#### Team

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Github: https://github.com/manikantavarmanadimpally/SVM-classification.git

**Project Report: Iris Flower Classification** 

### Introduction:

The goal of this project was to build a classification model to predict the species of iris flowers based on their sepal and petal measurements. The popular Iris dataset, containing measurements for three species of 1iris flowers (Setosa, Versicolor, and Virginica), was utilized for this task.

## **Data Exploration and Preparation:**

The Iris dataset consists of 150 samples with four features: sepal length, sepal width, petal length, and petal width.

There are no missing values in the dataset.

The class distribution shows that each class has an equal number of samples (50 samples each).

The dataset was split into training and testing sets with an 80:20 ratio.

## **SVM Implementation:**

A Support Vector Machine (SVM) classifier was implemented using Python's Scikit-learn library. The linear kernel was used for the SVM classifier.

The model was trained on the training data.

K-fold Cross-Validation:

K-fold cross-validation with K=5 was applied to assess the performance of the SVM model.

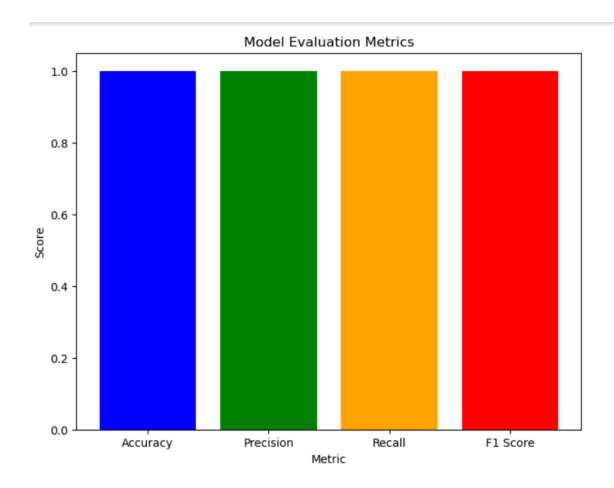
The dataset was shuffled before partitioning to prevent bias.

**Evaluation Metrics:** 

The performance of the model was evaluated using accuracy, precision, recall, and F1 score. The model achieved high scores for all evaluation metrics, indicating good classification performance.

# **Visualization of Results:**

Basic evaluation metrics (accuracy, precision, recall, and F1 score) were visualized using a bar plot.



## Conclusion:

The SVM model demonstrated strong performance in classifying iris flowers based on their sepal and petal measurements.

Further experimentation with different SVM kernels and hyperparameters could potentially improve the model's performance.

Recommendations:

Experiment with different SVM kernels (e.g., polynomial, radial basis function) to explore their impact on classification performance.

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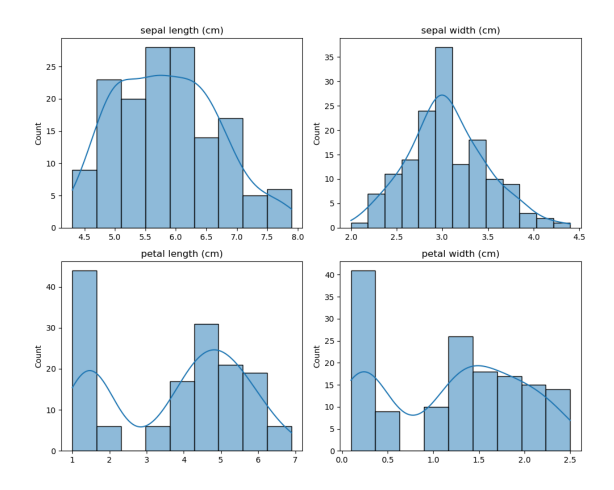
## April 26, 2024

```
[17]: # Importing necessary libraries
      from sklearn import datasets
      from sklearn.model_selection import train_test_split, cross_val_score
      from sklearn.preprocessing import StandardScaler, LabelEncoder
      from sklearn.svm import SVC
      from sklearn.metrics import accuracy score, precision score, recall score,

¬f1_score
      import numpy as np
      import matplotlib.pyplot as plt
      import seaborn as sns
[18]: # Load the Iris dataset
      iris = datasets.load_iris()
      X = iris.data
      y = iris.target
[19]: # Exploring the structure of the dataset
      print("Feature names:", iris.feature_names)
      print("Target names:", iris.target_names)
      print("Number of samples:", X.shape[0])
      print("Number of features:", X.shape[1])
      # Checking for missing values
      missing_values = np.isnan(X).sum()
      print("Missing values:", missing_values)
      # Checking class distribution
      print("Class distribution:", np.bincount(y))
     Feature names: ['sepal length (cm)', 'sepal width (cm)', 'petal length (cm)',
     'petal width (cm)']
     Target names: ['setosa' 'versicolor' 'virginica']
     Number of samples: 150
     Number of features: 4
     Missing values: 0
     Class distribution: [50 50 50]
```

```
[20]: # Visualizing feature distributions
      fig, axs = plt.subplots(2, 2, figsize=(10, 8))
      for i, feature in enumerate(iris.feature_names):
          sns.histplot(X[:, i], kde=True, ax=axs[i//2, i%2])
          axs[i//2, i%2].set_title(feature)
      plt.tight_layout()
      plt.show()
     C:\Users\nmkva\anaconda3\Lib\site-packages\seaborn\_oldcore.py:1119:
     FutureWarning: use_inf_as_na option is deprecated and will be removed in a
     future version. Convert inf values to NaN before operating instead.
       with pd.option_context('mode.use_inf_as_na', True):
     C:\Users\nmkva\anaconda3\Lib\site-packages\seaborn\ oldcore.py:1119:
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     future version. Convert inf values to NaN before operating instead.
```

with pd.option\_context('mode.use\_inf\_as\_na', True):



```
# Splitting the data into training and testing sets
X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2,u_orandom_state=42)

[22]: # Initializing the SVM classifier
svm = SVC(kernel='linear', C=1.0)

[23]: # Fitting the model
svm.fit(X_train, y_train)

[24]: # K-fold Cross-Validation
# Using 5-fold cross-validation
scores = cross_val_score(svm, X_train, y_train, cv=5)

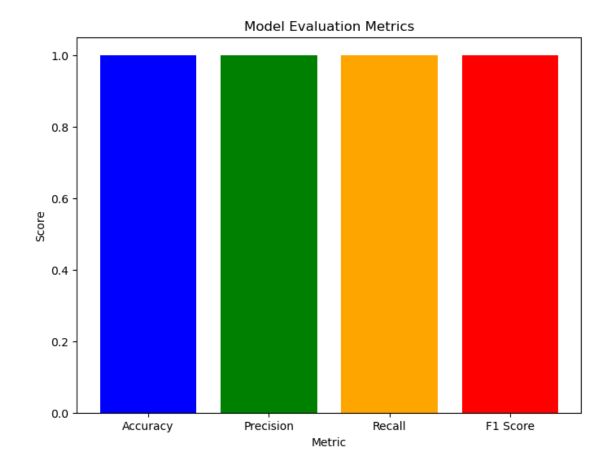
[25]: # Evaluation Metrics
# Making predictions
```

[21]: # SVM Implementation

```
y_pred = svm.predict(X_test)
[26]: # Calculating evaluation metrics
      accuracy = accuracy_score(y_test, y_pred)
      precision = precision_score(y_test, y_pred, average='weighted')
      recall = recall_score(y_test, y_pred, average='weighted')
      f1 = f1_score(y_test, y_pred, average='weighted')
[27]: # Printing evaluation metrics
      print("Accuracy:", accuracy)
      print("Precision:", precision)
      print("Recall:", recall)
      print("F1 Score:", f1)
     Accuracy: 1.0
     Precision: 1.0
     Recall: 1.0
     F1 Score: 1.0
[28]: # Visualize evaluation metrics
     metrics = {'Accuracy': accuracy, 'Precision': precision, 'Recall': recall, 'F1u

Score': f1}
      plt.figure(figsize=(8, 6))
      plt.bar(metrics.keys(), metrics.values(), color=['blue', 'green', 'orange', __

¬'red'])
      plt.title('Model Evaluation Metrics')
      plt.xlabel('Metric')
      plt.ylabel('Score')
      plt.show()
```



[]: