1_score_formula, '\n')

# Define a dictionary of models
models = {
    'Linear SVM': SVC(kernel='linear', random_state=0, probability=True),
    'RBF Kernel SVM': SVC(kernel='rbf', random_state=0, probability=True),
    'Poly Kernel SVM': SVC(kernel='poly', degree=3, random_state=0, ↴
        probability=True),
    'Sigmoid Kernel SVM': SVC(kernel='sigmoid', random_state=0, ↴
        probability=True),
    'Logistic Regression': LogisticRegression(max_iter=10000, random_state=0),
    'Decision Tree': DecisionTreeClassifier(criterion='entropy', ↴
        random_state=0),
    'K-Nearest Neighbors': KNeighborsClassifier(n_neighbors=5, ↴
        metric='minkowski', p=2),
    'Naive Bayes': GaussianNB(),
    'Random Forest': RandomForestClassifier(n_estimators=10, ↴
        criterion='entropy', random_state=0)
}

# Initialize plot
fig, axs = plt.subplots(len(models), 6, figsize=(24, len(models) * 5))
fig.subplots_adjust(hspace=0.5)

# Split the data
X_train, X_test, Y_train, Y_test = train_test_split(standard_scaled_X, Y, ↴
    test_size=0.2, random_state=0)

# Loop through models
for idx, (name, model) in enumerate(models.items()):
    # Train model
    model.fit(X_train, Y_train)

    # Predict results
    Y_pred = model.predict(X_test)
    Y_proba = model.predict_proba(X_test)[:, 1] if hasattr(model, ↴
        'predict_proba') else None
    accuracy = accuracy_score(Y_test, Y_pred)
    print(f"{name} Accuracy Score: {100 * accuracy:.2f}%")

    # Compute metrics
    cm = confusion_matrix(Y_test, Y_pred)

```

```

fpr, tpr, _ = roc_curve(Y_test, Y_proba) if Y_proba is not None else ([] ,[])
roc_auc = auc(fpr, tpr) if Y_proba is not None else None
precision, recall, _ = precision_recall_curve(Y_test, Y_proba) if Y_proba is not None else ([] ,[], [])
f1 = f1_score(Y_test, Y_pred)

# Plot Accuracy Score curve
if Y_proba is not None:
    thresholds = np.linspace(0, 1, 100)
    accuracy_scores = [accuracy_score(Y_test, (Y_proba > threshold).
    astype(int)) for threshold in thresholds]
    axs[idx, 5].plot(thresholds, accuracy_scores, color='green', lw=2)
    axs[idx, 5].set_xlim([0.0, 1.0])
    axs[idx, 5].set_ylim([0.0, 1.05])
    axs[idx, 5].set_xlabel('Decision Threshold')
    axs[idx, 5].set_ylabel('Accuracy Score')
    axs[idx, 5].set_title(f'{name} Accuracy Score Curve')
else:
    axs[idx, 5].text(0.5, 0.5, 'No Accuracy Curve',
horizontalalignment='center', verticalalignment='center', fontsize=12)
    axs[idx, 5].set_title(f'{name} Accuracy Score Curve')

# Plot confusion matrix
sns.heatmap(cm, annot=True, fmt='d', cmap='Blues', ax=axs[idx, 0],
cbar=False, xticklabels=['Malignant', 'Benign'], yticklabels=['Malignant',
'Benign'])
axs[idx, 0].set_xlabel('Predicted Label')
axs[idx, 0].set_ylabel('True Label')
axs[idx, 0].set_title(f'{name} Confusion Matrix')

# Plot ROC curve
if Y_proba is not None:
    axs[idx, 1].plot(fpr, tpr, color='darkorange', lw=2, label=f'ROC curve',
(area = {roc_auc:.2f}))
    axs[idx, 1].plot([0, 1], [0, 1], color='navy', lw=2, linestyle='--')
    axs[idx, 1].set_xlim([0.0, 1.0])
    axs[idx, 1].set_ylim([0.0, 1.05])
    axs[idx, 1].set_xlabel('False Positive Rate')
    axs[idx, 1].set_ylabel('True Positive Rate')
    axs[idx, 1].set_title(f'{name} ROC Curve')
    axs[idx, 1].legend(loc='lower right')
else:
    axs[idx, 1].text(0.5, 0.5, 'No ROC Curve',
horizontalalignment='center', verticalalignment='center', fontsize=12)
    axs[idx, 1].set_title(f'{name} ROC Curve')

```

```

# Plot Precision-Recall curve
if Y_proba is not None:
    axs[idx, 2].plot(recall, precision, color='blue', lw=2)
    axs[idx, 2].set_xlim([0.0, 1.0])
    axs[idx, 2].set_ylim([0.0, 1.05])
    axs[idx, 2].set_xlabel('Recall')
    axs[idx, 2].set_ylabel('Precision')
    axs[idx, 2].set_title(f'{name} Precision-Recall Curve')
else:
    axs[idx, 2].text(0.5, 0.5, 'No PR Curve', horizontalalignment='center',
                     verticalalignment='center', fontsize=12)
    axs[idx, 2].set_title(f'{name} Precision-Recall Curve')

# Plot feature importance for models that have it
if hasattr(model, 'feature_importances_'):
    importances = model.feature_importances_
    axs[idx, 3].barh(range(len(importances)), importances, align='center')
    axs[idx, 3].set_yticks(range(len(importances)))
    axs[idx, 3].set_yticklabels(bcancer.feature_names)
    axs[idx, 3].set_xlabel('Feature Importance')
    axs[idx, 3].set_title(f'{name} Feature Importance')
else:
    axs[idx, 3].text(0.5, 0.5, 'No Feature Importance', horizontalalignment='center',
                     verticalalignment='center', fontsize=12)
    axs[idx, 3].set_title(f'{name} Feature Importance')

# Plot F1 Score
axs[idx, 4].bar(name, f1, color='purple')
axs[idx, 4].set_ylim([0, 1])
axs[idx, 4].set_xlabel('Model')
axs[idx, 4].set_ylabel('F1 Score')
axs[idx, 4].set_title(f'{name} F1 Score')

# Add watermark to all subplots
for ax in axs[idx, :]:
    ax.text(0.5, 0.5, '23156022', fontsize=50, color='gray', alpha=0.3,
            rotation=45, ha='center', va='center', transform=ax.transAxes)

plt.tight_layout()
plt.show()

```

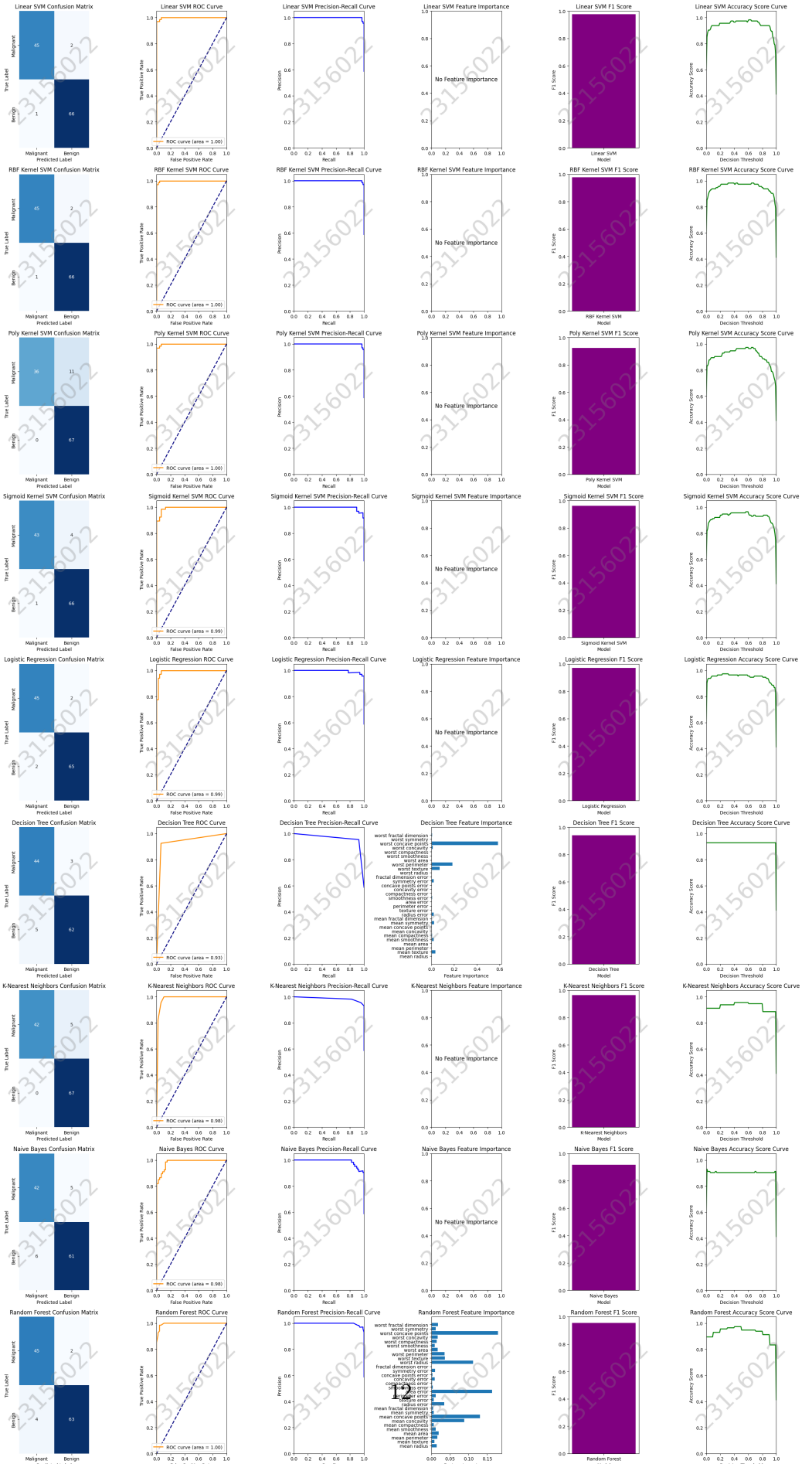
```

Precision = True Positives / (True Positives + False Positives)
Recall = True Positives / (True Positives + False Negatives)
F1 Score = 2 * (Precision * Recall) / (Precision + Recall)

```

Linear SVM Accuracy Score: 97.37%

RBF Kernel SVM Accuracy Score: 97.37%
Poly Kernel SVM Accuracy Score: 90.35%
Sigmoid Kernel SVM Accuracy Score: 95.61%
Logistic Regression Accuracy Score: 96.49%
Decision Tree Accuracy Score: 92.98%
K-Nearest Neighbors Accuracy Score: 95.61%
Naive Bayes Accuracy Score: 90.35%
Random Forest Accuracy Score: 94.74%



```
[7]: # Relative performance plots of all models - accuracy, ROC curve, PR curve, F1 score

# Function to add watermark
def add_watermark(ax, roll_number='23156022'):
    ax.text(0.5, 0.5, roll_number, fontsize=50, color='gray', alpha=0.3, rotation=45,
            ha='center', va='center', transform=ax.transAxes)

# Define a dictionary of models
models = {
    'Linear SVM': SVC(kernel='linear', random_state=0, probability=True),
    'RBF Kernel SVM': SVC(kernel='rbf', random_state=0, probability=True),
    'Poly Kernel SVM': SVC(kernel='poly', degree=3, random_state=0, probability=True),
    'Sigmoid Kernel SVM': SVC(kernel='sigmoid', random_state=0, probability=True),
    'Logistic Regression': LogisticRegression(max_iter=10000, random_state=0),
    'Decision Tree': DecisionTreeClassifier(criterion='entropy', random_state=0),
    'K-Nearest Neighbors': KNeighborsClassifier(n_neighbors=5, metric='minkowski', p=2),
    'Naive Bayes': GaussianNB(),
    'Random Forest': RandomForestClassifier(n_estimators=10, criterion='entropy', random_state=0)
}

# Split the data
X_train, X_test, Y_train, Y_test = train_test_split(standard_scaled_X, Y, test_size=0.2, random_state=0)

# Initialize dictionaries to store metrics
accuracy_scores = []
f1_scores = []
roc_data = []
prc_data = []

# Calculate metrics
for name, model in models.items():
    model.fit(X_train, Y_train)
    Y_pred = model.predict(X_test)
    Y_proba = model.predict_proba(X_test)[:, 1] if hasattr(model, 'predict_proba') else None
```

```

# Accuracy
accuracy = accuracy_score(Y_test, Y_pred)
accuracy_scores[name] = accuracy

# F1 Score
f1 = f1_score(Y_test, Y_pred)
f1_scores[name] = f1

# ROC Curve
if Y_proba is not None:
    fpr, tpr, _ = roc_curve(Y_test, Y_proba)
    roc_auc = auc(fpr, tpr)
    roc_data[name] = (fpr, tpr, roc_auc)

# Precision-Recall Curve
precision, recall, _ = precision_recall_curve(Y_test, Y_proba)
prc_data[name] = (precision, recall)

# Plot Accuracy Scores
plt.figure(figsize=(12, 6))
bars = plt.barh(list(accuracy_scores.keys()), list(accuracy_scores.values()), color='skyblue')
plt.xlabel('Accuracy')
plt.title('Model Accuracy')
plt.gca().invert_yaxis()
plt.grid(axis='x')
# Set x-axis ticks
plt.xticks(np.arange(0, 1.1, 0.1))
add_watermark(plt.gca())
# Add value labels to each bar
for bar in bars:
    plt.text(bar.get_width(), bar.get_y() + bar.get_height()/2,
             f'{bar.get_width():.2f}', va='center', ha='left')
plt.tight_layout()
plt.show()

# Plot ROC Curves
plt.figure(figsize=(12, 6))
for name, (fpr, tpr, roc_auc) in roc_data.items():
    plt.plot(fpr, tpr, lw=2, label=f'{name} (area = {roc_auc:.2f})')
plt.plot([0, 1], [0, 1], 'k--')
plt.xlim([0.0, 1.0])
plt.ylim([0.0, 1.05])
plt.xlabel('False Positive Rate')
plt.ylabel('True Positive Rate')
plt.title('ROC Curve for All Models')
plt.legend(loc='lower right')

```

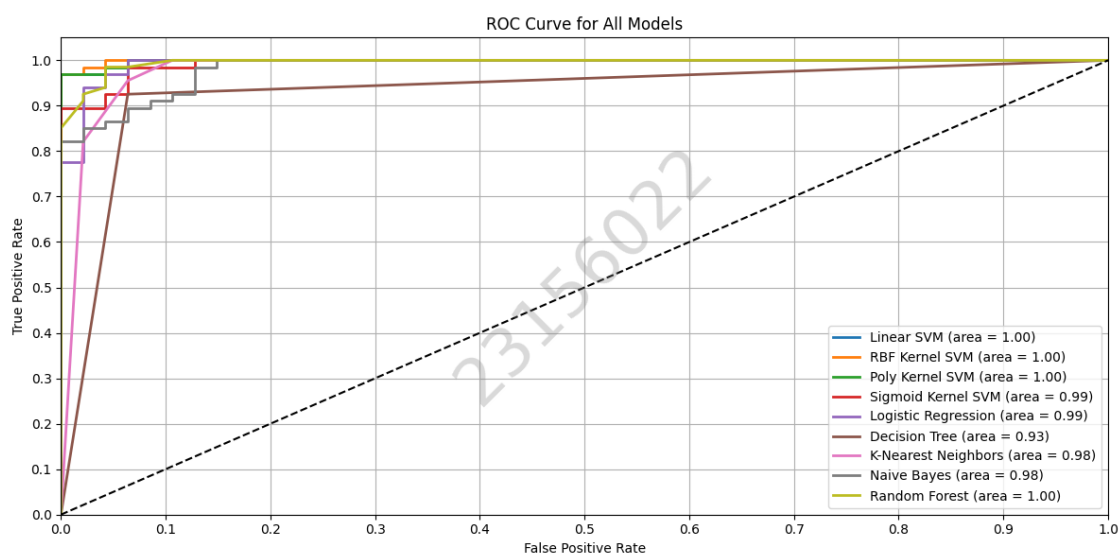
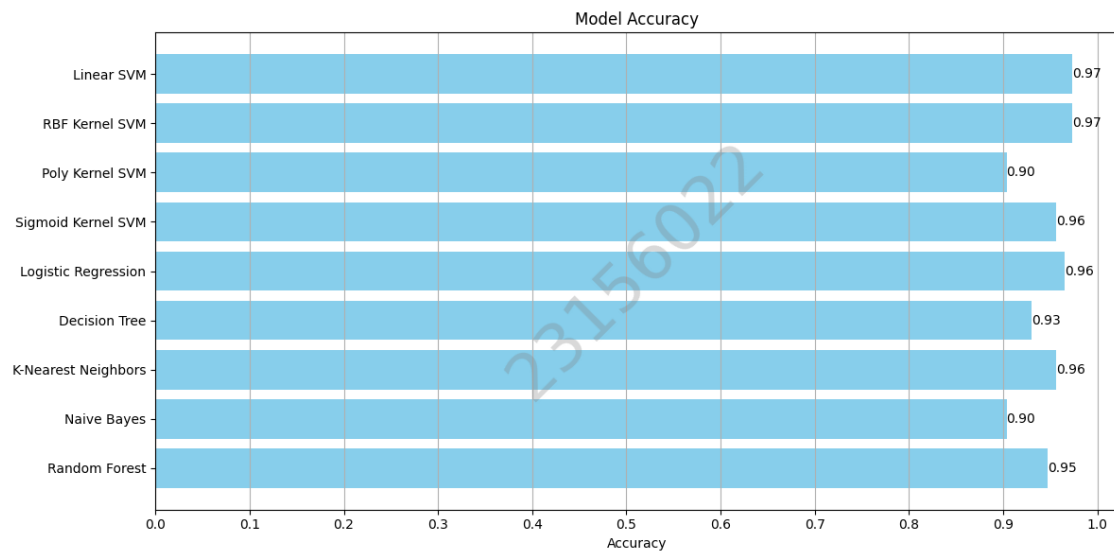
```

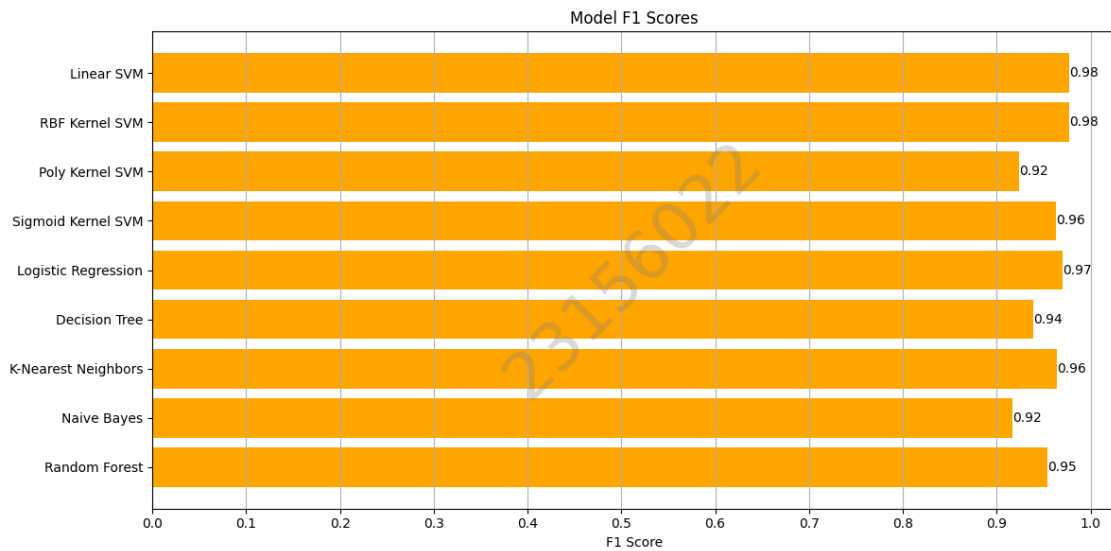
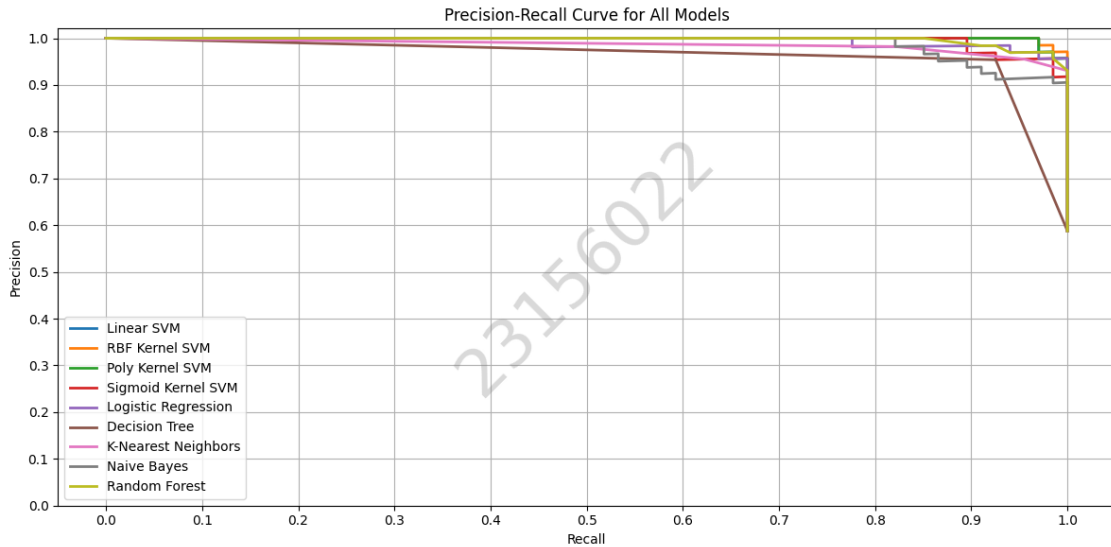
plt.grid()
# Set x-axis ticks
plt.xticks(np.arange(0, 1.1, 0.1))
# Set y-axis ticks
plt.yticks(np.arange(0, 1.1, 0.1))
add_watermark(plt.gca())
plt.tight_layout()
plt.show()

# Plot Precision-Recall Curves
plt.figure(figsize=(12, 6))
for name, (precision, recall) in prc_data.items():
    plt.plot(recall, precision, lw=2, label=name)
plt.xlabel('Recall')
plt.ylabel('Precision')
plt.title('Precision-Recall Curve for All Models')
plt.legend(loc='best')
plt.grid()
# Set x-axis ticks
plt.xticks(np.arange(0, 1.1, 0.1))
# Set y-axis ticks
plt.yticks(np.arange(0, 1.1, 0.1))
add_watermark(plt.gca())
plt.tight_layout()
plt.show()

# Plot F1 Scores
plt.figure(figsize=(12, 6))
bars = plt.barh(list(f1_scores.keys()), list(f1_scores.values()), color='orange')
plt.xlabel('F1 Score')
plt.title('Model F1 Scores')
plt.gca().invert_yaxis()
plt.grid(axis='x')
# Set x-axis ticks
plt.xticks(np.arange(0, 1.1, 0.1))
add_watermark(plt.gca())
# Add value labels to each bar
for bar in bars:
    plt.text(bar.get_width(), bar.get_y() + bar.get_height()/2,
             f'{bar.get_width():.2f}', va='center', ha='left')
plt.tight_layout()
plt.show()

```





```
[8]: # Plot curves for all classifiers - accuracy, confusion matrix, ROC curve, PR curve, feature importance, F1 score, decision boundaries
```

```
# Define the formulae as strings
precision_formula = "Precision = True Positives / (True Positives + False Positives)"
recall_formula = "Recall = True Positives / (True Positives + False Negatives)"
f1_score_formula = "F1 Score = 2 * (Precision * Recall) / (Precision + Recall)"

# Print the formulae
```

```

print(precision_formula)
print(recall_formula)
print(f1_score_formula, '\n')

# Define a dictionary of models
models = {
    'Linear SVM': SVC(kernel='linear', random_state=0, probability=True),
    'RBF Kernel SVM': SVC(kernel='rbf', random_state=0, probability=True),
    'Poly Kernel SVM': SVC(kernel='poly', degree=3, random_state=0,
                           probability=True),
    'Sigmoid Kernel SVM': SVC(kernel='sigmoid', random_state=0,
                               probability=True),
    'Logistic Regression': LogisticRegression(max_iter=10000, random_state=0),
    'Decision Tree': DecisionTreeClassifier(criterion='entropy',
                                             random_state=0),
    'K-Nearest Neighbors': KNeighborsClassifier(n_neighbors=5,
                                                 metric='minkowski', p=2),
    'Naive Bayes': GaussianNB(),
    'Random Forest': RandomForestClassifier(n_estimators=10,
                                             criterion='entropy', random_state=0)
}

# Initialize plot
fig, axs = plt.subplots(len(models), 7, figsize=(35, len(models) * 5))
fig.subplots_adjust(hspace=0.5)

# Split the data
X_train, X_test, Y_train, Y_test = train_test_split(standard_scaled_X, Y,
                                                    test_size=0.2, random_state=0)

# Select two features for decision boundary plot
feature1, feature2 = 0, 1 # example feature indices, change as needed

# Loop through models
for idx, (name, model) in enumerate(models.items()):
    # Train model on the full dataset
    model.fit(X_train, Y_train)

    # Predict results
    Y_pred = model.predict(X_test)
    Y_proba = model.predict_proba(X_test)[:, 1] if hasattr(model, 'predict_proba') else None
    accuracy = accuracy_score(Y_test, Y_pred)
    print(f"{name} Accuracy Score: {100 * accuracy:.2f}%")

# Compute metrics

```

```

cm = confusion_matrix(Y_test, Y_pred)
fpr, tpr, _ = roc_curve(Y_test, Y_proba) if Y_proba is not None else ([], []
[], [])
roc_auc = auc(fpr, tpr) if Y_proba is not None else None
precision, recall, _ = precision_recall_curve(Y_test, Y_proba) if Y_proba is not None else ([], [], [])
f1 = f1_score(Y_test, Y_pred)

# Plot Accuracy Score curve
if Y_proba is not None:
    thresholds = np.linspace(0, 1, 100)
    accuracy_scores = [accuracy_score(Y_test, (Y_proba > threshold).
astype(int)) for threshold in thresholds]
    axs[idx, 5].plot(thresholds, accuracy_scores, color='green', lw=2)
    axs[idx, 5].set_xlim([0.0, 1.0])
    axs[idx, 5].set_ylim([0.0, 1.05])
    axs[idx, 5].set_xlabel('Decision Threshold')
    axs[idx, 5].set_ylabel('Accuracy Score')
    axs[idx, 5].set_title(f'{name} Accuracy Score Curve')
else:
    axs[idx, 5].text(0.5, 0.5, 'No Accuracy Curve',
horizontalalignment='center', verticalalignment='center', fontsize=12)
    axs[idx, 5].set_title(f'{name} Accuracy Score Curve')

# Plot confusion matrix
sns.heatmap(cm, annot=True, fmt='d', cmap='Blues', ax=axs[idx, 0],
cbar=False, xticklabels=['Malignant', 'Benign'], yticklabels=['Malignant',
'Benign'])
axs[idx, 0].set_xlabel('Predicted Label')
axs[idx, 0].set_ylabel('True Label')
axs[idx, 0].set_title(f'{name} Confusion Matrix')

# Plot ROC curve
if Y_proba is not None:
    axs[idx, 1].plot(fpr, tpr, color='darkorange', lw=2, label=f'ROC curve
(area = {roc_auc:.2f})')
    axs[idx, 1].plot([0, 1], [0, 1], color='navy', lw=2, linestyle='--')
    axs[idx, 1].set_xlim([0.0, 1.0])
    axs[idx, 1].set_ylim([0.0, 1.05])
    axs[idx, 1].set_xlabel('False Positive Rate')
    axs[idx, 1].set_ylabel('True Positive Rate')
    axs[idx, 1].set_title(f'{name} ROC Curve')
    axs[idx, 1].legend(loc='lower right')
else:
    axs[idx, 1].text(0.5, 0.5, 'No ROC Curve',
horizontalalignment='center', verticalalignment='center', fontsize=12)

```

```

    axs[idx, 1].set_title(f'{name} ROC Curve')

# Plot Precision-Recall curve
if Y_proba is not None:
    axs[idx, 2].plot(recall, precision, color='blue', lw=2)
    axs[idx, 2].set_xlim([0.0, 1.0])
    axs[idx, 2].set_ylim([0.0, 1.05])
    axs[idx, 2].set_xlabel('Recall')
    axs[idx, 2].set_ylabel('Precision')
    axs[idx, 2].set_title(f'{name} Precision-Recall Curve')
else:
    axs[idx, 2].text(0.5, 0.5, 'No PR Curve', horizontalalignment='center',
                     verticalalignment='center', fontsize=12)
    axs[idx, 2].set_title(f'{name} Precision-Recall Curve')

# Plot feature importance for models that have it
if hasattr(model, 'feature_importances_'):
    importances = model.feature_importances_
    axs[idx, 3].barh(range(len(importances)), importances, align='center')
    axs[idx, 3].set_yticks(range(len(importances)))
    axs[idx, 3].set_yticklabels(bcancer.feature_names)
    axs[idx, 3].set_xlabel('Feature Importance')
    axs[idx, 3].set_title(f'{name} Feature Importance')
else:
    axs[idx, 3].text(0.5, 0.5, 'No Feature Importance',
                     horizontalalignment='center', verticalalignment='center', fontsize=12)
    axs[idx, 3].set_title(f'{name} Feature Importance')

# Plot F1 Score
axs[idx, 4].bar(name, f1, color='purple')
axs[idx, 4].set_ylim([0, 1])
axs[idx, 4].set_xlabel('Model')
axs[idx, 4].set_ylabel('F1 Score')
axs[idx, 4].set_title(f'{name} F1 Score')

# Train and plot decision boundary on two features
X_train_2d = X_train[:, [feature1, feature2]]
X_test_2d = X_test[:, [feature1, feature2]]
model_2d = model.__class__(**model.get_params()) # Create a new instance
with the same parameters
model_2d.fit(X_train_2d, Y_train)

X_set, y_set = X_train_2d, Y_train
X1, X2 = np.meshgrid(np.arange(start=X_set[:, 0].min() - 1, stop=X_set[:, 0].max() + 1, step=0.01),
                     np.arange(start=X_set[:, 1].min() - 1, stop=X_set[:, 1].max() + 1, step=0.01))

```

```

Z = model_2d.predict(np.array([X1.ravel(), X2.ravel()]).T).reshape(X1.shape)
axs[idx, 6].contourf(X1, X2, Z, alpha=0.75, cmap=ListedColormap(['red', ↴
'green']))
axs[idx, 6].scatter(X_set[:, 0], X_set[:, 1], c=y_set, ↴
cmap=ListedColormap(['red', 'green']), edgecolor='k')
axs[idx, 6].set_xlim(X1.min(), X1.max())
axs[idx, 6].set_ylim(X2.min(), X2.max())
axs[idx, 6].set_xlabel(bcancer.feature_names[feature1])
axs[idx, 6].set_ylabel(bcancer.feature_names[feature2])
axs[idx, 6].set_title(f'{name} Decision Boundary')

plt.tight_layout()
plt.show()

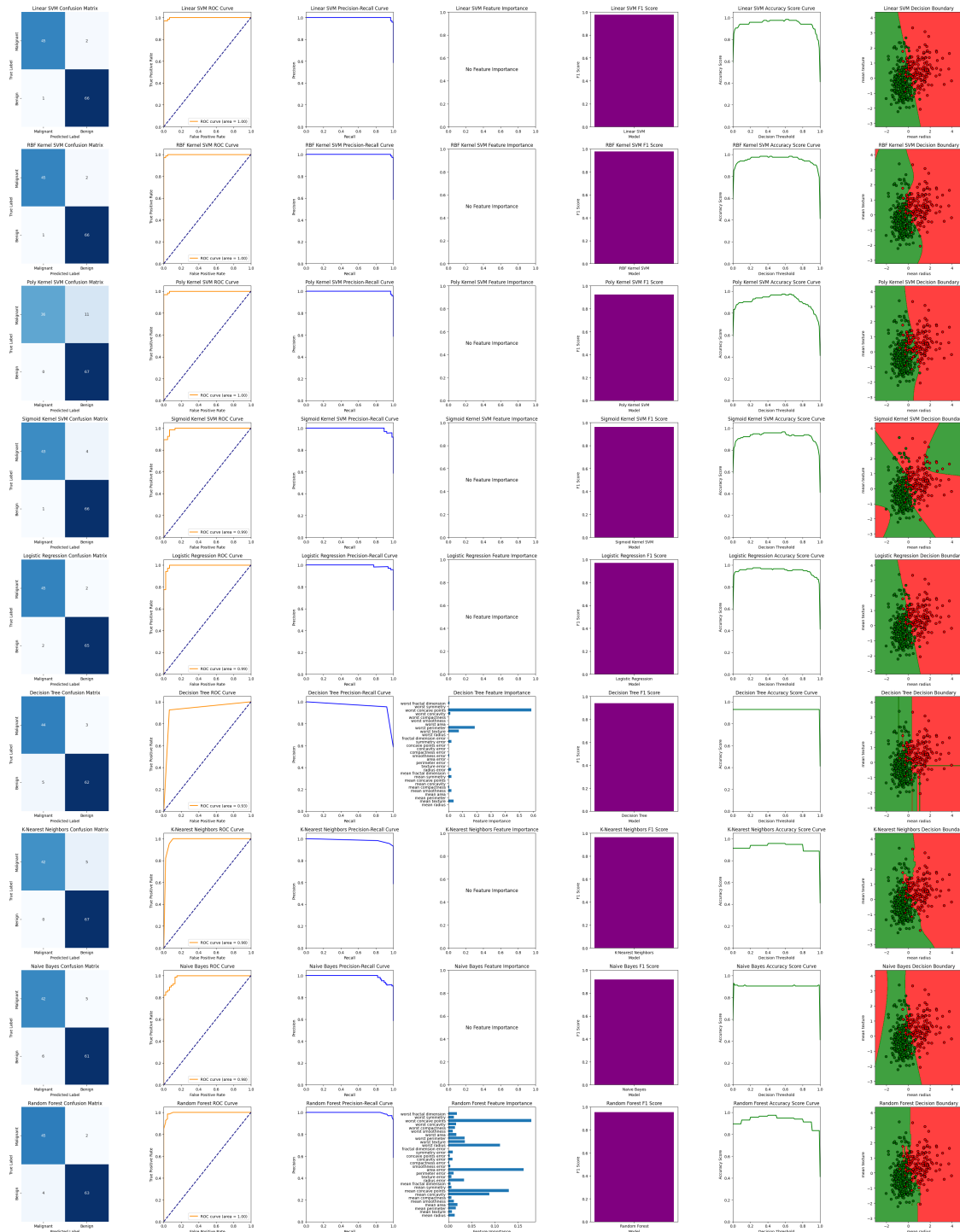
```

```

Precision = True Positives / (True Positives + False Positives)
Recall = True Positives / (True Positives + False Negatives)
F1 Score = 2 * (Precision * Recall) / (Precision + Recall)

```

Linear SVM Accuracy Score: 97.37%
 RBF Kernel SVM Accuracy Score: 97.37%
 Poly Kernel SVM Accuracy Score: 90.35%
 Sigmoid Kernel SVM Accuracy Score: 95.61%
 Logistic Regression Accuracy Score: 96.49%
 Decision Tree Accuracy Score: 92.98%
 K-Nearest Neighbors Accuracy Score: 95.61%
 Naive Bayes Accuracy Score: 90.35%
 Random Forest Accuracy Score: 94.74%



[9]: # Relative performance plots of all models - accuracy, ROC curve, PR curve, F_1 score etc.

Define the formulae as strings

```

precision_formula = "Precision = True Positives / (True Positives + False Positives)"
recall_formula = "Recall = True Positives / (True Positives + False Negatives)"
f1_score_formula = "F1 Score = 2 * (Precision * Recall) / (Precision + Recall)"

# Print the formulae
print(precision_formula)
print(recall_formula)
print(f1_score_formula, '\n')

# Define a dictionary of models
models = {
    'Linear SVM': SVC(kernel='linear', random_state=0, probability=True),
    'RBF Kernel SVM': SVC(kernel='rbf', random_state=0, probability=True),
    'Poly Kernel SVM': SVC(kernel='poly', degree=3, random_state=0,
                           probability=True),
    'Sigmoid Kernel SVM': SVC(kernel='sigmoid', random_state=0,
                               probability=True),
    'Logistic Regression': LogisticRegression(max_iter=10000, random_state=0),
    'Decision Tree': DecisionTreeClassifier(criterion='entropy',
                                             random_state=0),
    'K-Nearest Neighbors': KNeighborsClassifier(n_neighbors=5,
                                                 metric='minkowski', p=2),
    'Naive Bayes': GaussianNB(),
    'Random Forest': RandomForestClassifier(n_estimators=10,
                                             criterion='entropy', random_state=0)
}
}

# Initialize plot
fig, axs = plt.subplots(len(models) * 2, 4, figsize=(28, len(models) * 10))
fig.subplots_adjust(hspace=0.5)

# Split the data
X_train, X_test, Y_train, Y_test = train_test_split(standard_scaled_X, Y,
                                                    test_size=0.2, random_state=0)

# Select two features for decision boundary plot
feature1, feature2 = 0, 1 # example feature indices, change as needed

# Loop through models
for idx, (name, model) in enumerate(models.items()):
    # Train model on the full dataset
    model.fit(X_train, Y_train)

    # Predict results
    Y_pred = model.predict(X_test)

```

```

Y_proba = model.predict_proba(X_test)[:, 1] if hasattr(model, u
↪'predict_proba') else None
accuracy = accuracy_score(Y_test, Y_pred)
print(f'{name} Accuracy Score: {100 * accuracy:.2f}%')

# Compute metrics
cm = confusion_matrix(Y_test, Y_pred)
fpr, tpr, _ = roc_curve(Y_test, Y_proba) if Y_proba is not None else ([], u
↪[], [])
roc_auc = auc(fpr, tpr) if Y_proba is not None else None
precision, recall, _ = precision_recall_curve(Y_test, Y_proba) if Y_proba u
↪is not None else ([], [], [])
f1 = f1_score(Y_test, Y_pred)

# Plot Accuracy Score curve
if Y_proba is not None:
    thresholds = np.linspace(0, 1, 100)
    accuracy_scores = [accuracy_score(Y_test, (Y_proba > threshold).u
↪astype(int)) for threshold in thresholds]
    axs[2*idx + 1, 2].plot(thresholds, accuracy_scores, color='green', lw=2)
    axs[2*idx + 1, 2].set_xlim([0.0, 1.0])
    axs[2*idx + 1, 2].set_ylim([0.0, 1.05])
    axs[2*idx + 1, 2].set_xlabel('Decision Threshold')
    axs[2*idx + 1, 2].set_ylabel('Accuracy Score')
    axs[2*idx + 1, 2].set_title(f'{name} Accuracy Score Curve')
else:
    axs[2*idx + 1, 2].text(0.5, 0.5, 'No Accuracy Curve', u
↪horizontalalignment='center', verticalalignment='center', fontsize=12)
    axs[2*idx + 1, 2].set_title(f'{name} Accuracy Score Curve')

# Plot confusion matrix
sns.heatmap(cm, annot=True, fmt='d', cmap='Blues', ax=axs[2*idx, 0], u
↪cbar=False, xticklabels=['Malignant', 'Benign'], yticklabels=['Malignant', u
↪'Benign'])
axs[2*idx, 0].set_xlabel('Predicted Label')
axs[2*idx, 0].set_ylabel('True Label')
axs[2*idx, 0].set_title(f'{name} Confusion Matrix')

# Plot ROC curve
if Y_proba is not None:
    axs[2*idx, 1].plot(fpr, tpr, color='darkorange', lw=2, label=f'ROC_u
↪curve (area = {roc_auc:.2f})')
    axs[2*idx, 1].plot([0, 1], [0, 1], color='navy', lw=2, linestyle='--')
    axs[2*idx, 1].set_xlim([0.0, 1.0])
    axs[2*idx, 1].set_ylim([0.0, 1.05])
    axs[2*idx, 1].set_xlabel('False Positive Rate')

```

```

        axs[2*idx, 1].set_ylabel('True Positive Rate')
        axs[2*idx, 1].set_title(f'{name} ROC Curve')
        axs[2*idx, 1].legend(loc='lower right')
    else:
        axs[2*idx, 1].text(0.5, 0.5, 'No ROC Curve', horizontalalignment='center', verticalalignment='center', fontsize=12)
        axs[2*idx, 1].set_title(f'{name} ROC Curve')

# Plot Precision-Recall curve
if Y_proba is not None:
    axs[2*idx, 2].plot(recall, precision, color='blue', lw=2)
    axs[2*idx, 2].set_xlim([0.0, 1.0])
    axs[2*idx, 2].set_ylim([0.0, 1.05])
    axs[2*idx, 2].set_xlabel('Recall')
    axs[2*idx, 2].set_ylabel('Precision')
    axs[2*idx, 2].set_title(f'{name} Precision-Recall Curve')
else:
    axs[2*idx, 2].text(0.5, 0.5, 'No PR Curve', horizontalalignment='center', verticalalignment='center', fontsize=12)
    axs[2*idx, 2].set_title(f'{name} Precision-Recall Curve')

# Plot feature importance for models that have it
if hasattr(model, 'feature_importances_'):
    importances = model.feature_importances_
    axs[2*idx, 3].barh(range(len(importances)), importances, align='center')
    axs[2*idx, 3].set_yticks(range(len(importances)))
    axs[2*idx, 3].set_yticklabels(bcancer.feature_names)
    axs[2*idx, 3].set_xlabel('Feature Importance')
    axs[2*idx, 3].set_title(f'{name} Feature Importance')
else:
    axs[2*idx, 3].text(0.5, 0.5, 'No Feature Importance', horizontalalignment='center', verticalalignment='center', fontsize=12)
    axs[2*idx, 3].set_title(f'{name} Feature Importance')

# Plot F1 Score
axs[2*idx + 1, 0].bar(name, f1, color='purple')
axs[2*idx + 1, 0].set_ylim([0, 1])
axs[2*idx + 1, 0].set_xlabel('Model')
axs[2*idx + 1, 0].set_ylabel('F1 Score')
axs[2*idx + 1, 0].set_title(f'{name} F1 Score')

# Train and plot decision boundary on two features
X_train_2d = X_train[:, [feature1, feature2]]
X_test_2d = X_test[:, [feature1, feature2]]
model_2d = model.__class__(**model.get_params()) # Create a new instance with the same parameters
model_2d.fit(X_train_2d, Y_train)

```

```

X_set, y_set = X_train_2d, Y_train
X1, X2 = np.meshgrid(np.arange(start=X_set[:, 0].min() - 1, stop=X_set[:, 0].max() + 1, step=0.01),
                     np.arange(start=X_set[:, 1].min() - 1, stop=X_set[:, 1].max() + 1, step=0.01))
Z = model_2d.predict(np.array([X1.ravel(), X2.ravel()]).T).reshape(X1.shape)
axs[2*idx + 1, 1].contourf(X1, X2, Z, alpha=0.75,
                             cmap=ListedColormap(('red', 'green')))
axs[2*idx + 1, 1].scatter(X_set[:, 0], X_set[:, 1], c=y_set,
                           cmap=ListedColormap(('red', 'green')), edgecolor='k')
axs[2*idx + 1, 1].set_xlim(X1.min(), X1.max())
axs[2*idx + 1, 1].set_ylim(X2.min(), X2.max())
axs[2*idx + 1, 1].set_xlabel(bcancer.feature_names[feature1])
axs[2*idx + 1, 1].set_ylabel(bcancer.feature_names[feature2])
axs[2*idx + 1, 1].set_title(f'{name} Decision Boundary')

# Hide unused subplots
for ax in axs.flatten():
    if not ax.has_data():
        ax.axis('off')

plt.tight_layout()
plt.show()

```

```

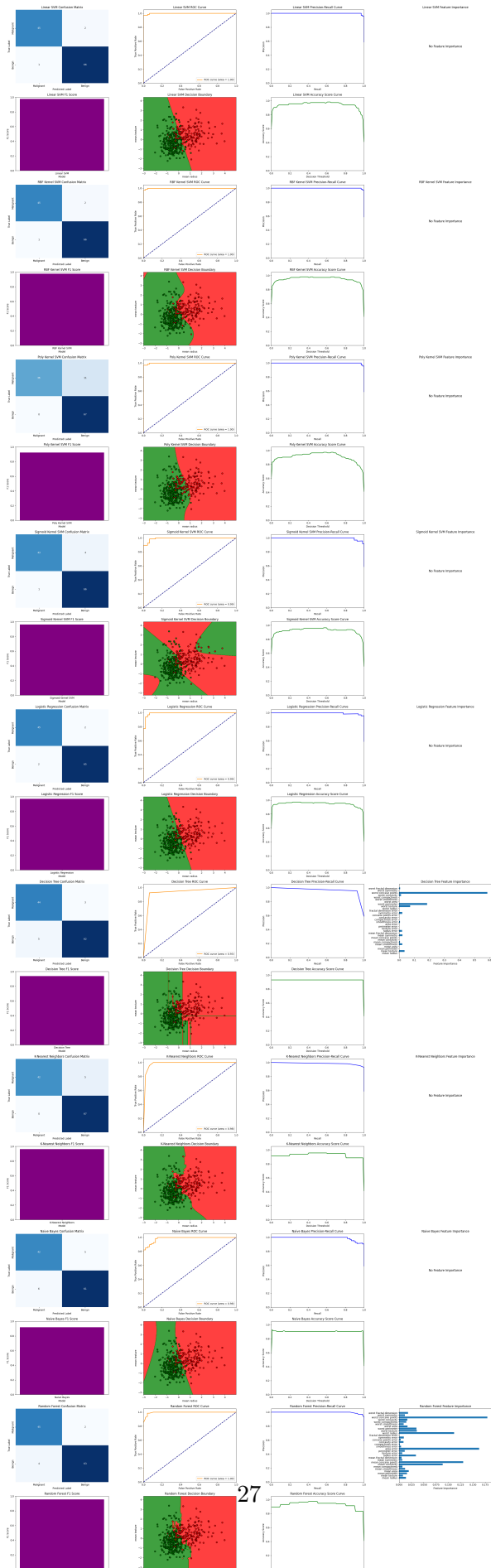
Precision = True Positives / (True Positives + False Positives)
Recall = True Positives / (True Positives + False Negatives)
F1 Score = 2 * (Precision * Recall) / (Precision + Recall)

```

```

Linear SVM Accuracy Score: 97.37%
RBF Kernel SVM Accuracy Score: 97.37%
Poly Kernel SVM Accuracy Score: 90.35%
Sigmoid Kernel SVM Accuracy Score: 95.61%
Logistic Regression Accuracy Score: 96.49%
Decision Tree Accuracy Score: 92.98%
K-Nearest Neighbors Accuracy Score: 95.61%
Naive Bayes Accuracy Score: 90.35%
Random Forest Accuracy Score: 94.74%

```



```
[1]: !jupyter nbconvert --to pdf Proj_04_Breast_Cancer_SVM_Full_4.ipynb
```

```
C:\Program Files\Python313\Scripts\jupyter-nbconvert.EXE\__main__.py:4:  
DeprecationWarning: Parsing dates involving a day of month without a year  
specified is ambiguous  
and fails to parse leap day. The default behavior will change in Python 3.15  
to either always raise an exception or to use a different default year (TBD).  
To avoid trouble, add a specific year to the input & format.  
See https://github.com/python/cpython/issues/70647.  
[NbConvertApp] Converting notebook Proj_04_Breast_Cancer_SVM_Full_3.ipynb to pdf  
[NbConvertApp] Support files will be in Proj_04_Breast_Cancer_SVM_Full_3_files\  
[NbConvertApp] Making directory .\Proj_04_Breast_Cancer_SVM_Full_3_files  
[NbConvertApp] Writing 146238 bytes to notebook.tex  
[NbConvertApp] Building PDF  
[NbConvertApp] Running xelatex 3 times: ['xelatex', 'notebook.tex', '-quiet']  
[NbConvertApp] Running bibtex 1 time: ['bibtex', 'notebook']  
[NbConvertApp] WARNING | b had problems, most likely because there were no  
citations  
[NbConvertApp] PDF successfully created  
[NbConvertApp] Writing 4595113 bytes to Proj_04_Breast_Cancer_SVM_Full_3.pdf
```