## Sri Sivasubramaniya Nadar College of Engineering, Chennai

(An autonomous Institution affiliated to Anna University)

Degree & Branch	B.E. Computer Science & Engineering	Semester	V
Subject Code & Name	ICS1512 & Machine Learning Algorithms Laboratory		
Academic year	2025-2026 (Odd)	Batch:2023-2028	Due date: 25/08/2025

## Experiment 3: Ensemble Prediction and Decision Tree Model Evaluation

## Aim and Objective:

To build classifiers such as Decision Tree, AdaBoost, Gradient Boosting, XGBoost, Random Forest, and Stacked Models (using SVM, Na¨ıve Bayes, Decision Tree) and evaluate their performance through 5-Fold Cross-Validation and hyperparameter tuning.

#### Libraries used:

- Numpy
- Pandas
- Matplotlib
- Seaborn
- sklearn
- xgboost

#### IMPORTING DATASET + EDA + FEATURE CORRELATION AND OTHER PLOTS:

```
# Importing dataset and required libraries
from ucimlrepo import fetch_ucirepo
import pandas as pd
import numpy as np
import matplotlib.pyplot as plt
import seaborn as sns

from sklearn.model_selection import train_test_split, GridSearchCV, StratifiedKFold, cross
from sklearn.preprocessing import StandardScaler
from sklearn.tree import DecisionTreeClassifier, plot_tree
from sklearn.pipeline import Pipeline
from sklearn.metrics import (
    accuracy_score, precision_score, recall_score, f1_score,
    roc_auc_score, roc_curve, precision_recall_curve, average_precision_score,
```

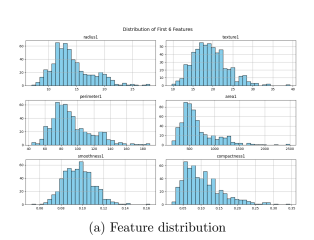
```
confusion_matrix, classification_report
from sklearn.utils import parallel_backend
# fetch dataset
breast_cancer_wisconsin_diagnostic = fetch_ucirepo(id=17)
# data (as pandas dataframes)
X = breast_cancer_wisconsin_diagnostic.data.features
y = breast_cancer_wisconsin_diagnostic.data.targets
# variable information
print(breast_cancer_wisconsin_diagnostic.variables)
# Adding columns headers
df = pd.DataFrame(X, columns=breast_cancer_wisconsin_diagnostic.variables.name)
# Removing unnecessary features
df.drop(columns=['ID','Diagnosis'], inplace=True)
# Malignant -> 1, Benign -> 0
df['Diagnosis'] = y
df['Diagnosis'] = df['Diagnosis'].map({'M': 1, 'B': 0})
# Splitting features and target
X = df.drop(columns=['Diagnosis'])
y = df['Diagnosis']
columns = df.columns.tolist()
print(columns)
# Check for missing values
y.isnull().sum()
''' Since there is no missing missing values in the dataset, we don't apply unwanted opera-
# Displaying first 5 records
print(df.head())
# 4. EDA
print("\nClass distribution:")
print(y.value_counts())
# Plot class distribution
sns.countplot(x='Diagnosis', data=df, palette = 'coolwarm')
plt.title("Class Distribution (Diagnosis)")
plt.show()
# Histograms of first 6 features
```

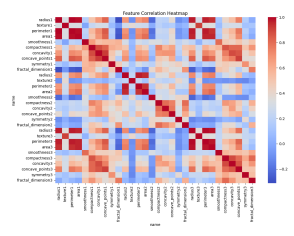
```
X.iloc[:, :6].hist(bins=30, figsize=(12, 8), color='skyblue', edgecolor='black')
plt.suptitle("Distribution of First 6 Features", fontsize=14)
plt.tight_layout()
plt.show()

# Feature correlation heatmap
plt.figure(figsize=(12, 8))
sns.heatmap(X.corr(), cmap='coolwarm')
plt.title("Feature Correlation Heatmap")
plt.show()

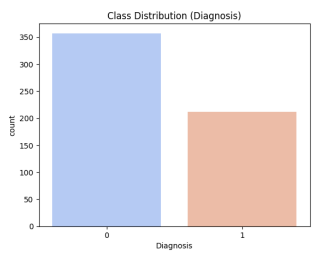
# Train test split
X_train, X_test, y_train, y_test = train_test_split(
    X, y, test_size=0.2, stratify=y, random_state=42
)
```

**Note:** This is common for all models





(b) Correlation Heatmap



(c) Class Distribution

Figure 1: Exploratory Data Analysis

### FUNCTIONS FOR MODEL TEST + PERFORMANCE METRICS + K - FOLD CV:

```
# Predict & Performance Analysis
def test_model(best_estimator, X_test,y_test):
    best_estimator = grid.best_estimator_
    y_pred = best_estimator.predict(X_test)
    y_proba = best_estimator.predict_proba(X_test)[:, 1]
    return performance_metrics(y_test, y_pred, y_proba)

# Performance metrics
def performance_metrics(y_true, y_pred, y_proba):
    acc = accuracy_score(y_test, y_pred)
    prec = precision_score(y_test, y_pred)
```

```
rec = recall_score(y_test, y_pred)
    f1 = f1_score(y_test, y_pred)
    roc_auc = roc_auc_score(y_test, y_proba)
    avg_precision = average_precision_score(y_test, y_proba)
    print("\nTest metrics")
    print(f"Accuracy: {acc:.4f}")
    print(f"Precision: {prec:.4f}")
    print(f"Recall: {rec:.4f}")
    print(f"F1: {f1:.4f}")
    print(f"ROC AUC: {roc_auc:.4f}")
    print(f"Avg Precision (PR AUC-ish): {avg_precision:.4f}")
    print("\nClassification report:\n", classification_report(y_test, y_pred))
    # Confusion matrix
    cm = confusion_matrix(y_test, y_pred)
    sns.heatmap(cm, annot=True, fmt='d', cmap='Blues')
    plt.xlabel("Predicted")
    plt.ylabel("Actual")
    plt.title("Confusion Matrix")
    plt.show()
    # 7. Curves: ROC & Precision-Recall
    fpr, tpr, _ = roc_curve(y_test, y_proba)
    precision, recall, _ = precision_recall_curve(y_test, y_proba)
    plt.figure(figsize=(6,4))
    plt.plot(fpr, tpr, label=f"ROC AUC = {roc_auc:.3f}")
    plt.plot([0,1],[0,1],'--', alpha=0.5)
    plt.xlabel("FPR"); plt.ylabel("TPR")
    plt.title("ROC Curve")
    plt.legend(); plt.show()
# Define multiple metrics
def kfoldCV(best_estimator, X, y, cv):
    scoring = {
        'precision': 'precision',
        'recall': 'recall',
        'f1': 'f1'
    }
    # Run cross-validation
    with parallel_backend('threading'):
        cv_results = cross_validate(
            best_estimator, X, y, cv=cv,
            scoring=scoring,
            return_train_score=False,
            n_{jobs}=-1
```

```
)
   n_splits = cv.get_n_splits()
    # Display per-fold results
    for i in range(n_splits):
        print(f"\nFold {i+1}: Precision: {cv_results['test_precision'][i]:.4f}
                                                                                  Recall: {
    # Display averages
    print("\n=== Average Metrics ===")
    print(f"Mean Precision: {cv_results['test_precision'].mean():.4f} + {cv_results['test_]
                          {cv_results['test_recall'].mean():.4f} ± {cv_results['test_recall']
    print(f"Mean Recall:
    print(f"Mean F1 Score: {cv_results['test_f1'].mean():.4f} + {cv_results['test_f1'].store: }
def feature_selection(estimator, X_train, y_train, method="model", top_k=None):
    feature_names = X_train.columns
    # ---- Model-based importance ----
    if method == "model":
        if hasattr(estimator, "feature_importances_"): # Trees, RF, XGB, GB
            importances = estimator.feature_importances_
        elif hasattr(estimator, "coef_"):
                                                          # Linear models
            importances = estimator.coef_.ravel()
        else:
            raise ValueError("Estimator does not support feature_importances_ or coef_. Us-
    # ---- Permutation importance (model-agnostic) ----
    elif method == "permutation":
        perm = permutation_importance(estimator, X_train, y_train, n_repeats=20, random_state)
        importances = perm.importances_mean
    else:
        raise ValueError("method must be either 'model' or 'permutation'.")
    # Build importance DataFrame
    feat_imp = pd.DataFrame({
        "Feature": feature_names,
        "Importance": importances
    }).sort_values(by="Importance", ascending=False).reset_index(drop=True)
    # If top_k specified
    if top_k:
        feat_imp = feat_imp.head(top_k)
    # Plot
    plt.figure(figsize=(8, 6))
   plt.barh(feat_imp["Feature"][::-1], feat_imp["Importance"][::-1])
   plt.title(f"Top {top_k if top_k else len(feat_imp)} Features ({method} importance)")
   plt.xlabel("Importance")
```

```
plt.ylabel("Feature")
plt.show()
return feat_imp
```

**Note:** These functions are common for all models

#### PIPELINE FOR DECISION TREES WITH GRIDSEARCHCV:

```
# Decision Tree Classifier with Hyperparameter Tuning
dtree = DecisionTreeClassifier(random_state=42)
param_grid = {
    'criterion': ['gini', 'entropy', 'log_loss'],
    'max_depth': [None, 5, 10, 15, 20],
    'min_samples_split': [2, 5, 10],
    'min_samples_leaf': [1, 2, 4],
    'max_features': [None, 'sqrt', 'log2']
}
# K-Fold Cross Validation
cv = StratifiedKFold(n_splits=5, shuffle=True, random_state=42)
grid = GridSearchCV(
    estimator=dtree,
    param_grid=param_grid,
    cv=cv,
    scoring='accuracy',
    n_{jobs=-1},
    verbose=2
)
with parallel_backend('threading'):
    grid.fit(X_train, y_train)
print("\nBest Parameters:", grid.best_params_)
print(f"Best CV Accuracy: {grid.best_score_ :.4f}")
# Test the decision tree model
best_estimator = grid.best_estimator_
test_model(best_estimator, X_test, y_test)
kfoldCV(best_estimator, X, y, cv)
```

#### PIPELINE FOR ADABOOST CLASSIFIER WITH GRIDSEARCHCV:

```
# AdaBoost Classifier with Hyperparameter Tuning
from sklearn.ensemble import AdaBoostClassifier
base_tree = DecisionTreeClassifier(random_state=42, max_depth=1) # stump
adaboost = AdaBoostClassifier(estimator=base_tree, random_state=42)
param_grid = {
    'n_estimators': [50, 100, 200],
    'learning_rate': [0.001, 0.01, 0.1, 1],
    'estimator__max_depth': [1, 2, 3], # tuning the base tree depth
    'estimator__criterion': ['gini', 'entropy']
}
cv = StratifiedKFold(n_splits=5, shuffle=True, random_state=42)
grid = GridSearchCV(
    estimator=adaboost,
    param_grid=param_grid,
    cv=cv,
    scoring='accuracy',
    n_{jobs=-1},
    verbose=2
)
with parallel_backend('threading'):
    grid.fit(X_train, y_train)
print("\nBest Parameters:", grid.best_params_)
print("Best CV Accuracy:", grid.best_score_)
# Test the decision tree model
best_estimator = grid.best_estimator_
test_model(best_estimator, X_test, y_test)
kfoldCV(best_estimator, X, y, cv)
```

#### PIPELINE FOR GRADIENT BOOSTING CLASSIFIER WITH GRIDSEARCHCV:

```
# Gradient Boosting Classifier with Hyperparameter Tuning
from sklearn.ensemble import GradientBoostingClassifier
gb = GradientBoostingClassifier(random_state=42)

param_grid = {
    'n_estimators': [100, 200, 300],  # number of boosting stages
    'learning_rate': [0.01, 0.1, 0.2],  # step size shrinkage
```

```
'max_depth': [3, 4, 5],
                                         # depth of individual trees
    'min_samples_split': [2, 5, 10],
                                         # minimum samples to split a node
    'min_samples_leaf': [1, 3, 5],
                                         # minimum samples per leaf
    'subsample': [1.0, 0.8]
                                         # fraction of samples for each tree (stochastic be
}
cv = StratifiedKFold(n_splits=5, shuffle=True, random_state=42)
grid = GridSearchCV(
    estimator=gb,
    param_grid=param_grid,
    cv=cv,
    scoring='accuracy',
    n_{jobs}=-1,
    verbose=2
)
with parallel_backend('threading'):
    grid.fit(X_train, y_train)
print("\nBest Parameters:", grid.best_params_)
print("Best CV Accuracy:", grid.best_score_)
# Test the decision tree model
best_estimator = grid.best_estimator_
test_model(best_estimator, X_test, y_test)
kfoldCV(best_estimator, X, y, cv)
```

#### PIPELINE FOR XGBOOST CLASSIFIER WITH GRIDSEARCHCV:

```
# XGBoost Classifier with Hyperparameter Tuning
xgb = XGBClassifier(
    eval_metric='logloss',
    use_label_encoder=False,
    random_state=42
)

param_grid = {
    'n_estimators': [100, 200],
    'learning_rate': [0.05, 0.1],
    'max_depth': [3, 4],
    'subsample': [1.0, 0.8],
    'colsample_bytree': [1.0, 0.8],
    'gamma': [0, 0.1]
}
```

```
cv = StratifiedKFold(n_splits=5, shuffle=True, random_state=42)
grid = GridSearchCV(
    estimator=xgb,
    param_grid=param_grid,
    cv=cv,
    scoring='accuracy',
    n_{jobs=-1}
    verbose=2
)
grid.fit(X_train, y_train)
print("\nBest Parameters:", grid.best_params_)
print("Best CV Accuracy:", grid.best_score_)
# Test the decision tree model
best_estimator = grid.best_estimator_
test_model(best_estimator, X_test, y_test)
kfoldCV(best_estimator, X, y, cv)
```

#### PIPELINE FOR RANDOM FOREST CLASSIFIER WITH GRIDSEARCHCV:

```
# Random Forest Classifier with Hyperparameter Tuning
from sklearn.ensemble import RandomForestClassifier
rf = RandomForestClassifier(random_state=42)
param_grid = {
     'n_estimators': [100, 200],
                                        # number of trees
    'max_depth': [None, 10, 20],  # depth of trees
'min_samples_split': [2, 5],  # min samples to split a node
'min_samples_leaf': [1, 2],  # min samples at a leaf
     'max_features': ['sqrt', 'log2'] # feature selection per split
cv = StratifiedKFold(n_splits=5, shuffle=True, random_state=42)
grid = GridSearchCV(
    estimator=rf,
    param_grid=param_grid,
    cv=cv,
    scoring='accuracy',
    n_{jobs=-1},
    verbose=2
```

```
)
   with parallel_backend('threading'):
        grid.fit(X_train, y_train)
   print("\nBest Parameters:", grid.best_params_)
   print("Best CV Accuracy:", grid.best_score_)
   # Test the decision tree model
   best_estimator = grid.best_estimator_
   test_model(best_estimator, X_test, y_test)
   kfoldCV(best_estimator, X, y, cv)
PIPELINES FOR STACKED ENSEMBLE MODELS:
    SVM + NAIVE BAYES + DECISION TREES (BASE MODELS):
  LOGISTIC REGRESSION (Final estimator):
   from sklearn.ensemble import StackingClassifier
   from sklearn.linear_model import LogisticRegression
   from sklearn.naive_bayes import GaussianNB
   from sklearn.svm import SVC
   # Define base learners
   svm_base = Pipeline([
        ("scaler", StandardScaler()),
        ("svc", SVC(probability=True, random_state=42))
   1)
   nb_base = GaussianNB()
   dt_base = DecisionTreeClassifier(random_state=42)
   # Define stack (example: Logistic Regression as final estimator)
    stack_clf = StackingClassifier(
        estimators=[
            ("svm", svm_base),
            ("nb", nb_base),
            ("dt", dt_base)
       ],
        final_estimator=LogisticRegression(max_iter=1000, random_state=42),
        n_{jobs}=-1
   )
   # Parameter grid (for meta-learner)
   param_grid = {
```

'final\_estimator\_\_C': [0.1, 1, 10], # Regularization strength

```
'final_estimator__penalty': ['12'],
                                              # Only 12 is supported with liblinear/saga
    'final_estimator__solver': ['lbfgs', 'saga']
}
# Cross-validation strategy
cv = StratifiedKFold(n_splits=5, shuffle=True, random_state=42)
# Grid search
grid = GridSearchCV(
    estimator=stack_clf,
    param_grid=param_grid,
    cv=cv,
    scoring='accuracy',
    n_{jobs}=-1,
    verbose=2
)
# Fit with parallel backend
with parallel_backend('threading'):
    grid.fit(X_train, y_train)
print("\nBest Parameters:", grid.best_params_)
print("Best CV Accuracy:", grid.best_score_)
# Test performance
best_estimator = grid.best_estimator_
test_model(best_estimator, X_test, y_test)
# K-fold CV evaluation
kfoldCV(best_estimator, X, y, cv)
# Feature importance (only works if final estimator supports it)
with parallel_backend('threading'):
    feat_imp = feature_selection(best_estimator, X_train, y_train, method="permutation", to
    print(feat_imp)
RANDOM FOREST (Final estimator):
from sklearn.ensemble import RandomForestClassifier
stack_clf_rf = StackingClassifier(
    estimators=[
        ("svm", svm_base),
        ("nb", nb_base),
        ("dt", dt_base)
    ],
```

```
final_estimator=RandomForestClassifier(random_state=42, n_jobs=-1),
     n_{jobs}=-1
 )
 param_grid_rf = {
     'final_estimator__n_estimators': [100, 200],
     'final_estimator__max_depth': [None, 10, 20],
     'final_estimator__min_samples_split': [2, 5],
     'final_estimator__min_samples_leaf': [1, 2],
     'final_estimator__max_features': ['sqrt', 'log2']
 }
 # Grid search
 grid_rf = GridSearchCV(
     estimator=stack_clf_rf,
     param_grid=param_grid_rf,
     cv=cv,
     scoring='accuracy',
     n_{jobs=-1},
     verbose=2
 )
 with parallel_backend('threading'):
     grid_rf.fit(X_train, y_train)
 print("\nBest Parameters:", grid_rf.best_params_)
 print("Best CV Accuracy:", grid_rf.best_score_)
 best_estimator_rf = grid_rf.best_estimator_
 test_model(best_estimator_rf, X_test, y_test)
 kfoldCV(best_estimator_rf, X, y, cv)
 with parallel_backend('threading'):
     feat_imp_rf = feature_selection(best_estimator_rf, X_train, y_train, method="permutation")
 print(feat_imp_rf)
SVM + KNN + DECISION TREES (BASE MODELS):
  LOGISTIC REGRESSION (Final estimator):
 from sklearn.neighbors import KNeighborsClassifier
 # Define KNN base
 knn_base = Pipeline([
     ("scaler", StandardScaler()),
     ("knn", KNeighborsClassifier())
 1)
```

```
# Define stack
stack_clf_knn = StackingClassifier(
    estimators=[
        ("svm", svm_base),
        ("dt", dt_base),
        ("knn", knn_base)
    ],
    final_estimator=LogisticRegression(max_iter=1000, random_state=42),
    n_{jobs}=-1
)
# Parameter grid (for meta-learner Logistic Regression)
param_grid_knn = {
    'final_estimator__C': [0.1, 1, 10],
    'final_estimator__penalty': ['12'],
    'final_estimator__solver': ['lbfgs', 'saga']
}
# Grid search
grid_knn = GridSearchCV(
    estimator=stack_clf_knn,
    param_grid=param_grid_knn,
    cv=cv,
    scoring='accuracy',
    n_jobs=-1,
    verbose=2
)
with parallel_backend('threading'):
    grid_knn.fit(X_train, y_train)
print("\nBest Parameters:", grid_knn.best_params_)
print("Best CV Accuracy:", grid_knn.best_score_)
best_estimator_knn = grid_knn.best_estimator_
test_model(best_estimator_knn, X_test, y_test)
kfoldCV(best_estimator_knn, X, y, cv)
# Logistic regression doesn't provide feature_importances_ directly
with parallel_backend('threading'):
    feat_imp_knn = feature_selection(best_estimator_knn, X_train, y_train, method="permuta"
    print(feat_imp_knn)
```

# CV TABLE FOR ALL MODELS:

MODEL	FOLD 1	FOLD 2	FOLD 3	FOLD 4	FOLD 5	AVG ACC
DECISION TREE	0.9386	0.8772	0.9298	0.9035	0.9649	0.9192
ADABOOST	0.9912	0.9035	0.9474	0.9561	0.9735	0.9543
GRADIENT BOOSTING	0.9825	0.9298	0.9561	0.9912	0.9823	0.9684
XGBOOST	0.9649	0.9474	0.9561	0.9649	0.9558	0.9578
RANDOM FOREST	0.9649	0.9386	0.9649	0.9649	0.9646	0.9596
STACKED MODEL	0.9912	0.9386	0.9737	0.9737	0.9823	0.9719

Table 1: Model performance across folds with average accuracy

### HYPERPARAMETER TRIALS - DECISION TREE MODEL

criterion	$\max_{-depth}$	Accuracy	F1 Score
log_loss	5	0.9385	0.9332
entropy	5	0.9319	0.9258
entropy	None	0.9253	0.9193
entropy	15	0.9187	0.9130
gini	10	0.9055	0.8977

Table 2: Hyperparameter trials - Decision Trees

### HYPERPARAMETER TRIALS - AdaBoost Model

$n_{-}estimators$	$learning_rate$	Accuracy	F1 Score
50	1	0.9758	0.9739
200	0.01	0.9604	0.9578
100	0.01	0.9451	0.9406
200	0.001	0.9319	0.9263
50	0.001	0.9011	0.8904

Table 3: Hyperparameter trials - AdaBoost

# HYPERPARAMETER TRIALS - Gradient Boosting Model

$n_{-}estimators$	learning_rate	$\max_{-depth}$	Accuracy	F1 Score
300	0.1	5	0.9824	0.9811
300	0.2	4	0.9692	0.9668
200	0.01	3	0.9560	0.9523
200	0.01	4	0.9407	0.9359
300	0.1	5	0.9143	0.9093

Table 4: Hyperparameter trials - Gradient Boosting

### HYPERPARAMETER TRIALS - XGBoost Model

$n_{-}estimators$	learning_rate	$\max_{-depth}$	gamma	Accuracy	F1 Score
200	0.1	3	0	0.9736	0.9718
100	0.1	3	0.1	0.9692	0.9670
100	0.05	3	0.1	0.9670	0.9648
100	0.05	3	0	0.9648	0.9625
100	0.05	4	0	0.9560	0.9531

Table 5: XGBoost Model

### **HYPERPARAMETER TRIALS - Random Forest Model**

$n_{-}estimators$	$\max_{-depth}$	criterion	Accuracy	F1 Score
200	10	gini	0.9670	0.9647
100	None	entropy	0.9648	0.9624
100	None	gini	0.9626	0.9600
200	20	entropy	0.9582	0.9552
200	20	$\log_{l}oss$	0.9538	0.9504

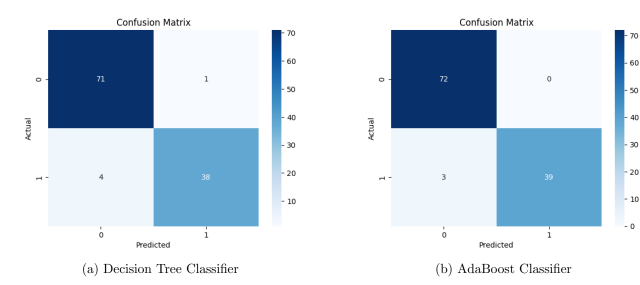
Table 6: Hyperparameter trials - Random Forest

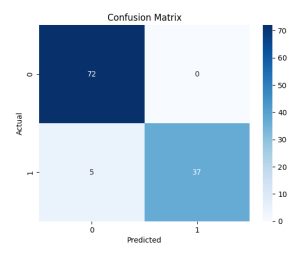
# STACKED ENSEMBLE MODEL TRIALS:

BASE MODELS	FINAL ESTIMATOR	ACC/F1
SVM + NB + DT	Logistic Regression	0.9825/0.9756
SVM + NB + DT	Random Forest	0.9649/0.9500
SVM + KNN + DT	Logistic Regression	0.9649/0.9500

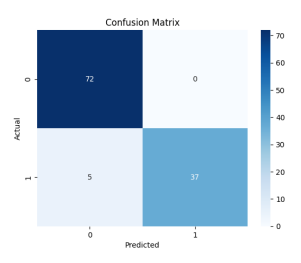
Table 7: Stack Ensemble Hyperparamter tuning

### **CONFUSION MATRICES:**

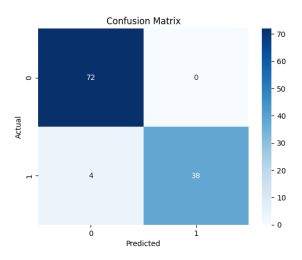




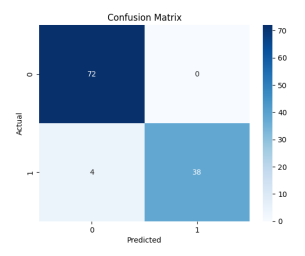
(a) Gradient Boosting Classifier



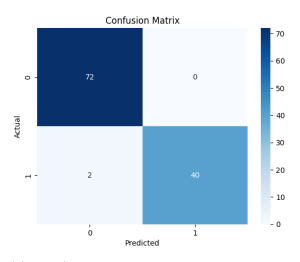
(c) Random Forest Classifier



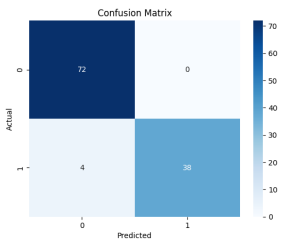
(e) SEM(SVM + NB + DT + Random Forest)



(b) XGBoost Classifier

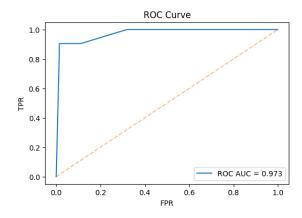


(d) SEM(SVM + NB + DT + Logistic Regression)

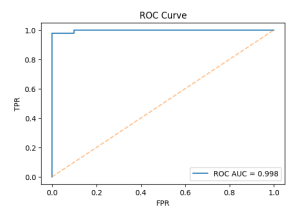


(f) SEM(SVM + KNN + DT + Logistic Regression)

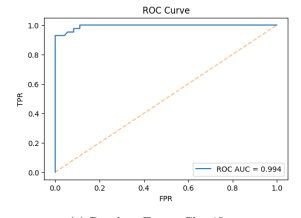
# ROC CURVES:



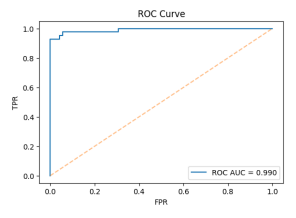
(a) Decision Tree Classifier



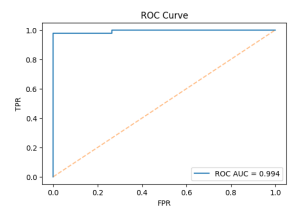
(c) Gradient Boosting Classifier



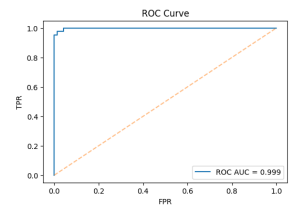
(e) Random Forest Classifier



(b) AdaBoost Classifier



(d) XGBoost Classifier



(f) SEM(SVM + NB + DT + Logistic)

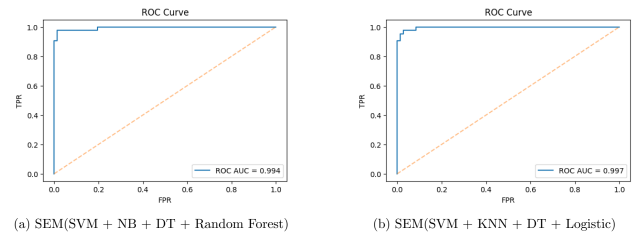
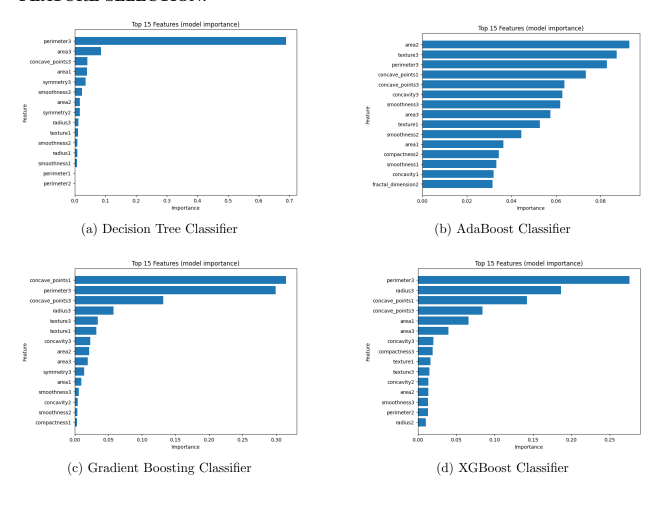


Figure 5: Comparison of different classifiers

#### FEATURE SELECTION:



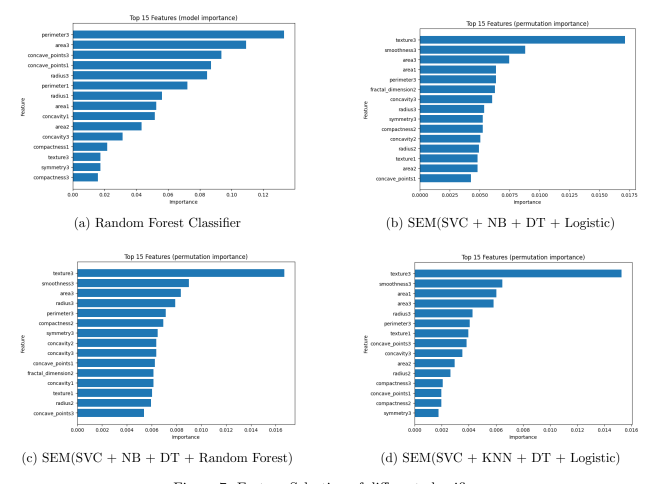


Figure 7: Feature Selection of different classifiers

#### Observations and Conclusion:

- 1. From the CV Table, we can see that **Stacked Ensemble Model with SVM**, **Naive Bayes** and **Decision Tree model as base and Logistic Regression as final estimator** achieved the best validation accuracy of **0.9719**.
- 2. Decision Tree performs underperformed compared to the ensemble models with average CV accuracy of 0.91 while others achieve an average of 0.95 accuracy
- 3. Random Forest achieved the best validation accuracy of **0.967** at **max depth of 10** and **n-estimator of 200**.
- 4. Stacked Ensemble Model with SVM, Naive Bayes and Decision Tree as base models and Logistic Regression as final estimator showed better generalization with best test accuracy of 0.9825 and roc auc of 0.999
- 5. From the above stacked ensembled models, it is evident that stacking improves the performance over base models.